USER GUIDE



# LifeScope<sup>™</sup> Genomic Analysis Software

## Command Shell

### DATA ANALYSIS METHODS AND INTERPRETATION

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# **About This Guide**

### Purpose

This guide is designed to help you quickly perform next-generation sequencing analyses using LifeScope<sup>™</sup> Software. These analyses support fragment, paired-end (PE) and long mate pair (LMP) library types of analyses. Specifically, the following workflows are supported:

- Whole genome resequencing
- Targeted resequencing
- Whole transcriptome resequencing
- Small RNA sequencing
- ChIP-Seq mapping
- MethylMiner<sup>™</sup> mapping

### **Prerequisites**

It is assumed that you have working knowledge of the:

- Linux<sup>®</sup> operating system
- Internet Protocol (IP) address of the LifeScope<sup>™</sup> Software cluster.
- Linux environment and know how to:
  - Navigate to directories.
  - Edit and save files in a text editor.
  - Run Linux shell scripts.
  - Run basic Linux commands such as chmod, ps, pwd, cd, echo, grep, and other commands.



# 1

# Introduction to LifeScope<sup>™</sup> Genomic Analysis Software

This introduction covers:

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### LifeScope<sup>™</sup> Software Overview

After years of development on analysis tools for SOLiD<sup>™</sup> data, in response to customer feedback, Life Technologies' LifeScope<sup>™</sup> Genomic Analysis Software enables fast translation of next-generation data for biologically meaningful results. LifeScope<sup>™</sup> Software matches the accuracy of the next generation 5500 Series SOLiD<sup>™</sup> Sequencers with Exact Call Chemistry (ECC) and streamlines your data analysis.

LifeScope<sup>TM</sup> Genomic Analysis Software is part of the LifeScope<sup>TM</sup> Genomic Analysis Features Solution. This informatics solution is comprised of genomics analysis software combined with a specified hardware platform. LifeScope<sup>™</sup> Software features: Seamless integration with the 5500 Series SOLiD<sup>TM</sup> Sequencer Performance-tuned algorithms for the 5500 Series and ECC Module Push-button workflows, intuitive user interface, and secure project management Optimized mapping and smaller file formats · Annotated variant reports, numerous charts, and select visualization tools for simple data interpretation Graphically driven configuration of multistep analysis workflows Ability to save and reuse workflows Ability to resume a workflow without repeating completed analyses Secure project-based data management specific to your data analysis Projects can be stored and data reanalyzed Complementing LifeScope<sup>TM</sup> Software features are the following types of data Data analyses analyses: Whole genome sequencing Targeted resequencing Whole exome sequencing Whole transcriptome RNA sequencing

• Small RNA sequencing

- MethylMiner<sup>TM</sup> mapping
- Detection of wide range of genomic variation, including:
  - SNP detection
  - Large and small indel detection
  - Copy number variation detection
  - Inversion detection
  - Fusion transcript detection
- Exon counting
- Splice finding

LifeScope<sup>™</sup> Software Components  $LifeScope^{TM}$  Software components include:

- LifeScope<sup>TM</sup> Server Software, the main software architecture that maintains the interaction of the graphical user interface (GUI) and compute engine. The server works with high performance cluster schedulers to maximize the computation demands of the finely tuned data analysis algorithms.
- The research GUI, which enables scientists to perform mapping and the detection of genomic variation from the convenience of a desktop computer, is comprised of a compute engine capable of mapping billions of reads of next-generation sequencing (NGS) data. The GUI provides auto-generated charts and plots for each genomic analysis run within the software.
- The command-line user interface, which gives bioinformaticians the power to customize workflows, and to manipulate every parameter available in each analysis module using the specific LifeScope<sup>TM</sup> command-line syntax.
- The Admin GUI Portal, which is an administration tool for managing user accounts and licensing permissions. Refer to *LifeScope<sup>™</sup> Genomic Analysis Software User Guide* (Part no. 4465676) for information on the Admin Portal.

### Workflows

This section describes a standard workflow, a project workflow, and several analysis workflows.

**Standard workflow** A standard workflow is a built-in series of commonly used analyses, which correspond to a common biological application. Using one of these workflows enables you to run an analysis with a minimum of setup required.

Standard workflows are provided for each of the analysis types listed above, on page 21.

In LifeScope<sup>™</sup> Software, running a standard workflow requires the following steps:

- 1. Create a project and an analysis.
- 2. Identify your input data (read set from your 5500 Series sequencing instrument).
- **3.** Identify the reference genome.
- 4. Choose the workflow to be executed on your data (from the above list).

- 5. Start and monitor your analysis.
- **6**. View the results of your analysis



**Analysis workflows** In LifeScope<sup>™</sup> Software, a workflow is a coordinated series of data analysis steps, with each step dependent only upon the steps that precede it. This section describes the flow of secondary and tertiary analyses in extensive detail, while primary analysis is performed on your 5500 Series sequencing instrument.

#### Secondary analysis

Secondary analysis begins with mapping. The input files required by mapping modules are eXtensible SeQuence files (XSQ) or converted color-space fasta (CSFASTA) files, and quality value (QV) files. Secondary analysis modules include:

SOLiD™ Accuracy Enhancement Tool (SAET)	Pairing
Mapping	Binary sequence Alignment Map statistics (BAMStats)

#### Tertiary analysis

The results of mapping and pairing in secondary analysis are used as input for tertiary analysis modules, which include:

SNP	Enrichment
CNV	Genomic annotations
Small indel	Gene and exon fusion
Large indel	Gene and exon count
Inversion	

Sequence Data (Transcriptomics) Epigenomics Genomics Whole Whole Targeted Small Genome ChIP-Seq Methylation Transcriptome Reseq RNA Reseg Mate-Pair Paired-End Paired-End Fragment Paired-End Fragment Fragment Fragment Fragment Fragment Paired-End SAET SAET SAET SAET SAET ¥ WT Mapping & Mapping & WT Mapping & Small RNA Mapping & Mapping Mapping Mapping & Pairing Pairing Pairing Mapping Mapping Mapping Mapping Pairing Pairing BAM BAM BAM BAM BAM BAM BAM BA<u>M</u> BAM 🛛 Stats BAM BAM 🚽 Stats 🚽 Stats 🕈 Stats Stats 🖌 Stats ↓ Stats 🕈 Stats Stats Stats Stats SNP SNP SNP SNP SNP Precursor Gene Gene Enrichment Enrichment Small RNA Fusion <u>Fusion</u> Exon CNV CNV CNV Mature Exon ¥ Small RNA Junction Junction Small Small Small Enriched BAM Small Small Enriched BAM Coverage Exon Exon Indel Indel Indel Indel Indel Count Count Large Large Gene Indel Gene Indel Count Count Inversions Inversions Coverage Coverage Annotations Annotations Annotations Annotations Annotations Legend Reporting Reporting Reporting Reporting Reporting Арр = Module Арр = Optional App = Off by default

File

Туре

= Output = Category

The illustration below shows the data analysis modules used in predetermined workflows.

Workflow	Analysis module	Library type	LifeScope <sup>™</sup> Software modules involved
ChIP-Seq	ChIP-Seq	Fragment	Secondary: • Fragment mapping • Mapping statistics-
Genomic resequencing	Genomic Resequencing	Fragment	Secondary: • Mapping • Mapping statistics
			Tertiary: • SNP Finding • CNV • Small indels • Annotations
		Mate-pair	Secondary: • Paired mapping • Mapping statistics
			Tertiary: • SNP Finding • CNV • Inversions • Large indels • Small indels • Annotations
		Paired-end	Secondary: • Paired mapping • Mapping statistics
			Tertiary: <ul> <li>SNP Finding</li> <li>Inversions</li> <li>Large indels</li> <li>CNV</li> <li>Small indels</li> <li>Annotations</li> </ul>
MethylMiner™	Methyl Miner	Fragment	Secondary: • Paired mapping • Mapping statistics
		Paired-end	Secondary: • Fragment mapping • Mapping statistics

The analysis workflows and the modules they execute are described in the following table.

Workflow	Analysis module	Library type	LifeScope <sup>™</sup> Software modules involved
Small RNA	Small RNA	Small RNA	Secondary:
			Small RNA mapping
			Tertiary:
			Small RNA count
			Small RNA coverage
Targeted resequencing	Targeted Resequencing	Fragment	Secondary:
			• SAET
			Fragment mapping
			Mapping statistics
			Tertiary:
			Enrichment
			SNP Finding
			Small indels
			Annotations
		Paired-end	Secondary:
			• SAET
			Paired mapping
			Mapping statistics
			Tertiary:
			Enrichment
			SNP Finding
			Small indels
			Annotations
Whole transcriptome	Whole Transcriptome	Fragment	Secondary:
			WT splice junction extractor
			WT fragment mapping
			Mapping statistics
			Tertiary:
			• WI counts
			WI coverage
			Splice finder
		Paired-end	Secondary:
			WI exon sequence extractor
			WI splice junction extractor
			Manning statistics
			Tartiany
			• Splice finder
			WT counts
			WT coverage

### Primary and secondary file types

Analysis type	File type	File name extension	File content	Used in
Primary	Raw reads file	*.xsq	Extensible sequence file (XSQ)	Secondary analysis
Secondary	BAM file	*.bam	Binary Alignment sequence Map (BAM), a generic file format used to store large numbers of nucleotide sequence alignments	Tertiary analysis

The following table lists the files used in secondary and tertiary analysis.

### Input and output file formats

All analysis modules use specific input files and produce specific output files. The input and output file requirements vary, depending on the type of analysis that you want to perform. The following table provides the names of the input and output file types.

Software or bioinformatics tool	Input file types	Output file type
Mapping tool	XSQ	ВАМ
MaToBAM tool	МА	Converts a MA file to a BAM file.
Mapping statistics (BAMStats)	ВАМ	Position error, probe error, multiple chart and statistics files
Small indel	ВАМ	GFFv3
diBayes	ВАМ	GFFv3, CSFASTA, consensus_calls
CNV	ВАМ	GFFv3, *.out
Large indel	ВАМ	GFFv3
Inversion	ВАМ	GFFv3, text
WT mapping	XSQ, FASTA, filter reference FASTA, WT GTF reference	ВАМ
WT counts	ВАМ	GTF
WT coverage	ВАМ	WIG
WT splice junction	ВАМ	WIG
Small RNA mapping	XSQ, GTF reference	ВАМ
Small RNA counts	ВАМ	GTF
Small RNA coverage	ВАМ	WIG

Introduction to LifeScope<sup>™</sup> Genomic Analysis Software Input and output file formats

# LifeScope<sup>™</sup> Genomic Analysis Software Installation

This chapter covers:

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### **Overview**

This chapter is intended for use by the LifeScope<sup>TM</sup> Software administrator. The chapter describes the LifeScope<sup>TM</sup> Genomic Analysis Software installation procedure, and system software and hardware requirements.

### Prerequisites

Some procedures in this chapter require that you:

- Know the Linux<sup>®</sup> operating system of the cluster on which LifeScope<sup>™</sup> Software is to be installed
- Know the Internet Protocol (IP) address of the cluster
- Have a login (ID) on the cluster Portions of the install require root access or your system administrator's assistance
- Know how to:
  - Navigate to directories in a Linux environment
  - Edit and save files in a text editor
  - Run Linux shell scripts
  - Run basic Linux commands such as chmod, ps, pwd, cd, echo, grep, and other commands

# Cluster hardware requirements

To successfully install and run the LifeScope<sup>™</sup> Software application, we recommend that your cluster meets the following hardware requirements:

CPU Speed	2 GHz Minimum	
Cores	8 Minimum	
Memory	24 GB Minimum per compute node	
Disk Space	500 GB of local or shared storage per node	
	200 MB for LifeScope <sup>™</sup> Software installation	
	225 GB for reference files installation <sup>‡</sup>	
	2 GB for examples installation	
	25 GB for performance verification data	

‡ You can use any type of shared storage to meet this requirement.

System LifeScope<sup>™</sup> Genomic Analysis Software requires a Linux<sup>®</sup> cluster with a TORQUE, Sun Grid Engine (SGE), or Load Sharing Facility (LSF) resource manager, with a requirements scheduler that can support scheduling policies and dynamic job priorities. Before you install LifeScope<sup>™</sup> Genomic Analysis Software, be sure that your cluster Server meets the following requirements: requirements • LifeScope<sup>™</sup> Genomic Analysis Software supports only Redhat 4.7 (or later version) and CentOS 4.7 (or later version) distributions on 64-bit platforms. PBS/TORQUE v2.3+, SGE v6.2+, or Platform LSF 7 Update 6. Note: If you use SGE, you must create a symmetric multiprocessing (SMP) parallel environment. Note: If you use LSF, it is highly recommended that LifeScope<sup>™</sup> Software jobs are non-preemptable by LSF. Before you install LifeScope<sup>™</sup> Software on the head node, pre-install compatible versions of these software packages on *all* compute nodes: Perl v.5.8.5 (or later version) Python 2.3 (or later version) GCC Compiler Ensure that the GNU Compiler Collection (GCC) compiler v3.4.6 or v4.1.2 installed and configured. While LifeScope<sup>™</sup> Software does not use the compiler itself, having the compiler installed and configured ensures that the necessary dynamically linked libraries are available. Ensure that there is no firewall restriction on the host running the LifeScope<sup>™</sup> Software Check firewall server. The LifeScope<sup>TM</sup> Software accesses the LifeScope<sup>TM</sup> Software license server on restrictions TCP/IP port 27000, by default. The LifeScope<sup>™</sup> Software UI clients and command shell clients access the LifeScope<sup>™</sup> Software server on TCP/IP port 9998 by default, although this port can be changed during the installation. Consult your system administrator to check the firewall access for LifeScope<sup>TM</sup> Software users.

#### Check DNS hostname

The DNS fully-qualified domain name and the local Linux hostname must match. Also, the forward and reverse lookup on the DNS against the hostname and its publicfacing IP address must be consistent. The LifeScope<sup>™</sup> Software licensing server may not work correctly if there are DNS discrepancies. Consult the system administrator to check this requirement.

### LifeScope<sup>™</sup> Software Administration

This section uses the acronyms in this list to describe installation roles and locations:

- **SA** The Linux system administrator for the cluster. The administrator's root access is required for user management and copying the data drive.
- LSA The LifeScope<sup>™</sup> Software administrator. The installation instructions recommend creating a new user account for the LSA. The LSA must run the installation script.
- LSU LifeScope<sup>™</sup> Software users. These users do not have admin permissions.
- LSBF The LifeScope<sup>™</sup> Software Binaries Folder. The LSBF is the installation directory.
- LSDF The LifeScope<sup>™</sup> Software Data Folder. The LSDF is a location for LifeScope<sup>™</sup> Software repositories, resources, and other files.

To install or run LifeScope<sup>TM</sup> Software, one person must be designated as the master LifeScope<sup>TM</sup> Software user. The master user also becomes the default LifeScope<sup>TM</sup> Software Administrator (LSA). The LSA and regular LifeScope<sup>TM</sup> Software Users (LSUs) must share a common UNIX<sup>®</sup> user group.

We recommend that the LSA not share the same user account as the UNIX System Administrator (SA), because of a requirement that all LSUs be members of the LSA's primary UNIX user group. The SA's primary UNIX user group is usually root. Non-administrator users typically are not allowed to be members of the root user group.

We recommend creating a new Linux user account, lifescope, for the LSA. The primary UNIX user group for the lifescope user should be the same group as most LifeScope<sup>TM</sup> Software Users (for example, the group users).

After installation, LifeScope<sup>™</sup> Software files and folders are owned by the LSA. All application jobs and processes are submitted by the LSA.

If you are not the LSA or if an LSA does not exist, please contact your System Administrator to have the master user and user group created. You must be logged in as the LSA to install LifeScope<sup>™</sup> Software.

Identify two shared folders for holding the software and data in a suitably common location, accessible to all LifeScope<sup>™</sup> Software users. Both locations should be accessible (mounted) from the compute nodes.

• Identify the LifeScope<sup>™</sup> Software Binaries Folder (LSBF) to install the lifescope software. This folder should be writable to LSA and readable to LSUs. Typically you may choose /opt/lifescope. If the LSBF does not exist, create it with appropriate permissions.

Identify the location of the LifeScope<sup>™</sup> Software Data Folder (LSDF) to hold all LifeScope<sup>™</sup> Software resources, repositories, pre-installation tar files, installer, examples, test data, and other files. This location should have read and write permissions for the LSA and all LSUs. This location also should be sufficiently large to hold the initial data and the results of future analysis. Typically you may choose /share/lifescope. If the LSDF does not exist, create it with appropriate permissions.

Make sure that you have the appropriate read and write permissions for the LSBF and LSDF folders, before proceeding with the installation.

### Installation workflow overview



### Copy data drive content

(System Administrator) Copy the data drive content to the LifeScope<sup>TM</sup> Software Data Folder (LSDF) on the LifeScope<sup>TM</sup> Software server, and change owner:group to those of the LifeScope<sup>TM</sup> Software Administrator (LSA).

The data drive's file system is ext3, which typically is supported only on Linux.

The drive content includes the following:

• Files for use with hg18 and hg19 genomes:

- Hash tables for the main mapping schemes
- Mappability files for use with the CNV analysis module
- Targeted regions files
- LifeScope<sup>TM</sup> Software reference repository files:
  - Hg18 and hg19 reference files and filter reference files
  - Genomic annotation files (dbSNP)
  - MiRNA database annotation files
- Data files for installation verification

The reference files downloaded in this step become the reference repository for your LifeScope<sup>™</sup> Software. The location of the reference files (the referenceData directory) is required during installation.

**IMPORTANT!** If you add your own reference files to the reference repository, you must follow the instructions in "Add new reference files" and "Prepare reference files" on page 520.

### Download LifeScope<sup>™</sup> Software

- (LSA) Visit the LifeScope<sup>™</sup> Software project page: http://solidsoftwaretools.com/gf/project/lifescope
- 2. On the project page, you must:
  - a. Read and accept the LifeScope<sup>™</sup> Software End User License Agreement.
  - **b**. Create an email with the following information:
    - Your acceptance of the LifeScope<sup>™</sup> Software End User License Agreement.
    - Your LifeScope<sup>™</sup> Software license key.
    - Your solidsoftwaretools account name.

Send the email to:

#### lifescope@lifetech.com

Instructions to download LifeScope<sup>™</sup> Software are emailed to you.

- 3. Go to the download page, as directed by the emailed instructions.
- **4.** Download LifeScope<sup>™</sup> Software (.tar.gz file, for example, *LifeScope-2.0r8380-2011040333223344.tar.gz*) to the LifeScope<sup>™</sup> Software Data Folder (LSDF).
- 5. Download the associated installer (install.sh file) to the LifeScope<sup>™</sup> Software Data Folder (LSDF).

### Install LifeScope<sup>™</sup> Software

This procedure provides typical, general instructions for installing LifeScope<sup>™</sup> Software. The actual procedures for your site might vary, depending on the configuration of the LifeScope<sup>™</sup> Software cluster and the installation options you select.

# Hardware configuration

This section explains the three types of installation based on hardware configuration. You must select your configuration when prompted by the installer in "Select the type of installation:", step 6 on page 35.

- Standalone Workstation LifeScope<sup>™</sup> Software is installed completely on the local workstation. There is no access to a cluster. Resource management software is still required on the workstation.
- **Cluster** LifeScope<sup>™</sup> Software is installed on a cluster with a headnode, a set of compute nodes, shared storage, node local storage, and resource management software, as described in "Cluster hardware requirements" on page 30.
- Workstation with Remote Submission The local workstation, instead of the cluster headnode, is used for running the LifeScope<sup>™</sup> Software server. The requirements for the workstation are:
  - A regular Linux 64-bit workstation with the following:
    - 250 GB harddrive
    - 4 GB RAM
    - Dual core
  - The operating system requirements are the same as those listed for the cluster in "Server requirements" on page 30.

LifeScope<sup>™</sup> Software jobs are submitted from the workstation onto the cluster. This requires that you *first* setup the workstation as described in "Set up a workstation with remote submission" on page 37.

You must be logged in as the LifeScope<sup>™</sup> Software administrator (LSA) on the local workstation, and not on the cluster. The installer (install.sh) also must be run on the local workstation and not on the remote cluster. You must provide the IP address of the cluster head node, and the LSA username and password, when asked by the installer. This information is required for the workstation to automatically login to the cluster for submitting jobs.

By default the password appears as plain text in the file <LSBF>/etc/ analysis/system.properties file. If the LSA is setup for password-less login as described in "Configure the workstation" on page 38, then you can give a blank password when asked by the installer, and prevent the password from appearing in the properties file.

# Authentication realm

This section explains the authentication realms supported by LifeScope<sup>™</sup> Software. You must supply your realm type when prompted by the installer in "Select an authentication realm to be used with LifeScope<sup>™</sup> Software:", step 3 on page 37.

- LifeScope This authentication realm refers to the LifeScope<sup>™</sup> Software application user database. If this realm is chosen, user passwords are maintained in the software database. Passwords are stored as one-way MD5 digests for security reasons. Users must be manually created in this realm. There is no relationship between the application users and Linux<sup>®</sup> host system users. User Account Management can be done by the LSA using the LifeScope<sup>™</sup> Software Admin module (included in this installation).
- LDAP This realm refers to an LDAP-compliant authentication server (OpenLDAP, Active Directory, etc.). If this realm is chosen, user passwords are *not* stored in LifeScope<sup>™</sup> Software. User credentials are authenticated against the configured LDAP server. Users need not be manually created in LifeScope<sup>™</sup>

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	<ul> <li>Software with this realm (though they may be). A valid LDAP server address and LDAP Bind DN must be provided in the configuration. User Account Management can be done by the LSA using the LifeScope<sup>™</sup> Software Admin module (included in this installation).</li> <li>Host – This realm refers to the Linux host machine on which the LifeScope<sup>™</sup> Software server is running. If this realm is chosen, user passwords are <i>not</i> stored in LifeScope<sup>™</sup> Software. User credentials are authenticated against the local machine using SSH (therefore uses whatever authentication scheme SSH uses, for example, NIS, /etc/passwd). There is a one-to-one relationship between the application users and Linux host system users. Users need not be manually created in LifeScope with this realm.</li> </ul>
	<b>Note:</b> If you change the authentication realm after installation, you must stop and restart the LifeScope <sup>TM</sup> Software server.
	<b>Note:</b> Changing a realm does not remove users from the previous realm. However, users might not be able to login in the new realm, unless they both exist in the new realm and are enabled in the new realm.
Installation instructions	<ol> <li>(LSA) Go to the downloaded folder LSDF: cd /path/to/<lsdf></lsdf></li> <li>Run this UNIX<sup>®</sup> command: chmod +x install.sh</li> </ol>
	to make the downloaded install.sh file executable.
	3. Run the installer: ./install.sh
	<ul> <li>4. Installation options include:</li> <li>1) Install the LifeScope<sup>™</sup> Genomic Analysis Software.</li> <li>2) Re-configure the LifeScope<sup>™</sup> Software.</li> <li>3) Request a LifeScope<sup>™</sup> Software license.</li> <li>4) Register a LifeScope<sup>™</sup> Software license.</li> <li>5) Exit this application.</li> <li>To install LifeScope<sup>™</sup> Software, type 1.</li> </ul>
	<b>5.</b> At the prompt, type <b>y</b> to begin installation.
	<ul> <li>6. Select the type of installation:</li> <li>1) Standalone Workstation Server (default)</li> <li>2) Cluster</li> <li>3) Workstation w/Remote Submission</li> <li>Review "Hardware configuration" on page 34 to determine your installation type.</li> </ul>
	<ul> <li>7. Enter the location where LifeScope<sup>™</sup> Software is to be installed. The default directory is /opt/lifescope.</li> <li>Accept the default or change the directory. Provide the location you have chosen for the LifeScope<sup>™</sup> Software Binaries Folder (LSBF):</li> <li>/path/to/<lsbf></lsbf></li> </ul>

If the directory you enter already exists, you are prompted "Do you still want to use this location?"

**8.** Enter the location for reference files. Provide the destination location of the data drive contents (from the "Copy data drive content" step on page 32):

```
/path/to/<LSDF>/referenceData
```

Note: LifeScope reference files are required for full functionality of LifeScope<sup>™</sup> Software. The default directory is /data/results/referenceData.

**9.** Enter the location for reads. This entry sets the location of the reads repository. You may choose

/path/to/<LSDF>/reads

The default directory is /data/results/reads.

**10.** Enter the location for projects. This entry sets the location of the projects repository. You may typically choose

/path/to/<LSDF>/projects
The default directory is /data/results/projects.

**11.** Enter the location for Binary Alignment Map (BAM) file repository. You may choose

/path/to/<LSDF>/bams

The default directory is /data/results/bams.

**12.** Enter the location for the scratch location for LifeScope. The default directory is /scratch.

**Note:** The scratch location is used as a temporary work space for submitted jobs. For best performance, cluster and remote server installations require a scratch location on the local file system of each *compute node* of the system, not on the file system of the *head node*.

**13.** Enter the IP address or Fully Qualified Domain Name (FQDN) for the LifeScope<sup>™</sup> Software server, for example: 192.168.1.27.

The address of the system where LifeScope<sup>™</sup> Software is installed is required to successfully access and run the software. The system address is the IP address or Fully Qualified Domain Name (FQDN) of this system or head node.

**Note:** Do not use the name localhost or IP address 127.0.0.1. The address should be the public-facing name or IP address of the LifeScope<sup>™</sup> Software server. The LSUs access the software using this name or IP address. Typically, the address is the domain name system (DNS) name of the server.

Test the connection to the IP address or FQDN to verify a connection to the system.

Enter the Web URL port address to be used to access the LifeScope Server. Default is 9998.

Configure the<br/>license server1. Enter the IP address or Fully Qualified Domain Name (FQDN) for the LifeScope<br/>License Server.<br/>The installer tests the connection to the given IP address or FQDN to verify a<br/>connection to the system.

2. Enter the port address for the license server to be used with LifeScope<sup>™</sup> Software.
- **3.** Select an authentication realm to be used with LifeScope<sup>TM</sup> Software:
  - 1) LifeScope<sup>TM</sup> Software (default)
  - 2) Lightweight Directory Access Protocol (LDAP)
  - 3) Host

Review "Authentication realm" on page 34 to determine your realm type.

**Note:** If you change the authentication realm after installation, you must stop and restart the LifeScope<sup>TM</sup> Software server.

**Note:** Changing a realm does not remove users from the previous realm. However, users might not be able to login in the new realm, unless they both exist in the new realm and are enabled in the new realm.

- **4.** Select the cluster resource manager to be used by LifeScope<sup>™</sup> Software:
  - 1) PBS/Torque (default)
  - 2) SGE
  - 3) LSF
- **5.** Enter the cluster-resource-manager-job-submission-queue-name to be used by LifeScope, for example, *lifescope*.
- 6. Enter the number of nodes available on the cluster. Default is 4.
- 7. Enter the number of cores for each compute node. Default is 8.
- 8. Enter the memory size (in GB) for each compute node. Default is 22.

**IMPORTANT!** Allow 2 GB for the compute nodes' OS. For example, for compute nodes with 24 GB, enter 22; for compute nodes with 48 GB, enter 46.

Continue installing<br/>LifeScope™1. At the prompt "Do you want to change any of these entries?" type n to accept the<br/>entries.<br/>The selected LifeScope™ Software configurations are saved.2At the prompt "Do you want to configurations are saved.

**2.** At the prompt, "Do you want to continue installing the LifeScope package?" type **y** to continue the installation.

#### Set up a workstation with remote submission

This section describes prerequisites for the install type *Workstation w/Remote Submission*. Skip this section if your installation type is either of these:

- 1) Standalone Workstation Server (default)
- 2) Cluster

In the *Workstation w/Remote Submission* scenario, LifeScope<sup>™</sup> Software is installed a workstation but submits analysis jobs to a remote cluster. The LifeScope<sup>™</sup> Software server and licensing server run on the local workstation. The analysis jobs are submitted to the remote cluster using SSH. LifeScope<sup>™</sup> Software services do *not* run on the cluster headnode.

The workstation and cluster nodes share common file system locations through NFS mounting or other means. The installation location <LSBF>/lifescope and the LifeScope<sup>TM</sup> Software data folder <LSDF> must be on the common shared file system. Both the <LSBF>/lifescope and the <LSDF> folders must first be created on the cluster. (See "LifeScope<sup>TM</sup> Software Administration" on page 31 for a description of <LSBF> and <LSDF>.)

The cluster headnode must be accessible (via SSH) from the workstation while the analysis is executing.

As with the normal cluster installation, the LifeScope<sup>™</sup> Software administrator account must exist on the cluster, as described in "LifeScope<sup>™</sup> Software Administration" on page 31.

### Configure the workstation

These instructions are only for Workstation w/Remote Submission, option 3) in "Select the type of installation:", step 6 on page 35. The IT system administrator performs these configuration steps. The commands given in this section are not applicable to all cluster and workstation combinations. The given commands are expected to serve as general guidelines to the system administrator to give an example of the procedures required. The actual commands to be used may vary depending on the system configuration.

- 1. Note down the UID and GID (user and group ids) of the LifeScope<sup>™</sup> Software administrator (LSA) on the cluster.
- **2.** Create a user and group on the workstation with the same UID and GID as that of the LSA on the cluster. If the LSA user account already exists on the workstation, the IT system administrate must modify the UID and GID of that user account on the workstation. For example:

usermod -u <UID> <LSA>

- 3. Set up password-less login for the LSA user account from the workstation to the cluster, using ssh-keygen, .ssh/authorized\_keys file on the destination, etc.
- **4.** On the workstation, mount the location of the *<LSBF>* and *<LSDF>* directories from the cluster, using the same path. The path used on the workstation must be the same as the path used on the cluster.

Example commands are:

mkdir -p /path/to/<LSBF>
mkdir -p /path/to/<LSDF>
mount -t nfs cluster\_ip:/path/to/<LSBF> /path/to/<LSBF>
mount -t nfs cluster\_ip:/path/to/<LSDF> /path/to/<LSDF>

These commands may require root permissions. These are example commands. The actual commands may vary according to the system administrator's choice.

You are now ready to proceed with the installation procedures described in these sections:

- "Hardware configuration" on page 34
- "Authentication realm" on page 34
- "Installation instructions" on page 35

### Activate the LifeScope<sup>™</sup> Software License Key

	(LSA) These instructions describe how to apply the License Key. The process includes (each of these steps is described below):
	<ul> <li>Obtaining the MAC address of the server that has LifeScope<sup>™</sup> Software</li> </ul>
	Activating your License Key and obtain a license file specific to your server
	<ul> <li>Applying the license file to activate your copy of LifeScope<sup>™</sup> Software</li> </ul>
	When applied to LifeScope <sup><math>TM</math></sup> Software, the core license file grants up to five users access to the software for one year.
Prerequisites	Before activating the LifeScope $^{\text{\tiny TM}}$ Software License Key, please ensure that:
	• You have downloaded and installed LifeScope <sup>™</sup> Software on computer hardware that meets the LifeScope <sup>™</sup> Software minimum hardware requirements (please see www.lifetechnologies.com/LifeScope for more details).
	<ul> <li>You have performed the DNS hostname check described in "Check DNS hostname" on page 31.</li> </ul>
	• You are familiar with using command-line Linux <sup>®</sup> , and you have the necessary permissions to log on to the server that has LifeScope <sup>™</sup> Software.
	• You have a LifeScope <sup><math>TM</math></sup> Software License Key.
Internet connectivity	Direct internet connectivity for the host computer is highly recommended for the license activation process.
Check the computer date	Check that the computer date matches the current date and time. Each license has a start date. If the computer date is earlier than the license start date, access to the software is disabled until the license start date. If you need to reset the computer date, only reset it <i>forward</i> . <b>Do not</b> reset the computer date backward; doing so prevents access to LifeScope <sup>™</sup> Software. Contact your system administrator (SA) if you need to change the date on the system.
	<b>IMPORTANT!</b> Resetting your computer date backwards disables your license key and prevents the operation of LifeScope <sup>™</sup> Software.
Obtain the MAC address	<b>IMPORTANT!</b> Run the Linux commands used in this procedure on the computer hardware that has LifeScope <sup>™</sup> Software installed on it. If you want to run the LifeScope <sup>™</sup> license server on a different machine, then you must obtain the MAC address for that machine.
	INDODTANTI The MAC eddress used for line size Life Compa <sup>TM</sup> Coffeen and the Life
	IMPURIANT! The MAC address used for licensing LifeScope <sup>™</sup> Software should be associated with a permanent Ethernet card on the host computer system. If there is a possibility that eth0 is not a permanent Ethernet card, consult your system administrator to determine which Ethernet card should be used instead. If the Ethernet card associated with the LifeScope <sup>™</sup> Software license is replaced on the host system, a new license file must be generated for your account.

To obtain the MAC address of the server that has LifeScope<sup>™</sup> Software installed on it:

2

- 1. Run the Linux command
  - /sbin/ifconfig A paragraph similar to the one shown below appears:

[corona@foshtddev02 ~]\$ /sbin/ifconfig

- eth0 Link encap:Ethernet HWaddr 00:1C:23:BB:E1:F3
  inet addr:10.1.1.1 Bcast:10.1.1.255 Mask:255.255.255.0
  inet6 addr: fe80::21c:23ff:febb:e1f3/64 Scope:Link
  UP BROADCAST RUNNING MULTICAST MTU:1500 Metric:1
  RX packets:259842678 errors:0 dropped:0 overruns:0 frame:0
  TX packets:296616310 errors:0 dropped:0 overruns:0 carrier:0
  collisions:0 txqueuelen:1000
  RX bytes:214032251167 (199.3 GiB) TX bytes:339284293393 (315.9 GiB)
  Interrupt:169 Memory:f4000000-f4012800
- eth1 Link encap:Ethernet HWaddr 00:1C:23:BB:E1:F1
  inet addr:167.116.6.72 Bcast:167.116.6.255 Mask:255.255.255.0
  inet6 addr: fe80::21c:23ff:febb:e1f1/64 Scope:Link
  UP BROADCAST RUNNING MULTICAST MTU:1500 Metric:1
  RX packets:25669852 errors:0 dropped:0 overruns:0 frame:0
  TX packets:82980132 errors:0 dropped:0 overruns:0 carrier:0
  collisions:0 txqueuelen:1000
  RX bytes:10648346642 (9.9 GiB) TX bytes:117855364190 (109.7 GiB)
  Interrupt:169 Memory:f8000000-f8012800

lo Link encap:Local Loopback inet addr:127.0.0.1 Mask:255.0.0.0 inet6 addr: ::1/128 Scope:Host UP LOOPBACK RUNNING MTU:16436 Metric:1 RX packets:11049203 errors:0 dropped:0 overruns:0 frame:0 TX packets:11049203 errors:0 dropped:0 overruns:0 carrier:0 collisions:0 txqueuelen:0 RX bytes:6069766283 (5.6 GiB) TX bytes:6069766283 (5.6 GiB)

2. In the paragraph labeled eth0, the first line should contain text that starts with HWaddr. The MAC address follows this text, and should look similar to 00:30:48:9F:72:76

Obtain the MAC Note: In this method, the MAC address is referred to as the "Computer ID." Address with the If you have difficulty finding the MAC address, the LifeScope<sup>TM</sup> Installer can find the LifeScope<sup>™</sup> the address for you. To use the LifeScope  ${}^{{}^{\mathrm{TM}}}$  Installer: Installer 1. From the software download directory, run install.sh The LifeScope<sup>™</sup> Installer information shown below appears: This is the LifeScope Genomic Analysis Software installation and licensing program Install the LifeScope Genomic Analysis Software. The following options require successful installation of LifeScope: 2) Re-configure the LifeScope Software. Request a LifeScope Software license. Register a LifeScope Software license. 5) Exit this application. Please select the operation you would like to perform (Q/q to quit) [1]: 3 To request a LifeScope product license for operating the application: Enter the following URL into a Web browser. This will present a form to request a LifeScope product application license. https://licensing.appliedbiosystems.com/activation/lifescope ۰ Enter the Computer ID listed below on the form. If this utility is being run on a system other than the system running LifeScope, this Computer ID is invalid. Stop and re-run this utility on the system where LifeScope is installed. \*\*\*\*\*\*\*\*\* ComputerID: 00:1C:23:BB:E1:F3 \*\*\*\*\*\*\*\*\*\*\*\*\*\* The License Key you must enter on the form is the license key you were given with the LifeScope system. When the form is completed and submitted, a LifeScope product license will be sent to the email address specified on the form. If multiple people are to receive this key their email addresses should be entered in the "CC'ed" field of the form. Once the LifeScope product license key is received, execute "Step 2" of this utility to register the license with the License Server. 2. Choose option 3 ("Request a LifeScope product license.") to display the information about product licensing, including the MAC address of the first Ethernet card.

#### Activate License Key and obtain license file

To activate the license and obtain the license file associated with your MAC address:

 Visit the LifeScope<sup>™</sup> License activation page, shown below, at https://licensing.appliedbiosystems.com/activation/lifescope



#### LifeScope License Activation

Complete the fields below. Fields with an \* are required.

*Computer ID		
*License Key		0
*Email Address		
Cc		
*Country	Choose One	*
*First Name		
Middle Name		
*Last Name		

- 2. Enter the MAC address in the **Computer ID** field.
- **3.** Enter the License Key included in the LifeScope<sup>™</sup> 2 Genomic Analysis Software license instructions.
- 4. Enter the email address where your license file (.lic) should be sent.
- Fill in the remaining information, then click Submit to activate your license.
   Note: Your license begins the moment you activate your License Key.

The License File "Lifescope.lic" is sent to you as an attachment in an email from "RightNow AB Technical Support <RightNow.ABTechnicalSupport@lifetech.com>".

#### Apply the License File to LifeScope<sup>™</sup> Software

(LSA) To apply the license to your copy of LifeScope<sup>™</sup> Software:

1. Download the license file (.lic) to the system that has LifeScope<sup>™</sup> Software has installed on it.

If you are using another computer for email, you can use tools such as ftp or winscp to transfer the .lic file to your LifeScope<sup>TM</sup> Software system. For more information on transferring files, contact your system administrator.

2. Move the .lic file to

<*lifescope-installed-dir*>/server/licenses/ Your LifeScope<sup>™</sup> Software is now licensed.

IMPORTANT! Keep a backup copy of your license file in a safe place.

Later in "Start the licensing server" on page 46, in Chapter 3, Before You Begin, you start the license server and confirm that the license has been correctly applied.

You have completed the LifeScope<sup>™</sup> licensing process.

After the license has been activated, refer to the *LifeScope<sup>™</sup> Genomic Analysis Software User Guide* (Part no. 4465696) administration appendix for instructions to make LifeScope<sup>™</sup> Software available to users.

LifeScope<sup>™</sup> Software installation is completed. However, you are recommended to do at least one installation verification step. These steps are optional but confirm that the software is functioning properly and that the LifeScope<sup>™</sup> Software Data Folder (LSDF) is properly configured. The optional installation verification steps are:

- The demo analyses
- The performance verification tests

See Chapter 4, "Verify the LifeScope<sup>TM</sup> Genomic Analysis Software Installation" on page 49 for instructions on optionally running example analysis workflows and optionally performing verification tests.

#### Download documentation and additional resources

To ensure you have the latest version of LifeScope<sup>™</sup> Software, please visit http:// solidsoftwaretools.com/gf/project/lscope\_release and look for the Download option. You also find user documentation and the latest information and technical resources. The full resources package includes pre-installation scripts, smaller data files, the installation wizard, and performance verification scripts. Additional links and online support resources are also available at this site.

Integrative<br/>Genomics Viewer<br/>(IGV)You can optionally install the Integrative Genomics Viewer (IGV), shown on page 44,<br/>available from the Broad Institute. The IGV is a visualization tool for interactive<br/>exploration of large, integrated datasets. It directly reads BAM files, which enables you<br/>to easily view and inspect alignments against the genome.

For more information about the IGV, go to www.broadinstitute.org/igv/.

A ICV				
File View Tracks Help				
Human hg 18	*	chr22 v chr22	60 👚 🏟 🔳	-
		p13 p12 p11.2 q	11.1 q11.21 q11.23 q12.1 q12.2 q12.3	q13.1 q13.2 q13.31 q13.3
	NAME DATA FILE DATA TYPF	nb 10 mb		40 mb 51
diBayesFrag-1.bam Cov erage diBayesFrag-1.bam		[D - 60] 	Zoom in to see alignments.	
diBayesFrag-1.bam Cove			Zoom in to see alignments.	
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RefSeq genes		▶	H HEREINEN HEREITEN KANNEN HEREITEN HEREITEN HEREITEN HEREITEN HEREITEN HEREITEN HEREITEN HEREITEN HEREITEN HER	GGAL TNRC6B TTLL1 RIBC2 FAM19A5
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## Before You Begin

This chapter covers:

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#### **Overview**

This chapter is used by both the LifeScope<sup>™</sup> Genomic Analysis Software administrators and also LifeScope<sup>™</sup> Genomic Analysis Software users.

For LifeScope<sup>™</sup> Software administrators, this chapter provides general prerequisites to complete after installation and before you run LifeScope<sup>™</sup> Genomic Analysis Software, depending on the LifeScope<sup>™</sup> Software cluster configuration.

Some pre-requisite administration procedures described in this section require that you:

- Know the IP address of the LifeScope<sup>™</sup> Software cluster
- Have a login ID on the LifeScope<sup>TM</sup> Software cluster
- Know how to:
  - Navigate to directories in a Linux environment
  - Edit and save files in a text editor
  - Run Linux shell scripts
  - Run Linux commands such as ls, pwd, cd, echo, grep, and other common utilities

See Chapter 2, "LifeScope<sup>™</sup> Genomic Analysis Software Installation" on page 29 for information on installing LifeScope<sup>™</sup> Software.

For LifeScope<sup>™</sup> Software users, this chapter describes the following procedures:

- Set your shell environment to use LifeScope<sup>™</sup> Software
- Start the LifeScope<sup>™</sup> Software servers
- Verify your browser version

#### Set the LifeScope<sup>™</sup> Software environment

To set your environment to run LifeScope<sup>™</sup> Software, add the LifeScope<sup>™</sup> Software bin directory to your PATH and export the new PATH variable. Enter this command every time you open a Linux shell window to run LifeScope<sup>™</sup> Software:

export PATH=<installdir>/bin:\$PATH

where *installdir* is the LifeScope<sup>TM</sup> Software installation directory.

The *installdir* is the *LSBF*>/lifescope directory defined in the administrator instructions ("LifeScope<sup>TM</sup> Software Administration" on page 31).

If BioScope<sup>™</sup> Software is not used on the same machine, you can make your PATH setting permanent by modifying your local .bashrc file or the global /etc/bashrc files. Contact your system administrator for more information.

BioScope<sup>™</sup> Software users' PATH variable If you install LifeScope<sup>TM</sup> Software on a machine that also has BioScope<sup>TM</sup> Software installed and the BioScope<sup>TM</sup> Software is actively being used, you must follow these instructions:

- LifeScope<sup>™</sup> Software users must *not* use their shell profile script to update the PATH variable with the LifeScope<sup>™</sup> Software bin directory. If the LifeScope<sup>™</sup> Software bin directory is added to the PATH, BioScope<sup>™</sup> Software cannot run correctly.
- **2.** Run BioScope<sup>™</sup> Software and LifeScope<sup>™</sup> Software in different Linux<sup>®</sup> shell windows.
- **3.** Set and export your PATH variable *every* time you open a Linux shell window to run LifeScope<sup>TM</sup> Software server or client software.

#### Start LifeScope<sup>™</sup> Software servers and check servers' status

This section describes starting the LifeScope<sup>™</sup> Software servers and checking their status. Commands described in this section are run in the Linux shell (not in the LifeScope<sup>™</sup> Software command shell).

The LifeScope<sup>™</sup> Software licensing server must be started first, before the LifeScope<sup>™</sup> Software web server. Only the LifeScope<sup>™</sup> Software administrator (LSA) may start the licensing server.

Start the licensing	To start the licensing server, follow these steps:
server	<ol> <li>Make sure LifeScope<sup>™</sup> Software binaries are in the PATH environment variable. If the PATH variable is not set properly, set PATH using this Linux<sup>®</sup> command: export PATH=<lifescope-installed-dir>/bin:\$PATH</lifescope-installed-dir></li> </ol>
	<b>2.</b> Start the license server, using this Linux command:

lscope-lmgrd.sh start

If the license server is already running, a second instance is not started.

Check the licensing server

1. Use the following command to check if the LifeScope<sup>™</sup> Software licensing server is running:

```
ps -ef | grep lmgrd
```

If the LifeScope<sup>TM</sup> Software licensing server is not running, the Linux command prompt returns (with no output). If the LifeScope<sup>TM</sup> Software licensing server is running, this command displays two processes, one for the license manager daemon (lmgrd) and one for the vendor daemon (lifetech). Example output when the licensing server is running is:

```
lifescope 20447 1 0 Mar04 00:00:00 ./lmgrd -c /opt/
lifescope/lifescope/server/licenses/LifeScope.lic
lifescope 20448 20447 0 Mar04 00:00:00 lifetech -T
foshtdvv12.na.ab.applera.net 11.9 4 -c /opt/lifescope/
lifescope/server/licenses/LifeScope.lic -lmgrd_port 6979
--lmgrd_start 4d59fb1f
```

**2.** Verify the license status and the available user licenses, shown below, using the Linux command:

lscope-lmgrd.sh status

```
[panakkj1@fospanakkj1d02 licenses]$ cd /share/apps/lifescope/server/licenses/
[panakkj1@fospanakkj1d02 licenses]$ ls -l
total 8
-rwxrw-r-- 1 panakkj1 panakkj1 1144 May 5 17:50 LifeScope.lic
[panakkj1@fospanakkj1d02 licenses]$ echo $PATH
/share/apps/lifescope/bin:/usr/lib64/qt-3.3/bin:/usr/kerberos/bin:/usr/local/bin:/usr/bin:/
sr/X11R6/bin:/home/panakkj1/bin
[panakkj1@fospanakkj1d02 licenses]$ lscope-lmgrd.sh start
Started License Server
[panakkj1@fospanakkj1d02 licenses]$ lscope-lmgrd.sh status
lmutil - Copyright (c) 1989-2010 Flexera Software, Inc. All Rights Reserved.
Flexible License Manager status on Thu 5/5/2011 17:53
License server status: 27001@fospanakkj1d02.ads.invitrogen.net
    License file(s) on fospanakkj1d02.ads.invitrogen.net: /share/apps/LifeScope-2.0.r0-8663
0412120300/server/licenses//LifeScope.lic:
fospanakkjld02.ads.invitrogen.net: license server UP (MASTER) v11.9
Vendor daemon status (on fospanakkj1d02.ads.invitrogen.net):
  lifetech: UP v11.9
Feature usage info:
Users of LTC.BIOSCP.USERS:
                            (Total of 5 licenses issued; Total of 0 licenses in use)
Users of LTC.BIOSCP.LAUNCH: (Total of 1 license issued; Total of 0 licenses in use)
Users of LTC.BIOSCP.CONCURRENT: (Total of 1 license issued; Total of 0 licenses in use)
```

	<b>3.</b> Check that you have activated the same number of licenses that you ordered by cross-verifying the number with the result of the status command.
	You must start the license server every time the system undergoes a reboot.
Start the LifeScope™	Only the LifeScope <sup>™</sup> Software administrator (LSA) must start the LifeScope <sup>™</sup> Software server. The command to start the LifeScope <sup>™</sup> Software server is:
Software server	lscope-server.sh start
	The server by default uses port 9998, but the default port number might be changed during installation. To run on a different port, use the following version:
	lscope-server.sh start -p port
	If you start the server on a port other than the default 9998, you must check that the firewall is also enabled for the new port number. See "Check firewall restrictions" on page 30.
	If the port is busy, the server start fails.
	If the server is already running, a second instance is not started. The server issues the following message:
	Web server already running (pid=nnnnn) on port nnnn
	You must start the LifeScope <sup>™</sup> Software server every time the system undergoes a reboot.
Check the LifeScope™	The following command checks if the LifeScope <sup>™</sup> Software server is running:
Software server	ps -ei   grep webserver
status	If the LifeScope <sup>™</sup> Software server is not running, the Linux command prompt returns (with no output). Example output when the web server is running is:
	<pre>lifescope 23521 1 0 13:26 pts/1 00:00:05 java -server -Xmx3072m -XX:PermSize=512m -XX:MaxPermSize=512m - XX:+UseParallelGC com.lifetechnologies.bioscope.webservice. WebServer -b /opt/lifescope/lifescope</pre>

#### Verify the browser version

 $LifeScope^{^{\rm TM}}\ Genomic\ Analysis\ Software\ supports:$ 

- Internet Explorer<sup>®</sup> versions 6 and 7
- Mozilla<sup>®</sup> 3.0.1

## Verify the LifeScope<sup>™</sup> Genomic Analysis Software Installation

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#### **Overview**

This chapter describes procedures and tests that the LifeScope<sup>™</sup> Software administrators should perform after the software has been installed. LifeScope<sup>™</sup> Software users may skip this chapter.



#### Workflow



#### Verify the installation

(LSA) Two mechanisms are provided to verify the installation:

- Demos A set of individual analysis modules with small data sets. The demos ٠ take approximately 30 minutes or less to complete.
- Performance verification A realistic computation scenario that takes ٠ approximately 24 hours to complete.

**IMPORTANT!** You are strongly recommended to run one or both of the verification procedures. The procedures are listed as optional because you are not required to run both of them. Run at least one of the optional verification procedures.

You may run the demo analyses and performance verification scripts as the LifeScope<sup>TM</sup> Software administrator or as a LifeScope<sup>TM</sup> Software user. See "Create user accounts" on page 53 create a LifeScope<sup>TM</sup> Software user account.

#### (Optional) Download and install demos files

(LSA) This step is required in order to run the optional sample analyses, "(Optional) Run demo analyses" on page 51.

Example scripts and data for demo LifeScope<sup>™</sup> Software analyses are available from the following site:

http://solidsoftwaretools.com/gf/project/lscope\_release

Included in this download are the required PLN and INI files for running individual LifeScope<sup>™</sup> Software analysis modules, and scripts to run the analyses in the LifeScope<sup>™</sup> Software command shell.

Follow these steps to install the demo files:

- 1. Download the .tar.gz file to the LifeScope<sup>™</sup> Software Data Folder (LSDF).
- 2. Go to the LSDF directory and untar the .tar.gz file:

cd <LSDF>
tar -xzf exampleWorkflows.tar.gz

**Note:** The demo analyses are not recommended as the first experience for users new to LifeScope<sup>™</sup> Software. The standard workflows (Chapter 7, Run a Standard Workflow Analysis) are recommended for new users to learn LifeScope<sup>™</sup> Software command-line analyses.

#### (Optional) Run demo analyses

(LSA) Run demo analyses, which takes approximately 30 minutes.

These instructions require the following (in addition to the software installation):

- The default reference repository, which includes hg18 and hg19 assemblies. The reference repository is created during the "Copy data drive content" on page 32 and "Install LifeScope<sup>TM</sup> Software" on page 33 procedures.
- The demos files and sample data installed during the "(Optional) Download and install demos files" on page 51.

Follow these steps to execute the demos in one run:

- Follow the instructions in "Set the LifeScope<sup>™</sup> Software environment" on page 46 to set the LifeScope<sup>™</sup> Software environment.
- **2**. Go to the demos directory:

cd <LSDF>/examples/demos

**3**. Execute this command in the Linux shell:

./runall.sh -u username -w password

LifeScope<sup>™</sup> Genomic Analysis Software Command Shell User Guide

where *username* and *password* must be for the LifeScope<sup>™</sup> Software administrator or for a valid LifeScope<sup>™</sup> Software user account. This script runs each demo one at a time.

**Check results** The results files are generated in the projects repository, at a location which is set during installation. Its default location is at *<LSDF>/projects*. Within the projects repository directory, the demos results are generated at the following directories:

username/examples/analysis\_name/outputs/sample\_name

Replace *username* with the user name used to run runAll.sh. The default is lifescope, the LSA account. The string examples is the project name used by all demos. *analysis\_name* is the name of the analysis module used by a demo, and is for example, cnv, dibayes, enrichment, etc. Each *analysis\_name* folder has an outputs subfolder. The final folder, *sample\_name*, is determined by the sample in the input data.

Each of the demo analyses creates a log file named summary.log in the lifescope/ examples/analysis\_name directory. Check for a success message in the last line of each log file.

The logs and output files are overwritten on subsequent runs executed by the same user.

#### (Optional) Download and install performance verification scripts

(LSA) This step is required in order to run the optional performance verification tests, "(Optional) Run the performance verification test" on page 52.

The performance verification test data file are in the LSDF directory, from the "Copy data drive content" step on page 32.

The performance verification test scripts are available from the following site:

http://solidsoftwaretools.com/gf/project/lscope\_release/ performanceVerificationTestScripts.tar.gz

Follow these steps to install the performance verification test scripts:

- Download the file performanceVerificationTestScripts.tar.gz to the LifeScope<sup>™</sup> Software Data Folder (LSDF).
- 2. Go to the LSDF directory and untar the .tar.gz file: cd <LSDF>

tar -xzf performanceVerificationTestScripts.tar.gz

#### (Optional) Run the performance verification test

(LSA) Run a performance verification test, which takes approximately 24 hours.

The performance verification data files are installed during the "Copy data drive content" step on page 32. The performance verification test scripts are installed during the "(Optional) Download and install performance verification scripts" step on page 52.

	Follow these steps to execute the performance verification test:
	<ol> <li>Follow the instructions in "Set the LifeScope<sup>™</sup> Software environment" on page 46 to set the LifeScope<sup>™</sup> Software environment.</li> </ol>
	2. Confirm that the performance verification test data is installed at this directory: <lsdf>/performanceVerificationData</lsdf>
	<b>3.</b> Go to the performance verification test scripts directory: cd < <i>LSDF</i> >/performanceVerificationTestScripts
	<b>4.</b> Execute this command in the Linux shell:
	./runAll.sh -u username -w password
	where username and password must be for t he LifeScope <sup>TM</sup> Software administrator or for a valid LifeScope <sup>TM</sup> Software user account.
Monitor progress	Monitor the progress of your run by checking the output log files and also through job status commands such as <code>qstat</code> .
Check results	The results files are generated in the projects repository, at a location which is set during installation. Its default location is at < <i>LSDF</i> >/projects. Within the projects repository directory, the performance verification results are generated at the following results directories:
	username/performanceVerification/analysis_name/outputs
	Replace <i>username</i> with the user name used to run runAll.sh. The default is lifescope, the LSA account. The string performanceVerification is the project name used by the performance verification test. The analysis names used by runAll.sh are genomicResequencing.20x, genomicResequencing.2x, and wholeTranscriptomePE. The outputs subfolder name is fixed.
	The performance verification tests creates a log file in the <i>analysis_name</i> directory. Check for a success message in the last line of each log file.
	The results files are overwritten on subsequent runs executed by the same user.

#### Create user accounts

(LSA or SA) Create user accounts.

For instructions in this section that require using the LifeScope<sup>TM</sup> Software Admin Portal, refer to the administration appendix in the LifeScope<sup>TM</sup> Genomic Analysis Software User Guide (Part no. 4465696).

The method for user account creation depends on the authentication realm for your installation:

 LifeScope realm – (LSA) The LifeScope<sup>™</sup> Software administrator creates LifeScope<sup>™</sup> Software user accounts using LifeScope<sup>™</sup> Software. The LifeScope<sup>™</sup> Software Admin Portal is recommended for user administration.

LifeScope<sup>™</sup> Software shell commands for user administration are described in Table 5 on page 77, in Chapter 6, Run a Command Shell Analysis. Although shell commands include some user administration commands, the Admin Portal is recommended for general user administration.

- LDAP realm (SA and LSA) The Linux system administrator creates user accounts on the LDAP server. The LifeScope<sup>™</sup> Software administrator must use the LifeScope<sup>™</sup> Software Admin Portal to enable each LifeScope<sup>™</sup> Software user.
- Host realm (SA) The Linux system administrator creates Linux user accounts on the local operating system.

**Enable named** (LSA) This step is not required if your installation uses the concurrent licensing scheme. (With concurrent licensing, all users in the authentication realm are automatically enabled for LifeScope<sup>™</sup> Software.)

If your installation uses a named licensing scheme, the LSA must use the Admin Portal to identify the users for the named license.

#### Distribute the URL and user credentials

(LSA) The LifeScope<sup>™</sup> Software URL follows this pattern:

http://<fqdn of host>:<port number>/LifeScope.html

For example, for an installation on host orange.intranet.company.com, using the default port 9998 for the LifeScope<sup>™</sup> Software server, the URL is:

http://orange.intranet.company.com:9998/LifeScope.html

With the LifeScope<sup>TM</sup> Software servers running, verify the LifeScope<sup>TM</sup> Software URL on a supported browser. The following page appears:



Notify LifeScope<sup>™</sup> Software users (LSUs) of the LifeScope<sup>™</sup> Software URL.

If new user accounts are created for LifeScope<sup>™</sup> Software users, notify them of the username and password credentials required to access the LifeScope<sup>™</sup> Software.

#### Maintenance

Stop the	The following cautions apply to stopping the LifeScope <sup>™</sup> Software server:
LifeScope <sup>™</sup> Software server	<ul> <li>The LifeScope<sup>™</sup> Software server should only be stopped for required maintenance such as license upgrades.</li> </ul>
	<ul> <li>Only the LSA account can stop the LifeScope<sup>™</sup> Software server.</li> </ul>
	• In order to stop both the LifeScope <sup>™</sup> Software server and the licensing server, you must stop the LifeScope <sup>™</sup> Software server first.
	• All running analyses are terminated when you stop the LifeScope <sup>™</sup> Software server.
	Follow these steps to stop the LifeScope <sup>™</sup> Software server:
	<ol> <li>If you have not already done so, set the LifeScope<sup>™</sup> Software environment. See "Set the LifeScope<sup>™</sup> Software environment" on page 46.</li> </ol>
	2. Check for LifeScope <sup>™</sup> Software jobs running on the cluster with a command such as qstat. Notify LifeScope <sup>™</sup> Software users that their jobs are affected.
	<b>3.</b> Execute this Linux command:
	lscope-server.sh stop
Stop the licensing	The following cautions apply to stopping the licensing server:
server	<ul> <li>The LifeScope<sup>™</sup> Software licensing server must not be stopped except for required maintenance such as license upgrades.</li> </ul>
	Only the LSA account can stop the licensing server.
	• You must stop the LifeScope <sup>™</sup> Software server first before stopping the licensing server.
	<ul> <li>Running LifeScope<sup>™</sup> Software analyses are affected if the licensing server is stopped.</li> </ul>
	<b>IMPORTANT!</b> If the LifeScope <sup>TM</sup> Software licensing server stops accidentally while the LifeScope <sup>TM</sup> Software server is running, the licensing server must be restarted as soon as possible. The licensing checks fail if the licensing server is not operating, and running analysis jobs may stop.
	Follow these steps to stop the LifeScope <sup>™</sup> Software licensing server:
	<ol> <li>If you have not already done so, set the LifeScope<sup>™</sup> Software environment. See "Set the LifeScope<sup>™</sup> Software environment" on page 46.</li> </ol>
	<b>2.</b> Execute this Linux command:
	lscope-lmgrd.sh stop
	<b>3.</b> Verify that the LifeScope <sup>™</sup> Software licensing server has stopped, with this command:

lscope-lmgrd.sh status

Upgrade	Upgrading to a new version of LifeScope <sup>™</sup> Software does not require you to uninstall your existing version of LifeScope <sup>™</sup> Software.
	Follow the installation instructions for the new version of LifeScope <sup>TM</sup> Software. This procedure installs the new version in the $$ alongside your existing version, and modifies the soft link $/lifescope$ to point to the new version. The older version of LifeScope <sup>TM</sup> Software is not deleted during installation.
	You must manually move your license files from the older installation to the newer one.
	Once the new installation has been verified, you may remove the older installation at your discretion.
Uninstall	This section describes the steps for the LifeScope <sup>™</sup> Software administrator (LSA) to uninstall LifeScope <sup>™</sup> Software. If you use an authentication realm that requires the IT system administrator (SA) to create user accounts, the SA deletes those user accounts as part of the uninstall process.
	Follow these steps to uninstall LifeScope <sup>™</sup> Software:
	<ol> <li>(LSA or SA) Backup your LifeScope<sup>™</sup> Software license file or files, from the directory <lsbf>/lifescope/server/licenses.</lsbf></li> </ol>
	<ul> <li>2. (LSA or SA) Backup any required files, from the <lsdf> directories (these directories hold the LifeScope<sup>™</sup> Software repositories):</lsdf></li> <li>Reads files (XSQ and BAM)</li> <li>Reference files</li> <li>Output files</li> <li>Data</li> <li>Projects and analyses</li> <li>Notify LifeScope<sup>™</sup> Software users to backup any files they might need from the <lsdf> directories.</lsdf></li> </ul>
	<b>3.</b> (LSA or SA) Delete the LifeScope <sup>™</sup> Software installation directory, with this Linux command:
	/bin/rm -rf < <i>LSBF</i> >/lifescope
	<ul> <li>4. (Optional) (LSA or SA) Delete the LifeScope<sup>™</sup> Software data folder, with this Linux command:</li> </ul>
	/bin/rm -ri <lsdf></lsdf>
	<ol> <li>(Optional) (SA) If you use host or LDAP authentication, delete the user accounts you created for LifeScope<sup>™</sup> Software users.</li> </ol>
	( <i>Optional</i> ) (LSA or SA) If you modified any bash profile scripts in /etc or in the local home directory to change the PATH variable, remove the lifescope location entries in the PATH variable.

## PART I Overview

## Understand The LifeScope<sup>™</sup> Software Shell

#### This chapter covers:

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Terminology and concepts	60
Common scenarios	66

#### **Overview**

This chapter describes the concepts and terminology required when working with LifeScope<sup>™</sup> Software command shell. New concepts used with the shell include the following list. These terms are explained in more detail in this chapter.

- **Project** A collection of analyses based on scientific experiments or computational methodologies.
- **Module** A module is a single step in an analysis workflow. For example, mapping, SAET, and CNV are modules.
- Analysis A set of modules within an analysis workflow. An analysis can contain one or more modules.
- Analysis workflow A pre-defined set of modules in a workflow based on common applications areas, such as targeted resequencing or small RNA analysis.
- Read-set Sequencing data associated with an indexed (barcoded) sample from one XSQ file.
- Read-set repository An indexed database of XSQ files based on individual readsets.
- **Group** A collection of data that can be analyzed together. Grouping allows for aggregrated data (for instance, data from different lanes or different runs) to be analyzed as one set of data.

#### Terminology and concepts

Table 1 defines terminology and concepts used to describe the LifeScope<sup>™</sup> Software command shell. Review these terms before proceeding to the shell syntax in Chapter 6. Table 1 Concepts related to the LifeScope<sup>™</sup> Software command shell

Term	Description
category	Category refers to the various types of information containers at the root level of the LifeScope <sup>™</sup> Software command shell. An 1s command run at the shell root level lists the following categories: bams, projects, reads, references, users, and workflow.
level, virtual directory	Levels refer to the virtual directory structure within the LifeScope <sup>™</sup> Software command shell. For example:
	The root level: /
	The workflow level: /workflows
	The project level: /projects
	The analysis level: /projects/proj1/run1
	These levels are a representation of your projects and other shell elements on the LifeScope <sup>™</sup> Software server.
prompt	The shell prompt displays your current level within the LifeScope <sup>™</sup> Software command shell environment. The following are examples of the prompt strings at various shell levels:
	At the root level: server:/>
	At the workflow level: server:/workflows>
	At the project level: server:/projects>
	At the analysis level: server:/projects/proj1/run1>
projects	A project is a collection of analyses based on scientific experiments or computational methodologies. In the LifeScope <sup>™</sup> Software command shell, projects appear at /projects virtual directory. You create your projects at the /projects level. Within each project, you create one or more analyses.
	You may use projects as you wish, to group your analyses in a way that is convenient and meaningful for you. Each project is private to an individual LifeScope <sup>™</sup> Software command shell user.
analysis	An analysis is a set of modules within an analysis workflow. An analysis can contain one or more LifeScope™ Software modules.
	An analysis is defined by the following configuration:
	<ul> <li>The analysis methodology (through the set workflow command, specifying which LifeScope<sup>™</sup> Software modules are executed).</li> </ul>
	<ul> <li>The source (input) reads (through the import, add xsq, and set groups).</li> </ul>
	<ul> <li>The genome reference to be used (with the set reference command).</li> </ul>
	In addition, for tertiary-only workflows you can specify source BAM files (through the add bam command).
module	A module is a single step in an analysis workflow. For example, mapping, SAET, and CNV are modules.

Term	Description
analysis workflow	A pre-defined set of modules in a workflow based on common applications areas such as small RNA analysis or targeted resequencing. Also known as a standard workflow. A workflow combines a set of analysis modules into one run of LifeScope <sup>™</sup> Software. See Chapter 7, "Run a Standard Workflow Analysis" on page 97 for more information.
read-set	Sequencing data associated with an indexed (barcoded) sample from one XSQ file.
group	A collection of data that can be analyzed together. Grouping allows for aggregrated data (for instance, data from different lanes or different runs) to be analyzed as one set of data.
	The implications of using groups with your read-sets include the following:
	• Reads that are grouped together are treated as one specimen, even if the reads are in different XSQ input files.
	<ul> <li>LifeScope<sup>™</sup> Software supports processing multiple groups in one analysis run, while treating each group's data as an independent run.</li> </ul>
	Groups are created through the following methods:
	• The group option of the $\operatorname{add}\operatorname{xsq}$ command.
	• The set groups command.
	<ul> <li>The -g option of the LifeScope<sup>™</sup> Software shell run command.</li> </ul>
	A read-set can belong to a maximum of one group.
groupsfile	A text file optionally specified with an analysis. The groups file is a tab- separated text file with multiple lines in the following format:
	groupname < <i>path</i> >/xsqorbamfile: <i>indexid</i> < <i>path</i> >/reference_file
	If no barcode <i>indexid</i> is specified, it is assumed to be "1" (corresponding to the non-barcoded data). If a reference column entry does not exist, the reference from the previous line in the file is used.
	All members of a group must use the same reference. A groupsfile supports only a single reference, and that reference must be specified on the <i>first</i> line of the groups file.
repository	A LifeScope <sup>™</sup> Software shell repositories are virtual containers that persist your reference files, reads files, and projects.
	The projects repository, at /projects, contains your projects.
	The read repository, at /reads, is an indexed database of XSQ files based on individual read-sets.
	The references repository, at /references, contains the reference genomes used in your analyses.
directory	The LifeScope <sup>™</sup> Software command shell's repositories and analyses are represented in a structure resembling directories, but these are in a virtual system that lets you browse a representation of your projects and analyses on the LifeScope <sup>™</sup> Software server.

Table 1	Concepts	related to	the	LifeScope <sup>™</sup>	Software	command	shell	(continued)

	Description
shell	This document discusses two different shells and uses the followin terms to distinguish between them:
	<ul> <li>LifeScope<sup>™</sup> Software command shell and LifeScope<sup>™</sup> Software shell—Refer to the LifeScope<sup>™</sup> Software command-line environn that is invoked with the lscope.sh command.</li> </ul>
	• Linux shell—Refers to the shell of the underlying operating sys
local file system	This document uses the phrase "on your local file system" to desc a file that is specified by a Linux absolute path, such as /data/ results/example.sh. Such a file could be on a different physi disk instead of the local file system, but is available to Linux throug absolute path. The file is navigable from or available to the local fi system.
INI and PLN files	( <i>Optional</i> ) INI and PLN (pronounced "plan" or "PLN") files define you LifeScope <sup>™</sup> Software experiment. The LifeScope <sup>™</sup> Software commashell creates these file for you, based on the projects and analyses create and on how you have configure your analyses.
	Each INI file usually corresponds to one LifeScope <sup>™</sup> Software mod (such as mapping, SNPs, or CNV), and defines the behavior of that module through a set of key-value parameters (for example, the parameters for allowing IUB codes in references or setting the cal stringency level).
	Each PLN file defines which LifeScope <sup>™</sup> Software modules execute perform your analysis. A PLN file lists the INI files corresponding to LifeScope <sup>™</sup> Software modules to execute. The PLN fie list defines order of execution and execution dependencies.
	You can define and run your analyses in the LifeScope <sup>™</sup> Software command shell, without directly editing INI and PLN files. It is also possible to define analyses outside of the command shell, using only and PLN files. See Appendix D, "Command Shell Control Files" on page 497 for information on INI and PLN files.
run in the projects repository, or run on the local file system	When "run in the projects repository", a LifeScope <sup>™</sup> Software analycreates its output files in the projects repository and the analysis ( its configuration, status, and results files) can be viewed through the LifeScope <sup>™</sup> Software UI. When "run on the local file system" (also can "run in the current directory"), the output files are created in the current directory on the local file system, and the analysis cannot be viewed through the LII

Table 1 Concepts related to the LifeScope<sup>™</sup> Software command shell (continued)

Within each project, you create one or more analyses. Through the configuration of an analysis, you define the executable run.

Projects and analyses

Table 2 lists example steps to create a run in the LifeScope<sup>™</sup> Software shell. These steps assume you have logged into the shell. (See Chapter 6, "Run a Command Shell Analysis" on page 71 for complete instructions.)

Command	Description
cd projects	Enter your projects area.
mk proj1	Create a project named proj1.
cd proj1	Enter the proj1 level.
mk run1	In your proj1 project, create an analysis named run1.
cd run1	Open the run1 analysis, to configure the reference and reads files, and the workflow (the analysis modules).

Table 2 Setting up an analysis run in the LifeScope<sup>™</sup> Software shell

The configuration of analysis includes these areas:

- The reference genome.
- The reads files.
- The type of analysis to be performed (which LifeScope<sup>™</sup> Software modules perform the analysis). The analysis type can be any of the following:
  - A pre-defined workflow, which is a built-in sequence of commonly-used analyses, corresponding to a common biological application.
  - An analysis sequence you define, by either creating your own or customizing one of the LifeScope<sup>™</sup> Software pre-defined workflows.

An example of a pre-defined workflow is the targeted resequencing fragment workflow. This workflow includes the following LifeScope<sup>™</sup> Software modules:

- 1. SAET
- 2. Enrichment
- **3.** Mapping
- 4. Mapping statistics (BAMStats)
- 5. SNPs
- **6.** Small indels
- 7. Annotations

One LifeScope<sup>™</sup> Software shell command defines this workflow for your analysis (the set workflow targeted.resequencing.frag command).

Once an analysis is configured, it is launched with a single shell command (the run command).

Your projects are private and cannot be seen by other users or shared with other users.

Naming restrictions

- Project and analysis names must conform to Linux filename conventions:
  - Contain only alphanumeric and underscore characters.
  - Do not contain spaces.
  - Must be unique (among siblings).

 Do not begin with an underscore character. (This requirement is a LifeScope<sup>™</sup> Software limitation, not a Linux rule.)

Reads Data The data from each lane of a 5500 Series SOLiD<sup>™</sup> Sequencers instrument results in a single XSQ output file. Different models of the 5500 support up to 6 or 12 lanes, so typically multiple XSQ files are generated per sequencing run. You can direct your LifeScope<sup>™</sup> runs to analyze data from specific barcodes in multiple lanes, or group the output of multiple lanes or multiple sequencing runs to be processed as a single specimen.

The type of data used in LifeScope<sup>TM</sup> Software is called a read-set, which is a collection of reads belonging to one index (barcode) from one XSQ file. The smallest set of data you can specify in the shell is one index from one XSQ file.

You can optionally use groups to combine more than one index into an analysis. Reads that have been combined into a group are processed as one specimen, even if the readsets come from different input XSQ files. Each group generates a combined set of output files.

In the LifeScope<sup>™</sup> Software command shell, input data is added at the analysis level. Input data is associated with individual analyses, not with entire projects.

Plan your analysis input carefully. The way you define your reads input affects the behavior of your analysis. Index IDs and group names give you flexibility in handling your input data. For example:

- Combine specific read-sets from one XSQ file or multiple XSQ files into one group to be analyzed together. All read-sets within a group must be of the same library type.
- Assign read-sets to different groups, to be analyzed separately within one LifeScope<sup>™</sup> Software run.

For more information and examples of LifeScope<sup>™</sup> Software shell commands to use with your input data, see "Define input data" on page 85.

For information about XSQ data and the XSQ file format, see the following sections:

- "XSQ file format" on page 456, in Appendix B, "File Format Descriptions" on page 455
- Appendix C, "XSQ Tools" on page 479

**Repositories** LifeScope<sup>™</sup> Software repositories organize and persist your projects, your input data, and your reference data files.

The reads repository is a storage place in LifeScope<sup>TM</sup> Software for instrument data intended to be input data for LifeScope<sup>TM</sup> Software analyses. Data imported into the reads repository are in unmapped XSQ-format files.

Importing reads and references file into LifeScope<sup>™</sup> Software repositories is optional, but preferred. The following are two advantages to using the shell's repositories:

- The shell is aware of barcoded readsets in the read repository. For reads files not in the read repository, you must manually specify barcodes.
- In your shell commands you can use repository names for files in the read and reference repositories. You do not have to use Linux paths to these files.

	Files imported into the read and reference repositories are visible to all LifeScope <sup>™</sup> Software users. However, each user has their own projects repository. The contents of your projects repository are private and cannot be seen by other users within the command shell.				
	<b>Note:</b> In the current release, one repository cannot be split across multiple partitions or file systems. (The various repositories can each be on a different partition or file system.)				
The reference repository	The reference repository is initially populated during the installation process with reference files for hg18 and hg19.				
	For more information (not required for most use), see Appendix G, "The Reference Repository" on page 515.				
Find reference files	To find reference files in the references repository, follow these steps:				
for an analysis	Log into the shell.				
	cd /references ls				
	The above example lists reference files that have previously been imported (registered) into the references repository. Reference files can alternatively be specified with an absolute path to the file on the local file system (or navigable from the local file system).				
	See also the shell command set reference <i>assembly_name</i> in Table 5 on page 77. When used with a known assembly such as hg18 or hg19, this command automates using the correct reference in your analysis.				
	One reference is supported per analysis run.				
Find data for an	To find data in the read-set repository, follow these steps:				
analysis	Log into the shell.				
	cd /reads ls				
	The above example lists reads files that have previously been imported (registered) into the reads repository. Reads files can alternatively be specified with an absolute path to the file on the local file system (or navigable from the local file system).				
View analysis information	To show the status of your jobs, run a shell status command in the Linux shell. The format of the command depends on whether the analysis is in the project repository or in the current directory.				
	For an analysis in the project repository, run this command:				
	lscope.sh status -u user -p projectname -a analysisname				
	For an analysis in the current directory, run this command:				
	lscope.sh status				
	An alternate method is to run an 1s command while in an analysis virtual directory, in the LifeScope <sup>™</sup> Software command shell. For example:				
	cd /projects				

```
cd proj1 (This command opens your project named proj1.)
cd run1 (This command opens your analysis named run1.)
ls
--ini--
secondary/bamstats.ini
secondary/global.ini
secondary/fragment.mapping.ini
--readsets--
lung:DH10B_ColorSpaceOnly_Unbarcoded_50_Frag.xsq:1:DH10B.fasta
--jobs--
run1 [70%]
```

The ls output lists the defined parameters and input read-sets. For running analyses, the output also lists the estimated percentage of completion.

#### View analysis read-sets

To show the read-sets in an analysis, cd to that analysis in the shell and run an ls command. Part of the ls output displays the read-sets which have been added to this analysis:

```
--readsets--(group:file:index:reference)
Group_1:helen_20100402_2_10103F_F3_1.3.1_2.0_converted.bam::
Group_1:helen_20100402_2_10103F_F3_1.3.1_2.0_converted.bam::
Group_1:WT_Chr17_Frag.xsq:1:
```

The output includes the group name, file name, index number, and reference file, separated by colons. Except for the file name field, some fields may be empty.

#### **Common scenarios**

This section describes how LifeScope<sup>™</sup> Software command shell handles common biologic analysis scenarios. New terms and concepts, such as project and analysis, are explained later in this chapter. Specific instructions and commands are given in Chapter 6, Run a Command Shell Analysis.

Basic

A basic scenario is to run a bioinformatics analysis, such as targeted resequencing, on sequence data. The steps to do this in LifeScope<sup>TM</sup> Software shell are:

- 1. Create a project.
- 2. In that project, create an analysis.
- 3. In that analysis:
  - a. Specify the type of analysis to be performed on the data (set the workflow).
  - **b.** Add the sequenced data file.
  - c. Specify the reference genome for this analysis.
  - **d**. Run the analysis.

A sample sequenced on multiple lanes	In this scenario one sample is sequenced in multiple lanes on the sequencing instrument. The output data from the sequencing instrument is contained in multiple XSQ files.						
	In the LifeScope <sup>™</sup> Software command shell, the multiple XSQ files are analyzed in one run and as one sample. LifeScope <sup>™</sup> Software treats the multiple data files as one set of data.						
	The steps to do this in LifeScope <sup>™</sup> Software shell are:						
	1. Create a project.						
	<b>2.</b> In that project, create an analysis.						
	<b>3.</b> In that analysis:						
	<b>a.</b> Specify the type of analysis (such as a standard workflow) to be performed on the data.						
	<b>b.</b> Add the XSQ data files. When you add the XSQ files, specify the same group name for each file.						
	<b>c.</b> Specify the reference genome for this analysis.						
	<b>d</b> . Run the analysis.						
Data from multiple samples	This scenario performs, in one run, the same analysis on sequence data from multiple specimens. The data can be in one or multiple XSQ files.						
•	LifeScope <sup>TM</sup> Software keeps the data from each specimen separate, and each specimen is analyzed separately. The steps to do this in LifeScope <sup>TM</sup> Software shell are:						
	1. Create a project.						
	<b>2.</b> In that project, create an analysis.						
	<b>3.</b> In that analysis:						
	<b>a.</b> Specify the type of analysis (such as a standard workflow) to be performed on the data.						
	<b>b.</b> Add the sequenced data file, specifying different grouping for the data.						
	<b>c.</b> Specify the reference genome for this analysis.						

**d.** Run the analysis.



## PART II Getting Started

## Run a Command Shell Analysis

This chapter covers:

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#### **Overview**

The LifeScope<sup>™</sup> Software command shell provides a way to quickly run bioinformatics experiments. The shell has its own working environment, which is organized around the concepts of projects and of analyses within projects. The LifeScope<sup>™</sup> Software command shell has elements similar to the Linux shell but with a different syntax and interpretation. If you are new to LifeScope<sup>™</sup> Software, please review Chapter 5, "Understand The LifeScope<sup>™</sup> Software Shell" on page 59 before reading this chapter.

New users are recommended to use the standard workflows built into LifeScope<sup>™</sup> Software. Standard workflows are developed with defaults optimized for common analyses. The applications supported by standard workflows are:

- Genomic resequencing
- Targeted resequencing
- Whole transcriptome
- Small RNA
- ChIP-Seq
- MethylMiner<sup>TM</sup>

#### Run a standard workflow

	A standard workflow is a built-in series of commonly-used analyses, which correspond to a common biological application. The pre-built standard workflows include genomic resequencing, targeted resequencing, small RNA, ChIP-Seq, MethylMiner <sup>™</sup> , and whole transcriptome. Using one of the workflows allows you to run an analysis with a minimum of setup.					
	In the LifeScope <sup>™</sup> Software command shell, running a standard workflow requires the following steps:					
	1. Create a project and analysis.					
	<b>2.</b> Identify the workflow to be executed on your data.					
	<b>3.</b> Identify your input data (your reads).					
	<b>4.</b> Identify the reference genome.					
	<b>5.</b> For targeted resequencing, identify the regions of interest file.					
	<b>6.</b> Issue the run command to start your analysis.					
Example steps to run a standard	These steps are an example of how to run a workflow in the LifeScope <sup>™</sup> Software command shell:					
workflow	<pre># log into the shell lscope.sh shell -u username -w password # cd to the projects repository cd /projects # make a project, and open it mk ecoli cd ecoli # make an analysis, and open it mk run1 cd run1 # define the analysis type set workflow targeted.resequencing.frag # define the input add xsq DH10B_ColorSpaceOnly_Unbarcoded_50_Frag.xsq lung # specify the reference to be used set reference /data/results/references/DH10B.fasta # define the any analysis-specific parameters set analysis.regions.file /data/results/readsdata/dh10b.bed # list the configuration of the analysis ls # start the run run # list progress information about the run ls</pre>					
# Explanation of<br/>example stepsTable 3 explains the purpose of the steps shown in "Example steps to run a standard<br/>workflow".

Table 3 Sample shell commands to run a workflow

Command	Purpose
lscope.sh shell -u username	Log into the LifeScope <sup>™</sup> Software command shell.
cd /reads import /data/xsq/DH10B_ColorSpaceOnly_ Unbarcoded_50_Frag.xsq	Go into the read repository and import an XSQ file (the input data file). This command in effect brings the reads file into the shell read repository.
	Once in the read repository, the reads file is available as input to your analysis runs.
cd /projects mk ecoli cd ecoli	In the projects repository, create a project named ecoli, and open that project.
mk run1	Create an analysis named run1. Open the run1 analysis.
cd run1	<b>Note:</b> The open analysis (cd run1, in this case) command is important. The data, reference, and other configuration commands which follow apply only to the current analysis.
<pre>set workflow targeted.resequencing.frag</pre>	Define the workflow used with this analysis. The workflow in this example includes SAET, mapping, mapping statistics, SNPs, inversion, large indel, small indel, and annotations.
add xsq DH10B_ColorSpaceOnly_Unbarcod ed_50_Frag.xsq	Associate the XSQ file with this analysis. This command directs LifeScope <sup>™</sup> Software to use this XSQ file as the input reads data for this analysis.
	This example uses the repository name for the XSQ file. The absolute path to the XSQ file on the local file system is also supported.
<pre>set reference /data/results/results/ references/DH10B.fasta</pre>	Define the reference file for this analysis.
<pre>setanalysis.regions.file/data/results /readsdata/dh10b.bed</pre>	Define the regions of interest file, for targeted resequencing.
ls	Display the configuration of your analysis, including input reads, reference files, workflow, and select parameter settings.
run	Start the analysis run.
wait	Wait for the analysis run to complete.
ls	Display configuration and status information about the analysis run.

# Use the command shell

The LifeScope<sup>™</sup> Software shell is a command-line tool for running LifeScope<sup>™</sup> Software data analyses. The shell also allows you to create and manage your LifeScope<sup>™</sup> Software data, projects, analyses, and users.

If you instead want to run the LifeScope<sup>TM</sup> Software user interface (UI), refer to the  $LifeScope^{TM}$  Genomic Analysis Software User Guide (Part no. 4465696).

There are four modes of operation for the LifeScope<sup>™</sup> Software command shell:

• **Run mode** – Runs a LifeScope<sup>™</sup> Software analysis or workflow from the Linux shell, without entering the LifeScope<sup>™</sup> Software command shell. The LifeScope<sup>™</sup> Software control file (*plnfile*) must be created and configured before executing a run mode command.

Example commands:

```
lscope.sh run plnfile -u marie -p proj1 -a run1 -g group1
lscope.sh resume plnfile -u joe
See "Run mode" on page 74.
```

• Status mode – Opens the LifeScope<sup>™</sup> Software shell status viewer, which shows the dynamic status of running and completed jobs for specified analysis.

Example commands:

```
lscope.sh status -p proj1 -a analysis1
lscope.sh status
See "Status mode" on page 75.
```

- Scripted mode You invoke the LifeScope<sup>™</sup> Software shell from commands listed in a file on your local system. See "Scripted mode" on page 76 for information on running in scripted mode.
- Shell mode Enters the LifeScope<sup>™</sup> Software command shell. In this shell, only LifeScope<sup>™</sup> Software shell commands are accepted. The LifeScope<sup>™</sup> Software shell commands are described in Table 5 on page 77. The LifeScope<sup>™</sup> Software shell commands allow you to create, manage, and run your LifeScope<sup>™</sup> Software analyses.

Example commands:

• To enter the shell: lscope.sh shell -u marie

Within the shell:

- set workflow whole.transcriptome.frag
- set reference DH10B.fasta

```
See "Shell mode" on page 77.
```

**Note:** Do not confuse the LifeScope<sup>™</sup> Software shell commands with Linux or UNIX commands. Some commands may have similar names, but the behavior of the LifeScope<sup>™</sup> Software shell commands is entirely different from the Linux or UNIX commands. LifeScope<sup>™</sup> Software shell commands also depend on the shell level (analysis, project, reads, references, or root) at which they are executed.

# **Run mode** With the run mode, you enter the LifeScope<sup>™</sup> Software command in the Linux shell. You remain in the Linux shell, and the LifeScope<sup>™</sup> Software command shell does not open. Run mode also includes the resume command.

The following are example run mode commands:

```
lscope.sh run ./splice.finder.pln -p wt_spliceFinder \
    -a wt_spliceFinder_PE_TC12 -u rjones
lscope.sh resume ./splice.finder.pln -p wt_spliceFinder \
    -a wt_spliceFinder_PE_TC12 -u rjones
```

The -p option specifies the project name. The -a option specifies the analysis name. The -u option specifies the username.

A –g groupsfile parameter is optional. When a groupsfile is provided, LifeScope<sup>™</sup> Software performs multi-group analysis, which involves analyzing the data for each group independently. See "Run a grouped analysis" on page 88 for more information about multi-group analyses.

#### When to use run mode

An lscope.sh run mode command is appropriate when you have already configured you analysis. Prerequisites to running an lscope.sh run mode command include:

- The LifeScope<sup>™</sup> Software web server and license server must be running.
- The LifeScope<sup>™</sup> Software user account must exist.
- The reads files and reference files required by the analysis must have been transferred to or available to the LifeScope<sup>™</sup> Software cluster or workstation.
- Reference files, reads files, and read-sets must either have been specified in a previous LifeScope<sup>™</sup> Software command shell session, or specified by absolute paths in the analysis PLN and INI files.
- The analysis PLN and INI files are configured correctly.

#### Run "in the projects repository" or "in the local file system"

A run command that includes the project and analysis options is considered to "run in the projects repository". The shell projects repository determines the output file location, and the analysis run is visible and trackable from the LifeScope<sup>™</sup> Software UI.

Without the project and analysis options, a run command is considered to "run in the local file system". Output files are created in the current directory on the file system, and the run is tied to the current directory for checking status.

Table 4 compares the two methods.

 Table 4
 Comparison of running in the projects repository and in the local file system

Category	Run in the projects repository	Run in the local file system
Command	lscope.sh run plnfile -u user -p proj -a analysis -g groupsfile	lscope.sh run plnfile -u user
format		Must be executed in the local file system in the same directory as plnfile.
Status command	lscope.sh status -u user -p proj -a analysis	lscope.sh status
		Must be executed in the same directory of the local file system.
Output file location	In the /projects repository	In the current directory of the local file system.
Visible in the UI?	Yes.	No.
	You can view the analysis configuration, reads files, parameters, status, and results files in the UI.	

# Status modeThe lscope.sh status command displays information about your running<br/>LifeScope™ Software analyses.

To check the status of the example run described in "Run mode", enter this command in the Linux shell:

lscope.sh status -u rjones -p wt\_spliceFinder \

-a wt\_spliceFinder\_WT\_SF\_PE\_TC0012

If your run command is running in the local file system, use one of the following commands to check status. The format depends on whether or not your analysis is in the projects repository:

- For an analysis in the project repository, run this command:
  - lscope.sh status -u user -p projectname -a analysisname
- For an analysis in the current directory, run this command: lscope.sh status

The following is an example of the lscope.sh status screen:

Status (t=toggle subjobs, arrows=scroll	l, q=quit view	i): [1	running]
TASK	STATUS	PROGRESS	INFO
human_hg18-pair.mapping	started	0%	
human_hg18-pair.mapping.1.splittin	completed	28	Pair End Mapping.doSpl
human_hg18-pair.mapping.2.run	started	78	mapreads
human_hg18-pair.mapping.3.run	started	88	mapreads
human_hg18-pair.mapping.4.run	started	88	mapreads
human_hg18-pair.mapping.5.run	started	78	mapreads
human_hg18-PairEndMapping.MarkDupl	not started	<b>1</b> 6	
human_hg18-PairEndMapping.BAMStats	not started	<del>1</del> 6	
solid0064_20101210_MP_2X60_T3_1_F3	not started	<b>1</b> 6	
solid0064_20101210_MP_2X60_T3_1_F3	not started	<b>1</b> 6	
solid0064_20101210_MP_2X60_T3_1_F3	not started	<b>1</b> 6	
solid0064_20101210_MP_2X60_T3_1_F3	not started	ale a	
solid0064_20101210_MP_2X60_T3_1_F3	not started	ale a	
solid0064_20101210_MP_2X60_T3_1_F3	not started	ale a	
solid0064_20101210_MP_2X60_T3_1_F3	not started	ale a	
solid0064_20101210_MP_2X60_T3_1_F3	not started	4	
solid0064_20101210_MP_2X60_T3_1_F3	not started	oło	

You must be in the correct directory on your local file system (the directory from which you executed the run command).

(For another status command, see Table 5. The shell 1s command, when executed with the shell in an analysis level, gives the status of an analysis run.)

**Scripted mode** The LifeScope<sup>™</sup> Software command shell accepts as input a text file containing shell commands.

1. Create a text file with one command per line. For example:

```
cd proj1
cd run2
set workflow analysis.pln
add xsq xsqfile
set reference hg19
run
ls
exit
```

2. Save the file, for example as projlscript.

#### **3.** Then run with this command:

lscope.sh -u user -w password < proj1script</pre>

**Note:** This example does not include the commands necessary to step up a new analysis.

Shell modeIn shell mode, you enter the LifeScope<sup>™</sup> Software command shell. In this shell, only<br/>LifeScope<sup>™</sup> Software shell commands are accepted. The LifeScope<sup>™</sup> Software shell<br/>commands allow you to create, manage, and run your LifeScope<sup>™</sup> Software analyses.

#### Terminology

Table 1 on page 60, in Chapter 5, Understand The LifeScope<sup>™</sup> Software Shell, defines concepts and terminology used with the LifeScope<sup>™</sup> Software command shell. Review these terms before proceeding to the shell commands and their syntax.

#### lscope.sh commands

Table 5 describes the commands available within the LifeScope<sup>TM</sup> Software command shell. The behavior of some shell commands depends on the level at which they are executed, as noted in Table 5.

**Table 5** LifeScope<sup>™</sup> Software shell commands

Command	Description		
Project and analysis	Project and analysis commands:		
mk <i>name</i>	Creates a new project or a new analysis within the current project. Must be executed at the /projects or /projects/ <i>projname</i> levels.		
	When executed at the /projects level, creates a new project named name.		
	When executed at the /projects/projname level, creates a new analysis named name, within your project projname.		
	Command examples:		
	• At the /projects level: mk testproj		
	• At the /projects/projname level: mk run1		
	See "Project and analysis naming rules" on page 85 for restrictions on name.		

## Table 5 LifeScope<sup>™</sup> Software shell commands (continued)

Command	Description
ls	Lists information about your project or analysis.
	When executed at the /projects level, lists the names of all your projects.
	At the /projects/projname level, lists the names of all analyses contained within your project named projname.
	At the /projects/projname/analysisname level, provides configuration and progress reporting for the analysis:
	• Lists INI files involved in the analysis, if set workflow has been given.
	<ul> <li>Lists read-sets involved in the analysis, if read-sets have been added. The read-set listing includes the group, file name, index, and reference.</li> </ul>
	Lists the status of jobs for the analysis.
	Example:
	ls ini fragmentMapping.ini global.ini
	readsets(group:file:index:reference) Group_1:case0030_Fragment.xsq:1:chr16.validated.fasta
	jobs run1 [100%] [JOB_SUCCESS] fragmentMapping [100%] [JOB_SUCCESS]
	Example two:
	<pre>lsini secondary/global.ini secondary/smallrnamapping.ini tertiary/global.ini tertiary/srCount.ini tertiary/srCoverage.ini</pre>
	readsets(group:file:index:reference) HumanFrag_smRNA:reads.xsq:1:hg1_1b.fasta
	jobs smallRNAworkflow [100%] [JOB_SUCCESS] secondary-hg1_1b-smallrnamapping [100%] [JOB_SUCCESS] tertiary-HumanFrag_smRNA-srCount [100%] [JOB_SUCCESS] tertiary-HumanFrag_smRNA-srCoverage [100%] [JOB_SUCCESS]

Command	Description
rm <i>name</i>	Permanently deletes a project or an analysis within the current project. Must be executed at the / projects or /projects/projname levels.
	When executed at the /projects level, deletes the project named name.
	When executed at the /projects/projname level, creates a new analysis named name, within your project projname.
	Command examples:
	• At the /projects level: rm testproj
	• At the /projects/projname level: rm analysis2
	<b>WARNING!</b> This command permanently deletes the underlying project and analysis directories on the server. Recovery is <i>not</i> supported.
set workflow <i>path</i>	Sets the workflow or PLN file for the current analysis. This command is a prerequisite to running an analysis. The set workflow command must also precede any set <i>param</i> value commands.
	Examples:
	set workflow targeted.resequencing.frag
	<pre>set workflow /share/lifescope/human/test1/my.pln</pre>
	The set workflow command replaces the manual configuration of PLN and INI files used in earlier versions of BioScope <sup>™</sup> Software. With this command the LifeScope <sup>™</sup> Software shell automatically creates the required PLN and INI files.
get workflow	Copies the current workflow's INI files, PLN files, and their directory structure to the local file system. These files are copied to the directory you are in when you give the <code>lscope.sh</code> command.
	You must have a shell analysis open (be in the analysis virtual directory) and that analysis must have a workflow already set (with the set workflow command).
ls	When executed at the level of a specific standard workflow, such as /workflows/small.rna, lists the INI files for the workflow. For example:
	server:/workflows> cd small.rna
	<pre>server:/workflows/small.rna&gt; ls</pre>
	secondary/bamstats.ini
	secondary/global.ini
	secondary/smallrnamapping.ini
	tertiary/filtermapping.bamstats.ini
	tertiary/global.ini
	tertiary/srCount.ini

 Table 5
 LifeScope<sup>™</sup> Software shell commands (continued)

#### Table 5 LifeScope<sup>™</sup> Software shell commands (continued)

Command	Description
import <i>path</i> [ <i>reponame</i> ]	When executed at the /reads level, imports a source XSQ file from the Linux file system into the reads repository (and updates the read repository index). The import command imports to the reads repository top folder, not to a subfolder.
	When executed at the /bams level, imports a BAM input data file into the BAM repository (for tertiary analysis). The /bams virtual directory is reserved for BAM files that have been converted from earlier versions of the software or imported from another installation of the same version of LifeScope <sup>™</sup> Software.
	The <i>path</i> argument is one of the following:
	• The absolute path to the file on the Linux file system (recommended), or
	• A relative path to the file, using as the current directory the Linux directory in which the lscope.sh shell command was given.
	The <i>reponame</i> argument is optional, and allows you to assign a repository name different from the filename. The file appears as <i>reponame</i> when the repository contents are displayed, and is accessed as <i>reponame</i> with commands such as $addxsq$ <i>reponame</i> .
	The import command may be one of these forms:
	<ul> <li>import-ln – Links the file (with ln -s) given by path to the repository directory on the LifeScope<sup>™</sup> Software server. Default.</li> </ul>
	• <b>import-cp</b> – Copies the file given by <i>path</i> to the repository directory on the LifeScope <sup>™</sup> Software server.
	• <b>import-mv</b> – Moves the file given by <i>path</i> to the repository directory on the LifeScope <sup>™</sup> Software server.
add xsq <i>name</i> [:id] [group]	The add $xsq$ command specifies input for secondary analysis. It adds the read-sets with index (barcode id) <i>id</i> in the XSQ file specified by <i>name</i> into the current analysis. add $xsq$ must be run at the analysis level.
	name must match either a reads file that has been imported into the reads repository or the full path to an XSQ file on the local file system. The ls command run at the /reads level lists available read repository files.
	<i>id</i> is optional. If no <i>id</i> is provided, all indices (barcodes) from the XSQ file are added. Each <i>id</i> refers to the data for one index.
	<i>group</i> is optional. If <i>group</i> is provided, all read-sets are grouped together, and analyzed together as one specimen. Read-sets from multiple XSQ files may be added the same <i>group</i> name.
	If group is not provided, the following occurs:
	Read-sets are grouped by index name from the XSQ file.
	The read-sets from each index (barcode) are processed independently.
	All read-sets in a group are analyzed together as one specimen. The group name is not required to match the specimen name or other information from the instrument run. Do not use spaces or special characters in a group name.
	Multiple $\operatorname{add} xsq$ commands are supported per analysis.
del xsq <i>filename</i>	Removes the XSQ file specified by <i>filename</i> from the current analysis. <i>filename</i> must appear in the listing of the read repository (as listed by the LifeScope <sup>™</sup> Software shell 1s command at the /reads level). This command does not physically delete the XSQ file from your file system.

Command	Description
rm xsq <i>name</i>	Removes the XSQ file <i>name</i> from the /reads repository. <i>name</i> is the file's repository name. If the XSQ file <i>name</i> was originally imported with <i>import-ln</i> , <i>rm</i> deletes the link in the repository directory on the local file system. If the XSQ file <i>name</i> was originally imported with <i>import-cp</i> or <i>import-mv</i> , <i>rm</i> deletes the file from the repository directory on the local file system.
set groups path	Creates read-set groups from the file given by <i>path</i> . Read-sets belonging to one group are analyzed together as one specimen. Each group generates an independent set of results files (containing data combined for the whole group).
	LifeScope <sup>™</sup> Software can process multiple groups in one analysis run, while treating each group's data as an independent run.
	The groups file must be a tab-separated text file with multiple lines in the following format:
	groupname <path>/XSQorBAMfile:indexid <path>/reference_file</path></path>
	<i>path</i> must be an absolute path to the groups file on (or available to) the local file system.
	If no index id is specified, it is assumed to be "1". All read-sets from the file are included.
	If the reference column entry does not exist, the reference from the previous line in the file is used.
	The XSQ, BAM, and reference files may be specified by their repository names, if the files have been imported into a LifeScope <sup>™</sup> Software shell repository. If these files are specified by a path on the local file system, an absolute path is recommended. Relative paths are interpreted relative to the directory where you open the LifeScope <sup>™</sup> Software command shell with the lscope.sh shell command.
add bam <i>path</i> [group]	( <i>Optional</i> ) Adds BAM file <i>path</i> into the current analysis. Use this command to specify input for tertiary analysis.
	<i>path</i> must match either a BAM file that has been imported into the bams repository or an XSQ file on the local file system. The ls command run at the /bams level lists available bams repository files.
	<i>group</i> is optional. If no <i>group</i> is provided, <i>group</i> is assumed to be "Default". All BAM files in a group are analyzed together as one specimen.
del bam <i>name</i>	Removes the BAM file <i>name</i> from read-sets to be analyzed.
	This command does not physically delete the BAM file from your file system.
rm bam <i>name</i>	Removes the BAM file <i>name</i> from the /bams repository. <i>name</i> is the file's repository name. If the BAM file <i>name</i> was originally imported with <i>import-ln</i> , <i>rm</i> deletes the link in the repository directory on the local file system. If the BAM file <i>name</i> was originally imported with <i>import-cp</i> or <i>import-mv</i> , <i>rm</i> deletes the file from the repository directory on the local file system.
cat bam <i>path</i>	Displays header information for the BAM file given by <i>path. path</i> must be the absolute path to a BAM file on the file system.
ls	When executed at the /reads level, lists the XSQ files in the read repository. When executed at the /bams level, lists the files in the BAM repository. The library type and data type (base space or color space) are included in the listing. Example output at the /reads level is:
	<pre>smallindel_simulation_2INS_22_25.xsq [Fragment] [color] AllMapBarcodes.xsq [MatePair] [mixed] [Barcoded] DH10B_ColorSpaceOnly_50_Frag.xsq [Fragment] [color] [Non-barcoded] DefaultLibrary_PE.xsq [PairedEnd] [color] [Non-barcoded] Fragment3panels.xsq [MatePair] [mixed] [Non-barcoded]</pre>

**Table 5** LifeScope<sup>™</sup> Software shell commands (continued)

#### Table 5 LifeScope<sup>™</sup> Software shell commands (continued)

Command	Description	
cat <i>xsqfile</i>	In /reads, the ls command with an XSQ file argument displays the libraries and barcodes contained in the XSQ file. For example:	
	<pre>server:/reads&gt; ls Indexing3panels.xsq</pre>	
	[lib:Library1] [idx:1] [MatePair] [50,50] [mixed]	
	[lib:Library1] [idx:13] [MatePair] [50,50] [mixed]	
	[lib:Library1] [idx:5] [MatePair] [50,50] [mixed]	
	[lib:Library1] [idx:9] [MatePair] [50,50] [mixed]	
	[lib:Library2] [idx:10] [MatePair] [50,50] [mixed]	
	[lib:Library2] [idx:14] [MatePair] [50,50] [mixed]	
	[lib:Library2] [idx:2] [MatePair] [50,50] [mixed]	
	[lib:Library2] [idx:6] [MatePair] [50,50] [mixed]	
	[lib:Library3] [idx:11] [MatePair] [50,50] [mixed]	
Reference file comman	ds:	
set reference assembly   path	assembly names an assembly in the LifeScope <sup>™</sup> Software shell reference repository. The LifeScope <sup>™</sup> Software shell sets reference-related parameters to appropriate default values. An example assembly name command is:	
	set reference hg18	
	<i>path</i> sets the genome reference to the reference file <i>path</i> in current analysis. <i>path</i> can either be an absolute filepath (on the Linux file system), or a reference name that exists in the shell reference repository (as listed by the LifeScope <sup>™</sup> Software shell 1s command at the /references level).	
	If <i>assembly</i> names a recognized assembly in the reference repository, then the following reference- related parameters are automatically set to default values for <i>assembly</i> :	
	• annotation.dbsnp.file	
	• annotation.gtf.file	
	• analysis.mirbase.precursor.file	
	• analysis.mirbase.mature.file	
	• analysis.filter.reference	
	• analysis.assembly.name	
	• analysis.species.name	
	These values can be changed with the set <i>param.key param.value</i> command, if necessary.	
	A set reference command (either <i>assembly</i> or <i>path</i> ) is mandatory before running an analysis. The use of <i>assembly</i> is recommended. A maximum of one reference is allowed per analysis.	
Parameter commands:		

Table 5 LifeScope	Software shell commands <i>(continued,</i>	)

Command	Description
set param.key param.value [inifile]	Sets a parameter for the analysis. The analysis workflow must be defined (with the set workflow arg command) before you set a parameter for that analysis. For example:
	<pre>set analysis.regions.file absolute_path/regions_bed_file set annotation.dbsnp.file absolute_path/dbsnp_file set annotation.gtf.file absolute_path/refgene_gtf_file set analysis.filter.reference absolute_path/filter_reference_file set analysis.mirbase.mature.file absolute_path/mature_forms_file set analysis.mirbase.precursor.file absolute_path/precursor_forms_file set analysis.assembly.name name_of_reference_assembly set analysis.species.name name_of_reference_species</pre>
	Some parameters are automatically pre-populated based on the reference chosen. See "List shell commands" on page 85.
	System parameters may also be set. For example:
	set scratch.dir /scratch/lifescope
	The <i>ini</i> argument is optional. When <i>ini</i> is specified, the parameter is set for an individual INI file within the analysis. For example, the following command sets the value of the parameter <code>bamstats.wig.primary.only</code> to 0, for only the BAMStats module:
	set bamstats.wig.primary.only 0 secondary/bamstats.ini
	The <i>ini</i> value is case sensitive. The secondary or tertiary location of the INI file is required.
	If the <i>ini</i> argument is not provided, the parameter is added to both the secondary and tertiary global.ini files.
	The <i>inifile</i> version of the command supports turning off certain analysis modules. This method is not recommended for a module that has dependencies in the PLN file. The following example turns off the bamstats module:
	set bamstats.run 0 secondary/bamstats.ini
	Note: The set workflow command must precede any set <i>param value</i> commands.
	Note: Do not use an equals sign '=' with the set command.
cat inifle	Lists parameters for the current analysis, including ones defined with the set param value [inifile] command. Example:
	<pre>cat pair.mapping.ini   default.maximum.insert.size=20000   default.minimum.insert.size=150   pair.mapping.run=1</pre>
del param inifile	Undefines a parameter from an INI file. The <i>inifile</i> value is case sensitive. If no INI file is specified, deletes the parameter from all global.ini files. If an INI file is specified, the secondary or tertiary location of the INI file is required. The following example deletes the definition of the parameter analysis.input.readset.file from a mapping INI file. The analysis.input.readset.file parameter must not be set if the SAET module is not run.
	del analysis.input.readset.file secondary/fragment.mapping.ini
Execution commands:	
run	Begins execution of the current analysis.
wait	Waits for the currently running analysis to complete. If the current analysis not running, returns immediately. Must be run in an analysis directory.
stop	Stops execution of the current analysis.

#### Table 5 LifeScope<sup>™</sup> Software shell commands (continued)

Command	Description
resume	Resumes execution of the current analysis, if that analysis or a portion of it is in a failed or stopped state. Modules that successfully completed execution are not re-run.
	Job resumption is at the module level. A module that partially completed but then failed is restarted from the beginning by the resume command.
	The shell resume command can also be issued in the Linux shell, as a run mode command. The syntax is:
	lscope.sh resume plnfile -u user -p project -a analysis
Navigation commands:	
cd /	Navigates to the root level of the LifeScope <sup>™</sup> Software command shell. May be executed at any shell level.
	<b>Note:</b> '/' is the only argument allowed. You may not specify multiple levels separated by one or more '/' characters.
cd	Navigates up one level (one virtual directory) in the shell.
	Note: The cd command navigates only one level at a time.
cd name	Navigates down one level (one virtual directory) in the shell.
	When executed at the shell root level (/), opens the category named $name$ . Allowed categories names are bams, projects, reads, references, users, and workflow.
	Note: Only admin users have access to the users category.
	When executed at the $/projects$ level, opens your project named <i>name</i> .
	At the /projects/projname level, opens your analysis named name.
	At the /workflows level, opens the standard workflow named name.
	Note: The cd command navigates only one level at a time.

#### Administrative user commands:

**Note:** Only admin users have access to the users category. Must be executed at the /users level. These commands require admin role privilege.

ls	Lists all users.
mk <i>name</i>	Creates a new user named <i>name</i> , with the password <i>name</i> and the role user.
rm name	Deletes the LifeScope <sup>™</sup> Software shell user named <i>name</i> .
loggedin	Shows logged in users.
set username <i>role</i> [password]	Sets a user's role and optionally set the user's password. Valid roles are: admin, user
makepasswdfile	Creates an encrypted password file for non-interactive login. The file is named <pre>lscope.passwd,</pre> and is created in the directory where your <pre>lscope.sh</pre> shell command.is originally issued.
	When <pre>lscope.sh shell is given without a password, LifeScope<sup>™</sup> Software looks for a <pre>lscope.passwd file in current directory. If the file exists, <pre>lscope.sh uses its password. This allows the command <pre>lscope.sh -u username</pre> to log the user in without asking for password.</pre></pre></pre>

#### Other administrative commands:

These commands require admin role privilege.

Table 5 LifeScope <sup>™</sup>	<sup>1</sup> Software shell	commands	(continued)
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Command	Description
rebuild	In /reads, rebuilds the read repository index, which is used to display the list of read-sets in the repository. Recent changes (additions or deletions) to the reads repository are not reflected in the ls output until the index is rebuilt.
Miscellaneous comm	ands:
config	Displays configuration information for the LifeScope <sup>™</sup> Software server.
help	Briefly lists the LifeScope <sup>™</sup> Software shell commands and syntax. The man page is more verbose. (The man page is access with the lscope.sh man command, issued in the Linux shell.)
exit	Logs out of the LifeScope <sup>™</sup> Software web server, and exits the shell. Running analyses are not affected.
!cmd	Runs a Linux shell command. Example: !pwd
	Do not use !cmd to run a Linux lscope.sh command from within the command shell.
	Project and analysis naming rules
	Project and analysis names must conform to Linux filename conventions:
	<ul> <li>Contain only alphanumeric and underscore characters.</li> </ul>
	Do not contain spaces.
	<ul> <li>Must be unique (among siblings).</li> </ul>
	<ul> <li>Do not begin with an underscore character. (This requirement is a LifeScope<sup>™</sup> Software limitation, not a Linux rule.)</li> </ul>
	Command completion and command history

#### Command completion and command history

The LifeScope<sup>™</sup> Software command shell supports command completion, with the keyboard Tab key. The command shell supports command history, with the keyboard up arrow.

List shell commands
 There are two methods to list shell commands and command syntax.
 On the Linux command line, enter this command: lscope.sh man This command lists the lscope.sh man page. This version is more verbose than the help output within the command shell.
 In the LifeScope<sup>™</sup> Software command shell, enter this command: help Help lists the shell commands, with their syntax and with a brief description.
 Define input data
 This section describes adding reads input data to one analysis. The reads input can either be in the read repository or specified by absolute path to a file on the local file system.
 Plan your analysis input carefully. The way you define your reads input affects the behavior of your analysis. Index IDs and group names give you flexibility in handling your input data.

#### Index IDs

The shell supports two methods of specifying index IDs with your reads data. Both the add xsq command and groupsfiles support index IDs. Using an index ID restricts the input to only the data for a specific index. For example:

```
add xsq xsqfile:62
```

#### Groups processing

Combining reads data into groups causes the data to be treated as one specimen, with one set of results files that represents the combined data for the group.

Groups are created with the optional group field of the add xsq command, or with a groupsfile and either the set groups command or the -g groupsfile option of the lscope.sh run command. See "Run a grouped analysis" on page 88 for an example of a groupsfile.

The following is an example of an add xsq command with a group field:

```
add xsq xsqfile:62 Human
```

**Example input data** The following sections describe scenarios for defining your input data.

#### scenarios

#### Define the input reads for a sequencing library instance

Suppose your specimen is sequenced on the instrument with three indices (barcodes). You require that LifeScope<sup>™</sup> Software treat the data from these three indices as one specimen and generate one set of results files for the specimen. You combine your reads data into a group to achieve this effect. The following are two methods to combine these indices into one group.

These examples imply the data is in one XSQ file, but that is not required (XSQfile does not have to be the same XSQ file for all the indices).

The add xsq command

If the indices for your specimen are 3, 4, and 9, and you choose the group name Human for your specimen's data, the following commands cause your data to be analyzed as one specimen:

add	xsq	XSQfile:3	Human
add	xsq	XSQfile:4	Human
add	xsq	XSQfile:9	Human

#### A groupsfile

A groupsfile also associates the specimen's indices with a group, and is more convenient for large numbers of indices. Here are example contents of the groupsfile:

```
Human <path>/XSQfile:3 <path>/reference_file
Human <path>/XSQfile:4
Human <path>/XSQfile:9
```

Use this groupsfile with either the set groups shell command or the -g option of the lscope.sh run command:

set groups <path>/groupsfilename
lscope.sh run ... -g <path>/groupsfilename

#### Define reads in separate groups

Contrast the following treatment of reads data with the example in "Define the input reads for a sequencing library instance" on page 86:

add xsq XSQfile:3 Human add xsq XSQfile:4 Rat add xsq XSQfile:9 Ecoli

In this example the analysis is run in three groups, with each group containing the data from one index. All groups receive the same analysis modules processing (for example, mapping, SNPs, and CNV). The data from each index is processed separately, and each index generates an independent set of results files.

#### Non-indexed data

When you use non-indexed XSQ files, all read-sets in those files become your input. (If you use a group name that includes other read-sets, the new read-sets are combined with the other read-sets already assigned to that group.)

Both the add xsq command and a groupsfile can add non-indexed data to your analysis:

add xsq xsqfile1 Human
add xsq xsqfile2 Human
Human <path>/XSQfile1 <path>/reference\_file
Human <path>/XSQfile2

#### The effect of not using the id index argument with indexed data

Consider what happens if you add multiple indexed XSQ files to an analysis and do not use the id argument to specify individual indices. The effect is very different when you assign a group and you do not assign a group.

Example with a group

add xsq xsq1 groupX add xsq xsq2 groupX add xsq xsq3 groupX

For example, the above add xsq commands, with a group specified, define as input all the indices in the three XSQ files. All read-sets (in all three XSQ files) are treated as one specimen, and in the output files the data from all the read-sets is combined.

Example without groups

If you do not specify the group name, the shell assigns the data to groups based on index name. For example, consider the following commands, assuming each XSQ file contains indexed data:

add xsq xsq1 add xsq xsq2 add xsq xsq3

The shell assigns the read-sets to groups based on index name, using group names of Group\_id. Each index is treated as a different specimen, the data for different indices are kept separate, and a separate output directory is used for each index. If the same index name occurs in more than one XSQ file, the read-sets for those indices are combined into one group. This combination might not be what you intend. For example, if files xsq1, xsq2, and xsq3 each contains indices 1–96, the data is processed as 96 separate groups. Group 1 contains index 1 from all 3 files, group 2 contains index 2 from all three files, and so on.

#### Granularity

The most specific set of data that can be defined as input to an analysis run is the data from one index in one XSQ file. For example:

add xsq *xsqfile*:62

## Run a grouped analysis

This section describes the behavior of a grouped analysis. If an analysis has read-set groups associated with it, the following jobs are run: One secondary sub-analyses: one job for all read-sets. A number of tertiary sub-analyses: one job per read-set group. The following mechanisms add read-set groups to an analysis: • The group option of the add xsq command. The set groups command. The -g groupsfile option of the lscope.sh run command. Example of a The following gives an example of a basic groupsfile: groupsfile Human /data/reads/xsqfile1.xsq:3 <path>/reference\_file1 Human /data/reads/xsqfile1.xsq:21 Human /data/reads/xsqfile2.xsq:51 The pattern is: groupname XSQfile:id referencefile. These field are separated by tabs. The *referencefile* field is optional, except for the first entry of a group. If the *referencefile* field is empty, the last reference specified is used. All read-sets within one group must use the same reference (and the groupsfile must contain only one reference). This section describes how to do a run that analyzes reads from more than one sample, Run a multi-group using a groupsfile that specifies multiple groups. All read-sets within the analysis analysis must use the same reference. Multi-group groupsfile With a multi-group analysis, a single run of LifeScope<sup>™</sup> Software processes the input data from more than one group. For each group, LifeScope<sup>™</sup> Software keeps the data, processing, and results separate. The following is an example groups file for a multi-group analysis:

```
lung /data/reads/xsqfile1.xsq:1 /refs/reference_file
lung /data/reads/xsqfile1.xsq:5
lung /data/reads/xsqfile1.xsq:13
brain /data/reads/xsqfile1.xsq:2
brain /data/reads/xsqfile1.xsq:6
brain /data/reads/xsqfile1.xsq:9
brain /data/reads/xsqfile1.xsq:10
brain /data/reads/xsqfile1.xsq:14
kidney /data/reads/xsqfile1.xsq:3
kidney /data/reads/xsqfile1.xsq:11
kidney /data/reads/xsqfile1.xsq:15
```

This read-set file defines three sample groups:

- lung barcodes 1, 5, and 13
- brain barcodes 2, 6, 9, 10, and 14
- kidney barcodes 3, 11, and 15

With this read-set file, LifeScope<sup>™</sup> Software in one run performs the processing of three separate runs, one for each group. All groups in the groupsfile must use the same reference, and the reference must be specified on the first line of the file.

#### Example commands for a multi-group analysis

This section describes the steps to run a multi-group analysis. The groupsfile used in "Multi-group groupsfile" is suitable for this example.

Follow these steps to run the multi-group analysis:

1. Login to lscope shell:

lscope.sh shell -u user

2. Create a project and analysis, and open the analysis:

```
cd /projects
mk proj1
cd proj1
mk run1
cd run1
```

**3.** Set the groups:

set groups path\_to\_groupsfile.

**4.** (*Optional*) Set these parameters:

set analysis.species.name species

set analysis.assembly.name name

An example of a species name is ecoli. An example of an assembly name is DH10B.

If the assembly name is not provided, an assembly name of unknown is used. If the species name is not provided, the name of the reference file is used.

5. Ensure that the INI files in your analysis *do not* set the analysis.sample.name parameter. If this parameter is set in your INI files, the LifeScope<sup>™</sup> Software shell overrides each sample group it processes with the value from the INI file.

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**6.** Launch the multi-sample analysis:

run

7. Optionally check the status of the analysis:

ls

An alternative method to check the is entered on the Linux shell:

lscope.sh status -u user

# Run a tertiary-only workflow

When you run a standard workflow and specify only BAM input files (not XSQ files), the framework automatically runs only the tertiary portion of the workflow. The secondary (mapping) portion of the workflow is not attempted.

Two examples are given, one with add bam commands and one with the set group command and a groupsfile.

With add bam The following is an outline of the shell commands to run only the tertiary portion of a workflow:

```
cd projects
mk tertiaryProj
cd tertiaryProj
mk tertiaryRun
cd tertiaryRun
set workflow genomic.resequencing.frag
add bam /path/to/bamfile1 group1
add bam /path/to/bamfile2 group1
add bam /path/to/bamfile3 group1
set reference /path/to/reference
run
```

The genomic.resequencing.frag workflow includes both secondary and tertiary analyses. The secondary analyses take as input XSQ files, and the tertiary analyses take BAM files as input. Because the above example has only BAM files as input, LifeScope<sup>™</sup> Software automatically skips the secondary analyses. the workflow runs only the tertiary analyses.

With a groupsfile A tertiary-only workflow can also be run using a groupsfile and the set groups command. For example, outside of the LifeScope<sup>TM</sup> Software shell, create a groupsfile named tertiarygroups with this content:

```
group1 /path/to/bamfile1 /path/to/reference
group1 /path/to/bamfile2
group1 /path/to/bamfile3
```

The following is an outline of the LifeScope<sup>™</sup> Software shell commands to run only the tertiary portion of a workflow, using the groupsfile:

cd projects mk tertiaryProj cd tertiaryProj mk tertiaryRun2 cd tertiaryRun2 set workflow genomic.resequencing.frag<br/>set groups ./tertiarygroups<br/>set reference /path/to/reference<br/>runWith these commands, the groupsfile tertiarygroups must in the current directory,<br/>which is the directory you are in when you give the lscope.sh command.Run a multi-group<br/>tertiary analysisThis section describes how to run a tertiary-only workflow using BAM file input and<br/>using groups. With a multi-group analysis, a single run of LifeScope™ Software<br/>processes the input data from more than one group. For each group, LifeScope™<br/>Software keeps the data, processing, and results separate.

BAM files with different library types can be combined as long as the workflow and tertiary modules can process them.

Two examples are given, one with add bam commands and one with the set group command and a groupsfile.

#### With add bam commands

This example uses add bam commands to create 2 groups:

```
set workflow yourworkflow
add bam /path/to/bamfile1 group1
add bam /path/to/bamfile2 group1
add bam /path/to/bamfile3 group1
add bam /path/to/bamfile4 group2
add bam /path/to/bamfile5 group2
set reference /path/to/reference
run
```

#### With a groupsfile

This example uses a groupsfile to create the same 2 groups. Outside of the LifeScope<sup>™</sup> Software shell, create a groupsfile with this content:

```
group1 /path/to/bamfile1 /path/to/reference
group1 /path/to/bamfile2
group1 /path/to/bamfile3
group2 /path/to/bamfile4
group2 /path/to/bamfile5
```

These example LifeScope<sup>™</sup> Software shell commands use the groupsfile:

```
set workflow yourworkflow
set groups /path/to/groupsfile
set reference /path/to/reference
run
```

# Run an individual analysis

The optional examples download includes an examples of how to run selected LifeScope<sup>™</sup> Software analyses individually, without being part of a standard workflow or other sequence of analyses. Users new to LifeScope<sup>™</sup> Software are recommended to use either a standard workflow or one of the optional examples, instead of the procedures described in the rest of this section.

This section describes how to run a single LifeScope<sup>™</sup> Software analysis module, using the command shell.

Some options required you to create or edit PLN and INI files. See Appendix D, "Command Shell Control Files" on page 497 for information about these files.

- If you have already run the analysis, and the command shell has created the underlying PLN and INI files. You can use these PLN and INI files to run the analysis.
- You can optionally configure the analysis independently outside of the shell, using INI and PLN files that you create yourself.

This section describes these methods of running a single module:

- In shell mode Open the LifeScope<sup>™</sup> Software command shell, make a project and analysis, configure the analysis, and run the analysis in the command shell.
- In run mode Use the lscope.sh run command to start the analysis. To use this command the PLN and INI file must already be created and customized. Two possibilities for creating the PLN and INI files are:
  - If you have already run the analysis in the command shell, then LifeScope<sup>™</sup> Software has created these files for you.
  - You can optionally create the PLN and INI manually, for instance with a text editor.

The run mode command you use to run the analysis depends on whether you are running in the project repository or in the local file system.

• Run in the project repository – The run command is in this format:

lscope.sh run plnfile -u user -p project -a analysis \
 -g groupsfile

When you run in the projects repository with the lscope.sh run command, the
results files are handled by the shell project repository. Where the results files
appear on your local file system is configured during installation, and is given by
the lifescope.projects parameter in the <installdir>/server/
server.properties file.

• Run in the local file system – The run command is in this format:

lscope.sh run plnfile -u user

When you run in the local file system with the lscope.sh run command, the results files appear in a subdirectory to your current directory on the file system. This command is executed in the same directory as *plnfile*.

# **Review job status**

How you started your run	How to check the status
In LifeScope <sup><math>M</math></sup> Software command shell, with the shell run command.	Open the LifeScope <sup>™</sup> Software shell, cd to your project and then to your analysis, and execute the shell ls command:
	Log in to the LifeScope <sup>™</sup> Software command shell. cd /projects cd proj1 cd run1 ls
	Output similar to the following is displayed:
	ini secondary/bamstats.ini secondary/fragment.mapping.ini secondary/global.ini tertiary/dibayes.genome.ini
	readsets lung:DH10B_ColorSpaceOnly_Unbarcoded_50_ Frag.xsq:1:DH10B.fasta
	jobs
	run1 [70%]
	Output for completed jobs also has a success or failure notice:
	run1 [100%] [JOB_FAILURE], or
	run1 [100%] [JOB_SUCCESS]
In the Linux shell, with either the <pre>lscope.sh</pre> run or <pre>lscope.sh</pre> resume command, and the analysis is	At the Linux prompt, enter a lscope.sh status command, with the -p project and -a analysis options:
running in the projects repository.	lscope.sh status -u rjones -p project \ -a analysis
In the Linux shell, with either the lscope.sh run or lscope.sh resume command, and the analysis is	At the Linux prompt, enter a lscope.sh status command. Do not specify the -p project and -a analysis options:
running in the current directory.	lscope.sh status -u rjones

The ways to check on your run vary according to how you started your run, as shown in the following table.

# **Review results**

Reviewing the output files generated by your the LifeScope<sup>™</sup> Software analyses is performed outside of the LifeScope<sup>™</sup> Software command shell. You can either inspect your results files on the Linux cluster or transfer the results files to your local machine to review.

If your analysis is run in the projects repository, you can view results and download results files using the LifeScope<sup>TM</sup> Software UI. Refer to the LifeScope<sup>TM</sup> Genomic Analysis Software User Guide (Part no. 4465696).

If your analysis is not run in the projects repository, you must find your results files on the local file system. The results files are under the following directory:

<projects\_repo\_dir>/<user\_name>/<project\_name>/<analysis\_name>/outputs

where *projects\_repo\_dir* is the location of the shell projects repository on the local file system, and *user\_name*, *project\_name*, and *analysis\_name* are your shell user id, project name, and analysis name. Contact your LifeScope<sup>TM</sup> Software administrator for the location of the shell projects repository. The projects repository directory is also given by the lifescope.projects parameter in the file <*installdir*/server/server/server.properties.

## **Review logs**

You can typically use the LifeScope<sup>™</sup> Software shell status commands (see "Review job status" on page 93) to determine your jobs success or failure status and to examine any error messages.

In the event you also want to look at your job's run logs, by default they are located in the LifeScope<sup>™</sup> Software shell projects repository. The location of this repository is set at installtime and is specific to your installation.

If you run your analysis in the current directory, your logs are in that directory.

For successful runs, the last line of summary.log ends in:

```
Analysis finished successfully.
```

When lscope.sh finishes, it prints the status of each job. If an error occurs, then the job's summary log is also printed.

## Error behavior and error messages

**Case sensitivity** LifeScope<sup>™</sup> Software shell levels, names, and commands are case sensitive. For example, the following sequence shows that a project named D157 exists, and the command cd d157 (with incorrect case) is not successful.

```
~/projects> ls
D152
D157
~/projects> cd d157
Server returned error status: 404, message: Project d157 does
not exist
~/projects> cd D157
~/projects/D157>
```

Unrecognized commands	The LifeScope <sup>™</sup> Software shell display an error message "Invalid command" for unrecognized commands. In the following examples, the 1s command is first mistyped and then given in the wrong case. The string mkdir proj1, while a correct Linux command, is not recognized by the LifeScope <sup>™</sup> Software command shell. (The correct LifeScope <sup>™</sup> Software shell command is mk proj1.)
	<pre>~/projects&gt; lss Invalid command ~/projects&gt; LS Invalid command ~/projects&gt; mkdir proj1 Invalid command ~/projects&gt;</pre>
Illegal usage	Error: String index out of range: -1 The message is thrown when the shell command is recognized but an illegal or unrecognized argument is used. Remember that shell commands do not behave like Linux commands. For example, the LifeScope <sup>™</sup> Software shell 1s command has a specific function, to list the contents of the current level. When mistakenly used to list the contents of a child or parent level, or used with wildcards, the error message appears:
	<pre>~/projects&gt; ls D152 D157 ~/projects&gt; ls D157 Error: String index out of range: -1 ~projects/D157/&gt; ls Error: String index out of range: -1 ~projects/D157/&gt; ls D1* Error: String index out of range: -1</pre>

# Run a Standard Workflow Analysis

#### This chapter covers:

Overview	. 97
Standard workflows	. 97
Run a standard workflow	103
Create a new workflow	106
Edit a workflow's control files	108
Run the tertiary portion of a workflow	109

## **Overview**

A standard workflow is a built-in sequence of experiments that combine multiple LifeScope<sup>™</sup> Software modules into a single command shell run. A workflow's analyses correspond to a common biological application, such as resequencing. The standard workflow directory also includes two file conversion utilities for XSQ and BAM conversion.

## Standard workflows

The standard workflows are provided for the following analysis types:

- ChIP-Seq
- Genomic resequencing
- MethylMiner<sup>™</sup>
- Small RNA
- Targeted resequencing
- Whole transcriptome

#### The workflows and the modules they execute are described in Table 6.

 Table 6
 Standard workflow libraries, locations, and modules

Analysis type	Library type	Workflow name	LifeScope <sup>™</sup> Software modules involved
ChIP-Seq	Fragment	chip.seq	Secondary: • Fragment mapping • Mapping statistics
Genomic resequencing	Fragment	genomic.resequencing.frag	Secondary: • Mapping • Mapping statistics Tertiary: • SNPs • Human CNV • Small indels • Annotations
Genomic resequencing	Mate-pair	genomic.resequencing.lmp	Secondary: • Paired mapping • Mapping statistics Tertiary: • SNPs • Human CNV • Inversions • Large indels • Small indels • Annotations
Genomic resequencing	Paired-end	genomic.resequencing.pe	Secondary: Paired mapping Mapping statistics Tertiary: SNPs Inversions Large indels Human CNV Small indels Annotations
Methyl Miner	Fragment	methyl.miner.frag	Secondary: • Paired mapping • Mapping statistics
Methyl Miner	Paired-end	methyl.miner.pe	Secondary: • Fragment mapping • Mapping statistics

Analysis type	Library type	Workflow name	LifeScope <sup>™</sup> Software modules involved
Small RNA		small.rna	Secondary: • Small RNA mapping Tertiary: • Small RNA count • Small RNA coverage
Targeted resequencing	Fragment	targeted.resequencing.frag	Secondary: • SAET • Fragment mapping • Mapping statistics Tertiary: • Enrichment
			<ul><li>SNPs</li><li>Small indels</li><li>Annotations</li></ul>
Targeted resequencing	Paired-end	targeted.resequencing.pe	Secondary: • SAET • Paired mapping • Mapping statistics
			Tertiary: • Enrichment • SNPs • Small indels • Annotations
Whole transcriptome	Fragment	whole.transcriptome.frag	Secondary: • WT splice junction extractor • WT fragment mapping • Mapping statistics
			Tertiary: • WT counts • WT coverage • Splice finder
Whole transcriptome	Paired-end	whole.transcriptome.pe	Secondary: • WT exon sequence extractor • WT splice junction extractor • WT paired-end mapping • Mapping statistics
			Tertiary: • Splice finder • WT counts • WT coverage

#### Table 6 Standard workflow libraries, locations, and modules (continued)

<b>Table 6</b> Standard workflow libraries, locations, and modules <i>(continu</i> )
--

Analysis type	Library type	Workflow name	LifeScope <sup>™</sup> Software modules involved
_	_	bam.convert	For internal use only.
_	—	xsq.convert	For internal use only.

**Note:** For improved accuracy in the SNPs module, turn on the BAMStats parameter bamstats.enable.probe.position, to generate position error files and probe error files:

```
set bamstats.enable.probe.position 1 secondary/bamstats.ini
```

#### Turn off SAET

If you turn off SAET in a workflow, make sure your mapping INI file does not contain the line:

```
analysis.input.readset.file = ${analysis.output.dir}/saet/*.rrs
```

Comment out this line if it appears in your mapping INI file.

List standard workflows	List workflows using the LifeScope <sup>™</sup> Software command shell	
	This section describes two methods of listing the standard workflows available on your installation of LifeScope <sup>TM</sup> Software.	
	Workflows are listed in the LifeScope <sup>™</sup> Software command shell by doing an 1s command in the /workflows virtual directory. To display a list of the standard workflows,	
	<b>1.</b> Open the LifeScope <sup><math>TM</math></sup> Software command shell:	
	lscope.sh -u username [-w password]	
	2. Enter the workflow level: cd workflows	
	<b>3.</b> List the available workflows:	
	ls	
	The workflows are listed.	
	List workflows using Linux commands	
	The LifeScope <sup>TM</sup> Software standard workflows are installed under the directory < <i>installdir</i> >/etc/workflows. To list the standard workflows, run this command in the Linux shell:	
	ls <installdir>/etc/workflows</installdir>	
List a workflow's INI files and parameters	The command shell provides commands to list a workflow's control files and parameters. The ls command, when run in at the level of a standard workflow, lists its INI files. The cat command is used to list the parameter settings of one of the workflow's INI files. The shell commands are shown in bold in the following example:	
	server:/workflows> <b>cd small.rna</b> server:/workflows/small.rna> <b>ls</b> secondary/bamstats.ini	

```
secondary/global.ini
secondary/smallrnamapping.ini
tertiary/filtermapping.bamstats.ini
tertiary/global.ini
tertiary/srCount.ini
tertiary/srCoverage.ini
server:/workflows/small.rna> cat secondary/bamstats.ini
bamstats.input.dir=${analysis.output.dir}/smallrnamapping
bamstats.run=1
```

# Workflow directory structure

All standard workflows follow a consistent directory structure, shown in Table 7. Table 7 Workflow directory and file layout

Contents of top-level workflow directory	Contents of sub-directories
analysis.pln	_
secondary	secondary.pln
	global.ini
	module1.ini
	module2.ini
	<i>moduleN</i> .ini
tertiary	tertiary.pln
	global.ini
	module1.ini
	module2.ini
	moduleN.ini

Do not delete or rename any of the following directories or files for your workflow:

- The secondary directory
- The tertiary directory
- The secondary.pln file
- The tertiary.pln file
- The analysis.pln file

# Workflow modules and parameters

The analysis parameters used in the workflow modules control the behavior of the modules' algorithms. Table 8 lists where the module parameters are described in this guide.

 Table 8
 Location of parameter descriptions

Workflow	Module	Chapter	Section	Page
Genomic resequencing	Mapping	Chapter 8	"Fragment mapping parameters"	123
	Mapping statistics	Chapter 8	"Mapping statistics"	141
	SNPs	Chapter 9	"SNPs runtime parameters"	164
	CNV	Chapter 10	"CNV module parameter descriptions"	189
	Small indel	Chapter 12	"Small indel parameter description"	217
	Annotations	Chapter 14	"Annotation parameters" See also the SNPs, CNV, and small Indel chapters.	277
Targeted resequencing	SAET	Chapter 16	"SAET parameters"	317
	Mapping	Chapter 8	"Fragment mapping parameters"	123
	Mapping Statistics	Chapter 8	"Mapping statistics"	141
	Enrichment	Chapter 15	"Enrichment parameters"	305
	SNPs	Chapter 9	"SNPs runtime parameters"	164
	Small indel	Chapter 12	"Small indel parameter description"	217
	Annotations	Chapter 14	"Annotation parameters" See also the SNPs, and small Indel chapters.	277
Small RNA	Small RNA mapping	Chapter 21	21 "Small RNA mapping parameters"	
	Mapping statistics	Chapter 21	"Mapping statistics"	395
	Small RNA coverage	Chapter 22	"Small RNA coverage parameters"	424
	Small RNA counts	Chapter 23	"Small RNA counts parameters"	429
Whole transcriptome	WT mapping	Chapter 17	"Single-read mapping parameters",	334,
			"Paired-end parameters"	346
	Mapping statistics	Chapter 17	"Mapping statistics"	354
	WT coverage	Chapter 18	"WT coverage parameters"	370
	WT counts	Chapter 19	"WT counts parameter description"	374
	WT splice finder	Chapter 20	"Splice finder parameters"	380
ChIP-Seq	Mapping	Chapter 8	"Fragment mapping parameters"	123
	Mapping statistics	Chapter 8	"Mapping statistics"	141

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Workflow	Module	Chapter	Section	Page
MethylMiner™	Mapping	Chapter 8	"Fragment mapping parameters"	123
	Mapping statistics	Chapter 8	"Mapping statistics"	141

 Table 8
 Location of parameter descriptions (continued)

# Run a standard workflow

Prepare to run a standard workflow	This section explains the preparation required to run a standard workflow.						
	Complete the prerequisites						
	Complete the applicable prerequisites described in Chapter 3, "Before You Begin" on page 45.						
	Determine your input and reference files						
	Before you can run the standard workflows you must know:						
	• The read and references files required by your analyses						
	<ul> <li>The regions of interest file, for targeted resequencing analyses</li> </ul>						
	Any non-default module parameter values						
Order commands	Shell commands must be ordered in the following sequence:						
correctly	1. Create the project.						
	<b>2.</b> Create the analysis.						
	<b>3</b> . Set the workflow.						
	4. Add the read-sets.						
	<b>5.</b> Set the reference.						
	<b>6.</b> Set the assembly name, species name, and other parameters, if required.						
	These rules apply to the analysis configuration commands:						
	• The set workflow command must precede all other analysis configuration commands.						
	• Read-sets must be added before the reference is set. (These commands add read- sets to your analysis: set groups <i>path</i> , add xsg <i>path</i> , and add bam <i>path</i> .)						
	<ul> <li>The set reference command must precede these two commands:</li> </ul>						
	- set analysis.assembly.name						
	- set analysis.species.name						
Specify the regions of interest file	Only for targeted resequencing workflows, you must provide the regions of interest file:						
	<pre>set enrichment.target.file path tertiary/enrichment.ini</pre>						

An absolute path is recommended for *path*. A relative path is interpreted relative to the directory you are in, in the Linux shell, when you execute the lscope.sh command.

Run a workflow in<br/>the command shellThis section describes how to run the fragment targeted resequencing workflow in the<br/>LifeScope<sup>TM</sup> Software command shell.

In the LifeScope<sup>™</sup> Software command shell, running a standard workflow requires the following steps:

- 1. Create a LifeScope<sup>™</sup> Software command shell project and analysis.
- 2. Identify the workflow to be executed on your data.
- **3.** Identify your input data (your reads).
- Identify the reference genome.
- 5. For targeted resequencing workflows, identify the regions of interest file.
- **6.** Issue the run command to start your analysis.

#### Example steps to run a standard workflow

This section shows the steps to run a workflow in the LifeScope<sup>TM</sup> Software command shell. These steps use the targeted resequencing fragment workflow as an example. A description of each step is given in Table 9.

```
# log into the shell
lscope.sh shell -u username -w password
# import your data into the reads repository
cd /reads
import /data/xsq/DH10B_ColorSpaceOnly_Unbarcoded_50_Frag.xsq
# cd to the projects repository
cd /projects
# make a project, and open it
mk ecoli
cd ecoli
# make an analysis, and open it
mk run1
cd run1
# define the analysis type
set workflow targeted.resequencing.frag
# define your input data
add xsq DH10B_ColorSpaceOnly_Unbarcoded_50_Frag.xsq
# specify the reference to be used
set reference hg19
# optionally change parameter defaults after this line or
# set analysis-specific parameters
set analysis.regions.file /data/results/readsdata/dh10b.bed
# list the configuration of the analysis
ls
# start the run
run
# list progress information about the run
ls
```

Paths (/data/results) and filenames are examples only. In your actual workflow runs you use paths to your own data and reference files.

Table 9 explains the purpose of the steps shown above. Information about your Table 9 Description of sample shell commands to run a workflow

Command	Purpose		
lscope.sh shell -u username	Log into the command shell. The shell prompts for your password if you do not provide it with the $-w$ option.		
cd /reads import /data/xsq/DH10B_ColorSpaceOnly_ Unbarcoded_50_Frag.xsq	Go into the reads repository and import an XSQ file (the input data file). This command brings the XSQ reads file into the LifeScope <sup>™</sup> Software reads repository.		
cd /projects mk ecoli cd ecoli	In the projects repository, create a project named ecoli, and in that project create an analysis named run1. Open the run1 analysis.		
mk run1 cd run1	<b>Note:</b> The open analysis (cd run1, in this case) command is important. The data, reference, and other configuration commands which follow apply only to the current analysis.		
set workflow targeted.resequencing.frag	Define the workflow used with this analysis. The workflow in this example includes SAET, mapping, mapping statistics, enrichment, SNPs, small indel, and annotations.		
add xsq DH10B_Unbarcoded_50_Frag.xsq	Define the XSQ file containing the input reads data for this analysis. Multiple input files are allowed.		
<pre>set reference /data/results/results/ references/DH10B.fasta</pre>	Define the reference file for this analysis.		
<pre>set enrichment.target.file /data/ results/readsdata/dh10b.bed tertiary/ enrichment.ini</pre>	Define the regions of interest file, for targeted resequencing.		
ls	Display the configuration of your analysis, including input reads, reference files, workflow, and select parameter settings.		
run	Start the analysis run.		
ls	Display status information about the analysis run.		

#### Information about your workflow job

The output of the ls command displays the analysis' INI files, read-sets, and run completion percentage or completion status. Example output is shown below:

```
--ini--
fragmentMapping.ini
global.ini
--readsets--(group:file:index:reference)
Group_1:case0030_Fragment.xsq:1:chr16.validated.fasta
--jobs--
run1 [100%] [JOB_SUCCESS]
fragmentMapping [100%] [JOB_SUCCESS]]
```

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Run a workflow in lscope.sh run mode	The lscope.sh run mode command executes the workflow without opening the LifeScope <sup>™</sup> Software command shell. This command is used at the Linux command prompt.					
	The following command runs the small RNA workflow in run mode:					
	lscope.sh run < <i>path</i> >/small.rna/analysis.pln -p rna \					

-a run1 -u sjones

where *path* is the path to the workflow's location in the LifeScope<sup>™</sup> Software installation directory. The project and analysis are created if they do not already exist.

As an alternative, you can copy an existing workflow to a directory of your choice (use a Linux command such as cp -p to preserve the directory structure, or use the command shell get workflow command). Then use the following version of the command:

lscope.sh run ./analysis.pln -p rna -a run1 -u sjones

See "Use the command shell" on page 73 for more information about run mode.

## Create a new workflow

This section describes how to create a new workflow. The new workflow appears in the shell 1s command (but not in the LifeScope<sup>™</sup> Software UI list of workflows). You run the new workflow following the directions for a standard workflow. The new workflow must have a unique name that both follows Linux file naming conventions and also does not start with an underscore character. Note: When you create your own workflow, you can view the analysis and results files in the LifeScope<sup>TM</sup> Software UI. But the LifeScope<sup>TM</sup> Software UI does not support other analysis activities, such as reuse, with a custom workflow created on the command line. This section describes two methods to create a new workflow, one with a Linux cp command, and the other with the shell get workflow command. With the Linux cp Using the Linux cp command method requires that you know the location of the LifeScope<sup>™</sup> Software installation directory on the local file system. You do not open command the LifeScope<sup>™</sup> Software command shell. The create a new workflow, follow these steps: 1. Identify an existing workflow similar to the one you want to create. 2. Change directory to the *<installdir>/etc/workflows* directory. **3**. Create a directory with the name of your new workflow: mkdir newworkflow You use this name to define your analysis type with the get workflow command, and the workflow appears as this name when you list the contents of the /workflows virtual directory. **4.** Copy an existing workflow into the new directory. Use a command such as cp - pin order to copy the directory structure as well as the files. cp -p small.rna/\* newworkflow

	5. Edit the PLN and INI files in newworkflow to customize it according to the requirements of your analysis.
	Do not rename the secondary or tertiary directory, and do not rename any of the PLN files (analysis.pln, secondary.pln, or tertiary.pln).
With the shell get workflow command	The shell get workflow command copies the current workflow's INI files, PLN files, and directory structure to your current directory on the local file system. This method requires you to have (or create) a LifeScope <sup>™</sup> Software command shell analysis configured with the shell set workflow command.
	The create a new workflow, follow these steps:
	1. Identify an existing workflow similar to the one you want to create.
	<ol> <li>Log into the LifeScope<sup>™</sup> Software command shell. Cd to your project and open the analysis. Or, if the project and analysis do not exist, create them with commands such as the following:</li> </ol>
	cd /projects
	mk myproj
	cd myproj
	mk run1

- cd run1
- **3.** If the analysis does not already have a workflow defined, use the shell set workflow command and name the workflow whose files you want to copy. For example:

set workflow genomic.resequencing.pe

4. Use the get workflow command. For example:

server:/projects/demo/getwf> get workflow

Analysis workflow saved to /home/dir/demo/getwf

This commands pulls the workflow files from the server and copies them to the your local file system. The destination directory is determined by the Linux current working directory when the command shell was opened (where the lscope.sh shell command was given).

The following files and directories are copied:

analysis.pln secondary (including its PLN and INI files) tertiary (including its PLN and INI files)

**5.** Edit the copied PLN and INI files to customize them according to the requirements of your analysis.

Do not rename the secondary or tertiary directory, and do not rename any of the PLN files (analysis.pln, secondary.pln, or tertiary.pln).

# Edit a workflow's control files

This section describes the files that define the behavior of a standard workflow. The fragment genomic resequencing workflow, genomic.resequencing.frag, is used as an example. This section describes the workflow's directory structure and the purpose of its PLN and INI files.

**PLN files** Each standard workflows contains PLN files of the same name and at the same relative location. These PLN files are described in Table 10.

Table 10 PLN files in a standard workflow

File name and relative location	File contents and explanation
<workflow_name>/analysis.pln</workflow_name>	secondary.pln tertiary.pln < secondary.pln
	First, runs the modules listed in the file secondary.pln.
	If the secondary.pln runs complete successfully, then the modules listed in the file tertiary.pln are run.
<workflow_name>/secondary/secondary.pln</workflow_name>	Lists the INI files involved in secondary analysis for this workflow.
	Runs the LifeScope <sup>™</sup> Software modules specified in these INI files.
<workflow_name>/tertiary/tertiary.pln</workflow_name>	Lists the INI files involved in tertiary analysis for this workflow.
	Runs the LifeScope <sup>™</sup> Software modules specified in these INI files.

# INI files Table 11 shows the INI files involved in the genomic resequencing standard workflow, along with the PLN files that direct the execution of the workflow. The global.ini files define parameters that are imported into one or more module INI files in the same directory. (These parameters are considered global for the single directory, not for the entire workflow.)

Table 11	PLN	and	INI	files	in	а	standard	workflow
----------	-----	-----	-----	-------	----	---	----------	----------

File name and relative location	File contents and explanation
<workflow_name>/analysis.pln</workflow_name>	secondary.pln tertiary.pln < secondary.pln
	First, runs the modules listed in the file secondary.pln.
	If the secondary.pln runs complete successfully, then the modules listed in the file tertiary.pln are run.
<workflow_name>/secondary/secondary.pln</workflow_name>	seat.ini fragment.mapping.ini < saet.ini bamstats.ini < fragment.mapping.ini
	Runs SAET using the parameters in the file saet.ini.
	If SAET completes successfully, runs fragment mapping using the parameters in the file fragment.mapping.ini.
	If the fragment mapping module completes successfully, runs bamstats mapping statistics.
File name and relative location	File contents and explanation
---	--
<workflow_name>/tertiary/tertiary.pln</workflow_name>	<pre>dibayes.genome.ini cnv.ini small.indel.ini annotation.dibayes.ini &lt; dibayes.genome.ini annotation.cnv.ini &lt; cnv.ini annotation.small.indel.ini &lt; small.indel.ini</pre>
	Runs, in order, the (1) SNPs, (2) CNV, and (3) small indel modules, using the parameters in the respective INI files.
	If the SNPs module completes successfully, runs SNPs annotations.
	If the CNV module completes successfully, runs CNV annotations.
	If the small indel module completes successfully, runs small indel annotations.

Table 11 PLN and INI files in a standard wo
---

# **.run parameters** The \*.run parameters control which LifeScope<sup>TM</sup> Software modules execute. The following are examples of run parameters used in standard workflows:

dibayes.run=1
small.indel.run=1
cnv.run=1

The allowed values for run parameters are 0 and 1:

- 0: Do not run the module.
- 1: Run the module analysis (default).

Because mapping is a prerequisite for all other runs, fragment.mapping.run may not be set to 0 unless it has already been successfully run to completion. You may set the .run parameters other modules to 0, under these conditions:

- You do not want to run that module's analysis.
- That module's analysis has already been completed (for instance, if you are restarting the workflow run and repeating a particular module is not necessary).

The run parameters are set automatically by the shell during standard workflows. Accept the default for most use.

## Run the tertiary portion of a workflow

This section describes running only the tertiary part of a standard workflow. With this method, you can optionally use input data with mixed library types.

First, set a standard workflow, such as targeted.resequencing.pe. You use one or more add bam commands to define the input data for the tertiary analysis. When run, LifeScope<sup>™</sup> Software automatically only runs the tertiary portion of the workflow, because there are no XSQ read-sets in the analysis.

The following are partial example steps:

```
set workflow targeted.resequencing.pe
add bam file1.bam
add bam file2.bam
add bam file3.bam
set reference referencefile1
run
```

In this scenario, you can optionally combine BAM files of different library types. The type of data you have determines which type of standard workflow you must use. Follow these guidelines for the workflow to specify:

- If your input includes mate-pair data, use a .lmp workflow. Only the genomic.resequencing.lmp workflow supports mate-pair libraries.
- If your input does not include mate-pair data, and does include paired-end data, use a .pe workflow.
- If your input does not include any paired libraries, use a . frag workflow.

# PART III Analysis Modules

# Run a Resequencing Mapping Analysis

This chapter covers:

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## **Overview**

This chapter describes how to perform resequencing mapping. The mapping analysis results files are required as input for tertiary analysis modules.

During secondary analysis LifeScope<sup>™</sup> Software performs the following steps:

- Maps or aligns reads to a reference genome.
- Accepts one or more input XSQ (eXtensible SeQuence) files to generate one or more sorted BAM files containing aligned reads.
- For mate-pair or paired-end libraries, pairs alignments that have the same bead ID.
- (Optional) Generates mapping statistics.

The 5500 Series SOLiD<sup>™</sup> Sequencer generates XSQ files, optionally with ECC (Exact Call Chemistry) reads. If an ECC primer round has been performed, the XSQ output also includes the sequence information in base space, in addition to color space. The ECC primer round is recommended during sequencing, for significant increases in sequencing accuracy.

XSQ is an extensible binary file format for storing sequence data, and supports multiple independent reads at the same position in a fragment. For information on the XSQ file format refer to this site:

http://solidsoftwaretools.com/gf/project/xsq

# Examples of how to run a mapping analysis

In a standardThe following standard workflows provide examples of running the mapping module:workflow• genomic.resequencing.frag

- genomic.resequencing.lmp
- genomic.resequencing.pe

The following are example shell commands using reads and references from the LifeScope<sup>TM</sup> Software repository. This example turns off tertiary modules in the workflow.

```
# log into the shell
lscope.sh shell -u username -w password
# cd to the projects repository
cd /projects
# make a project, and open it
mk ecoli
cd ecoli
# make an analysis, and open it
mk run1
cd run1
# define the analysis type
set workflow genomic.resequencing.pe
# define the input
add xsg /data/xsg/run0209a_50_PE.xsg
add xsq /data/xsq/run0209b_50_PE.xsq
# specify the reference to be used
set reference hg19
# these commands turn off the tertiary modules
set cnv.run 0 tertiary/cnv.ini
set dibayes.run 0 tertiary/dibayes.ini
set inversion.run 0 tertiary/inversion.ini
set large.indel.run 0 tertiary/large.indel.ini
set small.indel.run 0 tertiary/small.indel.ini
# optionally change mapping parameter defaults after this line
# list the analysis configuration
ls
# run the analysis
run
# list progress information for the analysis
ls
```

Dummy XSQ file names are used in this example.

In order to change a mapping parameter value in your analysis, use the set param shell command after the line # *optionally change mapping parameter defaults after this line*. For instance, to turn on the mapping.in.base parameter, use this shell command:

set mapping.in.base 1 secondary/pair.mapping.ini

This command shows how to turn on the mapping.in.base parameter for the genomic.resequencing.frag workflow:

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set mapping.in.base 1 secondary/fragment.mapping.ini

See Chapter 7, "Run a Standard Workflow Analysis" on page 97 for more information on running standard workflows.

**As a demo analysis** The optional examples download also includes an examples of how to run the mapping module by itself, as an individual analysis that is not part of a standard workflow. Fragment and paired examples are included. The use of a standard workflow is preferred.

See Appendix E, "Demo Analyses" on page 507 for more information on running a demo analysis.

### Input files

	The mapping module accepts as input one or more input XSQ files. The input reads can have different read lengths but they must be of the same library type. For example, LMP read-sets of length 50 and 60 can be processed together. The XSQ files processed in a single analysis can not have different library types. In one mapping analysis, the input must be of only one library type, either Fragment, LMP, or PE.
Plan your input read-sets	Plan your analysis input carefully. The way you define your reads input affects the behavior of your analysis. The following factors control how your input data is analyzed:
	• <b>Index (barcode) IDs</b> – Using an index ID restricts your input to the reads data of one or more indices.
	• <b>Grouping of reads</b> – Each group of reads is analyses together as one specimen. The output data for a group is combined into one set of results files.
	• <b>Multiple sample runs</b> – Unrelated reads can be processed together in one run of LifeScope <sup>™</sup> Software, but analyzed separately as separate input data.
	See "Define input data" on page 85 for more information on designing adding input data to your analysis.
	Mapping one tag of paired data is not supported.
Legacy data	If the data you want to process with LifeScope <sup>™</sup> Software is in CSFASTA and QUAL files, these files must be converted to the XSQ file format, through one of the following methods:
	<ul> <li>The LifeScope<sup>™</sup> Software UI handles converting these files to the XSQ format before mapping.</li> </ul>
	<ul> <li>If you are using the LifeScope<sup>™</sup> Software command shell, you must first convert the CSFASTA and QUAL files to the XSQ format. See Appendix C, "XSQ Tools" on page 479 for information on the standalone XSQ converter.</li> </ul>

# Stages of mapping

The mapping algorithm involves following stages:

- Scatter The process of distributing reads to the available compute nodes, to achieve parallelization.
- Mapping Each tag is aligned to the reference.
- **Pairing** With paired-end data, two tags are individually mapped using the same mechanism as fragment mapping. Then pairing is performed. Fragment data skips this stage.
- **BAM generation** The process of converting the output of mapping and pairing to BAM format. For color-space data, the base translation step is also included. Due to the scattering step, the BAM files at each compute node are a sub-set of the final BAM file and are referred to as baby BAMs.

BAM generation involves these steps:

- Ma2BAM A conversion step, from MA format to BAM format, for fragment data only.
- Pa2BAM A conversion step, from PA format to BAM format, for paired data only.
- Refcor Reference-assisted color-to-base translation.
- Sort BAM Sorting the BAM file from bead-id order to coordinate order.
- Merge BAM Merging the mini-BAM files created in the sort phase into the final output BAM files.

Figure 1 shows the phases involved in mapping fragment data. *Tag1* refers to the input data in the F3 strand.





Figure 1 The phases within the mapping module, when mapping fragment data



Figure 2 shows the phases involved in mapping paired data. *Tag1* and *Tag2* refer to the input data on the two different strands.

Figure 2 The phases within the mapping module, when mapping mate-pair or paired-end data

Scatter

The scatter stage distributes reads to the available compute nodes, in order to gain the efficiency benefits of parallel computing.

The mapping module can accept multiple read-sets, which are allowed to have different read lengths. The input set of read-sets from multiple XSQ files is first split based on read length. Fragment read-sets are partitioned into separate categories based on these read length categories:

- $0 \leq \text{Length} < 25$  (uses first | second.map.scheme.unmapped | repetitive.1)
- 25 ≤ Length < 35 (uses first | second.map.scheme.unmapped | repetitive.25)
- 35 ≤ Length < 50 (uses first | second.map.scheme.unmapped | repetitive.50)
- 50 ≤ Length < 60 (uses first | second.map.scheme.unmapped | repetitive.60)
- 60 ≤ Length < 75 (uses first | second.map.scheme.unmapped | repetitive.75)
- 75 ≤ Length (uses first | second.map.scheme.unmapped | repetitive.75)
- Trimmed reads, with variable read lengths (uses first|second.map.scheme.unmapped|repetitive.variable)

Mate-pair read-sets are partitioned into separate categories based on the pair length. Example categories are:

- $(0.0) \le (\text{Length1}, \text{Length2}) < (25, 25)$
- $(25.25) \le (\text{Length1}, \text{Length2}) < (50, 50)$

- $(50.50) \le (\text{Length1}, \text{Length2}) \le (60, 60)$
- $(60.25) \le (\text{Length1}, \text{Length2}) < (75, 35)$

The reads from each category are scattered into multiple jobs based on the parameters fragmap.minreads.per.node and fragmap.number.of.nodes. These parameters are described in the performance section (see "Mapping performance" on page 125).

# **Fragment mapping** The Map phase (also known as mapreads) includes map 1, map 2, and maMerge. Map combines information from reads in XSQ-formatted input files and a FASTA-format reference file, to generate alignment information in an intermediate file format known as match file format (.ma). Unmapped or poorly mapped reads from the first stage of mapping (map 1) can enter another mapping stage (map 2). Both these mapping stages have independent controls with respect to the mapping scheme and the gapped alignment algorithm. Since a poorly-mapped read can have alignments reported from map 1 and map 2, the maMerge is invoked to report the best alignments from the two mapping stages.

In the default setting, two mapping stages are invoked. The first one does ungapped alignment only. The second one does gapped alignments only. The second stage is switched off for read-sets with lengths less than the value of the parameter min.length.for.aggressive.gapped.mapping (default is 50). The maMerge step is not invoked if only one stage of mapping is run.

Alignment finding and gap finding use a proprietary algorithm. The presence of a gap can not be investigated in the anchor region. In general, for read length L and anchor defined by A.B.C,

- If A is zero, mapping looks for gap from position A+1 to L.
- If C is zero, mapping looks for gap only at position 0 to C.

Each map step is a multi-threaded executable that can handle one or more XSQ files of one library type. The unmapped and poorly reads from the first mapping stage are XSQ-formatted so that they can be processed by subsequent stages of mapping. The match file format is an internal file format that stores alignment and gap information. The MA files from two stages of mapping and maMerge are all generated in the scratch folder. The parameters to control mapping and gap finding algorithms are in "Mapping algorithm" on page 126. For details of the algorithm, see "Mapping schemes" on page 133.

**Mate-pair mapping** Each tag is mapped using a mapping protocol similar to fragment mapping. The match files (MA files) are provided to pairing stage.

PairingThe pairing module matches pairs of reads from the F3 and R3 mapping results of a<br/>mate-pair run. Pairing also matches reads from the F3 and F5-P2 files of a paired-end<br/>mapping run.

Pairing is divided in two stages. In the first stage, an estimate of the insert size distribution is made. Pairing module accepts the MA files, calculates the insert size from uniquely mapped beads and writes out a BINNED file (\*.binned) containing insert size distribution for each read-set. A perl script is invoked on the collection of BINNED files to calculate the insert size statistics (max, min, average, standard deviation) per read set. Output of the perl module is a tab-separated file (ISS, \*.iss) containing insert size statistics per read set. This ISS file is passed to pa2BAM module

that loads the average and standard deviation information to the appropriate BAM file. Pairing executable is invoked one more time with this ISS file and the MA files. This time pairing, rescue, and indels are called. Output of this pairing stage is a PA file, in an internal format.

The pairing algorithm performs the following steps for each pair of reads:

- 1. It finds all good AAA pairs based on the order, orientation, and distance between the two reads. (See the genomic classifications listed in Tables 20 and 21 on page 156 and 157 for information about AAA and other three-letter categories.)
- **2.** If no AAA pair is found, and a reference sequence is available, the algorithm performs mate-pair rescue using hits to one tag as an anchor, and then scanning for the "sister tag" in the region predicted by the library insert size.

**Note:** Either F3 or R3 tags can serve as the anchor, as long as the number of hits is below the Z threshold. The Z threshold is determined by the value entered for the pairing.gapped.max.hits parameter. If the F3 tag has x alignments, the R3 tag has y alignments, and both tags have fewer than Z hits, then the x+y anchor candidates are examined (see Figure 3, Mate-pair rescue example).



Figure 3 Mate-pair rescue example

- **3.** It determines if a AAA pair is unique. If multiple AAA pairing candidates exist for a given pair of tags, then the pair whose best pairing candidate scores at least *x* higher than the second-highest scoring pairing candidate is still considered unique, and all other AAA candidates with lower scores are discarded. The value of *x* is determined by the setting of the pair.uniqueness.threshold parameter.
- **4.** For non-AAA pairs where both tags have unique hits, the algorithm performs additional classifications based on the strand, distance, and orientation of the tags.

If either of the two tags has multiple mapping locations, but the highest mapping score is more than half of *x* higher than the second-highest score, then the location with the highest score is still considered unique and all other locations are discarded.

The pairing module supports paired-end experiments in addition to mate-pair experiments. In paired-end experiments, the actual sequencing is done on the same strand in opposite directions. However, because the representation in the sequencing output file follows the convention of representing the sequence from 5' to 3', the matching and pairing modules attempt to match the F5 and F3 tags on different strands, facing each other. In addition, a distance constraint determined by insert size is satisfied (see Figure 4). The pairing algorithm follows the same steps as the matepair algorithm. Steps 1 and 2 are modified to enforce the different order and

orientation requirement. If you convert matching positions for F3 tags to the opposite strand, then paired-end pairing is performed the same way as it is with mate-pair. The three-letter classification is defined the same way in mate-pair and pair-end. (See Table 20 on page 156 and Table 21 on page 157 for the genomic classification tables.)



Figure 4 Paired-end tags example

BAM file generation

This section describes the phases involved in BAM file generation.

#### Ma2BAM, pa2BAM

Ma2BAM and pa2BAM are format converters that change match files to a pro-BAM format. These pro-BAM files do not contain sequence or quality information but contain relevant BAM meta data. The pro-BAM files are sorted by incoming bead-id order. Relevant meta data from XSQ files is captured in pro-BAM files at this stage. The parameters bamgen.mqv.threshold and create.unmapped.bam.files control the functionality of the ma2BAM and pa2BAM utilities. The pro-BAM files are generated on the scratch folder.

Ma2BAM takes output of mapping (MA files) and converts them to pro-BAM format. The pa2BAM takes output of pairing (PA files) and converts them to pro-BAM format. During BAM generation, the mapping and pairing quality values are also computed.

For reads with multiple ungapped alignments, the read with the highest mapping or pairing quality value is chosen as the primary alignment for the read, and is reported to the BAM file. In cases where there are multiple alignments with the same quality value, the primary alignment is chosen at random from among the alignments with the same quality value.

#### Mapping and pairing quality values

This section describes mapping quality value (MQV) and pairing quality value (PQV) scores. Quality values are Phred-scaled quality scores. For any given alignment, an MQV is a measure of the confidence of a read aligning to the particular location, given all possible alignments for the read. A PQV represents the confidence in both alignments of a pair, and is a combined quality score for both reads. Quality values are within the range 0–100:

- 0 The highest probability of error
- 100 The lowest probability of error

The following factors increase confidence and increase an alignment's quality value:

- Longer alignment
- Fewer possible alignments
- · Fewer mismatches within the alignment

These factors reduce confidence and reduce an alignment's quality value:

- Shorter alignment
- More possible alignments
- More mismatches within the alignment

The pairing algorithm reports multiple sets of possible alignments for any given pair of reads (F3/R3 tags for a mate-pair run and F3/F5-P2 tags for a paired-end run). The pairing quality algorithm uses a Bayesian approach to calculate the quality of a given alignment for a pair of reads and the alignment with the highest pairing quality value (PQV) is chosen as the primary alignment for the pair of reads. The PQVs represent the Phred-scaled quality score, and are useful for downstream variant detection modules such as diBayes, small indels, large indels, and CNV.

In order to be consistent with the Phred quality score (-10\*log10[prob(error)]) used widely in literature, the quality is computed as the negative log odds of misaligning the read (or pairs of reads, for PQV). The resulting quality values are normalized by the maximum possible value to ensure that the quality values are within the range 0–100.

#### Gapped alignments

The pairing algorithm searches for gapped alignments (indels) when one of the tags (F3/R3/F5-P2/F5-BC) maps to the reference genome and the other tag does not map to the genome within the insert-size range, or it maps within the insert-range but the alignment is short. If both an ungapped and a gapped alignment are found for a given read, then, due to the low prior probability of 10^-4 assigned to the gapped alignments, the PQV for gapped alignments is zero.



Figure 5 Example of a gapped alignment and a partial alignment

In calculating the PQV for gapped alignments, the alternative hypothesis tested is the probability of finding the partial ungapped alignments. The read with the gapped alignment is treated as two partial reads on either side of the indel start point. The partial read with the greater length is used as the partial alignment length for the alternate hypothesis.

#### Refcor

Reference Assisted Color-to-Base Translation (also known as refcor) combines information from color calls, optional ECC extra primer round calls, and the reference sequence, to enhance the quality of base-space reads by detecting and fixing color errors that convert into multiple base-space errors. Refcor accepts a set of pro-BAM files, which contain alignments information without base or color sequence, and generates a corresponding set of BAM files containing alignments with base-space and color-space calls (on the scratch folder). These BAM files are sorted by bead-ID and contain dummy base calls and QVs, but are otherwise compatible with the BAM specification. The refcor module produces bead-ID-sorted BAM files that have valid calls and QVs.

For each alignment, the module constructs the primer-transition graph with vertices corresponding to all possible k-mers (k=4 for 2BE+4BE, k=2 for 2BE) in each read position. Two adjacent vertices are connected with an edge if they overlap in *k*–1 bases. A score corresponding to the (1–Pe), where Pe is the probability of being erroneous, is assigned to each of the vertices. Scores are derived from quality values corresponding to enumerated calls. When making a base call in a position, the current algorithm considers the cumulative probability of all sequences that have called base in that position. The evidence from the reference (base) is assigned to the vertex with *k*-mer ending in corresponding base only if the color transition between current and following reference bases matches the color call in a read. Such assignment reduces the reference over-correction in SNP positions. In order to reduce reference overcorrection in positions with color errors, we assign to reference base a very low weight corresponding to QV of 8. The weight can be reduced to QV of 0, completely reducing the reference guiding in the color-to-base translation. This results in a smaller skew between reference to non-reference allele ratio. Increasing the weight results in false positive variant calls reduction. This process restores the phase shift caused by color errors or completely eliminates the errors.

The parameters bamgen.refcor.softclip, bamgen.refcor.addcs, refcor.reference.weight, and refcor.base.filter.qv control the refcor program. The defaults provided are validated for a wide range of data.

- If the XSQ file contains base-space data only, the XSQ base call and quality value are transferred to BAM file. In this case, no color calls are present in BAM file irrespective of the value of bamgen.refcor.addcs.
- If the XSQ file contains 2BE color-space data only, refcor uses aligned 2BE and reference to generate base translation for the aligned part of the read. The base sequence and quality value are generated using a proprietary algorithm. For reads with no alignment, bases are determined by naively translating color to base. For these unaligned reads, the QV for the base at position *k* is the minimum color QV for all positions  $\leq k$ .
- If the XSQ file contains ECC data, then reference with 2BE and 4BE color are used to generate the new base sequence. The base sequence and quality value are generated using a proprietary algorithm. For reads with no alignment, the original base-space data from ECC is preserved.

#### Sort BAM

The output of refcor is a set of BAM files in bead-id sorted order. These files are sorted in this step using Picard sort into coordinate sorted order before merge. These BAM files are referred to as mini-BAM files and are stored in the analysis temporary directory.

#### Merge BAM

This phase accepts a set of mini-BAM files and outputs a set of BAM files. These BAM files are stored in the analysis output directory (which is typically a network-attached storage device).

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One BAM file is generated per read-set, which is defined as a barcode index in one XSQ file. For a 96-barcoded dataset in a single XSQ file, either 96 or 97 BAM files are created. The 97th BAM contains reads that have the default barcode index.

## Fragment mapping parameters

This section describes the runtime parameters which control the behavior of your fragment mapping analysis. Table 12 lists these parameters.

In order to change a parameter value, use the set param shell command to replace the line [optionally change mapping defaults after this line] in the example commands in "Examples of how to run a mapping analysis" on page 114. For instance, to have the unmapped BAM file generated, change the parameter create.unmapped.bam.files with this command:

set create.unmapped.bam.files 1 secondary/fragment.mapping.ini

# Fragment mapping<br/>parameters tableTable 12 lists the fragment mapping parameters.See Table 16 on page 141 for mapping statistics (BAMStats) parameters.

Parameter name	Default value	Description
Optional parallelization parameters	5	
fragmap.number.of.nodes	4	The number of compute nodes available for this analysis.
fragmap.max.number.of.jobs	100	The maximum number of jobs allowed for mapping.
		Allowed values: Integers 24-1000.
mapping.np.per.node	8	The number of processors per node use for mapping. The reads are divided into the number of chunks specified for this parameter.
fragmap.minreads.per.node	4000000	If the total number of fragments exceeds this amount, then the analysis is distributed across the available nodes.
		Allowed values: Integers 1-125000000 (125 Million)
fragmap.maxreads.per.node	150000000	The maximum number of fragments that can be processed on a single node.
		Allowed values: Integers 1–150000000 (150 Million)
Optional resource parameters		
wall.time	120	Total time for the process to complete.
java.heap.space	1500	Dynamic memory requirement.

Table 12 Mapping parameter description

#### Table 12 Mapping parameter description (continued)

Parameter name	Default value	Description
mapping.memory	15gb	The total memory, in gigabytes, that is available for map reads, per compute node. Include the units gb in the setting.
		This number is set to 15 GB because the minimum hardware requirement for memory is 16 GB.
		<b>Note:</b> Typically, the job scheduler will allocate on nodes that have the requested memory available. For mixed hardware, it is suggested that different queues are created based on the memory available. Mapping jobs should be launched on high memory systems. For human references (hg18, hg19) and a 25.2 scheme, 45 GB is recommended for fastest performance. The smallest recommended RAM for human reference is 19 GB.
Optional mapping parameters		
mapping.in.base	false	Whether to map in base space (true) or color space (false). Set to true if input data has base space available.
		Allowed values:
		• false: Map in color space.
		• true: Map in base space.
		If only color space is available, then the module fails when base space mapping is turned on.
second.map.gapped.algorithm	GLOBAL	Whether or not to do indel finding, and controls the behavior of indel finding.
		Allowed values:
		GLOBAL: Mapping reports global alignment up to one indel.
		• LOCAL: Mapping reports local alignment up to one indel.
Optional BAM generation paramete	ers	
bamgen.refcor.softclip	0	For cases when the unaligned portion of a read needs to be presented in the BAM record, we use soft clipping in the CIGAR string. If a base-space sequence generated by ECC decoder is available, we use that sequence to represent corresponding part of the read. If the ECC decoded base-space is not available, then we naively translate using 2BE and/or +4BE evidence (if available).
		The soft clipped part of the read can be permanently clipped if either of these conditions is true:
		• The ECC base QV is less than 4.
		<ul> <li>The ECC base and 2+4 naive propagation have less than 50% similarity.</li> </ul>
		Allowed values: 0,1.
bamgen.refcor.addcs	1	Whether or not to add the color sequence to BAM records. If the XSQ file does not contain color space, this parameter is ignored, and the resulting BAM file output does not contain color space. See "Refcor" on page 121 for information about refcor. Allowed values: 0,1.

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Table 12	Mapping	parameter	description	(continued)
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Parameter name	Default value	Description
refcor.reference.weight	8	Reference weight. This parameter is used during base translation. In the read reconstruction process, multiple signals are combined to generate the final base call. This parameter adds weight (in terms of Phred score) only to signals which are compatible with reference. Color combinations that result in a variant are considered compatible with reference. Additional weight helps to eliminate base propagation errors caused by color error(s) during base translation. Allowed values: Integers 0–100.
refcor.base.filter.qv	10	Bases with a quality value below the value of this parameter provided are replaced with 'N'. Allowed values: Integers 0–1000.
bamgen.mqv.threshold	0	Provides control over the contents written to the output BAM file depending on the quality value of the alignment. To preserve only high quality alignments, set this value to a positive integer. Allowed values: Integers 0–1000.
create.unmapped.bam.files	FALSE	If set to TRUE, creates an additional BAM output file containing unmapped reads. Allowed values:
		• FALSE: Do not create the unmapped reads output file.
		• <b>TRUE</b> : Create a BAM output file containing the unmapped reads.

MappingMapping speed depends on hardware properties as well as on the scattering logic.performanceMapping runs are split into multiple jobs for processing efficiency. The following<br/>parameters affect how a mapping run is split:

- **fragmap.number.of.nodes** Specifies the number of jobs that are created, only if a split is necessary. The size of each split job is approximately the total number of reads divided by the number of jobs.
- **fragmap.max.number.of.jobs** Determines the maximum number of jobs that can be launched per analysis. Setting this parameter to a very large value can cause the scheduler to behave incorrectly.
- **fragmap.minreads.per.node** Determines the threshold of beads beyond which splitting occurs.
- **fragmap.maxreads.per.node** Determines the largest number of beads that can be mapped per node. The default is based on the minimum scratch space requirements. If you scratch space is smaller or larger, this number can be made smaller or larger respectively.
- **processors.per.node** This is the number of cores available for mapping analysis on each node.
- mapping.memory For human mapping and 25.2.0 scheme, there are 3 schema lines. Simultaneous handling of schema lines reduces I/O and therefore improves mapping speed.

Memory requirements for mapping jobs are the following:

- Reference: ~6 GB
- 1 schema line: ~13 GB
- 1 schema line and reference: ~19 GB
- 2 schema lines and reference: ~32 GB

• 3 schema lines and reference: ~45 GB

Below 19 GB, the reference is split up and I/O increases significantly. Pre-built hash tables are not used for memory less than 24 GB. LifeScope<sup>™</sup> Software supports the reuse of hash tables for human genome reference files (hg18 and hg19) and the default mapping schemes 35.2, 25.2, and 20.1. Pre-calculated color-space hash tables are included in the reference directory available with LifeScope<sup>™</sup> Software. If system RAM is 24 GB or higher, the hash tables are loaded into memory. Depending on the pipeline, this hash table reuse typically saves 2–3 hours in analysis processing time.

In general, set the mapping.memory parameter to be 1 GB less than the total system memory (RAM) available.

Mapping only scales when the number of reads that are mapped together is greater than 100 M. Splitting such that the total number of fragments per node is significantly less than 100 M is not recommended. The splitting can be explained using the following pseudo-code.

```
if (num.reads <= fragmap.minreads.per.node ) { num.jobs = 1}
reads.per.job = num.reads / fragmap.number.of.nodes
if (reads.per.job > fragmap.maxreads.per.node) {
    reads.per.job = fragmap.maxreads.per.node}
num.jobs = celing (num.reads / reads.per.job)
if (num.jobs > fragmap.max.number.of.jobs ) {
    throw exception; }
```

In above pseudo-code, the input read-sets (possibly from multiple XSQ files) have num.reads that fall in one read length category. Num.jobs is the total number of jobs launched for mapping analysis of num.reads beads.

In general, the mapping parameters allow mixing and matching single-anchor and multiple-anchor schemes, and also mixing and matching different seed lengths and different numbers of mismatches. For example, an unmapped scheme of 25.2.0, 35.2.0:15:25, 20.1.0, with a repetitive scheme of 35.2.0:30, 45.2.0, 60.2.0, is supported, but not recommended. The impact on performance can be significant.

## Mapping algorithm

# Internal mapping parameters

Table 13 lists mapping parameters that we do not recommend changing. The "Mapping schemes" section which follows describes how these parameters affect the behavior of the mapping module.

Parameter name	Default value	Description
fragment.mapping.run	1	Indicates whether or not to run the mapping module. Allowed values:
		• <b>0</b> : Do not run the mapping module.
		• 1: Run the mapping module.
		The run parameter is set automatically by the shell during a standard workflow. Accept the default for most use.

Table 13 Internal mapping parameters

Parameter name	Default value	Description
mapping.use.iub.reference	FALSE	Whether or not to support reference sequences that contain IUB codes. Allowed values:
		• FALSE: Do not support IUB codes in the reference file.
		• <b>TRUE</b> : Mapping allows matching to either alleles of bi-allelic IUB codes in the reference file.
bamgen.primary.output.filter.type	primary_only	Filters the content written to the output BAM file depending on whether the alignment is gapped or not. Note that unmapped reads are considered primary. Allowed values:
		<ul> <li>no_filtering: Report all alignments.</li> </ul>
		• <b>primary_only</b> : Report all primary alignments (both gapped and ungapped).
		• primary_ungapped_all_gapped: Report all gapped alignments (even if not primary) and report ungapped alignments only if they are primary. Default.
fragmap.num.mapping.stages	2	The number of mapping passes to perform in this analysis. Allowed values:
		• 1: Perform mapping in one pass. Gapped alignment finding is off by default.
		• 2: Perform mapping with two passes. Recommended for increased accuracy of indel finding, but with an increase in processing time. The second stage of mapping performs only gap alignments.
		When set to 2, do not set second.map.gapped.algorithm to NONE.
mapping.fragment.tag	F3	When mapping XSQ files with more than one tag, this parameter selects the tag to map. The value must match one of the tag names in the TagDetails group of the XSQ file.
		Allowed values: F3, R3, F5-P2, F5-BC, F5-DNA, F5-RNA.
		This feature should not be used unless for data quality analysis. If a LMP/PE XSQ file is processed for fragment mapping, tertiary analysis is not supported.
mapping.schema.file	_	Specifies the full path to the file containing the mapping schema. If this parameter is not set, the default installed file is used.

Parameter name	Default value	Description
min.length.for.aggressive.gapped. mapping	50	Overrides some of other parameters' settings, depending on the relation between this value and read length.
		If the read length of tag1 is less than this value, the following parameters are set:
		<ul> <li>num.mapping.stages.tag1=1</li> </ul>
		<ul> <li>first.map.gapped.algorithm.tag1=NONE</li> </ul>
		<ul> <li>second.map.gapped.algorithm.tag1=NONE</li> </ul>
		<ul> <li>pairing.gap.max.mismatches.tag1=2</li> </ul>
		If the read length of tag2 is less than this value, the following parameters are set:
		<ul> <li>num.mapping.stages.tag2=1</li> </ul>
		<ul> <li>first.map.gapped.algorithm.tag2=NONE</li> </ul>
		<ul> <li>second.map.gapped.algorithm.tag2=NONE</li> </ul>
		<ul> <li>pairing.gap.max.mismatches.tag2=2</li> </ul>
First map parameters	1	
first.map.gapped.algorithm	NONE	Whether or not to do indel finding, and controls the behavior of indel finding. Allowed values:
		NONE: Turns off indel finding.
		• <b>GLOBAL</b> : Mapping reports global gapped alignments up to one indel.
		<ul> <li>LOCAL: Mapping reports local gapped alignments up to one indel.</li> </ul>
		first.map.gapped.algorithm is set to NONE if the read length is less than min.length.for.aggressive.gapped.mapping.
first.map.gapped.alignments.only	0	Whether or not the first pass outputs <i>only</i> gapped alignments. Allowed values: Integers 0 or 1
		• <b>0</b> : Do not restrict the output to only gapped alignments.
		• 1: Restrict the output to only gapped alignments.
first.map.gapped.deletion	19	The maximum deletion size in a gapped alignment in the first pass.
first.map.gapped.insertion	4	The maximum insertion size in a gapped alignment in the first pass. Allowed range: 0=< insertion size <= read length
		The module fails if the insertion size is not within this range.

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Parameter name	Default value	Description
first.map.gapped.penalty	10	The penalty for a single gap in alignment, for the first mapping pass. Allowed values: Integers >= 0.
		When the penalty is 10, either side of the gap must have a score of at least 10 for the gapped alignment to exist. Gapped alignment is effectively turned off if the penalty is set greater than the read length.
		For a gapped alignment, scoring is determined by three factors:
		The matching score, which is 1.
		• The mismatch score, which is set by first.map.mismatch.penalty and is -2, by default.
		• This parameter (first.map.gapped.penalty), which sets the score associated with a gap.
first.map.gapped.error.indel	3	The number of mismatches allowed for gap alignments on the first mapping pass. Reasonable range: 1 <= value <= 15% of the read length. Allowed values: Integers, from 1 to the read length. Only used if first.map.gapped.algorithm is set to GLOBAL.
first.map.gapped.edge.length. insertions	12	The minimum edge length required for insertions, on the first pass. Reasonable range: 1–15% of the read length. Lower values may lead to more false positives. Allowed values: Integers, from 1 to the read length. Only used if first.map.gapped.algorithm is set to GLOBAL.
first.map.gapped.edge.length. deletions	12	The minimum edge length required for deletions, on the first pass. Reasonable range: 1–15. Lower values may lead to more false positives. Only used if first.map.gapped.algorithm is set to GLOBAL.
first.map.gapped.seed.window. left	40	The window size allowed to the left of the anchor alignment, on the first pass. Reasonable range: 5–50. Only used if first.map.gapped.algorithm is set to GLOBAL.
first.map.gapped.seed.window. right	80	The window size allowed to the right of the anchor alignment, on the first pass. Reasonable range: (5 + read length) - (50 + read length). Only used if first.map.gapped.algorithm is set to GLOBAL.
first.map.gap.min.non.matched	9	For the first pass, the minimum number of non-matches required for a read to go to the second pass of mapping (if performed). For example, if this value is set to 9, for a 50-bp read we perform the second pass of mapping on this read if none of the alignments found in the first pass is longer than 50–9=41 bases (every alignment has at least 9 unmatched bases). Only poorly mapped (not fully mapped) reads need additional mapping, probably with indel allowed.

Parameter name	Default value	Description
first.map.mask.positions.	-	A string of 0s and 1s that indicate the read positions to be masked during the first mapping pass. There is no penalty in mapping towards positions that are masked.
		Default: blank
		Leaving the parameter blank results in no masking. Leave the parameter blank for classic style mapping. (Masking replaces read trimming.)
		If set, the length of the string must match the anchor length or seed length (the first value of scheme parameter such as 25.2.0).
		With a 25-mer read and a 25.2.0 scheme, an example mask is 11111111111111111100000 (20 1s, then 5 0s). The bases corresponding to the 0s are ignored. Under the 0s portion of the mask, any mismatches do not count toward the mismatch penalty.
first.map.max.hits	50	The maximum number of alignments for a read on the first pass. This setting gives the maximum number of hits that are reported in the mapping output (even if more hits are found).
		Allowed values: Integers >= 1.
first.map.mismatch.penalty	-2	A scoring penalty for mismatches. Used in local alignment mode. Must be a negative number or zero. Allowed values: Integers <= 0.
		If this penalty is set to a value greater than the read length, effectively no mismatches are allowed on the first pass.
first.map.scheme.unmapped.1	0.0.0	The scheme used for unmapped reads of a length less than 25, for the first mapping stage. See "Mapping schemes" on page 133.
first.map.scheme.repetitive.1	_	The scheme used for repetitive reads of a length less than 25, for the first mapping stage. See "Mapping schemes" on page 133.
first.map.scheme.unmapped.25	25.2.0	The scheme used for unmapped reads of a length 25–34, for the first mapping stage. See "Mapping schemes" on page 133.
first.map.scheme.repetitive.25	_	The scheme used for repetitive reads with a length 25–34, for the first mapping stage. See "Mapping schemes" on page 133.
first.map.scheme.unmapped.35	25.2.0	The scheme used for unmapped reads with a length 35–49, for the first mapping stage. See "Mapping schemes" on page 133.
first.map.scheme.repetitive.35	_	The scheme used for repetitive reads with a length 35–49, for the first mapping stage. See "Mapping schemes" on page 133.
first.map.scheme.unmapped.50	25.2.0:15	The scheme used for unmapped reads with a length 50–59, for the first mapping stage. See "Mapping schemes" on page 133.
first.map.scheme.repetitive.50	-	The scheme used for repetitive reads with a length 50–59, for the first mapping stage. See "Mapping schemes" on page 133.
first.map.scheme.unmapped.60	25.2.0:20	The scheme used for unmapped reads with a length 60–74, for the first mapping stage. See "Mapping schemes" on page 133.
first.map.scheme.repetitive.60	-	The scheme used for repetitive reads with a length 60–74, for the first mapping stage. See "Mapping schemes" on page 133.
first.map.scheme.unmapped.75	25.2.0:20	The scheme used for unmapped reads with a length of 75 or larger, for the first mapping stage. See "Mapping schemes" on page 133.

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Parameter name	Default value	Description
first.map.scheme.repetitive.75	-	The scheme used for repetitive reads with a length of 75 or larger, for the first mapping stage. See "Mapping schemes" on page 133.
first.map.scheme.unmapped. variable	25.2.0:20	Use for unmapped reads of variable length coming from ECC trimmed base-space reads, for the first mapping stage. See "Mapping schemes" on page 133.
first.map.scheme.repetitive. variable	_	Use for repetitive reads of variable length coming from ECC trimmed base-space reads, for the first mapping stage. See "Mapping schemes" on page 133.
Second map parameters		
second.map.gapped.alignments. only	1	Whether or not the second pass outputs <i>only</i> gapped alignments. Allowed values:
		• <b>0</b> : Do not restrict the output to only gapped alignments.
		• 1: Restrict the output to only gapped alignments.
second.map.gapped.deletion	19	The maximum deletion size in a gapped alignment in the second mapping pass.
second.map.gapped.insertion	4	The maximum insertion size in a gapped alignment in the second mapping pass. Allowed range: 0=< insertion size <= read length.
		The module fails if the insertion size is not within this range.
second.map.gapped.penalty	10	The penalty for a single gap in alignment, for the second mapping pass. Allowed values: Integers >= 0.
		When the penalty is 10, either side of the gap must have a score of at least 10 for the gapped alignment to exist. Gapped alignment is effectively turned off if the penalty is set greater than the read length.
		For a gapped alignment, scoring is determined by three factors:
		The matching score, which is 1.
		<ul> <li>The mismatch score, which is set by second.map.mismatch.penalty and is -2, by default.</li> </ul>
		<ul> <li>This parameter (second.map.gapped.penalty), which sets the score associated with a gap.</li> </ul>
second.map.gapped.error.indel	3	The number of mismatches allowed for gap alignments on the second mapping pass. Reasonable range: 1 <= value <= 15% of the read length. Maximum value: the read length. Only used if second.map.gapped.algorithm is set to GLOBAL.
second.map.gapped.edge.length. insertions	12	The minimum edge length required for insertions, on the second mapping pass. Reasonable range: 1–15% of the read length. Lower values may lead to more false positives. Only used if second.map.gapped.algorithm is set to GLOBAL.
second.map.gapped.edge.length. deletions	12	The minimum edge length required for deletions, on the second mapping pass. Reasonable range: 1–15. Lower values may lead to more false positives. Only used if second.map.gapped.algorithm is set to GLOBAL.

Parameter name	Default value	Description
second.map.gapped.seed.window. left	40	The window size allowed to the left of the anchor alignment, on the second pass. Reasonable range: 5–50. Only used if second.map.gapped.algorithm is set to GLOBAL.
second.map.gapped.seed.window. right	80	The window size allowed to the right of the anchor alignment, on the second pass. Reasonable range: (5 + read length) - (50 + read length). Only used if second.map.gapped.algorithm is set to GLOBAL.
second.map.gap.min.non.matched. length	9	For the second pass, the minimum number of non-matches required for a read to go to third pass of mapping (if performed). We currently do not perform a third mapping pass, and this parameter is not used.
second.map.masked.positions	_	A string of 0s and 1s that indicate the read positions to be masked during the second mapping pass. There is no penalty in mapping towards positions that are masked.
		Default: blank
		Leaving the parameter blank results in no masking. Leave the parameter blank for classic style mapping. (Masking replaces read trimming.)
		If set, the length of the string must match the anchor length or seed length (the first value of scheme parameter such as 25.2.0).
		With a 25-mer read and a 25.2.0 scheme, an example mask is 111111111111111111100000 (20 1s, then 5 0s). The bases corresponding to the 0s are ignored. Under the 0s portion of the mask, any mismatches do not count toward the mismatch penalty.
second.map.max.hits.tag1	50	The maximum number of alignments for a read on the second mapping pass. This setting gives the maximum number of hits that are reported in the mapping output (even if more hits are found). Allowed values: Integers >= 1.
		The module fails if the value is not within this range.
second.map.max.hits.tag2	50	The maximum number of alignments for a read on the second mapping pass. This setting gives the maximum number of hits that are reported in the mapping output (even if more hits are found). Allowed values: Integers >= 1.
		The module fails if the value is not within this range.
second.map.mismatch.penalty	-2	A scoring penalty for mismatches. Used in local alignment mode. Must be a negative number or zero. Allowed values: Integers <= 0.
		If the absolute of the penalty is set to greater than the read length, effectively no mismatches are allowed on this pass.
second.map.scheme.unmapped.1	0.0.0	The scheme used for unmapped reads of a length 1–24, for the second mapping stage. See "Mapping schemes" on page 133.
second.map.scheme.repetitive.1	-	The scheme used for repetitive reads of a length 1–24, for the second mapping stage. See "Mapping schemes".
second.map.scheme.unmapped.25	20.1.0	The scheme used for unmapped reads with a length 25–34, for the second mapping stage. See "Mapping schemes".

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Parameter name	Default value	Description
second.map.scheme.repetitive.25	_	The scheme used for repetitive reads with a length 25–34, for the second mapping stage. See "Mapping schemes".
second.map.scheme.unmapped.35	20.1.0	The scheme used for unmapped reads with a length 35–49, for the second mapping stage. See "Mapping schemes".
second.map.scheme.repetitive.35	_	The scheme used for repetitive reads with a length 35–49, for the second mapping stage. See "Mapping schemes".
second.map.scheme.unmapped.50	20.1.0:30	The scheme used for unmapped reads with a length 50–59, for the second mapping stage. See "Mapping schemes".
second.map.scheme.repetitive.50	-	The scheme used for repetitive reads with a length 50–59, for the second mapping stage. See "Mapping schemes".
second.map.scheme.unmapped.60	20.1.0:30	The scheme used for unmapped reads with a length 60 to 74, for the second mapping stage. See "Mapping schemes".
second.map.scheme.repetitive.60	-	The scheme used for repetitive reads with a length 60–74, for the second mapping stage. See "Mapping schemes".
second.map.scheme.unmapped.75	20.1.0:30	The scheme used for unmapped reads with a length of 75 or larger, for the second mapping stage. See "Mapping schemes".
second.map.scheme.repetitive.75	_	The scheme used for repetitive reads with a length of 75 or larger, for the second mapping stage. See "Mapping schemes".
second.map.scheme.unmapped. variable	20.1.0:30	The scheme used for unmapped reads of variable length coming from ECC trimmed base-space reads, for the second mapping stage. See "Mapping schemes".
second.map.scheme.repetitive. variable	_	The scheme used for repetitive reads of variable length coming from ECC trimmed base-space reads, for the second mapping stage. See "Mapping schemes".

 Table 13 Internal mapping parameters (continued)

#### Mapping schemes

The mapping scheme parameters, listed in Table 13, determine the behavior of the mapping module. The LifeScope<sup>™</sup> Software mapping module supports two types of mapping, classic mapping and seed-and-extend mapping. These two approaches are described in more detail below.

- **Classic mapping** With this approach the seed length matches the full read length, and the seed anchors at the beginning of the read. For example, classic mapping for 50-bp reads uses a scheme such as 50.6.0. (In this example, 50, the seed length, matches the read length, and 0, the seed start position, starts at the beginning of the read.)
- **Seed-and-extend** The initial alignment step that locates short matches between a read and the reference sequence. The seed specifies the length of the attempted match. This approach is also called local alignment.

#### **Classic mapping**

To map a 50 bp read using classic mapping, a typical scheme to use is 50.6.0. This allows for up to 6 mismatches. Because the seed length equals the length of the read, there is no extension.

#### Seed-and-extend

With the seed-and-extend approach, a mapping scheme parameter value such as 25.2.0 is interpreted as follows:

- **First value** The seed length (25).
- Second value The quantity of allowed mismatches in the seed (2).
- **Third value** The start site of the seed within the read (0).



Figure 6 Seeds for local alignments

For example, with a 25.2.0 seed extend scheme, the mapping module examines the first 25 bases of a 50 bp read, and if that portion aligns to the reference with two or fewer mismatches, then the mapping module attempts seed extension. For any alignment, we try two different extensions and report the one with higher score. The two extension approaches are described below:

- **Ungapped extension** Extends from the anchor alignment, to find the best ungapped local alignment with a score *S*. The alignment length in this case can be shorter than the read length.
- Gapped extension Looks for one indel, following two different algorithms that are controlled by the parameters first.map.gapped.algorithm and second.map.gapped.algorithm. Only one of the algorithms can be used at a time. These algorithms are:
  - Local indel finding extension Extends from the anchor alignment, doing gapped extension with a modified Smith-Waterman-like algorithm. In this algorithm, we use a matching score (same as in ungapped extension) and gap penalty (\*.map.gapped.penalty), to find a best local alignment within user specified size restriction (\*.map.gapped.insertion/ deletion). If this alignment has a score larger than S (the best ungapped alignment extended from the same anchor alignment), the alignment is reported. The alignment length in this case can be shorter than the read length.
  - Global indel finding extension Extends from anchor alignment, does a full-length gapped extension with the allowed number of mismatches (\*.map.gapped.error.indel), and within user-specified size restrictions (\*.map.gapped.insertion/deletion), window restrictions (\*.map.gapped.seed.window.\*), and edge length restrictions (\*.map.gapped.edge.length.insertions/deletions). If this alignment has a score larger than *S*, it is reported. This algorithm is the same as what was used with the frag indel module in earlier versions of the software.

For information on sliding periodic seeds, refer to the publication at this site:

http://bioinformatics.oxfordjournals.org/content/25/19/2514.full

#### Unmapped and repetitive schemes

Mapping schemes have two categories, a scheme for unmapped reads (\*.map.scheme.unmapped.\*) and a scheme for repetitive reads (\*.map.scheme.repetitive.\*). Either type of scheme can also be either single-anchor or multi-anchor.

A single-anchor scheme is represented by a comma-separated list of values, where each value is in the format *seed\_length.mismatches.seed\_start\_position* (see Figure 6 on page 134). The following apply to both unmapped and repetitive single-anchor schemes:

- Scheme 25.2.0,25.2.20 First maps using scheme 25.2.0 and only for those reads that do not have an alignment, maps a second round using the scheme 25.2.20.
- Scheme 25.2.0,25.2.20,25.2.30 First maps using scheme 25.2.0 and only for those reads that do not have an alignment, maps a second round using the scheme 25.2.20. The reads that do not map to 25.2.20 either are then tried with scheme 25.2.30.
- When both repetitive and unmapped are specified, the repetitive schemes are applied only to the repetitive reads from first unmapped scheme. Figure 7 depicts the logic followed for an unmapped scheme of 25.2.0, 25.2.20 and a repetitive scheme of 35.2.0, 35.2.30.

Multi-anchor schemes are represented by multiple colon-separated seed start positions, in the format *seed\_length.mismatches.seed\_start\_pos1:seed\_start\_pos2*. The following apply to both unmapped and repetitive multi-anchor schemes:

- Scheme 25.2.0:20 Simultaneously maps using scheme 25.2.0 and 25.2.20, and reports all alignments.
- Scheme 25.2.0:20:30 Simultaneously maps using scheme 25.2.0, 25.2.20 and 25.2.30, and reports all alignments.
- Figure 8 depicts the logic followed for an unmapped scheme of 25.2.0:20 and a repetitive scheme of 35.2.0:30.

The Figures 7 and 8 describe how the mapping scheme parameters are used in the mapping module. su1+su2 and sr1+sr2 are in the form of (L,m,b1)+(L,m,b2), where *L* is the seed length, *m* is number of mismatches allowed in the seed, *b1* and *b2* are the seed starting positions, *H* is the number of hits, and *Z* is the maximum number of alignments reported by the mapping program.

In these figures, *su* refers to a mapping scheme for matching unmapped reads. And *sr* refers to a mapping scheme for matching repetitive reads.

When H < Z, the read is not in a repetitive region. When  $H \ge Z$ , the read matches too many locations on the reference, so it is considered a repetitive read, or sequenced from a repetitive region.

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Figure 7 Single-anchor mapping for unmapped and repetitive reads



Figure 8 Multi-anchor mapping for unmapped and repetitive reads

#### Pairing parameters Table 15 lists parameters for the pairing algorithm.

In order to change a parameter value, use the set param shell command to replace the line [optionally change mapping defaults after this line] in the example commands in "Examples of how to run a mapping analysis" on page 114. For instance, to adjust the default minimum insert size, change the parameter default.minimum.insert.size with this command:

set default.minimum.insert.size 200 secondary/pair.mapping.ini.

#### Table 14 Pairing parameters

Parameter name	Default value	Description
default.minimum.insert.size	0	The minimum insert size allowed for the insert size range estimate. This value is used if this measurement fails. Allowed values: Integers 0–10000.
default.maximum.insert.size	20000 for LMP 2000 for PE	The maximum insert size allowed for the insert size range estimate. This value is used if this measurement fails. Allowed values: Integers 0–10000.

# Internal pairing parameters

Table 15 lists pairing parameters that we do not recommend changing.

The first stage of pairing does pairing without finding gaps. This stage is followed by 4 stages of pairing that allow gaps. Values in a format such as 0:3:14:14:0 are a list of one value for each indel pass, separated by colons (:).

Table 15 Internal pairing parameters

Parameter name	Default value	Description
pairing.gap.on	1	Enables indel searching in pairing analysis. Allowed values are:
		• <b>0</b> : Do not include indel search.
		• 1: Include indel search.
pairing.anchor.max.mismatches	4, if indel searching is off	Total mismatches for both reads of a pair. If indel searching is on, then this value is specified as a list of
	4:5:5:3:5, with indel searching	one number for each indel pass, separated by colons (:).
pairing.gap.min.non.matched.length	9	Used to determine whether the pair goes through an indel pass. If the regular pairing finds a good pair of alignments that are both longer than read length minus this value, then the pair does not need an indel finding run. Allowed values: Integers >= 0.
pairing.gap.max.mismatches.tag1	5	Number of mismatches allowed in the extension part of the tag 1. (The extension typically is 5% to 10% of the read length.)
		pairing.gap.max.mismatches.tag1 is set to 2 if the read length is less than the value of the parameter min.length.for.aggressive.gapped.mapping.

#### Table 15 Internal pairing parameters (continued)

Parameter name	Default value	Description
pairing.gap.max.mismatches.tag2	5	Number of mismatches allowed in the extension part of the tag 2.
		pairing.gap.max.mismatches.tag2 is set to 2 if the read length is less than the value of the parameter min.length.for.aggressive.gapped.mapping.
pairing.gapped.max.hits	50:10	For ungapped and gapped alignment determination during rescue, this value determines the maximum number of rescues attempted for each read. If a read has more than this number of hits, rescue is not performed on this bead.
		The first value is used for ungapped rescue. The second value is used for gapped rescue
pairing.output.uniqueness.type	-1	Whether to output non-unique hits. Allowed values:
		• -1: A clear zone determines uniqueness. 1 or 2 best pairs are output, depending on whether it is unique.
		• <b>0</b> : All good pairs are output.
		• 1: Only unique good pairs are output.
pairing.annotation.type		Whether to output non-AAA pairs. Allowed values:
		• <b>0</b> : Do not output non-AAA pairs.
		• 1: Output non-AAA pairs, only if pairing and rescue do not find good pairs.
pairing.mismatch.penalty	-2.0	A single penalty applied to alignments, to compare the significance between alignments. Helps evaluate the alignments. Allowed values: Floats <0.
pair.uniqueness.threshold	10.0	This value defines the clear zone. This value is considered only when pairing.output.uniqueness.type is set to -1. If the score of the best alignment is this threshold more than the score of the second best alignment, the best alignment is considered uniquely placed. A threshold of zero means that this uniqueness is strictly adhered to, regardless of the alignment's relative scores.

#### General pairing indel parameters

For these parameters each value is a colon-separated list of five numbers, which are used, in order, for the five pairing passes. The first pass of pairing does pairing without finding gaps. This pass is followed by four passes of pairing that allow gaps. These values are potentially overridden by size-specific indel parameters in the remaining sections of this table.

pairing.gap.max.deletion.size	0:11:0:0:500	The maximum deletion size allowed during pairing.
pairing.gap.max.insertion.size	0:4:15:21:0	The maximum insertion size allowed during pairing.
pairing.gap.edge.length.deletions	0:3:0:0:20	The gap edge length for deletions during pairing.
pairing.gap.edge.length.insertion	0:3:14:14:0	The gap edge length for insertions during pairing.

#### Pairing indel parameters for tag lengths 75–35

For pairs of these specific lengths, parameters in this section are used instead of the general pairing indel parameters. This section applies to pairs with F3 lengths of 75 and above, and R3 or F5 lengths of 35–59.

Parameter name	Default value	Description
pairing.gap.max.deletion.size.75-35	0:11:0:0:500	The maximum deletion size allowed for these lengths.
pairing.gap.max.insertion.size.75-35	0:4:15:21:0	The maximum insertion size allowed for these lengths.
pairing.gap.edge.length.deletions.75-35	0:3:0:0:20	The gap edge length for deletions for these lengths.
pairing.gap.edge.length.insertion.75-35	0:3:14:14:0	The gap edge length for insertions for these lengths.

#### Table 15 Internal pairing parameters (continued)

#### Pairing indel parameters for tag lengths 60-60

For pairs of these specific lengths, parameters in this section are used instead of the general pairing indel parameters. This section applies to pairs with both tags with lengths of 60–74.

pairing.gap.max.deletion.size.60-60	0:11:0:0:500	The maximum deletion size allowed for these lengths.
pairing.gap.max.insertion.size.60-60	0:4:15:21:0	The maximum insertion size allowed for these lengths.
pairing.gap.edge.length.deletion.size.60-60	0:3:0:0:20	The gap edge length for deletions for these lengths.
pairing.gap.edge.length.insertion.size.60-60	0:3:14:14:0	The gap edge length for insertions for these lengths.

#### Pairing indel parameters for tag lengths 50-50

For pairs of these specific lengths, parameters in this section are used instead of the general pairing indel parameters. This section applies to pairs with both tags with lengths of 50–59.

pairing.gap.max.deletion.size.50-50	0:11:0:0:500	The maximum deletion size allowed for these lengths.
pairing.gap.max.insertion.size.50-50	0:4:15:21:0	The maximum insertion size allowed for these lengths.
pairing.gap.edge.length.deletion.size.50-50	0:3:0:0:20	The gap edge length for deletions for these lengths.
pairing.gap.edge.length.insertion.size.50-50	0:3:14:14:0	The gap edge length for insertions for these lengths.

#### Pairing indel parameters for tag lengths 35–35

For pairs of these specific lengths, parameters in this section are used instead of the general pairing indel parameters. This section applies to pairs with both tags with lengths of 35–49.

pairing.gap.max.deletion.size.35-35	0:11	The maximum deletion size allowed for these lengths.
pairing.gap.max.insertion.size.35-35	0:4	The maximum insertion size allowed for these lengths.
pairing.gap.edge.length.deletions.35-35	0:3	The gap edge length for deletions for these lengths.
pairing.gap.edge.length.insertions.35-35	0:3	The gap edge length for insertions for these lengths.

#### Pairing indel parameters for tag lengths 25–25

For pairs of these specific lengths, parameters in this section are used instead of the general pairing indel parameters. This section applies to pairs with both tags with lengths of 25–34.

pairing.gap.max.deletion.size.25-25	0:11	The maximum deletion size allowed for these lengths.
pairing.gap.max.insertion.size.25-25	0:4	The maximum insertion size allowed for these lengths.
pairing.gap.edge.length.deletion.size.25-25	0:3	The gap edge length for deletions for these lengths.
pairing.gap.edge.length.insertion.size.25-25	0:3	The gap edge length for insertions for these lengths.

#### Pairing indel parameters for tag lengths 50-35

For pairs of these specific lengths, parameters in this section are used instead of the general pairing indel parameters. This section applies to pairs with F3 lengths of 50–74, and R3 or F5 lengths of 35–59.

The first pass of pairing does pairing without finding gaps. The second pass of pairing allows gaps.

pairing.gap.max.deletion.size.50-35	0:11	The maximum deletion size allowed for these lengths.
pairing.gap.max.insertion.size.50-35	0:4	The maximum insertion size allowed for these lengths.

#### Table 15 Internal pairing parameters (continued)

Parameter name	Default value	Description
pairing.gap.edge.length.deletions.50-35	0:3	The gap edge length for deletions for these lengths.
pairing.gap.edge.length.insertions.50-35	0:3	The gap edge length for insertions for these lengths.

#### Pairing indel parameters for tag lengths 50-25

For pairs of these specific lengths, parameters in this section are used instead of the general pairing indel parameters. This section applies to pairs with F3 lengths of 50–74, and R3 or F5 lengths of 25–34.

The first pass of pairing does pairing without finding gaps. The second pass of pairing allows gaps.

pairing.gap.max.deletion.size.50-25	0:11	The maximum deletion size allowed for these lengths.
pairing.gap.max.insertion.size.50-25	0:4	The maximum insertion size allowed for these lengths.
pairing.gap.edge.length.deletions.50-25	0:3	The gap edge length for deletions for these lengths.
pairing.gap.edge.length.insertions.50-25	0:3	The gap edge length for insertions for these lengths.

## Mapping output files

The resequencing mapping module generates a BAM file containing alignments in coordinate order. For information about the BAM file, see Appendix B, "File Format Descriptions" on page 455.

The number of BAM files generated depends on the input XSQ files. One BAM file is generated per read-set (a read-set is defined as data from one barcode, in one XSQ file). If the input data is an XSQ file containing 96 barcodes, then the mapping module generates at least 96 output BAM files. Beads that are unclassified in any barcode are output into a separate additional BAM file. If create.unmapped.bam.files is set to TRUE, then an additional 96 output BAM files corresponding to the unmapped reads are also generated.

The filenames for the output BAM files are created using information from the input XSQ file, including: file base name, file id, index name, and index id. The BAM files are named according to the following patterns:

- Non-indexed BAM files: xsqname-fileID-1.bam
- Indexed BAM files: xsqname-fileID-idx\_bcIndex-bcID.bam

The fields in the filenames are:

- **xsqname** The XSQ file base name, without the .xsq extension.
- **fileID** The file ID for internal XSQ file tracking.
- **bcIndex** The barcode index.
- **bcID** The barcode identifier for internal barcode tracking.

The directory structure for mapping output is as follows:

The directory name <*mappingIniFileBasename*> is the basename of the mapping INI file. The defaults are fragment.mapping and pair.mapping.

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The string *<sampleName*\*> is determined by the sample description for the particular read-set.

The mapping output files are used by the BAMStats mapping statistics module and by the LifeScope<sup>™</sup> Software tertiary analysis modules.

## **Mapping statistics**

Mapping statistics occur after mapping as an optional post-processing step named BAMStats. BAMStats accepts the output of the mapping step and generates statistics files to provide an in-depth understanding of the experimental data and to better detect the presence of anomalies. LifeScope<sup>TM</sup> Software shell users can display the mapping statistics output data as a chart with a spreadsheet program or other third-party program. A subset of the output from mapping can be visualized in the LifeScope<sup>TM</sup> Software UI as a series of line and bar charts, if the analysis is run in the projects repository.

# Mapping statisticsTable 16 lists the parameters which control the BAMStats output.parameters

Table 16 BAMStats parameter description

Parameter name	Default value	Description
Optional parameters		
bamstats.maximum.coverage	10000	Defines the maximum coverage allowed for locations in the reference. Locations with coverage more than the maximum coverage value are ignored during coverage calculations. Allowed values: Integers 1–10000.
bamstats.maximum.isize	100000	The maximum inset size for LMP and PE libraries. Reads with an insert size more then the specified value are ignored for Insert Range Report calculation.
		Allowed values: Integers 1–100000.
bamstats.wig.primary.only	1	Use only primary alignments for coverage in WIG file format. Allowed values:
		• <b>0</b> : Do not restrict coverage in WIG file format to only primary alignments.
		<ul> <li>1: Restrict coverage in WIG file format to only primary alignments.</li> </ul>
bamstats.combined.report.both. strands	0	Whether or not to combine data from both strands for coverage in WIG format. Allowed values: 0,1.
bamstats.wig.binsize	100	The bin size for coverage in WIG file format.
		Allowed values: Integers 1–100000.
bamstats.bin.isize	100	The bin size for insert range distribution.
		Allowed values: Integers 1–100000.

#### Table 16 BAMStats parameter description (continued)

Parameter name	Default value	Description
bamstats.wig.combined.report.both .strands	0	Whether to combine data from both the strands for coverage in WIG format. Allowed values:
		• <b>0</b> : Do not combine data.
		<ul> <li>1: Combine data from both the strands for coverage in WIG format.</li> </ul>
Optional resource parameters		
wall.time	120	Total time for the process to complete.
java.heap.space	13000	Dynamic memory requirement.

#### Table 17 lists BAMStats parameters that we do not recommend changing.

Table 17 BAMStats internal parameter description

Parameter name	Default value	Description
bamstats.run	1	Whether or not to run the BAMStats module. Allowed values:
		• <b>0</b> : Do not run the BAMStats module. BAMStats statistics are not generated.
		• 1: Run the BAMStats module during this analysis.
		The run parameter is set automatically by the shell during a standard workflow. Accept the default for most use.
bamstats.group.stats	1	Enable combined statistics covering all read-sets in the group or sample. Allowed values:
		• <b>0</b> : Only statistics for BAM files are generated.
		• 1: Also generate combined statistics for the group or sample.
bamstats.maximum.mismatches	100	The maximum mismatches allowed in the alignments. Any alignment with more then the specified number of mismatches is ignored while generating reports related to number of mismatches. Allowed values: Integers 0–100.
bamstats.maximum.baseqv	100	Max base quality values. Any base with base quality value more then the specified value is ignored while generating reports. Allowed values: Integers 0–100.
bamstats.maximum.mappingqv	255	Maximum mapping quality value. Any alignment with mapping quality value more than the specified value is ignored. Allowed values: Integers 0–255.
bamstats.wig.minimum.mappingqv	2	Defines the minimum mapping quality allowed for coverage in WIG format. Any alignment with a mapping quality value less than this value is ignored. Allowed values: Integers 0–100.
bamstats.enable.probe.position	0	Enable probe and position error reports. Allowed values:
		• <b>0</b> : Do not generate probe and position error reports.
		• 1: Also generate probe and position error reports.

Summary of mapping statistics output

#### File formats

For every input BAM file, a set of statistics files are generated. These files are in CHT, CSV, TXT, and WIG formats. Each CHT file corresponds to one displayed chart. A CHT file specifies the type of chart, the displayed range of each axis, and the data points, without using external references.

The CHT file format is an internal file format based on the CSV file format, with addition header information included. CHT header information is the following:

```
# name:
# type: scatter2d | pie | vbar | line
# title:
# xaxisname:
# yaxisname:
# yrange: <min>:<tickinterval>:<max>
# yrange: <min>:<tickinterval>:<max>
XAXISNAME, SERIES1NAME, SERIES2NAME, ...
x1, y1.1, y1.2, ...
x2, y2.1, y2.2, ...
x3, y3.1, y3.2, ...
```

The wiggle format (.wig) is a public format typically used for coverage. Visit their site for more information:

hgdownload.cse.ucsc.edu/goldenPath/help/wiggle.html

A genome browser such as the Integrative Genomics Viewer (IGV) can be used to visualize the coverage. For information is available from their site:

www.broadinstitute.org/igv/

For a collection of input BAM files that belong to a sample, a set of cumulative statistics files are generated. The cumulative statistics files are also in CHT, CSV, TXT, and WIG formats. The cumulative statistics can be visualized in the LifeScope<sup>™</sup> Software UI.

#### **Directory structure**

The output of the BAMStats module has the following directory structure:

```
bamstats/
    <sampleName>/*.cht, *.tbl
    <sampleName>/<bam>/.*cht
    <sampleName>/<bam>/Misc/*.csv, *.txt, *.wig
    <sampleName>/Misc/*.csv, *.txt
```

The position error files and probe errors files are created in the BAM file directories.

#### **Overview**

BAMStats generates a tab-separated summary file (BAMfilename-summary.tbl) that summarizes key mapping quality statistics per input BAM file. The summary report contains a snapshot of statistics in all BAM files present in this sample. This file contains one row for each input BAM file in the sample. The summary file is displayed in the LifeScope<sup>™</sup> Software UI. See "Summary file" on page 149.

<sampleName1>/BAMfilename-summary.tbl

Following directories and reports contain the cumulative statistics from all BAM files that belong to this sample. The Misc folder is not displayed in UI.

```
<sampleName1>/*.cht
<sampleName1>/Misc/*.csv, *.wig, *.txt
```

Following statistics are generated per BAM file in the mapping directory. These reports are not displayed in the UI.

```
<sampleName1>/<BAMfilename>/.*cht
<sampleName1>/<BAMfilename>/Misc/*.csv, *.txt, *.wig
```

In a sample, some of the BAM files possibly represent unhealthy DNA or RNA, causing the cumulative statistics to look poor. The summary tbl file is the unified location for examining the quality of data of all BAM files in the sample.

If the data for a particular BAM file is not as expected, look at the directory-level reports for that BAM file, for details.

# Mapping statistics output files

This section describes the mapping statistics files generated by the BAMStats module. When statistics reports are separated by tag type, the report's file name includes the tag in the file name. The tags used in file names are:

Library type	Tag
Fragment	F3
Mate-pair	F3
	R3
Paired-end	F3
	F5-P2

Table 18 Tags in mapping statistics output file names

The output files generated by BAMStats include the name of the BAM file. The file name pattern is:

BAMFileName-fileID-barcodeID.StatisticsReportName.tag.extension

This string is referred to by *prefix* in Table 19, the mapping statistics output table. In the description of mapping statistics output file names in Table 19, the string *tag* is used to refer to the tags in Table 18.

 Table 19
 Mapping statistics output files

Report	Description, axis information, filename	
Alignment Len	igth Distribution	
	A bar plot giving the distribution of alignment lengths found in various tags used in alignment. The report is separated by tag type to provide visibility into the accuracy of individual tags.	
	Only the primary alignment for each bead is considered in calculating the distribution.	
	Y axis: Frequency X axis: Alignment length (from 0 to the maximum read length)	
	Output file name: <i>prefix</i> .Alignment.Length.Distribution. <i>tag</i> .cht	
Alignment Len	gth Distribution for Unique Alignments	
The same report as the Alignment Length Distribution, except that only tags that have a primary alignment are considered in calculating the distribution.           By default BAM files contains primary alignments only. In this case, the reports           AlignmentLengthDistribution and AlignmentLengthDistribution for Unique Alignments are identical. However, users can also print all alignments or secondary alignments in BAM file. In this case, the two distributions are different.           Note: The title Unique Alignments is interpreted to mean primary alignments, for this report.           Y axis: Frequency           X axis: Alignment length (from 0 to the maximum read length)           Output file name: prefixAlignment.Length.Distribution.Unique.tag.cht           Base Mismatch Distribution           A bar plot giving the distribution of total number of mismatches found in various tags used in alignment. The bins are summed over all alignment lengths.           Only the primary alignment for each bead is considered in calculating the distribution.           Y axis: Frequency           X axis: Number of mismatches (from 0 to the maximum mismatches allowed)           Output file name: prefix.BaseMismatch.Distribution.tag.cht           Base Mismatch Distribution for Unique Alignments           Waxis: Frequency           X axis: Number of mismatches (from 0 to the maximum mismatches allowed)           Output file name: prefix.BaseMismatch.Distribution, except that only tags that have a primary alignment are considered in calculating the distribution.	Report	Description, axis information, filename
---	---------------	--
By default BAM files contain primary alignments only. In this case, the reports AlignmentLengthDistribution and Alignments or secondary alignments in BAM file. In this case, the two distributions are different.           Note: The title Unique Alignments is interpreted to mean primary alignments, for this report.           Y axis: Frequency           X axis: Alignment length (from 0 to the maximum read length)           Output file name: prefix.Alignment.Length.Distribution.Unique.tag.cht           Base Mismatch Distribution           A bar plot giving the distribution of total number of mismatches found in various tags used in alignment. The bins are summed over all alignment lengths.           Only the primary alignment for each bead is considered in calculating the distribution.           Y axis: Frequency           X axis: Number of mismatches (from 0 to the maximum mismatches allowed)           Output file name: prefix.BaseMismatch.Distribution.tag.cht           Base Mismatch Distribution for Unique Alignments           The same report as the Base Mismatch Distribution.           Note: The title Unique Alignments is interpreted to mean primary alignments, for this report.           Y axis: Frequency           X axis: Number of mismatches (from 0 to the maximum mismatches allowed)           Output file name: prefix.BaseMismatch.Distribution.Unique.tag.cht           Distribution of Alignment for 0 Mismatches in Tags           A report providing a simultaneous picture of the alignment length and number of mismatches distributio		The same report as the Alignment Length Distribution, except that only tags that have a primary alignment are considered in calculating the distribution.
Note: The title Unique Alignments is interpreted to mean primary alignments, for this report.           Y axis: Frequency           X axis: Alignment length (from 0 to the maximum read length)           Output file name: prefix Alignment.Length.Distribution.Unique.tag.cht           Base Mismatch Distribution           A bar plot giving the distribution of total number of mismatches found in various tags used in alignment. The bins are summed over all alignment lengths.           Only the primary alignment for each bead is considered in calculating the distribution.           Y axis: Frequency           X axis: Number of mismatches (from 0 to the maximum mismatches allowed)           Output file name: prefix.BaseMismatch.Distribution, except that only tags that have a primary alignment are considered in calculating the distribution.           Note: The title Unique Alignments is interpreted to mean primary alignments, for this report.           Y axis: Frequency           X axis: Number of mismatches (from 0 to the maximum mismatches allowed)           Output file name: prefix.BaseMismatch.Distribution.Unique.tag.cht           Note: The title Unique Alignments is interpreted to mean primary alignments, for this report.           Y axis: Frequency           X axis: Number of mismatches from 0 to the maximum mismatches allowed)           Output file name: prefix.BaseMismatch.Distribution.Unique.tag.cht           Parisitibution of Alignment Length and Number of Mismatches in Tags           A report providing		By default BAM files contain primary alignments only. In this case, the reports AlignmentLengthDistribution and AlignmentLengthDistribution for Unique Alignments are identical. However, users can also print all alignments or secondary alignments in BAM file. In this case, the two distributions are different.
Y axis: Frequency           X axis: Alignment length [from 0 to the maximum read length]           Output file name: prefix.Alignment.Length.Distribution.Unique.tag.cht           Base Mismatch Distribution           A bar plot giving the distribution of total number of mismatches found in various tags used in alignment. The bins are summed over all alignment lengths.           Only the primary alignment for each bead is considered in calculating the distribution.           Y axis: Frequency           X axis: Number of mismatches [from 0 to the maximum mismatches allowed]           Output file name: prefix.BaseMismatch.Distribution.tag.cht           Base Mismatch Distribution for Unique Alignments           The same report as the Base Mismatch Distribution.           Note: The title Unique Alignments is interpreted to mean primary alignments, for this report.           Y axis: Number of mismatches (from 0 to the maximum mismatches allowed)           Output file name: prefix.BaseMismatch.Distribution.Unique.tag.cht           Distribution of Alignment Length and Number of Mismatches in Tags           A report providing a simultaneous picture of the alignment length and number of mismatches distribution in various tags used in alignment. Only primary alignments for each tag are considered in calculating the distribution. The bins range from 0 to the maximum mismatches allowed]           Note: This report is located in the xisc folder, and is not displayed in the UI.           Y axis: Number of mismatches (from 0 to the maximum mismatches allowed] <td></td> <td>Note: The title Unique Alignments is interpreted to mean primary alignments, for this report.</td>		Note: The title Unique Alignments is interpreted to mean primary alignments, for this report.
Output file name: prefix.Alignment.Length.Distribution.Unique.tag.cht           Base Mismatch Distribution           A bar plot giving the distribution of total number of mismatches found in various tags used in alignment. The bins are summed over all alignment lengths.           Only the primary alignment for each bead is considered in calculating the distribution.           Y axis: Frequency           X axis: Number of mismatches (from 0 to the maximum mismatches allowed)           Output file name: prefix.BaseMismatch.Distribution.tag.cht           Base Mismatch Distribution for Unique Alignments           The same report as the Base Mismatch Distribution.           Note: The title Unique Alignments is interpreted to mean primary alignments, for this report.           Y axis: Frequency           X axis: Number of mismatches (from 0 to the maximum mismatches allowed)           Output file name: prefix.BaseMismatch.Distribution.Unique.tag.cht           Note: The title Unique Alignment of thismatches in Tags           A report providing a simultaneous picture of the alignment length and number of mismatches distribution in various tags used in alignment. Only primary alignments for each tag are considered in calculating thi distribution. The bins report is located in the Misc folder, and is not displayed in the UI.           Y axis: Alignment length (from 0 to the maximum mesmatches allowed)         Output file name: prefix.AlignmentLength.Mismatch.tag.csv           Distribution         Output file name: prefix.AlignmentLength.Mismatch.tag.csv		Y axis: Frequency X axis: Alignment length (from 0 to the maximum read length)
Base Mismatch Distribution           A bar plot giving the distribution of total number of mismatches found in various tags used in alignment. The bins are summed over all alignment lengths.           Only the primary alignment for each bead is considered in calculating the distribution.           Y axis: Frequency           X axis: Number of mismatches (from 0 to the maximum mismatches allowed)           Output file name: prefix.BaseMismatch.Distribution.tag.cht           Base Mismatch Distribution for Unique Alignments           The same report as the Base Mismatch Distribution, except that only tags that have a primary alignment are considered in calculating the distribution.           Note: The title Unique Alignments is interpreted to mean primary alignments, for this report.           Y axis: Number of mismatches (from 0 to the maximum mismatches allowed)           Output file name: prefix.BaseMismatch.Distribution.Unique.tag.cht           Distribution of Alignment Length and Number of Mismatches in Tags           A report providing a simultaneous picture of the alignment length and number of mismatches distribution in various tags used in alignment. Only primary alignments for each tag are considered in calculating this distribution. The bins range from 0 to the maximum mismatches allowed]           Output file name: prefix.AlignmentLength.Mismatch.tag.csv           Base QV Distribution           Quiptur file name: prefix.AlignmentLength.Mismatch.tag.csv           Base QV Distribution           Quiptu file name: prefix.AlignmentLength.Mismatch.tag.		Output file name: <i>prefix</i> .Alignment.Length.Distribution.Unique. <i>tag</i> .cht
A bar plot giving the distribution of total number of mismatches found in various tags used in alignment. The bins are summed over all alignment lengths.         Only the primary alignment for each bead is considered in calculating the distribution.         Y axis: Frequency         X axis: Number of mismatches (from 0 to the maximum mismatches allowed)         Output file name: prefix.BaseMismatch.Distribution.tag.cht         Base Mismatch Distribution for Unique Alignments         The same report as the Base Mismatch Distribution, except that only tags that have a primary alignment are considered in calculating the distribution.         Note: The title Unique Alignments is interpreted to mean primary alignments, for this report.         Y axis: Frequency         X axis: Number of mismatches (from 0 to the maximum mismatches allowed)         Output file name: prefix.BaseMismatch.Distribution.Unique.tag.cht         Distribution of Alignment Length and Number of Mismatches in Tags         A report providing a simultaneous picture of the alignment length and number of mismatches distribution: in various tags used in alignment. Only primary alignments for each tag are considered in calculating this distribution. The bins range from 0 to the maximum mismatches allowed)         Note: This report is located in the Misc folder, and is not displayed in the UI.         Y axis: Alignment length (from 0 to the maximum mismatches allowed)         Output file name: prefix.AlignmentLength.Mismatch.tag.csv         Base OV Distribution         A bar plot providin	Base Mismat	ch Distribution
Only the primary alignment for each bead is considered in calculating the distribution.           Y axis: Frequency           X axis: Number of mismatches [from 0 to the maximum mismatches allowed]           Output file name: prefix.BaseMismatch.Distribution.tag.cht           Base Mismatch           Distribution for Unique Alignments           The same report as the Base Mismatch Distribution, except that only tags that have a primary alignment are considered in calculating the distribution.           Note: The title Unique Alignments is interpreted to mean primary alignments, for this report.           Y axis: Frequency           X axis: Number of mismatches [from 0 to the maximum mismatches allowed]           Output file name: prefix.BaseMismatch.Distribution.Unique.tag.cht           Distribution of Alignment Length and Number of Mismatches in Tags           A report providing a simultaneous picture of the alignment length and number of mismatches distribution in various tags used in alignment. Only primary alignments for each tag are considered in calculating this distribution. The bins range from 0 to the maximum meal length, and from 0 to the maximum number of mismatches allowed.           Note: This report is located in the Misc folder, and is not displayed in the UI.           Y axis: Alignment length [from 0 to the maximum mismatches allowed]		A bar plot giving the distribution of total number of mismatches found in various tags used in alignment. The bins are summed over all alignment lengths.
Y axis: Frequency X axis: Number of mismatches [from 0 to the maximum mismatches allowed]           Output file name: prefix.BaseMismatch.Distribution.tag.cht           Base Mismatch Distribution for Unique Alignments           The same report as the Base Mismatch Distribution, except that only tags that have a primary alignment are considered in calculating the distribution. Note: The title Unique Alignments is interpreted to mean primary alignments, for this report.           Y axis: Number of mismatches [from 0 to the maximum mismatches allowed]           Output file name: prefix.BaseMismatch.Distribution.Unique.tag.cht           Distribution of Alignment Length and Number of Mismatches in Tags           A report providing a simultaneous picture of the alignment length and number of mismatches distribution in various tags used in alignment. Only primary alignments for each tag are considered in calculating this distribution. The bins range from 0 to the maximum mead length, and from 0 to the maximum number of mismatches allowed.           Note: This report is located in the Misc folder, and is not displayed in the UI.           Y axis: Number of mismatches (from 0 to the maximum mismatches allowed)           Output file name: prefix.AlignmentLength.Mismatch.tag.csv           Base QV Distribution           A bar plot providing a distribution of base quality values generated using the reference assisted error correction/base couversion algorithm. The base QV distributions are separated for each tag as quality of individual tags could be very different.           Y axis: Frequency X axis: Base QV (from 0 to the maximum QV)         Output file name:		Only the primary alignment for each bead is considered in calculating the distribution.
Output file name: prefix.BaseMismatch.Distribution.tag.cht           Base Mismatch Distribution for Unique Alignments           The same report as the Base Mismatch Distribution, except that only tags that have a primary alignment are considered in calculating the distribution.           Note: The title Unique Alignments is interpreted to mean primary alignments, for this report.           Y axis: Frequency           X axis: Number of mismatches (from 0 to the maximum mismatches allowed)           Output file name: prefix.BaseMismatch.Distribution.Unique.tag.cht           Distribution of Alignment Length and Number of Mismatches in Tags           A report providing a simultaneous picture of the alignment length and number of mismatches distribution in various tags used in alignment. Only primary alignments for each tag are considered in calculating this distribution. The bins range from 0 to the maximum read length, and from 0 to the maximum number of mismatches allowed.           Note: This report is located in the Misc folder, and is not displayed in the UI.           Y axis: Number of mismatches (from 0 to the maximum mismatches allowed)           Output file name: prefix.AlignmentLength.Mismatch.tag.csv           Base QV Distribution           A bar plot providing a distribution of base quality values generated using the reference assisted error correction/base could be very different.           Y axis: Base QV (from 0 to the maximum QV)         Output file name: prefix.BaseQV.tag.cht		Y axis: Frequency X axis: Number of mismatches (from 0 to the maximum mismatches allowed)
Base Mismatch Distribution for Unique Alignments           The same report as the Base Mismatch Distribution, except that only tags that have a primary alignment are considered in calculating the distribution.           Note: The title Unique Alignments is interpreted to mean primary alignments, for this report.           Y axis: Frequency           X axis: Number of mismatches (from 0 to the maximum mismatches allowed)           Output file name: prefix.BaseMismatch.Distribution.Unique.tag.cht           Distribution of Alignment Length and Number of Mismatches in Tags           A report providing a simultaneous picture of the alignment length and number of mismatches distribution in various tags used in alignment. Only primary alignments for each tag are considered in calculating this distribution. The bins range from 0 to the maximum read length, and from 0 to the maximum number of mismatches allowed.           Note: This report is located in the Misc folder, and is not displayed in the UI.           Y axis: Alignment length (from 0 to the max read length)           X axis: Number of mismatches (from 0 to the maximum mismatches allowed)           Output file name: prefix.AlignmentLength.Mismatch.tag.csv           Base QV Distribution           A bar plot providing a distribution of base quality values generated using the reference assisted error correction/base conversion algorithm. The base QV distributions are separated for each tag as quality of individual tags could be very different.           Y axis: Frequency         X axis: Base QV (from 0 to the maximum QV)         Y axis: Base QV (from 0 to the maximum		Output file name: <i>prefix</i> .BaseMismatch.Distribution. <i>tag</i> .cht
The same report as the Base Mismatch Distribution, except that only tags that have a primary alignment are considered in calculating the distribution.           Note:         The title Unique Alignments is interpreted to mean primary alignments, for this report.           Y axis:         Frequency           X axis:         Number of mismatches (from 0 to the maximum mismatches allowed)           Output file name:         prefix.BaseMismatch.Distribution.Unique.tag.cht           Distribution of Alignment Length and Number of Mismatches in Tags         A report providing a simultaneous picture of the alignment length and number of mismatches distribution in various tags used in alignment. Only primary alignments for each tag are considered in calculating this distribution. The bins range from 0 to the maximum read length, and from 0 to the maximum number of mismatches allowed.           Note:         This report is located in the Misc folder, and is not displayed in the UI.           Y axis:         Number of mismatches (from 0 to the max read length)           X axis:         Number of mismatches (from 0 to the maximum mismatches allowed)           Output file name:         prefix.AlignmentLength.Mismatch.tag.csv           Base QV Distribution         A bar plot providing a distribution of base quality values generated using the reference assisted error correction/base conversion algorithm. The base QV distributions are separated for each tag as quality of individual tags could be very different.           Y axis: Base QV (from 0 to the maximum QV)         Output file name: prefix.BaseQV.tag.cht  <	Base Mismat	ch Distribution for Unique Alignments
Note:         The title Unique Alignments is interpreted to mean primary alignments, for this report.           Y axis:         Frequency           X axis:         Number of mismatches (from 0 to the maximum mismatches allowed)           Output file name:         prefix.BaseMismatch.Distribution.Unique.tag.cht           Distribution of Alignment Length and Number of Mismatches in Tags         A report providing a simultaneous picture of the alignment length and number of mismatches distribution in various tags used in alignment. Only primary alignments for each tag are considered in calculating this distribution. The bins range from 0 to the maximum read length, and from 0 to the maximum number of mismatches allowed.           Note:         This report is located in the Misc folder, and is not displayed in the UI.           Y axis:         Number of mismatches (from 0 to the maximum mismatches allowed)           Output file name:         prefix.AlignmentLength.Mismatch.tag.csv           Base OV Distribution         A bar plot providing a distribution of base quality values generated using the reference assisted error correction/base conversion algorithm. The base QV distributions are separated for each tag as quality of individual tags could be very different.           Y axis:         Frequency           X axis:         Base QV (from 0 to the maximum QV)           Output file name:         prefix.BaseQV.tag.cht		The same report as the Base Mismatch Distribution, except that only tags that have a primary alignment are considered in calculating the distribution.
Y axis: Frequency X axis: Number of mismatches (from 0 to the maximum mismatches allowed)           Output file name: prefix.BaseMismatch.Distribution.Unique.tag.cht           Distribution of Alignment Length and Number of Mismatches in Tags           A report providing a simultaneous picture of the alignment length and number of mismatches distribution in various tags used in alignment. Only primary alignments for each tag are considered in calculating this distribution. The bins range from 0 to the maximum read length, and from 0 to the maximum number of mismatches allowed.           Note: This report is located in the Misc folder, and is not displayed in the UI.           Y axis: Alignment length (from 0 to the max read length) X axis: Number of mismatches (from 0 to the maximum mismatches allowed)           Output file name: prefix.AlignmentLength.Mismatch.tag.csv           Base QV Distribution           A bar plot providing a distribution of base quality values generated using the reference assisted error correction/base conversion algorithm. The base QV distributions are separated for each tag as quality of individual tags could be very different.           Y axis: Frequency X axis: Base QV (from 0 to the maximum QV)         Output file name: prefix.BaseQV.tag.cht		Note: The title Unique Alignments is interpreted to mean primary alignments, for this report.
Output file name: prefix.BaseMismatch.Distribution.Unique.tag.cht           Distribution of Alignment Length and Number of Mismatches in Tags           A report providing a simultaneous picture of the alignment length and number of mismatches distribution. In various tags used in alignment. Only primary alignments for each tag are considered in calculating this distribution. The bins range from 0 to the maximum read length, and from 0 to the maximum number of mismatches allowed.           Note: This report is located in the Misc folder, and is not displayed in the UI.           Y axis: Alignment length (from 0 to the max read length)           X axis: Number of mismatches (from 0 to the maximum mismatches allowed)           Output file name: prefix.AlignmentLength.Mismatch.tag.csv           Base QV Distribution           A bar plot providing a distribution of base quality values generated using the reference assisted error correction/base conversion algorithm. The base QV distributions are separated for each tag as quality of individual tags could be very different.           Y axis: Frequency         X axis: Base QV (from 0 to the maximum QV)           Output file name: prefix.BaseQV.tag.cht		Y axis: Frequency X axis: Number of mismatches (from 0 to the maximum mismatches allowed)
Distribution of Alignment Length and Number of Mismatches in Tags         A report providing a simultaneous picture of the alignment length and number of mismatches distribution in various tags used in alignment. Only primary alignments for each tag are considered in calculating this distribution. The bins range from 0 to the maximum read length, and from 0 to the maximum number of mismatches allowed.         Note: This report is located in the Misc folder, and is not displayed in the UI.         Y axis: Alignment length (from 0 to the max read length)         X axis: Number of mismatches (from 0 to the maximum mismatches allowed)         Output file name: prefix.AlignmentLength.Mismatch.tag.csv         Base QV Distribution         A bar plot providing a distribution of base quality values generated using the reference assisted error correction/base conversion algorithm. The base QV distributions are separated for each tag as quality of individual tags could be very different.         Y axis: Frequency       X axis: Base QV (from 0 to the maximum QV)         Output file name: prefix.BaseQV.tag.cht		Output file name: <i>prefix</i> .BaseMismatch.Distribution.Unique. <i>tag</i> .cht
A report providing a simultaneous picture of the alignment length and number of mismatches distribution in various tags used in alignment. Only primary alignments for each tag are considered in calculating this distribution. The bins range from 0 to the maximum read length, and from 0 to the maximum number of mismatches allowed.         Note: This report is located in the Misc folder, and is not displayed in the UI.         Y axis: Alignment length (from 0 to the max read length)         X axis: Number of mismatches (from 0 to the maximum mismatches allowed)         Output file name: prefix.AlignmentLength.Mismatch.tag.csv         Hase QV Distribution         A bar plot providing a distribution of base quality values generated using the reference assisted error correction/base conversion algorithm. The base QV distributions are separated for each tag as quality of individual tags could be very different.         Y axis: Frequency       X axis: Base QV (from 0 to the maximum QV)         Output file name: prefix.BaseQV.tag.cht       Output file name: prefix.BaseQV.tag.cht	istribution o	of Alignment Length and Number of Mismatches in Tags
Note: This report is located in the Misc folder, and is not displayed in the UI.         Y axis: Alignment length (from 0 to the max read length)         X axis: Number of mismatches (from 0 to the maximum mismatches allowed)         Output file name: prefix.AlignmentLength.Mismatch.tag.csv         Base QV Distribution         A bar plot providing a distribution of base quality values generated using the reference assisted error correction/base conversion algorithm. The base QV distributions are separated for each tag as quality of individual tags could be very different.         Y axis: Frequency         X axis: Base QV (from 0 to the maximum QV)         Output file name: prefix.BaseQV.tag.cht		A report providing a simultaneous picture of the alignment length and number of mismatches distributions in various tags used in alignment. Only primary alignments for each tag are considered in calculating this distribution. The bins range from 0 to the maximum read length, and from 0 to the maximum number of mismatches allowed.
Y axis: Alignment length (from 0 to the max read length)         X axis: Number of mismatches (from 0 to the maximum mismatches allowed)         Output file name: prefix.AlignmentLength.Mismatch.tag.csv         Base QV Distribution         A bar plot providing a distribution of base quality values generated using the reference assisted error correction/base conversion algorithm. The base QV distributions are separated for each tag as quality of individual tags could be very different.         Y axis: Frequency         X axis: Base QV (from 0 to the maximum QV)         Output file name: prefix.BaseQV.tag.cht		Note: This report is located in the Misc folder, and is not displayed in the UI.
Output file name: prefix.AlignmentLength.Mismatch.tag.csv         Base QV Distribution         A bar plot providing a distribution of base quality values generated using the reference assisted error correction/base conversion algorithm. The base QV distributions are separated for each tag as quality of individual tags could be very different.         Y axis: Frequency         X axis: Base QV (from 0 to the maximum QV)         Output file name: prefix.BaseQV.tag.cht		Y axis: Alignment length (from 0 to the max read length) X axis: Number of mismatches (from 0 to the maximum mismatches allowed)
Base QV Distribution         A bar plot providing a distribution of base quality values generated using the reference assisted error correction/base conversion algorithm. The base QV distributions are separated for each tag as quality of individual tags could be very different.         Y axis: Frequency         X axis: Base QV (from 0 to the maximum QV)         Output file name: prefix.BaseQV.tag.cht		Output file name: <i>prefix</i> .AlignmentLength.Mismatch. <i>tag</i> .csv
A bar plot providing a distribution of base quality values generated using the reference assisted error correction/base conversion algorithm. The base QV distributions are separated for each tag as quality of individual tags could be very different. Y axis: Frequency X axis: Base QV (from 0 to the maximum QV) Output file name: <i>prefix</i> .BaseQV. <i>tag</i> .cht	Base QV Dist	ribution
Y axis: Frequency X axis: Base QV (from 0 to the maximum QV) Output file name: <i>prefix</i> .BaseQV. <i>tag</i> .cht		A bar plot providing a distribution of base quality values generated using the reference assisted error correction/base conversion algorithm. The base QV distributions are separated for each tag as quality of individual tags could be very different.
Output file name: <i>prefix</i> .BaseQV. <i>tag</i> .cht		Y axis: Frequency X axis: Base QV (from 0 to the maximum QV)
		Output file name: <i>prefix</i> .BaseQV. <i>tag</i> .cht

Table 19	Mapping	statistics	output	files	(continued)
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## Table 19 Mapping statistics output files (continued)

Report	Description, axis information, filename
	A report providing a distribution of base quality values by individual base positions. This report identifies if certain base positions (particularly towards the end of the read) have poor base quality values. All the tags (F3/R3/F5-P2) used in the alignment are combined into a single bin.
	Y axis: Base QV X axis: Base position (from 0 to read length)
	Output file name: <i>prefix</i> .BaseQV.by.Position.csv
Distribution of	f Mismatches by Base QV
	A 3D surface plot providing a distribution of errors (mismatches to reference) by base quality values bins. This report measures whether the base QVs generated are well calibrated to the probability of error in that particular base position.
	Y axis: Frequency (error rate) X axis: Base QV (from 0 to maximum QV)
	Output file name: <i>prefix</i> .Mismatches.By.BaseQV. <i>tag</i> .cht
Distribution of	f Mismatches by Position
	A bar plot providing a distribution of errors (mismatches to reference) by position within the read. Only the primary alignments for each bead are used to generate this distribution.
	Y axis: Frequency X axis: Position (from 0 to maximum read length)
	Output file name: <i>prefix</i> .Mismatches.By.Position.BaseSpace. <i>tag</i> .cht
Distribution L	y Mapping QVs by Tag
	A bar plot providing a distribution of mapping quality values for individual tags (F3/R3/F5-P2). Only the primary alignment for each bead is used in calculating this distribution.
	Y axis: Frequency X axis: Mapping QV (from 0 to maximum mapping QV)
	Output file name: <i>prefix</i> .MappingQV. <i>tag</i> .cht
Distribution b	y Pairing QVs
	A bar plot providing a distribution of pairing quality values. Only the primary alignment for each bead is used in calculating this distribution.
	Y axis: Frequency X axis: Pairing QV (from 0 to maximum pairing QV)
	Output file name: <i>prefix</i> .PairingQVs.cht
Coverage Re	oort
	A line plot providing a distribution of coverage obtained after mapping/pairing. Only the primary alignment for each bead is used in this calculation.
	Y axis: Frequency X axis: Coverage (from 0 to number of reads)
	Output file name: <i>prefix</i> .Coverage. <i>tag</i> .cht
Coverage Re	port by Chromosome (Contig) and Base Windows

Report	Description, axis information, filename
	A line plot providing a distribution of coverage within each reference window. The coverage is calculated within each window along a reference chromosome. The window size can theoretically be anywhere from a single base to the contig length. However the calculations of base level coverage are computationally expensive and less interpretable as the window size increases. Only the primary alignment for each bead is used in this calculation.
	Y axis: Frequency X axis: Coverage (from 0 to number of reads)
	Output file name: <i>prefix</i> .Coverage.By.Chromosome.contig <i>nn</i> .cht
Coverage by	Strand
	A line plot providing the distribution of coverage within each reference window separated by reference strand (+/-). Only the primary alignment for each bead is used in this calculation.
	Y axis: Frequency X axis: Coverage (from 0 to number of reads)
	Output file name: <i>prefix</i> .Coverage.By.Strand. <i>tag</i> .cht
Coverage file	is a second s
	Coverage reports in wiggle format. For each genome position, reports the number of reads that cover (map to or span) the position. Because reporting coverage for each position results in very large files, coverage is reported for bins, with each bin spanning a user-defined number of bases. For each bin, the mean coverage of all the positions in that bin is reported. The parameter <code>bamstats.wig.binsize</code> controls the size of the bins in this file.
	By default one coverage file is generated, per chromosome, per strand. If the parameter bamstats.combined.report.both.strands is set to true, then one file per chromosome is generated, combining coverage from both strands into one file per chromosome.
	Each coverage file includes a header as the first line. The header lines follow this pattern:
	track type=wiggle_0 name=< <i>chrname&gt;</i> description=< <i>coverage from positive/negative/both strand&gt;</i> visibility=full color=0,0,255 fixedStep chrom=< <i>chrname&gt;</i> start=< <i>startpos&gt;</i> step=< <i>binsize&gt;</i> span=< <i>binsize&gt;</i>
	Output file name: coverage_chrnn.POS.wig, coverage_chrnn.NEG.wig
Insert Range	Distribution
	A line plot providing the distribution of insert sizes for paired data (PE and LMP library types).
	Y axis: Frequency X axis: Insert Range bins
	Output file name: <i>prefix</i> .Insert.Range.Distribution.cht
Distribution (	of Read Pair Types
	For paired data, this report calculates the distribution of read pair types (AAA, AAB, C**, etc.).
	Y axis: Frequency X axis: Read pair type
	Output file name: <i>prefix</i> .ReadPair.Type.cht
Pairing Stat	stics

## Table 19 Mapping statistics output files (continued)

#### Table 19 Mapping statistics output files (continued)

Report	Description, axis information, filename				
	A pie chart showing the percentages of the following F3, R3 (or F5) combinations, for paired data,:				
	Mapped, Mapped				
	Mapped, Unmapped				
	Unmapped, Mapped				
	Unmapped, Unmapped				
	Mapped, Missing				
	Missing, Mapped				
	Unmapped, Missing				
	Missing, Unmapped				
	Output file name: <i>prefix</i> .Pairing.Stats.cht				
Unique Start	Position				
	A text report with the following statistics about the start position on the genome, based only on primary alignments. The report contains:				

- 1. The number of starting points in uniquely placed tags:
  - Reports the positions in the reference with at least one uniquely placed alignment starting at that position. Unique here does not mean primary.
  - Also reports the percentage of this number within the total number of positions in the reference.
- 2. The average number of uniquely mapped reads per starting point:

Reports the total number of unique alignments divided by the number of starting points in uniquely placed tags.

**3.** An estimated number of starting points for all mapped tags:

Reports the total number of primary alignments divided by the number of starting points in uniquely placed tags.

Output file name: prefix.Unique.Start.Positions.txt

#### Position error files

One or more tab-delimited text files generated in the directory containing the input BAM file. This file is used by the diBayes module for SNP analysis. The file records the frequencies of dicolor mismatches between reads and the reference at different positions in a read.

The following position error files are generated, based on library type:

- Fragment runs An F3 position error file is created.
- Mate-pair runs Both F3 and R3 position error files are created.
- Paired-end runs Both F3 and F5 position error files are created.

Output file names: *BAMfilename\_*positionErrors\_*tag\_*basespace.txt, *BAMfilename\_*positionErrors\_*tag\_*colorspace.txt

Probe error files

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Table 19	Mapping	statistics	output	files	(continued)
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Report	Description, axis information, filename				
	One or more tab-delimited text files generated in the directory containing the input BAM file. This file is used by the diBayes module for SNP analysis. The probe error file records the frequencies of dicolor mismatches between the reads and the reference as a function of different 6-mer probes.				
	The following probe error files are generated, based on library type:				
	• Fragment runs – An F3 probe error file is created.				
	• Mate-pair runs – Both F3 and R3 probe error files are created.				
	• Paired-end runs – Both F3 and F5 probe error files are created.				
	Output file names: <i>BAMfilename_</i> probeErrors_ <i>tag_</i> colorspace.txt				

## Mapping statistics example output

This section provides example output of some mapping statistics output files.

## Summary file

The summary file provides statistics for each BAM file in the sample. This list describes labels used in the summary file:

- **NumUnFilteredBeads** The total number of beads, before any filtering on the instrument. This value is also the fragment count in the input XSQ file.
- NumFilteredBeads The number of beads that pass filtering on the instrument.
- NumMapped The number of reads with primary alignment.
- % filtered that mapped The number of primary reads divided by the number of beads passing instrument filtering (NumMapped / NumFilteredBeads).
- % total that mapped The number of primary reads divided by the number of total beads (NumMapped / NumUnFilteredBeads).

The number of beads filtered out in the instrument is found by subtracting the number of beads that pass filtering from the total number of beads:

NumUnFilteredBeads - NumFilteredBeads

The following are example contents for a summary file:

#title: BAMStats Summary

```
BamFileName,IsColorInBam,IsBaseInXSQ,IsECC,LibraryType,ReadLeng
th,PredictedInsertSize,NumFilteredBeads,NumUnFilteredBeads,Tag1
-NumMapped,Tag1- % total Mapped,Tag1- % filtered mapped,Tag2-
NumMapped,Tag2- % total Mapped,Tag2- % filtered mapped,(Tag1-
AlignmentLength;Min;Max;Avg;Median;StdDev),(Tag1-
NumMismatches;Min;Max;Avg;Median;StdDev),(Tag1-
MappingQV;Min;Max;Avg;Median;StdDev),(Tag1-
BaseQV;Min;Max;Avg;Median;StdDev),(Tag2-
AlignmentLength;Min;Max;Avg;Median;StdDev),(Tag2-
NumMismatches;Min;Max;Avg;Median;StdDev),(Tag2-
BaseQV;Min;Max;Avg;Median;StdDev),(Tag2-
MappingQV;Min;Max;Avg;Median;StdDev),(Tag2-
StdDev),Min;Max;Avg;Median;StdDev),(Tag2-
StdDev),Min;Max;Avg;Median;StdDev),(Tag2-
StdDev)
```

```
hg19_Simulated_GR_CS_BC16_PE50X25_320Mil-1-Idx_BC15-
15.bam,Y,N,N,PE,50x25,200,18000000,18000000,16785960,93.26,93.2
6,14720544,81.78,81.78,(F3;25;50;46.19;50;6.20),(F3;0;25;1.64;1
;2.03),(F3;0;60;34.37;43;22.61),(F3;1;41;34.66;41;11.02),(F5-
P2;25;25;25.00;25;0.00),(F5-P2;0;16;0.61;0;0.91),(F5-
P2;0;60;33.35;41;23.21),(F5-
P2;1;41;36.15;41;9.56),(0;9921;0.34;0;7.11)
```

## **Unique Start Position**

The following is an example of the output of a Unique Start Position report:

```
#
Starting Points within Placed Tags
Number of Starting Points in Uniquely placed tag 109,068,047
(2.005740% of reference)
Average Number of Uniquely Mapped reads per Start Point
1.992770
Estimate number of starting point for all mapped tags
152,843,627 (2.810765% of reference)
```

## Coverage files

Example contents for the file coverage\_chr1\_positive.wig are:

```
browser position chr1:1-20000000
browser hide all
browser pack refGene encodeRegions
# minimumMapq=25, minimumCoverage=1,
alignmentFilteringMode=PRIMARY, filterOrpha-nedMates=false,
track name="BAM Coverage positive strand" description="BAM
Coverage positive strand" visibility="full color 0,0,255"
priority=10 yLineMark=0 type=wiggle 0 yLineOnOff=on
variableStep chrom=chr1 span=1
336171
336181
336191
336201
336211
336221
336231
336241
336251
336261
336271
. . .
```

Example contents for the file coverage\_chr1\_negative.wig are:

browser position chr1:1-20000000

browser hide all browser pack refGene encodeRegions # minimumMapg=25, minimumCoverage=1, alignmentFilteringMode=PRIMARY, filterOrpha-nedMates=false, track name="BAM Coverage negative strand" description="BAM Coverage negative strand" visibility="full color 0,0,255" priority=10 yLineMark=0 type=wiggle\_0 yLineOnOff=on variableStep chrom=chr1 span=1 57432 57442 57452 57463 57473 57483 57493 57503 57513 57523 57533

•••

#### Position error file

The following is a truncated example of position error file contents:

```
# Generated by: BAM Stats
Plugin(20110415_F12_KF_FC1_1x50frag_AN_02-1-1.bam-t f3,r3 -o /
data/whist/Lifescope_Results/projects/cafe/pHRECC88020/a1/
outputs/bamstats -r /share/lifescope/Reference Data/2.0/
lifetech/hg18/reference/human_hg18.fa ) (Base space)
# Version: 1.0
# Date: May 2, 2011
#? Total read positions = 201701987
#? Missing base calls = 34330
#? Unresolved reference positions = 4
##position_[refCall][readCall] nErrors nReadOccurrences Erro
r frequency
                         0.0044
1 AC
      93033 1272759
1 AG
      25231 1255139
                         0.0059
      184621 37994592
1 AT
                         0.0049
1_CA
      66929 37196998
                         0.0018
```

#### Probe error files

The following is an example of probe error file contents:

```
# Generated by: CountErrorsByPrefix(-t,F3,-d,-1,-k,6,--
mask1=01111,--maskn=11111,--tmfile=/share/apps/corona-r33455M/
etc/analysis/6mers.Tm,/data/results/RegressionDriver/
CaseManager/results/latestBuild/2011-02-19_101137_r3
```

	3455M3.5-trunk_sanityTest/mutationST/case0009/						
	F3_alignment.v2.gff)						
	# Versic	on: 1.0.	29869				
	<pre># Date:</pre>	02/19/2	2011 06:	41 PM			
	##ref cc	ntext	num errors num		num oc	curences	err frequency
	Tm						
	АААААА	0	16	0.0000	00	-24.04	
	CAAAAA	0	6	0.0000	00	-24.53	
	GAAAAA	0	18	0.0000	00	-25.87	
	TAAAAA	0	0	0.0000	00	-37.81	
	ACAAAA	0	0	0.0000	00	-24.77	
Run BAMStats standalone	This section describes requirements to generate mapping statistics outside of the context of the mapping module. These requirements are:						
	• <b>Directory structure</b> – The input directory for BAMStats module must mimic the output directory structure of mapping:						
	Input BAMStat dir/						
	<samplenamel>/* ham. * hai</samplenamel>						
	<samplename2>/*.bam, *.bai</samplename2>						
	• Index	files – Fo	or every B	AM file_th	ere must h	e a correspor	nding BAL-formatted
	index	ed file. B	AI files car	i be genera	ted using t	he following	; command:
	samtools index <i><bamfile></bamfile></i>						
	This command generates index sorted alignment for fast random access. The index file < bamfile > . bai is created. For details, please refer to the samtools site:						

http://samtools.sourceforge.net/samtools.shtml

• Write permission – The user must also have write permission for the input directory, because the BAMStats module generates position and probe error files in that directory.

## FAQ – Mapping

## How does the local alignment approach affect mapping?

Using the local alignment approach removes the constraint of a whole-read alignment, and mapping rate is significantly increased. It is not uncommon for a poor dataset with a 30% mapping rate at 50\_6, to reach more than a 60% mapping rate with the mapping algorithm used in LifeScope<sup>™</sup> Software.

While the majority of alignments are full length, some alignments can vary in length. If a read has many errors towards the end, the final alignment will not include these positions if a shorter alignment receives a better overall score.

## 2

1

How do l increase mapping rate?

Mapping rate increases with each additional round of mapping. However, the gain in rate comes at the cost of increased runtime and disk requirement. Testing has found sets of mapping schemes that strike a good balance between mapping rate and runtime for 25-, 35,- and 50-bp reads. These schemes are shipped as LifeScope<sup>™</sup> Software defaults and should work well for most purposes.

For 50-bp reads, two keys in the mapping.ini file define how many rounds of mapping are run, and what happens in each round. The default values of the keys are shown below, and their meanings are further explained in Table 12 on page 123.

Note: Repetitive schemes are empty by default.

- mapping.scheme.unmapped.50 = 25.2.0,25.2.15
- mapping.scheme.repetitive.50 = 38.3.0,25.2.0

The repetitive scheme does not affect the number of mapped reads, and setting it only increases the chance of finding better alignments for repetitive reads. If your main goal is to improve throughput, first try to add more iterations at the unmapped scheme. For example, add another step of 25.2.25 at the end of the unmapped scheme above.

#### What is a good seed for my application?

Consider the following factors when picking seed parameters:

- Mapping is slower when more mismatches are allowed in the seed. However, allowing more mismatches improves the mapping rate.
- Shorter seeds have higher mapping rates, but also lead to more spurious alignments.
- Color-calls at the beginning of the read are more reliable than those at the end. Therefore, you have the option to anchor the seed near the front of the read. However, applications such as transcriptome sequencing are scenarios where it is an advantage to anchor the seed near the end of the read.

For 50-bp reads, the default setting of 25.2.0 for the first round, followed by 25.2.15 in the second round, delivers good results in most cases. For a single round, 30.3.0 is recommended.

3

#### What reads scenario achieves the best accuracy?

The best performance is achieved by considering 2BE and 4BE reads, their alignment, and reference. Typically, base-space reads generated by ECC have lower mapping throughput. Base-space reads generated from combined 2BE and 4BE calls and no reference have lower accuracy, compared to ECC calls.

#### How do I interpret a mapping quality value?

First of all, it is worth noting that the mapping quality value (QV) is not like the p-value in BLAST. In BLAST, the p-value is used to estimate the likelihood that the aligned sequences are related. In general, reads come from the same source and it is known that the two are related.

The purpose of mapping QV is to estimate the probability that the read originates from the mapped genomic location. The estimate is determined mainly by the difference in significance between the best- and second-best hits.

The numeric interpretation of mapping QV is the same as the base call QV. A mapping QV of 10 means that there is a 90% chance that the alignment is correct. A mapping QV of 20 means that there is a 99% chance that the alignment is correct.

## 6

#### How much RAM do I need to analyze human samples?

For optimal performance, 24 GB of RAM per cluster node is recommended for human samples. If the cluster node has less than 24 GB of available RAM, mapreads can split the genome into smaller segments, and align to each segment sequentially. Splitting the genome into smaller segments is implemented inside mapreads and is completely transparent to the user.

Mapping with less than the recommended amount of RAM slows down performance. If the cluster node has 16 GB of RAM, mapping human samples will be roughly 30% slower compared to a machine with 24 GB of RAM.

## 7

## In the mapping statistics genome coverage calculation report, how are Ns in the reference counted?

Ns in the reference are counted as missing coverage. For example, in the human genome, about 7% of the reference sequence consists of Ns. This means that the coverage calculation will never be reported as higher than 93%. If the frequency distribution says 7.49% of the genome is uncovered, this means that about 0.34% of the non-N reference sequence does not have reads mapped to it, and that also the 7.15% of the genome that consists of N does not have reads mapped to it.

## 8

#### What should I do to ensure balanced allele ratios at heterozygous positions?

The base-translation algorithm, refcor, uses instrument color-space data (ECC or non-ECC) and the reference sequence to produce the most likely base calls, along with a Phred-scale quality value. The degree to which the reference influences the base call is controlled by the refcor.reference.weight parameter. A position is converted to reference only if the refcor.reference.weight setting is higher than the difference between the quality values of a correct color call and of an adjacent erroneous color call that supports reference.

The refcor.reference.weight default setting, 8, is chosen to be low enough to ensure balanced allele ratios, and also be large enough both to distinguish correct from incorrect color calls and to maximize base accuracy.

In the event an incorrect reference base call is made, the disagreement between evidence sources causes that call's QV to be low. Incorrect calls can be filtered out based on QV threshold, and this filtering improves allele ratios.

Another way of filtering low quality base-calls is to adjust the parameter refcor.base.filter.qv, that converts all base calls with a quality value less than refcor.base.filter.qv into Ns. This filtering has the effect of setting QV to zero in the calls with low confidence. To decrease the number of N base calls, reduce this threshold (for example, from the default of 10 to 5).

**Note:** Setting the refcor.reference.weight parameter to zero completely eliminates the reference as a guide for base-translation (completely eliminates reference bias), but also considerably reduces the quality of base-translation.

## FAQ – Pairing

#### How is the uniqueness of a pair determined?

A pair of reads is unique when there is exactly one good AAA pair. With the advent of local alignment, the likelihood of finding only one good pair is decreased. As a result, a different heuristic is used to determine "uniqueness".

Consider an alignment of length *L* with *M* mismatches. If the local score is defined as L+(m-1)M, where m<0 is the mismatch penalty, then the score of each good pair candidate is the sum of its two constituent local scores.

A pair of tags is considered unique if it has only one good pair, or if the score of the best pair is at least X greater than the score of the second best pair. X is specified by the pair.uniqueness.threshold key, and has a default value of 10.0. See Table 15 on page 137 for a description of this and all pairing parameters.

## 2

1

#### What do the 3-letter pair classifications mean?

Table 20 lists a summary of genomic code classifications for mate-pair pairing runs. See Table 21 on page 157 for a summary of genomic code classifications for paired-end runs.

Class name	Strand	Orientation	Insert size	Preference
AAA	Same	R3 to F3	Normal	1
AAB	Same	R3 to F3	< min	2
AAC	Same	R3 to F3	> max	2
ABA	Same	F3 to R3	Normal	2
ABB	Same	F3 to R3	< min	3
ABC	Same	F3 to R3	> max	3
BAA	Different	Outward	Normal	3
BAB	Different	Outward	< min	4
BAC	Different	Outward	> max	4
BBA	Different	Inward	Normal	3
BBB	Different	Inward	< min	4
BBC	Different	Inward	> max	4

 Table 20
 Genomic code classifications, for mate-pair pairing

The first letter determines whether the two tags match the same strand:

- A Match the expected strand.
- **B** Match different strands.

The third letter determines if the distance between two hit positions is within the correct range:

- **A** Within the correct range.
- B The distance is less than the insert size measurement minimum (set with default.minimum.insert.size and denoted by "< min" in the table above).</li>
- C The distance is greater than the insert size measurement maximum (set with default.maximum.insert.size and denoted by "> max" in the table above).

The default minimum insert size is 0. For paired-end and long mate pair workflows, the default maximum insert sizes are 2000 and 20,000, respectively. An insert size measurement automatically determines the actual values using the defaults as the largest interval, and the defaults are used if the insert size measurement fails.

The middle letter has different meanings, depending on whether the first letter is A or B. If the first letter is A, the second letter indicates if the order of the two tags is correct:

- A The order is correct, and R3 is upstream of F3.
- **B** The order is incorrect, and F3 is upstream of R3.

When the two tags are on different strands, the second letter indicates whether the two tags are pointing toward or away from each other

- A Pointing away from each other.
- **B** Pointing toward each other.

In addition, C\*\* classification is assigned to a pair of reads that have one hit each, but are on different chromosomes. If both F3 and R3 tags are present, but only one has a unique hit while the other one is unmapped, the pair is labeled D\*\*. Finally, if one tag from a pair has a unique hit but the other one is missing, as opposed to unmapped, then the pair is classified as E\*\*.

The Preference value indicates likelihood of the variation can occur in nature. Larger values indicate the variation is less likely to occur in nature.

Table 21 shows the classifications for paired-end runs. Summary statistics of these categories are calculated in the optional BAMStats Read Pair Type Statistics Report (.ReadPair.Type.cht).

Class name	Strand	Orientation	Insert size	Preference
AAA	Different	Inward	Normal	1
AAB	Different	Inward	< min	2
AAC	Different	Inward	> max	2
ABA	Different	Outward	Normal	2
ABB	Different	Outward	< min	3
ABC	Different	Outward	> max	3
BAA	Same	F3 to F5	Normal	3
BAB	Same	F3 to F5	< min	4
BAC	Same	F3 to F5	> max	4
BBA	Same	F5 to F3	Normal	3
BBB	Same	F5 to F3	< min	4
BBC	Same	F5 to F3	> max	4

Table 21 Genomic code classifications, for paired-end pairing

# Run an SNPs Analysis

This chapter covers:

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(Optional) Genomic annotation parameters	169
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## **Overview**

The LifeScope<sup>™</sup> Genomic Analysis Software SNPs analysis module uses the diBayes algorithm to find Single Nucleotide Polymorphisms (SNPs). The diBayes package performs independent SNP analysis at each position in the reference, using either a Bayesian or Frequentist algorithm.

The SNPs module is used to call Single Nucleotide Polymorphisms (SNPs) from mapped and processed SOLiD<sup>™</sup> System reads. The module takes the reads, quality values, the reference sequence, and error information of each SOLiD<sup>™</sup> System slide as its input, and calls SNPs.

The SNPs module takes the mapped and processed SOLiD<sup>™</sup> System reads, quality values, the reference sequence, and error information of each SOLiD<sup>™</sup> System slide as its input, and calls SNPs.

The SNPs module creates these results files:

- A list of SNPs.
- A quartile file with coverage as well as color and base quality value distribution.
- (*Optional*) A consensus FASTA file with the same number of bases as the reference sequence.
- (*Optional*) A consensus calls file with a list of all covered positions.
- (Optional) A collection of annotated files.

The consensus calls file and the quartile file are each generated as one file per contig. The list of SNPs and the consensus FASTA file are each generated both as one file per contig and also as a consolidated file for the entire run. Annotated files are generated for the entire run, not for individual contigs. See "SNPs output files" on page 171.

## **SNPs** input files

The diBayes input files include the reference file, the regions of interest file (BED), the BAM file (the output of the mapping-pairing or enrichment modules), and position error and probe error files (the output of the BAMStats mapping statistics module). When you run a standard workflow that includes a SNP analysis, the wiring of these input files is handled automatically by the LifeScope<sup>™</sup> Software command shell. These input files are described in Table 22.

Table 22 diBayes input file description

Input File	Description
Reference genome	The sequence to which the reads are aligned and mapped. The recommended format for the reference file is the FASTA format, for example, chr20.validated.fasta.
	<b>Note:</b> When you run the SNPs module, you must use the same reference that was used for mapping. If different reference files are used, during the SNPs module analysis the coordinates in the BAM file might not correctly represent the coordinates in the reference file. The SNPs module might reject the input BAM files if the reference information in headers of BAM files is not matched to the reference provided.
	IUB codes at positions of known SNPs.
Regions of interest file	The regions of interest file, in BED format, contains a list of defined regions. This file is
(BED file)	resequencing workflows.
	If users want to use a BED file in whole genome resequencing workflows, the file must be sorted based on the start positions for defined regions. The BED files for target resequencing workflows do not have this requirement. Instead the enrichment module within the target resequencing workflow sorts the BED file and passes the sorted BED file to the SNPs module. We recommend users to choose target resequencing workflows for analyses that include a BED file.

Table 22	diBayes	input	file	descriptio	n (continued)
----------	---------	-------	------	------------	---------------

Input File	Description
F3 (R3/F5) position error file	F3 (R3/F5) position error files are tab-delimited text files created during secondary analysis. The files record the frequencies of color/base mismatches between reads and the reference at different positions in a read.
	For fragment runs, only an F3 position error file is needed. Both F3 and R3(F5) position error files are created for mate-pair (paired-end) runs, and both are needed to run diBayes. Position error files are automatically generated during mapping statistics if BAMStats is enabled (recommended).
	If position error files are not available, the SNPs module uses internal default position errors for SNP calling. Using default position error files might slightly affect the module's accuracy.
	Position error files are located in the mapping results folder, and share the same name prefix as the output BAM files. You do not specify their names or location when you run SNPs in a standard workflow. If you run SNPs outside of a workflow and also move the mapping output BAM files, always copy the position and probe error files to the same directory as the BAM files.

## Table 22 diBayes input file description (continued)

Input File	Description			
F3 (R3/F5) probe error file	F3 (R3/F5) probe error files are tab-delimited text files that record the frequencies of dicolor mismatches between the reads and the reference as a function of different 6-mer probes. LifeScope <sup>™</sup> Software calculates the probe error files.			
	For fragment runs, only F3 probe error files are needed. Both F3 and R3(F5) probe error files are created for mate-pair (paired-end) runs, and both are needed to run diBayes on paired data. Different SOLiD <sup>™</sup> system runs of the same sample might generate different F3 (R3/F5) probe error files for each run, because of the random nature of probe errors.			
	Probe error files are automatically generated during mapping statistics if BAMStats is enabled (recommended). However, probe error files are not generated for base space data. If the probe error file is not found, the SNPs module uses internal default probe error files for SNP calling. Using default probe error files might slightly affect the module's accuracy.			
	Probe error files are located in the mapping results folder, and share the same name prefix as the output BAM files. You do not need to specify their name or location when you run SNPs in a standard workflow. If you run SNPs outside of a workflow and also move the mapping output BAM files, always copy the position and probe error files to the same directory as the BAM files.			
BAM file input	The BAM input files are generated by LifeScope <sup>™</sup> Software mapping or enrichment modules. The SNPs module accepts one or more BAM files as input. Each BAM file must have only one read group and only one library type. The SNPs module does not support merged BAM files with multiple read groups or with different library types (within one BAM file).			
	For reads containing Ns, the SNPs module uses all information in the reads except for the missing bases.			
	The SNPs module accepts both color space BAM files (from regular two base color- encoding runs) and base space BAM files (from ECC runs), but does <i>not</i> support combining both color space and base space BAM files in one run. The module determines what algorithms to use automatically based on the input files. Multiple base space BAM files can be processed as combined input using base space algorithms. Multiple color space BAM files are processed as combined input using color space algorithms. If color space BAM files are missing color space information (for instance when users choose not to output color information into the BAM files.), then base space algorithms are used.			
	<b>IMPORTANT!</b> Do not concatenate BAM files from different run types.			
	<b>IMPORTANT!</b> Do not merge the BAM files before running the SNPs module. The module can accept multiple BAM files of different run types as its input.			
	<b>Note:</b> If you move the input BAM file from its default location (not recommended), you must also move its corresponding position and probe error files to the same directory (the BAM file and its corresponding position and probe error files have the same file name prefix).			

## Examples of running a SNPs analysis

Examples in a standard workflow

The following standard workflows provide examples of running the SNPs module:

- genomic.resequencing.frag
- genomic.resequencing.lmp
- genomic.resequencing.pe
- targeted.resequencing.frag
- targeted.resequencing.pe

The following are example shell commands using reads and references from the LifeScope<sup>™</sup> Software repository.

```
# log into the shell
                      lscope.sh shell -u username -w password
                      # cd to the projects repository
                      cd /projects
                      # create a project and open it
                      mk ecoli
                      cd ecoli
                      # create an analysis and open it
                      mk run1
                      cd run1
                      # define the analysis type
                      set workflow genomic.resequencing.pe
                      # define the input data
                      add xsg run0209a_50_PE.xsg
                      add xsq run0209b_50_PE.xsq
                      # specify the reference
                      set reference hg19
                      # optionally change SNPs defaults after this line
                      # list the analysis configuration
                      ls
                      # start the analysis
                      run
                      # list progress information for the run
                      15
                      Dummy file names for XSQ files are shown here.
                      See for Chapter 7, "Run a Standard Workflow Analysis" on page 97 for more
                      information on running standard workflows. Chapter 6, "Run a Command Shell
                      Analysis" on page 71 for information on shell commands and syntax.
As a demo analysis
                      The optional examples download includes an example of how to run the SNPs module
                      by itself, as a single analysis that is not part of a standard workflow. The demo
                      example is not recommended for general use.
```

See Appendix E, "Demo Analyses" on page 507 for information on running the SNPs module as a standalone analysis.

## **SNPs runtime parameters**

Table 23 describes the SNPs module runtime parameters.

In order to change a parameter value in your analysis, use the set param shell command to replace the line # *optionally change SNPs defaults after this line* in the example commands in "Examples in a standard workflow". For instance, to change the call stringency, use this shell command:

set dibayes.call.stringency high /tertiary/dibayes.ini

These workflows use the INI file name dibayes.ini:

- genomic.resequencing.lmp
- genomic.resequencing.pe
- targeted.resequencing.pe

For the workflow targeted.resequencing.frag, use dibayes.targeted.frag.ini. For the workflow genomic.resequencing.frag, use dibayes.genome.ini.

Certain parameters affect only the module's base-space or color-space algorithm. These parameters are noted in the table. Changes to a base-space parameter's setting have no effect if the module runs its color-space algorithm. Similarly, changes to a color-space parameter's setting have no effect if the module runs its base-space algorithm.

Table 23 SNP (diBayes) parameter description

Parameter name	Default	Description			
Required parameters	Required parameters				
dibayes.call.stringency	medium	Specifies the SNPs call stringency. Each call stringency setting specifies a set of filters. An individual filter parameter setting overrides the call stringency setting, in case of conflict. Allowed values:			
		• <b>highest</b> : Recommend when a very low false positive tolerance is allowed.			
		• <b>high</b> : Requires the allele on both strands.			
		• medium: No both-strand requirement.			
		low: Very aggressive.			
		lowest: Even more aggressive. Very minimal filtering.			
		Lower stringency settings result in more SNP calls, but more false positives are also expected. Higher settings result in fewer SNP calls, with fewer false positives. See "The call stringency parameter" on page 167.			
Optional general parameters	•				
dibayes.het.skip.high.coverage	1	Do not call SNPs when the coverage of position is too high compared to the median of the coverage distribution of all positions. Allowed values:			
		• <b>0</b> : Call a position a SNP under these conditions.			
		• 1: Do not call a position a SNP under these conditions.			
		<b>Note:</b> Enable this filter for whole genome resequencing application. Disable it for whole transcriptome or target resequencing.			
		See "The skip high coverage filter" on page 168.			

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7

Table 23	SNP	(diBayes)	parameter	description	(continued)
----------	-----	-----------	-----------	-------------	-------------

Parameter name	Default	Description
dibayes.reads.min.mapping.qv	8	Requires that the mapping quality value of the read be higher than this minimum mapping/pairing QV. Allowed values: Integers 0–100.
		See "The reads mapping QV parameter" on page 168.
dibayes.detect.2.adjacent.snps	0	Whether or not to detect two adjacent SNPs. Allowed values:
		• <b>0</b> : Do not call a second adjacent SNP as a SNP.
		• 1: If two adjacent SNPs are found, call both as SNPs.
dibayes.polymorphism.rate	0.001	The expected frequency of heterozygotes in the population, for example, 0.001 in humans. Allowed values: Float 0.0–1.0.

## **Optional read filters:**

The entire read is excluded from analysis if a read fails to pass one of these filters.

dibayes.reads.with.unmapped. mate.include	1 for PE data; 0 for other data	<ul> <li>Whether or not to include reads that only have one tag mapped (their mate tags are either unmapped or missing). Allowed values:</li> <li>0: Do not include these reads.</li> <li>1: Include these reads.</li> </ul>
dibayes.reads.with.indel.exclude	1	<ul> <li>Whether or not to exclude reads that have indels. Allowed values:</li> <li>0: Do not exclude reads that have indels.</li> <li>1: Exclude reads that have indels.</li> </ul>
dibayes.reads.only.uniquely. mapped.allow	0	<ul> <li>Whether or not to require that reads be uniquely mapped. A very stringent filter. Allowed values:</li> <li>0: Do not require that reads be uniquely mapped.</li> <li>1: Require uniquely mapped reads. Filter out reads that are not uniquely mapped.</li> </ul>
dibayes.reads.max.mismatch. alignlength.ratio	1	The threshold of mismatch/alignment-length ratio. The reads whose mismatch/alignment length ratio is higher than this specified threshold are filtered (are ignored). Allowed values: Float 0.0–1.0.
dibayes.reads.min.alignlength. readlength.ratio	0	The threshold of alignment-length/read-length ratio. The reads whose alignment-length/read-length ratio is less than this specified threshold are filtered (are ignored). Allowed values: Float 0.0–1.0.

#### **Optional general position filters:**

The set of pile-up color/base calls at a given position is excluded from both heterozygous and homozygous SNP analysis if the position pile-up fails to pass one of these filters.

dibayes.snp.both.strands	0	<ul> <li>Whether or not to require that the novel allele is present on both strands and statistically similar represented on both strand for both heterozygous and homozygous SNPs. Allowed values:</li> <li>0: Do not make the above requirement.</li> <li>1: Make the above requirement.</li> </ul>
dibayes.snps.min.base.qv	26	Require that the candidate allele have at least this base quality value. A base call that fails this filter is excluded from the analysis, but the base pile-up is still considered for SNP calling. Allowed values: Integers 0–41. Base space only. See "The minimum base qv filters" on page 168.

Table 23	SNP	(diBayes)	parameter	description	(continued)
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Parameter name	Default	Description
dibayes.snps.min.nonref.base.qv	26	Require that the non-reference allele have at least this base quality value. A base call that fails this filter is excluded from the analysis, but the base pile-up is still considered for SNP calling. Allowed values: Integers 0–41. Base space only. See "The minimum base qv filters" on page 168.

#### Optional heterozygous position filters:

The set of pile-up color/base calls at a given position is excluded from only heterozygous SNP analysis if the position pileup fails to pass one of these filters.

dibayes.het.min.allele.ratio	0.15	The less-common allele must be at least this proportion of the reads of the first two most common valid alleles. Allowed values: Float 0.0–0.5.
dibayes.het.min.coverage	2	Require at least this coverage to call a heterozygous SNP. Allowed values: Integers >= 0.
dibayes.het.min.start.pos	2	The minimum number of unique start positions required to call a heterozygote. Allowed values: Integers >= 0.
dibayes.het.min.nonref.color.qv	7	Requires the non-reference allele to have at least this average color quality value to call a heterozygous SNP. Allowed values: Integers 0–62. Color space only.
dibayes.het.min.nonref.base.qv	26	Requires the non-reference allele to have at least this average base quality value to call a heterozygous SNP. Allowed values: Integers 0–41. Base space only.
dibayes.het.min.validreads. totalreads.ratio	0.65	The proportion of the total reads containing either of the two candidate alleles. Filters positions with high raw-error rates. Allowed values: Float 0.0–1.0.
dibayes.het.min.adjacentsnp. allele.count	2	Minimum valid tricolor counts (Het). Allowed values: Integers >= 0. Color space only.

## Optional homozygous position filters:

The set of pile-up color/base calls at a given position is excluded from only homozygous SNP analysis if the position pileup fails to pass one of these filters.

dibayes.hom·min.coverage	1	Requires at least this coverage to call a homozygous SNP. Allowed values: Integers >= 0.
dibayes.hom.min.nonref.allele. count	2	Requires at least this non-reference allele count to be called a homozygous SNP. Allowed values: Integers >= 0.
dibayes.hom.min.nonref.color.qv	7	Requires the non-reference allele to have at least this average color quality value to call a homozygous SNP. Allowed values: Integers 0–62. Color space only.
dibayes.hom.min.nonref.base.qv	26	Requires the non-reference allele to have at least this average base quality value to call a homozygous SNP. Allowed values: Integers 0–41. Base space only.
dibayes.hom.min.nonref.start. pos	2	Requires the non-reference allele to have at least this minimum number of unique start positions, to call a homozygote. Allowed values: Integers >= 0.

Table 23	SNP	(diBayes)	parameter	description	(continued)
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Parameter name	Default	Description
Output file processing		
dibayes.write.fasta	1	Whether or not to create FASTA files. Allowed values:
		• <b>0</b> : Do not create the output FASTA files.
		• 1: Create the individual and combined output FASTA files of all contigs.
dibayes.write.consensus	1	Whether or not to create individual consensus calls files of each contig. Allowed values:
		• <b>0</b> : Do not create individual consensus calls files of each contig.
		• 1: Create individual consensus calls files of each contig.
		Neither setting creates a combined consensus calls file of all contigs.
dibayes.compress.consensus	0	Whether or not to zip the generated consensus files. Allowed values:
		• <b>0</b> : Do not compress the consensus files.
		• 1: Compress the individual consensus file of each contig (within the individual contig folders).
Resource parameters		
memory.request	15gb	Memory request for process. Include the units gb in the setting.
java.heap.space	14000	Dynamic memory requirement.
maximum.workers	24	Maximum number of nodes that can be used for this analysis.

# The call stringency<br/>parameterThe dibayes.call<br/>setting represents a c<br/>stringency parameter

The dibayes.call.stringency parameter controls the SNP calling results. Each setting represents a different combination of filters. Table 24 explains the call stringency parameter settings.

Stringency setting	Stringency	SNPs	False positives	Recommended coverage	Comments
highest	_			>80x	Recommend when very low false positive tolerance is allowed.
high		Λ	$\wedge$	20x ~ 80x	Requires the allele on both strands.
medium				1x ~ 25x	No both-strand requirement.
low	$\bigvee$			-	Very aggressive.
lowest	۷			_	Very aggressive. Minimal filtering.

 Table 24
 Empirical dibayes.call.stringency settings for diBayes

Table 25 shows the non-default filter parameter values for each level of stringency. The medium stringency level is not listed because it calls for default values of the filtering parameters. If you explicitly set one of the listed filter parameters, the value that you specify overrides the value called for by the call stringency level.

 Table 25
 Filter parameters affected by call stringency level

Call stringency			Parameter se	tting change		
setting		Filter parameter	Default value	New value		
highest	dibaye	es.snp.both.strands	0	1		
	dibaye	es.het.min.start.pos	2	3		
	dibaye	es.het.min.validreads.totalreads.ratio	0.65	0.70		
	dibaye	es.het.min.adjacentsnp.allele.count	2	3		
	dibaye	es.hom.min.nonref.start.pos	2	3		
high	dibaye	es.snp.both.strands	0	1		
low	dibaye	es.het.min.start.pos	2	1		
	dibaye	es.hom.min.nonref.start.pos	2	1		
lowest	dibaye	es.het.min.adjacentsnp.allele.count	2	1		
	dibaye	es.het.min.nonref.color.qv	7	0		
	dibaye	es.het.min.start.pos	2	1		
	dibaye	es.het.min.validreads.totalreads.ratio	0.65	0.0		
dibaye dibaye		es.hom.min.nonref.allele.count	2	1		
		es.hom.min.nonref.base.qv	26	0		
	dibayes.hom.min.nonref.color.qv		7	0		
	dibaye	es.snps.min.nonref.base.qv	26	0		
The skip high coverage filter		Enable the dibayes.het.skip. positions with high coverage. If yo coverage of a position is too high o of all positions. Enable the dibay genome resequencing. Disable the The filter is enabled by default.	high.coverage ou enable this filter compared to the n es.het.skip.h e filter for transcrip	e filter to skip fal r, the SNP position nedian of the cov igh.coverage ptome or targete	se-positive SNP on is skipped if the rerage distribution filter for whole d resequencing.	
The reads mapping Th QV parameter ma to		The parameter dibayes.reads.min.mapping.qv is a read-level filter. Reads with mapping QVs lower than this value are filtered out. The empirical default of the mapping QV is 8, which gives a balanced performance between sensitivity (more SNPs to be called) and specificity (fewer false positive SNPs) on multiple known data sets.				
		Raising the mapping QV reduces because more reads are filtered ou	the number of fals it, the total numbe	se positive SNP o er of SNP calls de	calls. However, ecreases also.	
The minimum b qv filters	ase	The dibayes.snps.min.base.qv parameters work like a read filter	and dibayes.sr	ps.min.nonref rel. These param	.base.qv eters are very	

ers parameters work like a read filter at the position level. These parameters are very effective in removing bad base calls for SNP analysis. Increasing the value of these parameters causes only high-quality base calls to be considered, and results in higher SNP calling accuracy with a slight sacrifice of the total number of SNPs. When data coverage is high, we recommend using a higher value for higher accuracy.

Lowering the value of the dibayes.snps.min.base.qv and dibayes.snps.min.nonref.base.qv parameters results in more SNP calls.

Table 30 on page 175 also describes how the optional tuning parameters are used in filtering reads.

## (Optional) Genomic annotation parameters

Table 26 describes the annotation parameters you can optionally add to your analysis, to have annotation files generated.

All annotations options that are turned on must be fulfilled by the variant entry, for the variant to be printed in the filtered output file. See Chapter 14, "Add Genomic Annotations to Analysis Results" on page 267 for information about annotations.

Parameter name Default Description Annotation general parameters annotation.run Whether or not to add annotations. Allowed values: • 0: Do not generate annotations. • 1: Generate annotations. The input variant GFF file (see "Annotation input files" on annotation.input.gff.file page 274, in Chapter 14, Add Genomic Annotations to Analysis Results). This file is the output of the SNPs module. Expected file extensions: .gff, .gff3. Example: /data/HuRef\_chr22/DB\_out/HuRef\_rmm13\_SNP.gff3 Must be a valid path, and the file must be readable. If this file is not found or is problematic, or if the parameter is missing, the annotations output file is not created. Annotation source parameters annotation.dbsnp.annotate.snp Whether or not to annotate variants with SNP entries from the dbSNP file. Allowed values: • 0: Do not annotate variants with SNP entries from the dbSNP Note: Applies only to SNPs and small file indels modules • 1: Annotate variants with SNP entries from the dbSNP file. annotation.dbsnp.annotate.indel Whether or not to annotate variants with indel entries from the dbSNP file. Allowed values: • 0: Do not annotate variants with indel entries from the dbSNP Note: Applies only to SNPs and small file. indels modules • 1: Annotate variants with indel entries from the dbSNP file.

Table 26 Optional annotation parameters

Table 26	Optional	annotation	parameters	(continued)
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Parameter name	Default	Description
annotation.indel.border.slack	5	An indel is considered to be matched to a dbSNP indel if the two overlap or if the distance between them is less than or equal to the value of annotation.indel.border.slack.
		Allowed values: Integers >= 0.
Annotation filtering parameters (See Chapter 14, "Add Genomic Annotat	ions to Analysi	s Results" on page 267, for more information on annotation output.)
annotation.show.only.variants.in.exons	0	When set to 1, filters the annotated output: restricts the Filtered Variant annotation output file to only the variants that overlap any exon. Allowed values:
		• <b>0</b> : Do not filter the output for variants that overlap any exon.
		• 1: Restrict output to variants that overlap any exon.
annotation.show.only.variants.in.genes	0	When set to 1, filters the annotated output: restricts the Filtered Variant annotation output file to only the variants that overlap any gene, even if it does not overlap any exon. Allowed values:
		• <b>0</b> : Do not filter the output for variants that overlap any gene.
		• 1: Restrict output to variants that overlap any gene.
		<b>Note:</b> if the parameter annotation.show.only.variants.in.exons is set to 1, then the setting for the parameter annotation.show.only.variants.in.genes is ignored.
annotation.show.only.variants.in.dnsnp	0	When set to 1, filters the annotated output: restricts the Filtered Variant annotation output file to only SNPs or small indels that exist in dbSNP.
		Allowed values:
		• <b>0</b> : Do not filter the output for variants that <i>appear</i> in dbSNP.
<b>Note:</b> Applies only to SNPs and small indels modules		• 1: Restrict output to SNPs or small indels variants that <i>appear</i> in dbSNP.
annotation.show.only.variants.not.in. dnsnp	0	When set to 1, filters the annotated output: restricts the Filtered Variant annotation output file to only SNPs or small indels that do not exist in dbSNP.
		Allowed values:
Note: Annlies only to SNDs and small		• <b>0</b> : Do not filter the output variants that <i>do not appear</i> in dbSNP.
indels modules		• 1: Restrict output to SNPs or small indels variants that <i>do not appear</i> in dbSNP.
Resource parameters		
memory.request	22gb	Memory request. Include the units gb in the setting.
java.heap.space	21500	Dynamic memory request.

If you are creating you own INI files, the annotation parameters are by convention defined in the file tertiary/annotation.dibayes.ini.

Table 27 describes SNPs internal parameters that we do not recommend changing.

Table 27 SNPs internal parameter description

dibayes.run	1	Enables the SNPs module. Allowed values:		
		• <b>0</b> : Do not run a SNPs analysis.		
		• 1: Run the SNPs module during this analysis.		
		The run parameter is set automatically by the shell during a standard workflow. Accept the default for most use.		

## SNPs output files

There are multiple output subfolders (for example, chr\_1, chr\_2, ... chr\_n), corresponding to individual chromosomes or chromosome/contigs. Each subfolder contains up to 4 files:

- <prefix>\_SNP.gff
- <prefix>\_quartiles.txt
- (Optional) <prefix>\_Consensus\_Calls.txt
- (Optional) <prefix>\_Consensus\_Basespace.fasta

The remainder of this section describes these files and provides examples.

In the root of output folder, the individual GFF and FASTA file in the chromosome subfolders are concatenated into the final GFF and FASTA file for the whole genome. Because of the large size of individual consensus calls files, we do not consolidate them into a summarized copy to save space.

The output layout is listed below. The first two files are consolidated data for the entire run. The files in contig subfolders are data for an individual contig.

```
<prefix>_SNP.gff3
<prefix>_Consensus_Basespace.fasta.
contig1/
<prefix>_SNP.gff3
<prefix>_Consensus_Calls.txt
<prefix>_Consensus_Basespace.fasta
<prefix>_quartiles.txt
contig2/
<prefix>_SNP.gff3
<prefix>_Consensus_Calls.txt
<prefix>_Consensus_Calls.txt
<prefix>_Consensus_Basespace.fasta
<prefix>_quartiles.txt
```

GFF file format

The <prefix>\_SNP.gff file is the list of output SNPs. This file is described in Table 28.

Table 28	<dibaves< th=""><th>s.output.</th><th>prefix&gt;.</th><th>aff file</th><th>format</th><th>descri</th><th>ption</th></dibaves<>	s.output.	prefix>.	aff file	format	descri	ption
			p. e				

Column name	Description	Example
##	Header comment lines.	Input files and algorithm parameters
#	Header of the results.	_
seqid	The string ID of the sequence to which the start and end coordinates refer.	chr1
source	The source of the data.	S0LiD_diBayes
type	Sequence ontology derived type for this variation. For diBayes, this is always SNP.	SNP
start	Start position of the SNP.	420
end	End position of the SNP.	420
score	Calculated p-value of the SNP.	0.000000
strand	-	_
phase	-	—
Attributes:		
reference	The base of the reference sequence at the current position.	reference=c
• allele-call	Genotype in alleleA/alleleB (in alphabet order) format for bases observed in all valid reads that cover the position.	allele-call=A/C
coverage	The number of valid reads that cover the current position.	coverage=52
<ul> <li>rawCoverage</li> </ul>	The number of raw reads that cover the current position.	rawCoverage=52
refAlleleCounts	The number of reads of the reference allele at the current position.	refAlleleCounts=22
refAlleleStarts	The number of different start positions of reads having the reference allele at the current position.	refAlleleStarts=15
refAlleleMeanQV	The mean of quality values of all reference allele reads at the current position.	refAlleleMeanQV=15
novelAlleleCount	The number of reads of the most abundant non-reference allele at the current position.	novelAlleleCounts=24
novelAlleleStarts	The number of different start positions of reads having the most abundant non-reference allele at the current position.	novelAlleleStarts=14
novelAlleleMeanQV	The mean of quality values of all novel allele reads.	novelAlleleMeanQV=17
• diColor1	The most abundant allele in the reads (not necessarily the reference allele) in dicolor encoding (for example, one of 00, 01, 32, 33 of 16 possible dicolors).	diColor1=00
	Does not apply to the results of base-space BAM files.	
• diColor2	The second-most abundant allele in the reads.	diColor2=22
	Does not apply to the results of base-space BAM files.	

Column name	Description	Example
• zygosity	Heterozygosity flag. Values are:	zygosity=heterozygous
	homozygous: Homozygous SNP	
	heterozygous: Heterozygous SNP	
• flag	Reports why the SNP is not called as a homozygous SNP or as a heterozygous SNP, when information is available. Values are:	flag=t4,t10
	• <b>m*</b> : Flags beginning with the letter "m" report why the SNP is not called as a Hom.	
	<ul> <li>t*: Flags beginning with the letter "t" report why the SNP is not called as a Het.</li> </ul>	
	• —: No information. The SNP can be either a Het or a Hom.	
	The m* and t* flag values are listed in Table 30 on page 175.	
	A position that is not called as a heterozygous SNP (due to failing current filter settings) may still be called as a homozygous SNP. In this case, the flag reports the reason for not calling the position a heterozygous SNP. Similarly, a position that is not called as a homozygous SNP (due to failing current filter settings) may still be called as a heterozygous SNP. In this case, the flag reports the reason for not calling the position a homozygous SNP.	
	See also the consensus calls Flag field, which reports the reason why a position is not called a SNP, and adds which filter causes this result.	

Table 28	<dibayes.out< th=""><th>put.prefix&gt;.g</th><th>ff file format</th><th>description</th><th>(continued)</th></dibayes.out<>	put.prefix>.g	ff file format	description	(continued)
	,				

**GFF file example** The following is example content of the GFF file generated by the SNPs module.

##gff-version 3
##source-version LifeScope(tm) 2.0 diBayes
<pre>##feature-ontology http://song.cvs.sourceforge.net/*checkout*/</pre>
<pre>song/ontology/sofa.obo?revision=1.141</pre>
##type DNA
##date 2011-03-15
##time 15:56:41
<pre>##genome-build /local/xux1/validataion/HuRef/hg18/reference/</pre>
chr22.validated.fasta
<pre>##input-files /local/xux1/validataion/HuRef/hg18/input/</pre>
<pre>solid0107_20100514_PE_launch_verification_F3.sowmi-1_0309.bam</pre>
<pre>##options call-stringency: medium; dibayes-base-space: 0;</pre>
experiment-name:
<pre>Test_huref_4.0_base_newbam0309_1of3_color2.0_newout_chr22; het-</pre>
<pre>skip-high-coverage: 1; index: 1; log-dir: /local/xux1/</pre>
<pre>validataion/HuRef/hg18/log/newbam0309_lof3_color2.0; output-</pre>
dir: /local/xux1/validataion/HuRef/hg18/outputs/

	<pre>newbam0309_lof3_color2.0; poly-rate: 0.003; reference: /local/ xux1/validataion/HuRef/hg18/reference/chr22.validated.fasta; working-dir: /local/xux1/validataion/HuRef/hg18/tmp/ newbam0309_lof3_color2.0; #ChrSourceTypePos_StartPos_EndScoreStrandPhaseAttributes chr22SOLiD_diBayesSNP14432513144325130.000984 reference=G;allele-call=C/ G; coverage=6; rawCoverage=9; refAlleleCounts=3; refAlleleStarts=3; refAlleleMeanQV=22; novelAlleleCounts=3; novelAlleleStarts=3; nove lAlleleMeanQV=29; diColor1=01; diColor2=32; zygosity=heterozygous; flag=- chr22SOLiD_diBayesSNP14433730144337300.002050 reference=C; allele-call=A/ C; coverage=8; rawCoverage=8; refAlleleCounts=5; refAlleleStarts=4; refAlleleMeanQV=20; novelAlleleCounts=3; novelAlleleStarts=3; nove lAlleleMeanQV=20; novelAlleleCounts=3; novelAlleleStarts=3; nove lAlleleMeanQV=20; novelAlleleCounts=3; novelAlleleStarts=3; nove lAlleleMeanQV=20; novelAlleleCounts=3; novelAlleleStarts=3; nove lAlleleMeanQV=20; novelAlleleCounts=3; novelAlleleStarts=3; nove lAlleleMeanQV=26; diColor1=20; diColor2=31; zygosity=heterozygous; flag=-##</pre>
Combined GFF for all contigs	The combined GFF file is a consolidated version of individual GFF files.
	é(tagk output dir) (é(analygig nama) CND gff?
	The first of the f
	The following is an example of the summarized header of the combined GFF file:
	<pre>##gff-version 3 ##List of SNPs. Date Thu Oct 7 14:32:53 2010 Stringency: medium Mate Pair: 0 Read Length: 50 Polymorphism Rate: 0.001000 Bayes Coverage: 60 Bayes_Single_SNP: 1 Filter_Single_S NP: 1 Quick_P_Threshold: 0.997000 Bayes_P_Threshold: 0.040000 Minimum_Allele_Ratio: 0.150000 Minimum_Allele_Ratio_Multiple_of_Dicolor_Error: 100 ##Data source: Experiment: test_case0003 ##chr1 1// chr\${index} chromosome name ##chr2 2// chr\${index} chromosome name ##chr3 3// chr\${index} chromosome name</pre>
	The portion in bold is the chromosome index and name table, which combine the information from individual GFF files.
Consensus_Calls file format	The <prefix>_Consensus_Calls.txt is a tab-separated file that covers all positions that have coverage and provides general information about each position (see Table 29). Its flag column shows a list of codes of filters (see Table 30 on page 175) that the position fails to pass to be called as a SNP. It is very useful for identifying the reason of false negative SNP calls (the known SNPs that are not called by diBayes). By adjusting the value of keys listed in Table 30, user can loosen the filter stringency and make the positions that originally fail the filters to be considered in the SNP analysis. Consensus calls files are created with file names according to the following pattern:</prefix>

\${task.output.dir}/contig\${index}/\${analysis.name}\_ Consensus\_Calls.txt
Table 29 <diBayes.output.prefix>\_Consensus\_Calls.txt file format description

File Name/Column	Description	Example
Chr	Chromosome/Contig number.	chr1
Position	Location of the SNP on the reference sequence.	442
Allele_DiColor1	The most abundant allele in the reads (not necessarily the reference allele) in dicolor encoding (for example 00, 0132, 33) of 16 possible dicolors. Color space only.	03
Allele_DiColor2	The second most abundant allele in the reads in dicolor encoding (for example 00, 0132, 33) of 16 possible dicolors. Color space only.	03
Reference	The base of the reference sequence at the current position.	С
Genotype	Genotype in the form of IUB codes for bases observed in all the reads.	С
P-value	Calculated p-value of the SNP. (P-values are generated from the Bayesian and Frequentist algorithms.)	1.00000
Coverage	The number of the reads that cover the current position.	2
Unfiltered Coverage	The number of all unfiltered reads that cover the current position.	2
nCounts_1st_allele	The number of the most abundant allele at the current position.	2
nCounts_Reference_allele	The number of reads having the reference allele at the current position.	2
nCounts_NonReference_allele	The number of reads having the most abundant non-reference allele reads at the current position.	0
Ref-Avg-QV	The mean of quality values of all reference allele reads at the current position.	29
Novel-Avg-QV	The mean of quality values of all novel allele reads.	0
Heterozygous	Heterozygosity flag. Values are:	0
	<b>0</b> : Homozygous SNP.	
	1: Heterozygous SNP.	
Algorithm	The algorithm used to call the current SNP. Values are:	-1
	• -1: Not a SNP.	
	• <b>0</b> : Homozygous position.	
	• 1: Heterozygous SNP, called by the Bayesian algorithm.	
	• 2: Heterozygous SNP, called by the Frequentist algorithm.	
Flag	Flag indicating why a location is not called a SNP.	m4
	See Table 30 for a description of Flag values.	

## Table 30 <diBayes.output.prefix>\_Consensus\_Calls.txt flag column description

Flag	Filter meaning	Related parameter
Heterozygote		
t1	Insufficient coverage for a heterozygous position.	dibayes.het.min.coverage.

Flag	Filter meaning	Related parameter	
t2	Insufficient unique start positions for a heterozygous position.	dibayes.het.min.start.pos	
t3	The coverage is too high, compared to the coverage distribution of all positions.	dibayes.het.skip.high.coverage	
t4	The fraction of the second-most common <i>valid</i> color (or base allele) in the total of top two valid colors (or the top two base alleles) is higher than the threshold (usually a function of raw error squared).	dibayes.het.min.allele.ratio	
t5	Genome positions with sufficient coverage (20x) at which there is only 1 unique read position for all the reads. It could be a PCR error.	_	
t6	The candidate SNP is evenly distributed over positions (not used).	-	
t7	The non-reference allele is not on both strands (counting the reads).	dibayes.snp.both.strands	
t8	Both alleles not evenly represented on both strands (doing statistical test on read distribution of the both strands).	dibayes.snp.both.strands	
t9	The second-most common valid dicolor is not more frequent than the third-most. Color space only.	_	
t10	There are no other valid SNPs, or the second-most common valid SNP (as a proportion of all valid SNPs) is less than half the 2-dibase error frequency. Color space only.	dibayes.het.min.allele.ratio	
t11	Sum of the first and second-most common alleles must be at least this proportion, for example, 0.5, of all reads at this position.	dibayes.het.min.ratio.validreads. totalreads.ratio	
t12	The dicolor alleles are inconsistent to the reference allele. Color space only.	_	
t13	The difference of quality values of the non-reference allele and the reference allele is much lower the quality value of the reference allele.	_	
t14	The average of p-values of color/base quality values of the non- reference allele is lower than one standard deviation from the mean of p-value distribution, when the non-reference allele has to be at low frequency (rare variant).	_	
t15	The average of p-values of color/base quality values of the position is lower than one standard deviation from the mean of the p-value distribution.	_	
t16	Insufficient coverage of the reference allele.	-	
t17	Insufficient coverage of the non-reference allele.	-	
t18	Zero coverage.	_	
t19	Insufficient number of start positions of non-reference allele.	_	
t20	Insufficient coverage for either of two alleles, when neither allele is same as the reference.	_	
t21	Quality value of non-reference allele too low (lower than a relative threshold depend on the distribution of all color quality values).	_	
t22	Quality value of non-reference allele too low (lower than an absolute threshold).	dibayes.het.min.nonref.color.qv	
Homozygote			
m1	Insufficient coverage for a homozygous SNP.	dibayes.hom.min.coverage	

## Table 30 <diBayes.output.prefix>\_Consensus\_Calls.txt flag column description (continued)

Flag	Filter meaning	Related parameter
m2	Insufficient proportion of the count of the most common valid dicolor (base) allele to all valid reads.	_
m3	Non-reference allele not on both strands.	dibayes.snp.both.strands
m4	Reserved for future filters (not used).	_
m5	Second-most common allele too close in coverage to the first most common allele.	_
m6	Insufficient coverage (as a fraction of the median of the coverage distribution of all positions) for a homozygous call.	_
m7	Dicolor inconsistent with reference. Color space only.	_
m8	Insufficient number of non-reference alleles	dibayes.hom.min.nonref.allele. count
m9	Same as the m2 flag.	-
m10	Insufficient number of start positions of non-reference alleles.	dibayes.hom.min.nonref.start.pos
m11	No coverage.	-
m12	Quality value of non-reference allele too low (lower than an absolute threshold).	_
m13	Quality value of non-reference allele too low (lower than a relative threshold depend on the distribution of all color quality values).	_

		<i>с</i> 0	<u> </u>		()
Table 30	<dibayes.output.pr< th=""><th>etix&gt;_Consensus</th><th>_Calls.txt flag</th><th>column descri</th><th>ption <i>(continued)</i></th></dibayes.output.pr<>	etix>_Consensus	_Calls.txt flag	column descri	ption <i>(continued)</i>

Consensus calls	The following is example output of a SNPs consensus calls file.
file example	##consensus-call
	##source-version LifeScope(tm) 2.0 diBayes
	##type DNA
	##date 2011-03-10
	##time 11:38:51
	<pre>##genome-build /local/xux1/validataion/HuRef/hg18/reference/ chr1.validated.fasta</pre>
	##input-files /local/xux1/validataion/HuRef/hg18/input/
	solid0107_20100514_PE_launch_verification_F3.sowmi-1_0309.bam
	<pre>##options call-stringency: medium; dibayes-base-space: 0;</pre>
	experiment-name:
	Test_huref_4.0_base_newbam0309_1of3_color2.0_chr1; het-skip-
	high-coverage: 1; index: 1; log-dir: /local/xux1/validataion/
	<pre>HuRef/hg18/log/newbam0309_1of3_color2.0; output-dir: /local/</pre>
	<pre>xux1/validataion/HuRef/hg18/outputs/newbam0309_1of3_color2.0;</pre>
	<pre>poly-rate: 0.003; reference: /local/xux1/validataion/HuRef/</pre>
	hg18/reference/chr1.validated.fasta; working-dir: /local/xux1/
	<pre>validataion/HuRef/hg18/tmp/newbam0309_1of3_color2.0;</pre>
	#Chr Position Allele_DiColor1 Allele_DiColor2 Reference
	Genotype P-value Coverage Unfiltered_Coverage
	nCounts_1st_allele nCounts_Reference_allele
	nCounts_NonReference_allele Ref_Avg_QV NonRef_Avg_QV
	Zygousity Algorithm Flag

	chrl1     50262     10     10     T     T     1.000000     1       1     1     1     0     25     0     homozygous     -
	1 t4,t10,t1,t2,t21,t22,m4 cbr11 50263 00 00 T T 1.000000 1
	1 1 1 0 25 0 homozygous -
	1 t4,t10,t1,t2,t21,t22,m4
Contig base space FASTA file	The <i><prefix>_</prefix></i> Consensus_Basespace.fasta file is a base space translation of the genotype column of its corresponding consensus call file. The bases are encoded with IUB codes. All reference positions that have 0 coverage are replaced by "N".
	The individual base space FASTA files are created with file names according to the following pattern:
	\${task.output.dir}/contig\${index}/\${analysis.name}_\ Consensus_Basespace.fasta
	The following is example content of a contig base space FASTA file. The title line is always a ">" sign followed by the chromosome name. The length of each sequence line is 70.
	>chromosome_name NNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNN
	NNNNNNNNNNNNNNACGGGGGGGGGGGGGGGGGGGGGGAAAAAAAA
Combined FASTA for all contigs	The combined FASTA file for all contigs is a consolidated version of individual base space FASTA files. The combined file is named according to the following pattern:
j	<pre>\${task.output.dir}/\${analysis.name}_Consensus_Basespace.fasta</pre>
	For field information and example content, see "Contig base space FASTA file".
Quartiles file	The <prefix>_quartiles.txt file lists the quartile and percentile information about the coverage and color quality value distribution of all positions of the chromosome. Quartiles files are created with file names according to the following pattern:</prefix>
	<pre>\${task.output.dir}/contig\${index}/\${analysis.name}_ quartile.txt</pre>
	The following shows an example output of a SNPs quartiles output file.
	25 percentile of the coverage distribution (the first quartile): 4 50 percentile of the coverage distribution (median): 11 75 percentile of the coverage distribution (the third quartile): 23
	0.1 percentile of the average color quality value per position distribution: 4
	0.2 percentile of the average color quality value per position distribution: 4
	1 percentile of the average color quality value per position distribution: 5

50 percentile of the average color quality value per position distribution (median): 19

## SNPs algorithm description

LifeScope<sup>™</sup> Software uses the diBayes package to find Single Nucleotide Polymorphisms (SNPs). The diBayes package performs independent SNP analysis at each position in the reference, using either a Bayesian or Frequentist algorithm (see Figure 9 on page 180 and Figure 10 on page 181).

FrequentistThe Frequentist algorithm is used when coverage is high for a given position, for<br/>example, 60x. Given the assumption that the errors follow a Poisson distribution, the<br/>Frequentist algorithm calculates the probability of a null hypothesis that the observed<br/>valid dicolor base mismatches are errors. If the probability of the null hypothesis is too<br/>low, the hypothesis is rejected and the position is called a SNP.

**Bayesian algorithm** The SNP finding problem is to predict the most likely genotype (*G*) at a genome position from a list of reads (*R*) that are mapped to that position. To do that, for each genome type *G*, we want to estimate the posterior probability of P(G|R). Using Bayesian theorem, P(G|R)=P(R|G)P(G)/P(R), where P(G) is the prior probability of the genotype *G*, P(R|G) is the posterior probability of observing the given read set at the position assuming the true genome type is *G*, and P(R) is the prior probability of the read sets. P(R) is a constant for a given set of *R* and can be ignored from further consideration. To estimate P(R|G), the algorithm takes into consideration the color (or base) quality values, position error profiles, and probe error profiles. (The Frequentist algorithm also uses these information.)

Define S(G)=P(R|G)P(G) and  $T = \sum_{G} S(G)$ , the likelihood of each genotype, L(G),

can be estimated by S(G)/T. To simplify the computation, our Bayesian algorithm compares two hypotheses; the first, that the true genotype is homozygous for the most common allele ( $G_0$ ) in the reference, and the second, that the true genotype is heterozygous (SNP) for the top two alleles ( $G_a$ ) in the reference. Therefore, *T* can be simplified to  $S(G_0)+S(G_a)$ , and  $L(G_0)=S(G_0)/T$ , and  $L(G_a)=S(G_a)/T$ . The algorithm reports the genotype with the highest likelihood.

Data flow Figure 9 shows how diBayes can take multiple BAM files that are sorted by genome positions as input to make SNP calls at positions with coverage. Running diBayes requires the position error and probe error files of each BAM file. The position error files and probe error files are generated by the mapping statistics module. In a multi-CPU computing cluster, diBayes automatically distributes parallel jobs for individual chromosomes to different computing nodes. The final results for the whole genome are merged after the individual jobs are complete.





Internal moduleThe SNPs module first validates the input BAM files, then determines whether to use<br/>its internal color-space or base-space processing.

#### **BAM** file validation

The SNPs module requires the following:

- Each BAM file must have only one read group and only one library type.
- Each BAM file is sorted by genomic coordinates.
- The reference sequence lengths in BAM file header and in reference FASTA file are identical.
- The BX and BY fields in the BAM headers of all input BAM files must be consistent. (The BX and BY fields record whether the original XSQ reads file had base space present on tag 1 and tag 2, respectively.)

If BAM files that do not meet these criteria are included in the analysis, the analysis fails.
#### Color-space and base-space processing

The SNPs module uses its color-space processing algorithm if the following are both true:

- For paired data, BX and BY are both 0, or, for fragment data, BX is 0.
- Color space information is present in the BAM data (CS and CQ fields are present in the first 10 read records).

For other conditions, the SNPs module uses its base-space processing algorithm. (The exception is BAM files with inconsistent BX and BY fields. These BAM files are rejected previously during BAM file validation.)

The type of processing selected is important because certain filtering and other parameters are only effective on either base space or color space, but not both. The parameters that only apply to one type of processing are noted in the parameters table, Table 23 on page 164.

Both types of processing follow the steps shown in Figure 10.



Figure 10 diBayes algorithm flowchart

# FAQ – SNPs

#### SNP calling is taking too long. What can I do?

Parallelization is the best solution for running diBayes on large data sets. LifeScope<sup>™</sup> Software has implemented the parallel distribution of diBayes jobs for individual chromosomes. It can help the users to achieve the best running efficiency. The diBayes uses BAM files as its input and removes all large temporary files (improving run times).

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#### I seem to be missing SNPs — how do I troubleshoot?

First, try repeating the analysis with a lower stringency level (for example, changing dibayes.call.strengency from medium to high). Look at the positions that you expect to be SNPs in the consensus calls file. Typically, you should find a list of flags that describe the reasons the position was not called as a SNP (see Table 28 on page 172). Look at the properties of this position. Visualizing the reads might be helpful here.

Is the coverage much higher than average? The filter

dibayes.het.skip.high.coverage can remove heterozygous SNPs at positions of extremely high coverage because these have previously been observed to be variants in repeat regions rather than truly heterozygous SNPs at a single position.

Are the reads strongly biased towards one strand, or is the non-reference allele missing from one strand? You probably want to perform a run with dibayes.call.stringency=medium, or switch off the "both strands" requirement (dibayes.snp.both.strand=0).

Do all the reads have the same start position because of the way the sample was prepared? You need to reduce the number of unique start positions required to call a SNP (dibayes.het.min.start.pos and dibayes.hom.min.start.pos).

#### I seem to be finding too many SNPs — how do I troubleshoot?

Look at the properties of the SNPs and compare them to the properties of all the positions in the consensus\_calls.txt file. Is the coverage of these SNPs much lower or much higher than average? Is the color quality value of the non-reference allele much lower than average? You might want to post-filter the results if you find these kinds of patterns. For example, you might want to remove SNPs with very low coverage, or very low color quality values, or p-values close to one. You might want to repeat the analysis with a more stringent setting for example, changing the parameter dibayes.call.stringency from medium to high.

Filtering by p-values is not recommended for homozygous SNPs. See also FAQ 4, "How do I interpret the SNPs p-values?" on page 183.

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#### How do I interpret the SNPs p-values?

P-values close to 0 mean that the algorithm has strong confidence that the genotype call is correct. P-values close to 1 mean that the algorithm does not have very strong confidence that the genotype call is correct.

Note that the p-value refers to the probability that the *genotype* call is correct. This meaning is not the same as the probability as "there is a SNP at this position". A homozygote SNP can have a p-value close to 1, but it is still highly likely that a SNP is present at this location.

Most commonly, a p-value close to 1 occurs when there is good evidence that a SNP is present, *but* it is difficult to confidently decide whether the genotype is heterozygote or homozygote. For example, take a case where there is coverage=3 and all of these are high QV non-reference allele calls (e.g., the reference is A and there are three C's observed). In this case, it is clear there is a SNP in this location. However, it is possible this might really be a heterozygous SNP, and if we observed a fourth allele, it might be an A. So the p-value on this homozygous SNP call might be close to 1, since we are not very confident of the genotype call, even though we are confident that there is a SNP in this location. In this case, we are still confident that a SNP is present at this location.

For this reason, it is recommended not to filter out homozygous SNPs with high pvalue, as this will result in removing many real SNPs.

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#### Can the SNPs module find two adjacent SNPs?

Yes, in some circumstances. You must set the dibayes.detect.2.adjacent.snps parameter to 1. With this setting, diBayes detects and properly calls these scenarios:

- Two adjacent heterozygous SNPs
- Two adjacent homozygous SNPs
- A homozygous SNP adjacent to a heterozygous SNPs (base-space algorithm only)
- A heterozygous SNP adjacent to a homozygous SNPs (base-space algorithm only)

However, diBayes does not call a homozygous SNP adjacent to a heterozygous SNP in color-space BAM files (even with dibayes.detect.2.adjacent.snps on). In general, adjacent SNPs calls may have more false positives than single SNPs calls do.

#### Why does SNPs output sometimes contain a blank dicolor value?

Occasionally, the color sequence is not reported and the base sequence is reported as an 'N', before and after a SNP. This reporting usually occurs when the non-reference allele is more abundant than the reference allele. In these cases, the bases flanking the SNP are the normal reference sequence. (The color sequence appears as a space character in the output file. The space character may affect pattern-matching tools.)

#### How can I control the sensitivity and specificity of SNP calling?

Different call stringency settings and filter settings may help users to achieve different sensitivity and specificity requirements. The more stringent the filters are, the less sensitivity the SNP call and the higher specificity in general. Two filters that are very flexible for users' different needs are the dibayes.reads.min.mapping.qv (MQV) and dibayes.reads.min.alignlength.readlength.ratio (ARR) parameters. Changing one or both of these parameters filters out low confident reads and gets SNP calls with high accuracy. The following figures show that changing MQV and ARR may affect the total number of SNP calls and the dbSNP concordance of the predictions.



Figure 11 Total SNP calls on chromosome 1 of a HuRef long mate-pair data as a function of mapping QV cutoffs and ratios of alignment length and read length (ARR)



Figure 12 dbSNP concordance of SNP calls on chromosome 1 of a HuRef long mate-pair data as a function of mapping QV cutoffs and ratios of alignment length and read length (ARR)

# Why are the coverages reported by diBayes for individual genomic positions sometimes different from the coverage derived from input BAM files?

The difference happens because diBayes reports the coverage of filtered reads instead of raw coverages derived from input BAM files. The diBayes module calls SNPs only for these reads that are:

- Based on reads that are primary alignments.
- Are classified as an AAA pair (See "FAQ Pairing" on page 155 for a description of the 3-letter genomic code classifications).
- Are not duplicates.
- Do not contain indel.
- Have high mapping quality values (default is 8).
- Pass other read level filters (such as optional tuning parameters such as dibayes.reads.max.mismatch.alignlength.ratio and others described in Table 23 on page 164).

The SNPs module reports both coverage and raw coverage at every position, for users' convenience.

Users can find detailed information about the numbers of reads filtered by read level filters and the total number of raw reads at the end of each algorithm log file.

#### Why is a SNPs analysis not recommended with whole transcriptome data?

The reverse transcription process that is part of the WT library prep intrinsically adds a percentage of sequencing errors. In addition, alternative splicing and other alignment complexities around exon junctions may cause local misalignments, which can lead to false positive SNP calls. Although the SNPs module can run successfully on BAM files generated from the WTA module, using those BAM files may cause a higher number of false positive SNPs.

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# 10

# Run a Human CNVs Analysis

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# **Overview**

The LifeScope<sup>™</sup> Software Copy Number Variation (Human CNV) module detects copy number variations in a data sample that is mapped to the reference genome. The analysis method is based on depth of coverage. The module performs these phases in the analysis:

- Samples the genome sequence into non-overlapping windows
- Identifies the regions with significantly higher or lower coverage
- Assigns copy number calls to the identified regions, based on the scale of coverage

The CNV module requires predicted mappability files, which are provide with the LifeScope<sup>™</sup> Software reference repository. These pre-computed files are available for the human reference sequences hg18 and hg19. The module currently supports only human CNV detection. By default, the module does not call CNVs within 1 MBase of the chromosomes' centromeres and telomeres. These regions are highly repetitive and tend to contain many apparent CNVs that are not of general interest. The size of the blackout regions is a user-configurable parameter, cnv.trim.distance. The module is designed to detect larger CNVs (which can take up whole chromosomes or large portions of chromosomes) as well as smaller CNVs, by doing either global or local normalization, respectively. By default, and if the number of valid chromosome arms is greater than 6, the module does global normalization. The parameter cnv.local.normalization is provided to switch to local normalization.

The major components of the CNV module include the following:



- Coverage calculation
- Sampling into windows
- Normalization
- Segmentation
- Post-processing

# Examples of running a CNV analysis

#### In a standard workflow

The following standard workflows provide examples of running the CNV module:

- genomic.resequencing.frag
- genomic.resequencing.lmp
- genomic.resequencing.pe

The following are example shell commands using reads and references from the LifeScope<sup>™</sup> Software repository.

```
# log into the shell
lscope.sh shell -u username -w password
# cd to the projects repository
cd /projects
# create a project and open it
cd /projects
mk proj1
cd proj1
# create an analysis and open it
mk run1
cd run1
# define the workflow
set workflow genomic.resequencing.pe
# define the input (dummy file names are used in the example)
add xsq run0209a_50_PE.xsq
add xsg run0209b_50_PE.xsg
# define the reference
set reference hq19
# optionally change CNV defaults here
# list the configuration of the analysis
ls
# start the run
run
# list progress information about the run
ls
See for Chapter 7, "Run a Standard Workflow Analysis" on page 97 for more
information on running standard workflows. Chapter 6, "Run a Command Shell
Analysis" on page 71 for information on shell commands and syntax.
```

# **As a demo analysis** The optional examples download includes an example of how to run the CNV module by itself, as a single analysis that is not part of a standard workflow. The demo example is not recommended for general use.



See Appendix E, "Demo Analyses" on page 507 for information on running the CNV module as a standalone analysis.

# **CNV** module parameter descriptions

This section describes the parameters that control the runtime behavior of the CNV module. See also Table 32 on page 191, which lists the effect of the cnv.stringency.setting value on other stringency parameters.

In order to change a parameter value in your analysis, use the set param shell command to replace the line # *optionally change CNV defaults here* in the example commands in "In a standard workflow". For instance, to change the call stringency, use this shell command:

set cnv.stringency.setting High /tertiary/cnv.ini

 Table 31
 CNV module parameter descriptions

Parameter name Default value		Description				
CNV parameters						
cnv.window.size	5000	Size of the window block to be considered as a region. Allowed values: Integers >= 100.				
cnv.trim.distance	1000	Distance in kilo bases to be trimmed from the extreme ends of the chromosome arms. Allowed values: Integers 0–100000.				
cnv.min.quality	2	Minimum mappability quality value of the alignments. Allowed values: Integers 0–100.				
cnv.ploidy	2	General ploidy of the genome. Allowed values: Integers >= 1.				
cnv.ploidy.exception	None	List of all the contigs whose ploidy is different to the general ploidy of the genome. Entries in the list are in the format {contig id: ploidy of the contig}, and are separated by commas. Use the string "None" to indicate no entries.				
cnv.local.normalization 0 Whether or not genome-wide normalizat be performed. Allowed values:		Whether or not genome-wide normalization or local normalization should be performed. Allowed values:				
		O: Perform genome-wide normalization.				
		1: Perform chromosome-arm local normalization.				
cnv.write.coverage	0	Whether or not to create coverage output files. Allowed values:				
		O: Do not create coverage output files.				
		1: Create coverage output files, in WIG format				
cnv.coverage.wsize	1000	Size of the window block to be considered as a region for writing coverage output. The mean coverage of all bases in each of these windows is output. Allowed values: Integers 0–100000.				
cnv.gender	Male	Set the ploidy of sex chromosomes in Human correctly. Allowed values: Male, Female.				
CNV stringency parameter	Ś					



#### Table 31 CNV module parameter descriptions

Parameter name	Default value	Description	
cnv.stringency.setting	Medium	Set the stringency setting for calling CNVs. Allowed values:	
		• <b>High</b> : Recommend when a very low false positive tolerance is allowed.	
		Medium: Default values.	
		Low: Aggressive CNV calling.	
		Lower settings results in more CNV calls, but with more false positives. Higher settings result in fewer CNV calls, but with fewer false positives.	
		See also Table 32 on page 191.	
cnv.deletions.min. mappability	50	Minimum mappability percentage for regions to be shown as copy number deletions. Allowed values: Floats 0.0–100.0.	
		If this parameter is not specified, and	
		• cnv.stringency.setting is set to <i>High</i> , then this parameter is set to 75.	
		• cnv.stringency.setting is set to <i>Low</i> , then this parameter is set to 10.	
cnv.insertions.min. mappability	10	Minimum mappability percentage for the regions to be shown as copy number insertions. Allowed values: Floats 0.0–100.0.	
		If this parameter is not specified, and	
		• cnv.stringency.setting is set to <i>High</i> , then this parameter is set to 25.	
		• cnv.stringency.setting is set to <i>Low</i> , then this parameter is set to 0.	
cnv.deletions.min. windows	2	Minimum number of windows for the regions to be shown as copy number deletions. Allowed values: Integers >= 0.	
		If this parameter is not specified, and	
		• cnv.stringency.setting is set to <i>High</i> , then this parameter is set to 4.	
		• cnv.stringency.setting is set to <i>Low</i> , then this parameter is set to 1.	
cnv.insertions.min. windows	2	Minimum number of windows for the regions to be shown as copy number insertions. Allowed values: Integers >= 0.	
		If this parameter is not specified, and	
		• cnv.stringency.setting is set to <i>High</i> , then this parameter is set to 4.	
		• cnv.stringency.setting is set to <i>Low</i> , then this parameter is set to 1.	
cnv.deletions.max.pval	1.0	Maximum p-value for regions to be shown as copy number deletions. Allowed values: Floats 0.0–1.0.	
		If this parameter is not specified, and cnv.stringency.setting is set to <i>High</i> , then this parameter is set to 0.1.	
cnv.insertions.max.pval	1.0	Maximum p-value for regions to be shown as copy number insertions. Allowed values: Floats 0.0–1.0.	
		If this parameter is not specified, and cnv.stringency.setting is set to <i>High</i> , then this parameter is set to 0.1.	
cnv.deletions.max.ratio	0.5	Maximum ratio between the coverage of the region and the expected coverage, for a region to be called as CNV deletion. Allowed values: Floats 0.0–1.0.	
		If this parameter is not specified, and	
		• cnv.stringency.setting is set to <i>High</i> , then this parameter is set to 0.3.	
		• cnv.stringency.setting is set to <i>Low</i> , then this parameter is set to 0.6.	

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Table 31 CNV module	parameter	descriptions
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Parameter name	Default value	Description
cnv.insertions.min. 1.25 ratio		Minimum ratio between the coverage of the region and the expected coverage, for a region to be called as CNVs insertion. Allowed values: Floats >= 1.0.
		If this parameter is not specified, and
		• cnv.stringency.setting is set to <i>High</i> , then this parameter is set to 1.5.
		• cnv.stringency.setting is set to <i>Low</i> , then this parameter is set to 1.1.
Resource parameters		
memory.request	15gb	Memory request. Include the units gb in the setting.
java.heap.space	3000	Dynamic memory request.
processors.per.node	8	Number of processors per node. The maximum number of CPUs that all simultaneous processes are expected to use to full capacity.

Table 32 shows the non-default filter parameter values for each level of stringency that is set with the cnv.stringency.setting parameter. The Medium stringency level is not listed because it calls for default values of the filtering parameters. If you explicitly set one of the listed filter parameters, that value overrides the value called for by the call stringency level.

Table 32	Filter	parameters	affected	by	call	stringency	level	L
----------	--------	------------	----------	----	------	------------	-------	---

Call stringency	Filter perspector	Parameter setting change			
setting	Filler parameter	Default value	New value		
High	cnv.deletions.min.mappability	50	75		
	cnv.insertions.min.mappability	10	25		
	cnv.deletions.min.windows	2	4		
	cnv.insertions.min.windows	2	4		
	cnv.deletions.max.pval	1.0	0.1		
	cnv.insertions.max.pval	1.0	0.1		
	cnv.deletions.max.ratio	0.5	0.3		
	cnv.insertions.min.ratio	1.25	1.5		
Low	cnv.deletions.min.mappability	50	10		
	cnv.insertions.min.mappability	10	0		
	cnv.deletions.min.windows	2	1		
	cnv.insertions.min.windows	2	1		
	cnv.deletions.max.ratio	0.5	0.6		
	cnv.insertions.min.ratio	1.25	1.1		



# (Optional) Genomic annotation parameters

Table 33 describes the annotation parameters you can optionally add to your CNV analysis, to have annotation files generated. See Chapter 14, "Add Genomic Annotations to Analysis Results" on page 267 for information about annotations.



Parameter name	Default value	Description			
Annotation general parameters					
annotation.run	-	Specifies whether or not to add annotations. Allowed values:			
		• <b>0</b> : Do not generate annotations.			
		• 1: Generate annotations.			
annotation.input.gff.file	_	The input variant GFF file, generated by CNV analysis module. (See "Annotation input files" on page 274.)			
		Expected file extensions: .gff, .gff3.			
		Example: /data//HuRef_chr22/DB_out/HuRef_rmm13_SNP.gff3			
		Must be a valid path, and the file must be readable. If this file is not found or is problematic, or if the parameter is missing, annotations output is not created.			
Annotation source parameters					
annotation.indel.border.slack	5	An indel is considered to be matched to a dbSNP indel if the two overlap or if the distance between them is less than or equal to the value of annotation.indel.border.slack.			
		Allowed values: Integers >= 0.			
Annotation filtering parameters					
annotation.show.only.variants.in.exons	0	When set to 1, filters the annotated output: restricts the Filtered Variant annotation output file (see "Annotation output files" on page 282) to only the variants that overlap any exon.			
		Allowed values:			
		<ul> <li>0: Do not filter the output for variants that overlap any exon.</li> <li>1: Restrict output to variants that overlap any exon.</li> </ul>			
annotation.show.only.variants.in.genes	0	When set to 1, filters the annotated output: restricts the Filtered Variant annotation output file (see "Annotation output files" on page 282) to only the variants that overlap any gene, even if it does not overlap any exon.			
		Allowed values:			
		• <b>0</b> : Do not filter the output for variants that overlap any gene.			
		• 1: Restrict output to variants that overlap any gene.			
		<b>Note:</b> if the parameter annotation.show.only.variants.in.exons is set to 1, then the setting for the parameter appotation show only variants in genes is ignored			
Resource narameters					



#### Table 33 Annotation parameters description (continued)

Parameter name	Default value	Description
memory.request	22gb	Memory request. Include the units gb in the setting.
		For hg18 or hg19 sources, use memory.request=15gb.
java.heap.space	21500	For hg18 or hg19 sources, use java.heap.space=14000.

If you are creating your own INI files, annotations parameters for CNV are by convention defined in the file <*analysis*>/tertiary/annotation.cnv.ini.

Table 34 describes CNV parameters that we do not recommend changing.

Table 34 Annotation internal parameters description

Parameter name	Default value	Description
cnv.run	1	Enables the CNV module. Allowed values:
		• <b>0</b> : Do not run a CNV analysis.
		• 1: Run the CNV module during this analysis.
		The run parameter is set automatically by the shell during a standard workflow. Accept the default for most use.
cnv.mappability.dir	\${analysis.	Path to the directory of mappability files.
	mappability.dir}	This parameter is set automatically by the shell when you use the default hg18 or hg19 assemblies in the reference repository.

# Human CNVs results file format description

This section describes the file format of the \*.out files and the GFF file created by the Human CNVs module run.

\*.out files

The \*.out files and their contents are:

- AllSegments.out Contains all the segments, including segments with normal copy number.
- AllCNVs.out Contains all the segments with copy number deletions or duplications (CN != 2).
- **OutputCNVs.out** Contains all the segments with copy number deletions or duplications which pass the specified filtering criteria. These segments are the output CNVs, and are the same as those reported in a different format in the GFF file.

The three \*.out files share a common file format, which is described in Table 35. You can view the files in a text editor or a spreadsheet application.

Wiggle fileThe CNV module optionally creates, per chromosome, one wiggle format file named<br/>Coverage\_chr\*\*.wig. This file contains the coverage in specified window sizes.<br/>You can visualize the file in a browser such as the Integrative Genomics Viewer (IGV).<br/>You can download the IGV browser from the Broad Institute website:

www.broadinstitute.org/igv

The wiggle file specification is available from this site:

#### http://genome.ucsc.edu/goldenPath/help/wiggle.html

**GFF file** The module creates one OutputCNVs.gff file. The file formats are described in Table 36. You can view the OutputCNVs.gff file in a text editor or a spreadsheet application. You can also visualize the file in a browser such as the Integrative Genomics Viewer (IGV), or in a browser such as the UC Santa Cruz genome browser. You can download the IGV browser from the Broad Institute website. For more information, go to these sites:

#### www.broadinstitute.org/igv genome.ucsc.edu

Table 35 CNV \*.out file format descriptions

Column Title	Description	Example
Chrom	Chromosome number.	chr1
start	Start location of the CNV region.	1636395
end	End location of the CNV region.	1780200
mappability	Fraction of mappable bases in the CNV region.	83.745453
coverage	Mean coverage of the windows in the CNV region.	4.080000
log2Ratio	Mean of Log2Ratios of the windows in the CNV region.	-1.126154
copy number	Copy number of the region.	The copy number is relative to a diploid genome, with a normal copy number of 2.
numWindows	Number of windows in the CNV region.	22
p-val	p-value of the CNV call for the region.	0.00001 is very confident. 0.99 is not at all confident.

#### Table 36 OutputCNVs.gff file format

Column Title	Description	Example
Chr	The ID of the sequence to which the start and end coordinates refer.	chr1
Source	Free text-qualifier indicating the algorithm or method that generated the feature.	AB_CNV_PIPELINE
Туре	Sequence ontology derived type for this variation.	repeat_region
Pos_Start	Start position of the CNV Region.	1144255
Pos_End	End position of the CNV Region.	0.04223
Score	p-Value of the CNV Region.	_
Strand	Not used for this output.	_
Phase [attribute]	Not used for this output.	_
Attributes		
copynumber	Copy number of the region.	1

1	$\cap$
	U

	<b>_</b>	
Column Title	Description	Example
logRatio	Mean of Log2Ratios of the windows in the CNV region.	-1.258771
numWindows	Number of windows in the Human CNV region.	23
		Usually, the larger this number is, the more confident the CNV call is.
coverage	Mean coverage of the windows in the CNV region.	4.080000
mappability	Fraction of mappable bases in the Human CNV	91.421745
	region.	The maximum value is 100. A low number may indicate that this region is difficult to map, increasing the likelihood of a false positive CNV call.

Table 36 OutputCNVs.gff file format (continued)

# Human CNVs results file examples

This section provides examples of the \*.out files and the GFF file created when you run the Human CNVs module. This output can be viewed in a text editor or a spreadsheet application.

The following is example \*.out file output, from an OutputCNVs.out file.

chrom p-val	start	end	mappak	oility	coverage	logRatio	С	оруі	number	numWindows
chr12	50772857	507	98807	59.6667	6.13873	-1.23401	1	3	0.000122	201
chr12	51001962	510	67993	98.3077	23.8285	0.628887	4	13	1.53379	e-16
chr12	52032318	520	51850	76.6667	7.27728	-1.09105	1	3	0.12058	3
chr12	52756738	528	06997	30	5.85307	-1.3758	1	3	1.4977e	-05
chr12	62211177	624	34738	31.5385	5.84658	-1.21177	1	13	0.25666	б

The following is example output of the file OutputCNVs.gff:

##gff-version 3 ##source-version LifeScope(tm) 2.0 {CNVcaller} ##feature-ontology http://song.cvs.sourceforge.net/\*checkout\*/song/ontology/ sofa.obo?revision=1.141 ##type DNA ##date 2011-02-10 ##time 15:44:31 ##genome-build NCBI B36 ##input-files /local/gottimrk/cnv\_test\_datasets/solid0122\_20100412\_chr12Q\_chr16Q\_CNVs-Paired.bam #Chr Pos\_Start Pos\_End Score Strand Phase Source Type Attributes 50772857 50798807 AB\_CNV\_PIPELINE chr12 repeat\_region 0.00012201 . copynumber=1;numWindows=3;coverage=6.13873;logRatio=-1.23401;mappability=59.6667 51001962 51067993 chr12 AB\_CNV\_PIPELINE repeat\_region 1.53379e-16 . copynumber=4;numWindows=13;coverage=23.8285;logRatio=0.628887;mappability=98.3077 52032318 52051850 chr12 AB\_CNV\_PIPELINE repeat\_region 0.120583 copynumber=1;numWindows=3;coverage=7.27728;logRatio=-1.09105;mappability=76.6667 52756738 52806997 chr12 AB\_CNV\_PIPELINE repeat\_region 1.4977e-05 . copynumber=1;numWindows=3;coverage=5.85307;logRatio=-1.3758;mappability=30

LifeScope<sup>™</sup> Genomic Analysis Software Command Shell User Guide



# Algorithm for the Human CNV module

The Human CNV module algorithm has six steps:

- Preprocessing
- Coverage calculation
- Sampling into windows
- Normalization
- Segmentation
- Post processing
- **Preprocessing** During preprocessing, user-defined configuration parameters are validated and initialized. The BAM file headers are validated against the reference sequence. Information about the chromosomes (including chromosome name, start and end locations of arms) is loaded from the mappability file. The chromosomes that are present in both the mappability file and in the reference sequence (with exactly the same name) are selected for the analysis.
- Coverage A position's coverage is the number of alignments spanning the position in the chromosome. The coverage at every position is computed from the BAM file. By default, all alignments with a quality value of at least 8 are used to calculate coverage. Low quality alignments can be filtered out by modifying the mapping quality threshold parameter (cnv.min.quality).

Optionally, the coverage computed for every chromosome in this step is output in WIG file format. The cnv.coverage.wsize parameter controls the window sizes for this coverage output.

Sampling into The algorithm divides the chromosomal region into windows of variable size, depending upon the mappability of the region. The window sizes are determined dynamically so that exactly the same number of mappable positions are in each window. The program distinguishes between mappable and unmappable positions using the precomputed mappability files.

The coverage mean for every window is computed by taking the average of the coverage values from all the mappable positions in each window. The log ratios between the coverage mean of every window and the expected coverage are computed. The mean of all windows in the whole chromosome arm is used as the expected coverage.

#### **GC** Correction

GC content is the number of G or C bases compared to the total number of bases in a particular region. In the regions of the genome where the percentage of GC content is either high or low, the coverage is observed to be decreased (see Figure 13 on page 197).





Figure 13 Extreme GC contact reduces coverage

In Figure 13, the X-axis is the log ratio (coverage of the window or coverage of the whole chromosome arm). The Y-axis is the percentage of GC content in each window.

The algorithm normalizes this effect of GC contact by scaling the coverage of the windows with extreme GC content to the median coverage. The scaling factors for the windows are computed inline for every chromosome arm during runtime by the algorithm.

**Normalization** The previous section explained how the log ratios computed in the previous step use the mean coverage of the local chromosomal arm as the expected value. However, if the algorithm is used with these settings, it cannot detect large CNV regions spanning more than half the length of the chromosome arm. To detect large CNVs, which can take whole chromosome arms or large portions of chromosome arms, the algorithm performs global normalization. The algorithm computes log ratios by using the median of the coverage means of all chromosome arms as the expected value (see Figure 14).

To skip the normalization step, set the local normalization parameter cnv.local.normalization to 1. By default, the algorithm performs global normalization if the number of valid chromosome arms provided for the analysis is nine or higher. In some cases, you might perform an initial run using Global Normalization, and then perform a second run using local normalization to detect more fine-grained Human CNVs.





**Segmentation** The algorithm uses the Finite First Order Bayesian Hidden Markov Model ("the model") to take the normalized log ratios of the windows as input and convert the continuous log ratio values into discrete copy number states (see Figure 15).

The model consists of 10 states {0, 1, 2, 3, 4,...9}, where each state represents a copy number. For diploid species, state 2 is the normal state, states less than 2 are copy number deletions, and states greater than 2 are copy number amplifications.

The prior probabilities and transition probability matrix for the model are trained using the Baum-Welch EM algorithm.

A copy number state with a p-value is assigned to each window using the Viterbi decoding algorithm. The p-value is a measurement of statistical significance. In general, the lower the p-value, the more likely it is that the prediction will be true.



X-Axis : Window IDs (windows of size of 5000 mappable bases)

Figure 15 Segmentation example



**Post processing** Adjacent windows with the same copy number are merged into a segment.

The algorithm applies filtering criteria on the Human CNV calls according to parameter settings (see Table 31 on page 189 for descriptions of these parameters). The filtering criteria can include parameters defined for minimum mappability, number of continuous blocks with copy number greater than two, and so forth. The algorithm uses the GFF format for structural variants to format the Human CNV segments that pass all of the filtering criteria. See Table 36 on page 194 for a description of the CNV GFF file format.

# FAQ – Human CNVs

1

#### Does the Human CNVs module work on any species?

No. The current version of the module works only for the human hg18 and hg19 reference. The module cannot work on other species because LifeScope<sup>™</sup> Software does not have predicted mappability files for any species other than human.

2

#### How do various parameters affect the sensitivity and specificity of the CNV calls?

The default configuration is designed such that CNV calls are made with a balance between sensitivity and specificity.

To increase the specificity, increase the cnv.window.size value, use only high quality alignments by filtering out the low quality ones using cnv.min.quality, and change the cnv.stringency.setting to High.

To increase the sensitivity, work at higher resolution by using a smaller value for cnv.window.size, by using a lower value for cnv.min.quality, and by changing the cnv.stringency.setting to Low.

#### 3

#### What do the mappability files contain?

The mappability files are essentially a representation of reads from the reference that are mapped back to the reference. The mappability files have one row per every position, indicating whether that position is or is not uniquely mappable. A *mer* is the number of the bases per read. These files are in a binary format (HDF).

#### 4

#### What mappability files are provided?

LifeScope<sup>™</sup> Software provides mappability files for 50.4 fragment mapping.



# 5

#### Can users generate mappability files if they have FASTA files?

LifeScope<sup>™</sup> Software does not provide any applications that allows users to generate their own customized mappability files. Contact your Life Technologies BioInformatics FAS for additional questions.

#### 6

#### When should we change the default value of the cnv.window.size parameter?

The size of Human CNV segments detected by the module is directly dependent on the value of cnv.window.size. Typically, the smallest Human CNV segment that can be detected is at least twice the value of the cnv.window.size setting. The smaller the window size, the smaller the CNV that can be detected. However very small CNVs may be more likely to be false positives (or at least, under-represented in existing public databases).

As the cnv.window.size value decreases, the time taken for the Human CNV analysis and the sizes of the files generated increases significantly.

# 7

#### When should we use the cnv.local.normalization parameter?

To detect smaller Human CNVs (kbp scale) with tumor samples, normalize by the local chromosome context to detect these smaller Human CNVs. If you do not use cnv.local.normalization, large perturbations in ploidy across the whole genome might confuse detection. Ploidy is the number of complete sets of chromosomes in a biological cell. In humans, the somatic cells that compose the body are diploid (containing two complete sets of chromosomes, one set derived from each parent).

The module applies various user-configurable filtering criteria, such as minimum mappability, number of continuous blocks, and so forth on the Human CNV calls.

#### 8

#### How are the p-values for Human CNV segments computed?

The p-values for Human CNV segments are computed using probabilities from the Finite First Order Bayesian Hidden Markov Model (the model). Sequence of log ratios of coverage per window are given as input to the model for segmentation. The module calculates the most probable Human CNV-state, that is, the hidden state, for each window using "Forward-Backward" Algorithm on the model. For example, if window 'W' is assigned Human CNV state 'c' with a probability 'p', then the p-value of that window is '1-p'. The calculations control Human CNV states and p-values for all windows. In the next step, the CNV algorithm merges the neighboring windows with similar copy number states into a Human CNV segment. The p-value of that Human CNV segment is given by the minimum p-value of all merged windows.



#### How do I interpret a CNV with a copy number value of 2?

9

CNVs with a copy number value of 2 can occur in male samples because these samples have only one X chromosome and one Y chromosome. Because the normal copy number is 1 for these chromosomes in males, a CNV with copy number 2 on these chromosomes is a gain in copy number. All CNVs that have a copy number value of 2 are present on either chromosome X or Y, and not on other chromosomes.



**Chapter 10** Run a Human CNVs Analysis *FAQ – Human CNVs* 



# Run an Inversions Analysis

#### This chapter covers:

Overview	203
Examples of running an inversion analysis	204
Inversion module parameters	204
Input files	206
Inversion output files	207
Inversion algorithm	210

### **Overview**

An inversion is defined by its two breakpoints. The numbers of mate pairs supporting occurrence of the starting and ending inversion breakpoints are counted for each base pair. The genomic ranges corresponding to local peaks of these counts, if above a score threshold, are called as candidate breakpoint ranges. To define an inversion, its starting and ending breakpoints are paired up only if they are the reciprocal nearest neighbor of each other in the correct order. The score for the inversion is the harmonic mean of its two breakpoints. Each breakpoint range can be scanned for coverage of normal (AAA) mate pairs to identify a sub-range with the lowest normal mate pair coverage as the most probable location of a breakpoint, and to differentiate homozygous inversions from heterozygous ones. The supporting evidence of all inversions can be visually inspected using a genomic browser.



# Examples of running an inversion analysis

In a standard The following standard workflows provide examples of running the inversion module:

genomic.resequencing.lmp

genomic.resequencing.pe

The following are example shell commands using reads and references from the LifeScope<sup>TM</sup> Software repository.

```
lscope.sh shell -u username -w password
cd /projects
mk ecoli
cd ecoli
mk run1
cd run1
set workflow genomic.resequencing.pe
add xsq run0209a_50_PE.xsq
add xsq run0209b_50_PE.xsq
set reference hg19
# optionally change inversion defaults here
ls
run
```

See for Chapter 7, "Run a Standard Workflow Analysis" on page 97 for more information on running standard workflows. Chapter 6, "Run a Command Shell Analysis" on page 71 for information on shell commands and syntax.

**As a demo analysis** The optional examples download includes an example of how to run the inversion module by itself, as a single analysis that is not part of a standard workflow. The demo example is not recommended for general use.

See Appendix E, "Demo Analyses" on page 507 for information on running the inversion module as a standalone analysis.

# Inversion module parameters

Table 37 describes the parameters that control the inversion algorithm.

In order to change a parameter value in your analysis, use the set param shell command to replace the line # *optionally change inversion defaults here* in the example commands in "In a standard workflow". For instance, to change recover small inversion to on, set the inversion.recover.tiny.inversions parameter to 1 with this command:

set inversion.recover.tiny.inversions 1 /tertiary/inversion.ini

Table 37 Inversion parameter description

Parameter name	Default value	Description		
Optional parameters				



 Table 37 Inversion parameter description (continued)

Parameter name	Default value	Description
library.type	matepair	The library type for alignment data. Use when the BAM LB: header field is not populated. Allowed values: matepair, pairedend.
inversion.calculate.mp.coverage	0	<ul> <li>Whether to calculate normal mate-pair coverage around inversion breakpoints. Allowed values:</li> <li>0: Do not use this calculation.</li> <li>1: Use this calculation.</li> </ul>
inversion.number.of.chromosomes	25	The number of chromosomes used to distribute jobs on a cluster (one job per chromosome). Allowed values: Integers >= 0.
inversion.abx.score	0	<ul> <li>The ABX score. ABX is ABA/ABB/ABC, which are mates with both tags on the correct strands but in reverse order. Allowed values:</li> <li><b>0</b>: Do not use ABX types as candidate inversions.</li> <li><b>1</b>: Use ABX types as candidate inversions.</li> </ul>
inversion.force.update.intermediate. files	0	<ul> <li>Whether to force updating of all intermediate files. Allowed values:</li> <li>0: Do not force updating of all intermediate files.</li> <li>1: Force updating of all intermediate files.</li> </ul>
inversion.down.weight.mp. mismatches	0	<ul> <li>Whether to down-weight mate-pairs with mismatches exponentially.</li> <li>Allowed values:</li> <li><b>0</b>: Do not down-weight mate-pairs with mismatches exponentially</li> <li><b>1</b>: Down-weight mate-pairs with mismatches exponentially.</li> </ul>
inversion.max.bxx.mp.length	3000000	The maximal mapped length of BXX mate-pairs. Allowed values: Integers >= 0.
inversion.max.inversion.length	100,000	The maximum mapped length of inversions. Allowed values: Integers >= 0.
inversion.min.pairing.quality	10	Minimum pairing quality threshold. Allowed values: Integers 0–100.
inversion.score.run.individually	0	<ul> <li>Whether to score every run individually. The parameter is used when rerunning the inversions module with a different breakpoint score threshold, and reusing the inversion evidences and score calculated from previous run. Allowed values:</li> <li>0: Do not score every run individually.</li> <li>1: Score every run individually.</li> </ul>
inversion.breakpoint.rescue	0	<ul> <li>Whether to pair breakpoints with rescue. Allowed values:</li> <li>0: Do not pair breakpoints with rescue</li> <li>1: Pair breakpoints with rescue.</li> </ul>
inversion.recover.tiny.inversions	0	<ul> <li>Whether to recover small inversions. Allowed values:</li> <li><b>0</b>: Do not recover small inversions.</li> <li><b>1</b>: Recover small inversions.</li> </ul>
inversion.max.length.tiny.inversions	0	The maximum mapped length of small inversions. Allowed values: Integers >= 0.
inversion.breakpoint.score.threshold	4	The break point score threshold. Allowed values: Integers 0–100.



#### Table 37 Inversion parameter description (continued)

Parameter name	Default value	Description
inversion.sab.gff.score.threshold	0	The output score threshold. Allowed values: Integers 0–100.
inversion.breakpoint.peak.width	peak.width 100 The break point peak width. Allowed values: Integers 0–100.	
Resource parameters		
memory.request	4gb	Memory request. Include the units gb in the setting.
java.heap.space	3000	Dynamic memory requirement.

Table 37 describes the inversion parameters that we do not recommend changing.

Table 38 Inversion internal parameter description

Parameter name	Default value	Description
inversion.run	1	Whether or not to run the inversion module.
		• <b>0</b> : Do not run the inversions module.
		• 1: Run the inversions module during this analysis.
		The run parameter is set automatically by the shell during a standard workflow. Accept the default for most use.
inversion.max.anchor.mismatch	0	Filter out mates with more anchor mismatches on either tag. Allowed values: Integers 0–10.
inversion.min.alignment.length	0	Filter out mates with shorter alignment length on either tag. Allowed values: Integers 0–100.
inversion.max.alignment.start	0	Filter out mates with farther alignment start position on either tag.
		Allowed values: Integers 0–15.

# Input files

The inversion module takes the BAM file output of the mapping module as input. Because the inversion module specifically selects for pairs that are in the incorrect orientation, it is important to know the correct orientation. As a result, either the library type information in the LB field or the library.type parameter must be properly set.

Libraries in the input file or files must be either:

- All mate-pair
- All paired-end

When different library types are provided as input on the same run, the inversion module fails with an error.

#### **Read lengths**

Input BAM files may contain multiple read lengths.



# Inversion output files

Table 39 describes t	the inversion	GFF output	file format.
----------------------	---------------	------------	--------------

 Table 39
 The inversion output file format

Column title or number	Description	Example
1	Chromosome.	chr10
2	Method.	AB_SOLID
3	Feature keywords.	inversion
4	Inversion start coordinate.	46443097
5	Inversion end coordinate.	46479578
6	Inversion score.	129.8
7	Not used.	_
8	Not used.	_
9 Attributes, see below:		
left <sup>‡</sup>	Starting breakpoint range.	left=chr10:46443097-46443161
right	Ending breakpoint range.	right=chr10:46479540-46479578
leftscore	Left breakpoint range score.	leftscore=185
rightscore	Right breakpoint range score.	rightscore=100
count_AAA_further_left	The number of AAA mate-pairs spanning the genomic region to the immediate left of the starting breakpoint range.	count_AAA_further_left=1
count_AAA_left	The number of AAA mate-pairs spanning the starting breakpoint range.	count_AAA_left=1
count_AAA_right	The number of AAA mate-pairs spanning the ending breakpoint range.	count_AAA_right=2
count_AAA_further_right	The number of AAA mate-pairs spanning the genomic region to the immediate right of the ending breakpoint range.	count_AAA_further_right=1
left_min_count_AAA	Sub-range within starting breakpoint range that has the minimal AAA coverage.	left_min_count_AAA=chr10:46443 097-46443112
count_AAA_min_left	Minimal AAA coverage in starting breakpoint range.	count_AAA_min_left=0
count_AAA_max_left	Maximal AAA coverage in starting breakpoint range.	count_AAA_max_left=4
right_min_count_AAA	Sub-range within ending breakpoint range that has the minimal AAA coverage.	right_min_count_AAA=chr10:4647 9576-46479578
count_AAA_min_right	Minimal AAA coverage in ending breakpoint range.	count_AAA_min_right=4
count_AAA_max_right	Maximal AAA coverage in ending breakpoint range.	count_AAA_max_right=11
homozygous	Whether AAA coverage at both breakpoints ranges are lower than one-fifth of their neighboring ranges.	homozygous=YES

‡ This and the remaining entries are allowed values for Attributes in column 9.

The following is an example of the file inversions.s2.w100.100000.GFF, whose format is described in Table 43 on page 209:

```
##gff-version 3
##generated by SOLiD inversion tool
chr10 AB_SOLiD
                  inversion
                              46442583
                                           46479737
                                                        4
left=chr10:46442583-46443522;right=chr10:46479431-
46479737; leftscore=6.0; rightscore=3.0; count_AAA_further_left=0; count_AAA_left
=0;count_AAA_right=0;count_AAA_further_right=0;left_min_count_AAA=chr10:46442
583-
46443522;count_AAA_min_left=0;count_AAA_max_left=0;right_min_count_AAA=chr10:
46479431-46479737; count_AAA_min_right=0; count_AAA_max_right=0; homozygous=YES
chr21 AB SOLiD
                  inversion
                              26295366
                                           26297167
                                                       2.7
left=chr21:26295366-26296041;right=chr21:26296491-
26297167; leftscore=2.7; rightscore=2.7; count_AAA_further_left=0; count_AAA_left
=0;count_AAA_right=0;count_AAA_further_right=0;left_min_count_AAA=chr21:26295
366 -
26296041; count_AAA_min_left=0; count_AAA_max_left=0; right_min_count_AAA=chr21:
26296491-26297167;count_AAA_min_right=0;count_AAA_max_right=0;homozygous=YES
chr6 AB_SOLiD
                  inversion
                              168834702
                                           168837413
                                                       2.1
                                                              .
left=chr6:168834702-168835567;right=chr6:168836564-
168837413;leftscore=2.2;rightscore=2.1;count_AAA_further_left=0;count_AAA_lef
t=0;count_AAA_right=0;count_AAA_further_right=0;left_min_count_AAA=chr6:16883
4702-
168835567; count_AAA_min_left=0; count_AAA_max_left=0; right_min_count_AAA=chr6:
168836564-
168837413; count AAA min right=0; count AAA max right=0; homozygous=YES
```

Column	Description	Example
1	Display label. For internal use.	_
2	GFF name, type of breakpoint.	InvStart
3	GFF type.	exon
4	Range start coordinate.	1297
5	Range end coordinate.	1362
6	Breakpoint range score, number of supporting mate- pairs.	1
7	Strand.	-
8	-	_
9	-	g=.1
10	Chromosome.	chr1

Table 40 Inversion all.chr, all.chr\*, pair.orphan, pair.txt file format description

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Column	Description	Example
1	Chromosome.	chr1
2	Inversion start coordinate.	1632874
3	Inversion end coordinate.	1706252
4	Inversion score.	4.8
5	Inversion length.	73379
6	Starting breakpoint range.	chr1:1632874-1633166
7	Ending breakpoint range.	chr1:1705726-1706252
8	Left breakpoint range score.	6.0
9	Right breakpoint range score.	4.0

#### Table 41 inversion rank.txt file format description

Table 42 Inversion AAA GFF file format description

Column	Description	Example
1	Mate-pair bead ID.	1_11_215_288
2	GFF name.	LEFT
3	GFF type.	exon
4	Mate-pair start coordinate.	3971
5	Mate-pair end coordinate.	4020
6	Score, number of mismatches.	1
7	Strand.	+
8	-	-
9	-	g=

 Table 43
 Inversion coords.inversions.s\*.\* and coords.orphan.\* file format description

Column	Description	Example
1	The genomic coordinate of an inversion.	chr10:46443097-46479578
2	Inversion score.	129.8

The following is an example of the file coords.inversions.s2.w100.100000-, whose format is described in Table 43:

chr10:46442583-46479737	4
chr21:26295366-26297167	2.7
chr6:168834702-168837413	2.1



Column	Description	Example
1	Chromosome.	chr10
2	Inversion start coordinate.	46443097
3	Inversion end coordinate.	46479578
4	Inversion score.	129.8
5	Inversion length.	36482
6	Starting breakpoint range.	chr10:46443097-46443161
7	Ending breakpoint range.	chr10:46479540-46479578
8	Left breakpoint range score.	185
9	Right breakpoint range score.	100
10	The number of AAA mate-pairs spanning the genomic region to the immediate left of the starting breakpoint range.	1
11	The number of AAA mate-pairs spanning the starting breakpoint range.	1
12	The number of AAA mate-pairs spanning the ending breakpoint range.	2
13	The number of AAA mate-pairs spanning the genomic region to the immediate right of the ending breakpoint range.	1

#### Table 44 Inversion AAA/\*.txt and AAA.txt file format description

#### Table 45 Inversion rescore.txt file format description

Column	Description	Example
1-13	Same as AAA.txt	See Table 44.
14	Sub-range within starting breakpoint range that has the minimal AAA coverage.	chr10:46443097-46443112
15	Minimal AAA coverage in starting breakpoint range.	0
16	Maximal AAA coverage in starting breakpoint range.	4
17	Sub-range within ending breakpoint range that has the minimal AAA coverage.	chr10:46479576-46479578
18	Minimal AAA coverage in ending breakpoint range.	4
19	Maximal AAA coverage in ending breakpoint range.	11

# Inversion algorithm

**Overview** 

The inversion module exploits the large insert size of SOLiD<sup>TM</sup> System mate-pair libraries to detect inversion polymorphisms that are important but often poorly characterized. The large insert size is critical for spanning the repeat regions that are associated with inversion break points.



In its simplest form, the algorithm collects evidence for an inversion by observing accumulations of pairs with correct relative positioning, but with an improper orientation. For SOLiD<sup>™</sup> System mate-pair libraries, this algorithm requires tags to be mapped to opposite strands. For example, if the R3 tag is to the left of an F3 tag, but the R3 tag maps to the top strand and the F3 tag maps to the bottom strand, this pair provides inversion evidence, specifically that the R3 tag is to the left of the inversion starting break point and that the F3 tag is to the right of the break point. Once the evidence is collected, candidate inversion break points are scored, paired, and ranked. Additional evidence is provided by a scan for drops in the coverage by normal matepairs (see Figure 16).



Figure 16 Inversions and break point ranges

T: 47	1	.1	C 11	•	1 .
HIGHING 16	domicto	tho	tollo	MATIN CT	alamonte
Figure to	Gerneis	THE	111111	100 1119	elements.
1 1 2 2 2 2 2 2	e.ep.1000			· · · · · · · · · · · · · · · · · · ·	010111011001

Thick black line	The sequenced genome.
Thin black lines	Mate-pairs.
Red and green arrows	Two ends of an inverted mate-pair.
Blue and orange bars	The maximum distance separating the two ends of a normal mate-pair (AAA mates).

By definition, an inversion has two breakpoints: a starting breakpoint and an ending breakpoint. The inversion module plug-in uses SOLiD<sup>™</sup> System mate files or matepair BAM files as input (using paired-end BAM files is not recommended due to their small insert size). The number of mate-pairs supporting an occurrence (of both starting and ending inversion breakpoints) are counted for each base pair as breakpoint scores. Candidate breakpoint ranges are genomic ranges corresponding to local peaks of counts above a predetermined score threshold. The clone insert size constraints specify a minimum and a maximum (blue and orange horizontal bars) distance separating the

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	two ends (small green or red arrows) of a mate-pair (thin black lines) in the sequenced genome (thick black line). Each green mate-pair suggests a starting breakpoint of an inversion occurring to the left, and an ending breakpoint between its two tags. The green pairs then define the range of the starting breakpoint by contributing positive counts to all base pairs to the left of their left tags within 1 max. The green pairs also help refine the ending breakpoint range by contributing negative counts to all base pairs 1 max to the left and 1 max to the right. The red mate-pairs contribute counts in a similar fashion.
	When small inversion detection is enabled (with the parameter key inversions.recover.tiny.inversions), a second pass of the algorithm is performed, but with a smaller window for the scoring function, allowing the detection of inversions of 200 base pairs, or even lower, depending on coverage.
	Like other SOLiD <sup>™</sup> System modules, the inversion module outputs results in GFF format for easy viewing in common genome browsers. Because of the inherently ambiguous nature of the detection, the start and end locations represent the widest possible range. More details are provided by the GFF attributes, including the break point ranges and scores.
	While the same algorithm can be applied to SOLiD <sup>™</sup> System paired-end libraries, in practice, the smaller insert size makes inversions more difficult to detect. Without the large insert size of mate-pair libraries, both tags would place into repeat regions that typically flank inversions, making tag placement ambiguous.
Input data	The first step in the detection of inversions is the collection of inverted mates. These are filtered from the BAM file by selecting for mates of opposite orientation. In pairing category terms, this filtering includes BA*, BB*, and AB*. These records are written to an intermediate directory, so that the filtered set can be reused. It is important that either the library.type parameter be set or that the LB field be properly constructed to indicate a mate-pair library.
	Non-redundant mates are selected from BAM file input using the PCR duplicates flag (0x0400). Pairing quality values can be used to select unique records via the inversion.min.pairing.quality parameter key. Values greater than 20 should be unique, but criteria may be different for different applications.
	The Pairing "SV" output filter is specifically designed to capture more of the deviant pairs used for structural variant programs such as the inversion module. A lower inversion.min.qv value (~10) and a BAM file generated with the "SV" output filter can potentially find a larger number of candidate inversions.
Workflow	Figure 17 shows the inversion module workflow. The inversion module begins with a long mate-pair BAM file and proceeds through scoring, pairing, ranking, rescoring, and GFF output.





Figure 17 Inversion module algorithm workflow

Scoring	Candidate breakpoints are scored by first determining the tag of a pair that is properly oriented. The inverted mate of the pair is then used to select reference locations that are in the vicinity of the inversion breakpoint. All reference locations that are between the inverted mate and the maximum possible insert size accumulate a positive score, resulting in a list of high scoring locations that represent candidate breakpoints.
Pairing	The high scoring candidate breakpoints are then recombined into inversion candidates by matching start and end locations that are defined by an analysis of the high scoring locations. A user-definable window size (inversion.breakpoint.peak.width) is used to determine regions with a peak score greater than the threshold (inversion.breakpoint.score.threshold). These peaks are combined with their reciprocal nearest neighbor to create a set of possible inversions.
	In some cases there will be a peak whose nearest neighbor is slightly below the threshold. A rescue analysis will retrieve those that are significant peaks, but below the threshold cutoff. This is controlled by the inversion.breakpoint.rescue flag.
	A correct threshold for inversion breakpoint scoring depends on clone coverage of sequencing data, its nature, and its mapping quality. The default value 4 is set for 100–500X clone coverage of sequencing from a normal genome. As clone coverage increases, the inversion score threshold must be raised to filter out mismapping noise. Because more inversions can be expected in cancer samples, the threshold may be lowered when running on those samples.



Ranking	Inversions are ranked based on the harmonic mean of the scores of the start and end breakpoints. Additionally, inversions are separated out by the user specified maximum inversion length (inversion.max.inversion.length).
Inversion length thresholds	Table 46 describes the how the parameters inversion.max.inversion.length and inversion.max.bxx.mp.length affect inversion calling. The parameter inversion.max.inversion.length also affects the output file name, according to these patterns:
	<ul> <li>coords.inversions.s4.w100.<max.inversion.length>-</max.inversion.length></li> </ul>

• coords.inversions.s4.w100.<max.inversion.length>+

The example output file names in this table are based on the parameter setting inversion.max.inversion.length=100000.

	Length	Inversion calling behavior
	From 1 to inversion.max.inversion.length	Inversions are called, into an output file named, for example, coords.inversions.s4.w100.100000
		Note: The minus sign ("-") is part of the file name.
	From inversion.max.inversion.length to inversion.max.bxx.mp.length	Inversions are called, into an output file named, for example, coords.inversions.s4.w100.100000+.
		Note: The plus sign ("+") is part of the file name.
	Greater then inversion.max.bxx.mp.length	Inversions are not called.
Tiny inversions	If the tiny inversions flag is set (inve pairing and ranking components are small to detect inversions with a smal inversion.max.length.tiny.ir	rsion.recover.tiny.inversions), the rerun with a window size that is sufficiently ler size (controlled by the parameter nversions).

 Table 46
 Inversion threshold parameters

Normal pair coverage Additional evidence is provided for breakpoints by examining the relative coverage of unique proper pairs. These pairs are selected from the BAM input using the proper pair flag (0x0002), and they are used to reduce the score of reference positions covered by proper pairs.



# Run a Small Indels Analysis

#### This chapter covers:

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### **Overview**

Small indel detection is a tertiary module in LifeScope<sup>™</sup> Genomic Analysis Software. Data in BAM format from the fragment and paired mapping modules serve as direct inputs for small indel discovery. Gap alignments, those with a CIGAR string containing D or I, serve as the primary evidence for this. In the second.map part of the mapping module, these evidences are found by taking partial read anchor alignments, and then either extending allowing for a gap, called LOCAL alignment, or as seed placement for a full read, called GLOBAL, very similar to what is done in the pairing phase. Gap alignments are also found separately in the pairing phase of the mapping module. Here, these evidences are found in mate pairs by using the placement of one tag such that the other tag can be used to search for the presence of a small indel. Specifically, with the unplaced tags, the algorithm finds all possible placements of the beginning and ends of the read. Then, these are joined together only if the resulting gap size is within the limits set in the pairing module. All successful joinings form the set of gap alignments.

The LifeScope<sup>™</sup> Software small indel module allows flexible processing of these gap alignments. LifeScope<sup>™</sup> Software takes input gap alignments from one or more input BAM files and collects them (based on their proximity from each other) to form pileups. The module, in the process of making the pileups or in evaluating these pileups, employs a number of heuristics, including map QV filtering, insertion sequence determination, and assessment of zygosity using ungapped alignment read counting and clustering of gap sizes. In the end, the small indel module produces quality indel calls in a public GFF version 3 format.



# Examples of running a small indels analysis

In a standardThe following standard workflows provide examples of running the small indelsworkflowmodule:

- genomic.resequencing.frag
- genomic.resequencing.lmp
- genomic.resequencing.pe
- targeted.resequencing.frag
- targeted.resequencing.pe

The following are example shell commands using reads and references from the LifeScope<sup>™</sup> Software repository.

```
# log into the shell
lscope.sh shell -u username -w password
# cd to the projects repository
cd /projects
# make a project, and open it
mk proj1
cd proj1
# make an analysis, and open it
mk run1
cd run1
# define the analysis type
set workflow genomic.resequencing.pe
# define the input (this example uses dummy XSQ filenames)
add xsq run0209a_50_PE.xsq
add xsg run0209b_50_PE.xsg
# specify the reference to be used
set reference hg19
# optionally change parameter defaults here
# list the analysis configuration
ls
# run the analysis
run
# list progress information for the analysis
ls
In order to change a parameter value in your analysis, add a set param shell
command to after the line # optionally change parameter defaults here.
For instance, to change the filtering parameter, use this shell command:
set small.indel.perform.filtering 0 /tertiary/small.indel.ini
See for Chapter 7, "Run a Standard Workflow Analysis" on page 97 for more
information on running standard workflows. Chapter 6, "Run a Command Shell
Analysis" on page 71 for information on shell commands and syntax.
```

# **As a demo analysis** The optional examples download includes an example of how to run the small indel module by itself, as a single analysis that is not part of a standard workflow. The demo example is not recommended for general use.
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See Appendix E, "Demo Analyses" on page 507 for information on running the small indel module as a standalone analysis.

# Small indel parameter description

Table 47 describes the small indel module parameters.

Table 47 Small indel module file parameters

Parameter name	Default value	Description	
Optional input parameters			
analysis.regions.file	-	BED format file indicating the target regions for the analysis. Used for targeted analysis.	
		In targeted resequencing workflows, this parameter is set in the both secondary and tertiary global.ini files.	
General options	ł		
small.indel.detail.level	3	For BAM file inputs, the level of detail in output:	
		• <b>0</b> : Keeps only position information about the anchor read and no information for the ungapped alignment.	
		• <b>1–8:</b> Keeps only some of the alignment's anchor alignment but none of the ungapped alignment.	
		• 9: Is most detailed, but also the slowest.	
small.indel.zygosity.profile.name	max-mapping	Zygosity profile name.	
		• <b>classic</b> : Profile for classically (full read) mapped reads.	
		• <b>max-mapping</b> : Profile for seed and extend ungapped alignments.	
		• max-mapping-v2: Reserved for future use.	
		• <b>gap-align-only</b> , and <b>no-calls</b> : Force all zygosity calls to be homozygous calls.	
small.indel.genomic.region	_	Names a specific genomic region to be selected from the BAM file. Only full chromosomes are guaranteed not to alter results. Specifying partial chromosomes is allowed but may result in the loss of indels near the edges of that region. For example, chr1:2945-9659 causes a reduction of coverage for approximately a read length after position 2945.	
small.indel.display.base.qvs	0	Whether or not to display base quality value scores in the GFF file. Allowed values:	
		<b>0</b> : Do not add QV scores in the output file.	
		<b>1</b> : Displays the FASTQ base QV scores for all of the reads used for each indel in the GFF file. FASTQ strings contain semi-colons, so adding these strings may produce a GFF file that is not compatible with certain applications.	

Parameter name	Default value	Description
small.indel.num.aligns.per.pileup	1000	For pileups with more than this number of alignments, sets the expected number of alignments per pileup.See FAQ about this setting regarding rare variant detection. Values much higher than 1000 may result in a significant increase in computational time.
		Allowed values: Integers >= 0.
small.indel.random.seed	94404	The random seed value used to determine which pseudo random set of reads to use when there are greater than 1000 reads in a pileup. The random number generator used is the Mersenne Twister MT19937 algorithm.
		Allowed values: Integers >= 0.
Pileup options		
small.indel.min.num.evid	2	Minimum number of evidences required for an indel call. This parameter does not have an upper limit, but a value higher than the average coverage level in most cases causes a significant reduction in sensitivity.
small.indel.max.num.evid	-1	Maximum number of evidences. Use -1 for no maximum. Setting this value to some multiple of the average coverage could remove indels found in abnormally high coverage areas.
small.indel.consGroup	1	Indel grouping method. Allowed values:
		1: Conservative grouping of indels with 5bp max between consecutive evidences.
		<b>2</b> : Lax grouping. Groups indels that are at maximum the higher of 15 or 7 times the indel size.
		<b>9</b> : No grouping. Makes every indel evidence a separate pileup.
Mapping quality filtering	1	·
small.indel.max.reported.alignments	-1	Only uses those alignments where the NH field (the number of reported alignments; from the BAM record) is this value or lower. A value of $-1$ is to have no upper limit. The range where this is effective depends on the input BAM file's range of values of the NH tag.
small.indel.min.mapping.quality	8	Keeps only reads that have this or higher pairing qualities. For paired tags, mapping quality is for the pair (pairing quality), and for fragment, it is the single tag's map quality.
		Reads that are lower than this value are filtered out. Allowed values: Integers 0–100.
small.indel.min.best.mapping.quality	10	For a particular indel called with a set of reads, at least one pairing quality in this set must be higher than this value. Allows for supporting evidences to have a lower mapping quality threshold then the best read. Allowed values: Integers 0–100.
small.indel.min.anchor.mapping. quality	-1	Minimum mapping quality for a non-indel (anchor) tag. Effective only for paired reads, for the number of anchors queried as defined by small.indel.detail.level.

1	2
	<u> </u>

Parameter name	Default value	Description
small.indel.ungapped.bam.flag.filter	ProperPair,Primary	<ul> <li>For ungapped alignments, specifies the BAM flag properties that a read must have to be included. Allowed values: A comma-separated string of one or more of these values:</li> <li>ProperPair</li> <li>UniqueHit (not a BAM flag; requires BAM NH=1)</li> <li>NoOptDup</li> <li>Primary</li> </ul>
		None
		If None is present in the string, then all filters, even those explicitly set, are turned off.
small.indel.gapped.bam.flag.filter	Primary	<ul> <li>For gapped alignments, specifies the BAM flag properties that a read must have to be included. Allowed values: A comma-separated string of one or more of these values:</li> <li>ProperPair</li> <li>UniqueHit (not a BAM flag; requires BAM NH=1)</li> <li>NoOptDup</li> <li>Primary</li> <li>None: All filters, even those explicitly set, are turned off. If None is present in the string, then all filters, even those explicitly set.</li> </ul>
	0	Con clier result that do not have this minimum longth on
small.indel.edge.length.deletions	U	either side of the indel will not be considered. Allowed values: Integers >= 0.
small.indel.edge.length.insertions	0	Gap alignments that do not have this minimum length on either side of the indel will not be considered. Allowed values: Integers >= 0.
Heuristic filtering		
small.indel.perform.filtering	1	Whether or not to perform filtering in each pileup. Allowed values:
		• <b>0</b> : Do not perform filtering on pileups.
		• 1: Perform heuristic filtering on pileups.
		Parameters that change the makeup of pileups contained in the PAS.SUM file, such as small.indel.min.num.evid are still active.

Parameter name	Default value	Description
small.indel.indel.size.distribution. allowed	can-cluster	Indel sizes in a pileup are allowed to have certain indel size distributions.
		Allowed values:
		• <b>similar-size</b> : 75% of the reads of a pileup must have exactly the same size.
		• <b>similar-size-any-large-deletions</b> : Any pileups with at least 2 large deletion alignments, the other pileups must have similar sizes.
		<ul> <li>can-cluster: Allowed if at least one cluster of any indel size is found. [Default].</li> </ul>
		• <b>can-cluster-any-large-deletions</b> : Any pileups with at least 2 large deletion alignments; other pileups must be able to cluster (will have indels with two more reads with larger deletions, even if they don't form good clusters).
		<ul> <li>any: Can have any size distribution (same as small.indel.require.called.indel.size=false).</li> </ul>
small.indel.remove.singletons	1	Whether or not to remove the singletons that occur when different alignment methods are combined based on identical bead ids and read sequence. Allowed values:
		• <b>0</b> : Do not remove the singletons.
		• 1: Remove singletons.
small.indel.alignment.compatibility. filter	1	Alignment compatibility level. Checks color space compatibility around the gap. Allowed values:
		• <b>0</b> : No alignment compatibility filtering.
		• 1: The small indel module determines whether the data contains base-space or color-space sequence.
		• 2: Force the use of base-space sequence, if present.
		• <b>3</b> : Force the use of color-space sequence, if present
small.indel.max.coverage.ratio	12	Maximum allowed value for the ratio of number of reference alignments allowed by the ungapped.bam.flag.filter over the number of non-redundant indel variant reads selected with the gapped.bam.flag.filter.
		Use -1 for no limit (no coverage ratio filtering).
		Allowed values: Integers.
small.indel.max.nonreds-4filt	2	Maximum number of non-redundant reads where read position filtering is applied. Allowed values: Integers.
small.indel.min.from.end.pos	9.1	Minimum average number of base pairs from the end of the read required of the pileup, when there are at most a certain number of reads defined by small.indel.max.nonreds-4filt. Allowed values: Floats.
Indel size filtering	1	
small.indel.min.insertion.size	0	Minimum insertion size to include. Allowed values: Integers.

Parameter name	Default value	Description
small.indel.min.deletion.size	0	Minimum deletion size to include. Allowed values: Integers.
small.indel.max.insertion.size	100000000	Maximum insertion size to include. Allowed values: Integers.
small.indel.max.deletion.size	100000000	Maximum deletion size to include. Allowed values: Integers.
Resource parameters		
memory.request	4gb	Memory request. Include the units gb in the setting.
java.heap.space	30000	Dynamic memory requirement.
processors.per.node	1	On a node, the number of parallel threads used to run this process.
wall.time	336	The maximum total time in hours for the process to complete. The default is equivalent to 14 days.

## Small indel module output file formats

The main output file of this module is the produced GFF\_3 file. This list describes the other file types generated by the small indel module:

- **.pas.sum** An internal pileup format which represents collections of gap alignments extracted from the BAM file.
- **.align** Displays the alignments of reads of the gaps in a human readable format.
- **.ungapped** For each pileup, contains the list of beads ids from reads that are aligned without gaps but also span the location of the indel.
- .pas A legacy file produced but not used by the caller.

#### **Small indel GFF** format As shown in the small indel output file example below (after Table 48), the GFF file created by the small indel module begins with the General File Format Version 3 headers. The section following the GFF\_3 headers displays BAM header and read group information. The lines containing information about each indel follows. Table 48 describes each column and attribute contained in the file. Table 49 on page 227 describes the text format, which contains similar information as the GFF file.

 Table 48
 Small indel GFF file format descriptions

Column/Attribute tag name	Description	Example
seqid	The ID of the sequence to which the start and end coordinates refer, such as a chromosome number.	chrV
source	Free-text qualifier that indicates the algorithm or method that generated the feature.	AB_SOLiD Small Indel Tool



Table 48	Small ir	ndel GFF	file	format	descriptions	(continued)
						(,

Column/Attribute tag name	Description	Example	
type	SOFA feature. For indels, possible values are:	deletion	
	insertion_site		
	deletion		
	combination		
start	Start position of the feature.	200587060	
end	End position of the feature.	200587063	
	1-based integer coordinates of the feature, relative to the sequence in column 1. For zero-length features, such as insertion sites, "start" equals "end" and the implied site is to the right of the indicated base in the direction of the landmark. For deletions, the start and end indicate the positions in the reference that are not present in the sample.		
score	Floating-point value representing the quality of the evidence for the feature.	1	
	"Score" is not currently calculated, so all indels are reported with a score of 1.		
strand	The strand of the feature. The type of indels detected are not stranded, because they are found with sequence reads that are on either or both strands.		
phase	Translation frame; relevant only for CDS features.		
Attributes			
ID	Unique indel id. Non-sequential ids are due to filtering.	21	
ins_len	Number of bases inserted relative to the reference.	7	
del_len	Number of bases missing from the reference.	2	
len	The size of the indel, if both an insertion and a deletion are detected. Deletions are indicated by a negative value.	7, -2	
clustered-indel-sizes	The average indel size of each cluster rounded to the nearest integer. Deletions are indicated by a negative value.	4, -298	
reference	The reference sequence at the position of the allele call. An event that is an insertion and is not a substitution, is indicated with a	ag	
allele-call	The allele sequences detected. The reference sequence is only present if sufficient reads span the first breakpoint of the indel. The string ag/- indicates a hemizygous deletion where one allele is the reference, while -/- indicates a homozygous deletion. A third allele is possible if sufficient reads indicate its presence.	ag/	

Column/Attribute tag name	Description	Example
context-pos	Position of the first base of the sequences reported with additional surrounding sequence context. If the first allele is missing, it is the position immediately before the insertion.	713660
context-reference- seq	Reference sequence at the context position.	acagagagaag
(ref part of alleles in v1.3)		
context-variant-seq	A complete list of non-reference variant sequences	aCAGAGAAG/ acagagagagaag
(var part of alleles in 1.3)	found by all gap alignments in the pileup. Using the ambiguity information from the aligner (XA tag), this contains the context around the indel representing a possible short tandem repeat. "NO_CALL" indicates reads where a misalignment is likely (that is, color- space alignment is not in agreement with base- space alignment).	
reference-reads	Number of reads detected to span the first indel breakpoint location by 5bp.	5
context-variant- reads (was allele-counts in v1.3)	The number of indel reads found for each of the above variants. If the number of gap alignments found is greater than the parameter small.indel.num.aligns.per.pileup, then an estimated counts is displayed followed by the read counts of the sample in parentheses.	3,1 15464,1497(899,87)
gap-total-reads	Total number of gap alignments in the pileup.	4
gap-nonred-reads	Total number of reads that are not marked as PCR duplicates in the BAM file.	4
tight_chrom_pos	Conservative estimate of the chromosome start position range of the feature.	713662–713667
loose_chrom_pos	Estimate of the maximum chromosome start position range of the feature.	713662–713667
no_nonred_reads	Number of reads with unique start positions (non-redundant reads).	3
coverage_ratio	Clipped ungapped coverage divided by the number of non-redundant gap alignment reads.	1.6667
	Clipped coverage means the last 5 bp at either end are not counted as a part of coverage.	

 Table 48 Small indel GFF file format descriptions (continued)



#### Table 48 Small indel GFF file format descriptions (continued)

Column/Attribute tag name	Description	Example
zygosity	Experimental zygosity call. Allowed values:	HEMIZYGOUS-REF
(v1.3 name: experimental- zygosity)	• <b>HEMIZYGOUS-RE</b> F: enough ungapped alignments overlap the indel break point to indicate that the reference allele is present. Also, all the indels have the same size and the same sequence.	
	• <b>HOMOZYGOUS-NON-REF</b> : occurs when there is no indication that the reference allele is present. Also, all the indels have the same size and the same sequence.	
	• <b>HEMIZYGOUS-REF-MULTI-INDEL-ALLELE</b> : occurs if the reference is detected, the variant has a single clustered indel size, but multiple variant sequences are detected.	
	• HETEROZYGOUS-NON-REF: occurs if the reference is not detected, the variant has a single clustered indel size, but multiple variant sequences are detected.	
	• <b>MULTI-HEMIZYGOUS-REF</b> : occurs if the reference, multiple clustered indel sizes, and multiple variant sequences are all detected.	
	• <b>HEMIZYGOUS-NON-REF</b> : occurs if the reference is not detected, but multiple clustered indel sizes and multiple variant sequences are.	
zygosity-score (v1.3 name: experimental- zygosity-score)	Experimental zygosity score. The range is 0–1. The closer to 1, the more likely the zygosity is hemizygous with the reference allele, and the closer to 0, the less likely that it is.	0.9656
rg_id_nums (was contained in run_names in v1.3)	The list of runIdNum's that correspond to the ##@HD header line of the GFF file. This information indicates which input BAM file the particular read came from.	1,1,2,1
bead_ids	Bead IDs for each read. The run_names and all the comma-separated attributes below are in the same order as these.	984_536_1054, 1431_2007_1567, 116_364_1582
overall_qvs	Alignment quality values for each bead.	26,47,66
no_mismatches	List of number of mismatches for each read.	-1,-1,-1
read_pos	Position in each read at which there was a gap in the alignment.	20,17,10
from_end_pos	Same as above, except that the value is the number of base pairs from the end of the read.	30,33,40
strands	Strand for each read.	+,+,+
tags	Tags where the indel was found. Possible values are F3, R3, and FRAG.	F3,F3,F3
indel_sizes	List of indel sizes found for each evidence.	-2,-2,-2

1	5
	L

Column/Attribute tag name	Description	Example
non_indel_no_ mismatches	Number of mismatches of the other tag that was matched without a gap. Values of NIL occur if that particular bead-id is from a fragment analysis.	1,3,3
unmatched-lengths	For a particular bead-id, the length of the read that was left unmatched (equal to the read length if no ungapped match was found). This read length is relevant in extended read alignments.	50,50,50
ave-unmatched	Average of the unmatched lengths.	50
anchor-match- lengths	Anchor tag's match lengths in extended read alignments. If the detail level is not set to 9, some of these values may be reported as 99.	49,44,49
ave-anchor-length	Average of the anchor match lengths.	47.3333
read_seqs	The read sequence where the indel was found for each bead.	T302000, T120320, T232132
base_qvs	The color call QVs for the read sequence where the indel was found. This value is not currently displayed.	_
non_indel_seqs	Sequences of the non-indel anchor tag.	G100220, G313000, G203330
non_indel_qvs	The color call QVs for the non-indel anchor tag sequence. This value is not currently displayed.	_

Table 48	Small in	del GEE	file form	at description	ons (continued)
	Jinattin		nice for fin	at description	JIIS (CONTINUEU)

# **Example output file** A composite example of a small indel output file is shown below. The file contents are as follows:

- 1. The general GFF version 3 headers.
- 2. The BAM header and read group information.
- 3. Lines containing information about each indel call.

Table 48 on page 221 describes each tag of the attributes column.

```
##gff-version 3
##solid-gff-version 0.3
##source-version LifeScope(tm) 2.0 SmallIndel
```

#### Header Example:

##other-source-version small-indel-tool.pl/splitread-pileup/process-small-indels v 1.4.0, 2011-03-11 17:23:21 ##type DNA ##date 2011-04-09 ##time 21:16:33 ##feature-ontology http://song.cvs.sourceforge.net/\*checkout\*/song/ontology/ sofa.obo?revision=1.141 ##options reference=/reference/human\_hg18.fa output.dir=. output.prefix=baseFilename bam.file=output/bam/bamA.bam,output/bam/bamB.bam,output/bam/bamC.bam ##reference /reference/human\_hg18.fa ##input-files output/bam/bamA.bam,output/bam/bamB.bam,output/bam/bamC.bam



```
##run-path /path/of/ini/ini-filename/
##genomic-region chr1
##Filter-settings: max-ave-read-pos=none,min-ave-from-end-pos=9.1,max-nonreds-4filt=2,min-
insertion-size=none,min-deletion-size=none,max-insertion-size=none,max-deletion-
size=none, indel-size-distribution-allowed=CAN-CLUSTER, max-coverage-ratio=12, min-mapping-
quality=none,min-best-mapping-quality=5,remove-singletons?=T
##BAM header:
##CODE_VERSION splitread-pileup 1.4.0
##BAM-input='output/bam/bamA.bam' bam_file_num:1
##@HD VN:1.0
               SO:2
                                ID:20100629040351402
##@RG rg_id_num:1
                        LT:MP
                                                        SM:HuRef
                                                                        LB:50x50MP
PU:bioscope-pairing
                      PI:1645 DT:2010-06-28T21:03:51-0700
                                                                PL:SOLiD
              LN:247249719
                                UR:file:/reference/human_hg18.fa
##@SQ SN:chr1
. . .
##@SQ SN:chrM LN:16571
                                UR:file:/reference/human_hg18.fa
##BAM-input='output/bam/bamB.bam' bam_file_num:2
##@HD_ VN:1.0 SO:2
##@RG_ rg_id_num:2
                        LT:MP
                                ID:20100617182840485
                                                        SM:HuRef
                                                                        LB:50x50MP
PU:bioscope-pairing
                        PI:1575 DT:2010-06-17T11:28:40-0700
                                                                PL:SOLiD
##@SQ SN:chr1 LN:247249719
                             UR:file:/reference/human_hg18.fa
. . .
##@SQ SN:chrM LN:16571
                                UR:file:/reference/human_hg18.fa
##BAM-input='output/bam/bamC.bam' bam_file_num:3
##@HD____ VN:1.0 SO:2
##@RG____rg_id_num:3
                                ID:20100627170940231
                        LT:MP
                                                        SM:HuRef
                                                                        LB:50x50MP
                        PI:1645 DT:2010-06-27T10:09:40-0700
                                                                PL:SOLiD
PU:bioscope-pairing
##@SQ SN:chr1 LN:247249719
                              UR:file:/reference/human_hg18.fa
. . .
##@SQ SN:chrM LN:16571
                                UR:file:/reference/human_hg18.fa
##Annotation Apr 13, 2011; LifeScope_resources//lifetech/hg18/GTF/refGene.20090513.gtf;
LifeScope_resources/lifetech/hg18/dbSNP/dbSNP_b130.tab
##dbSNP function codes: Locus region = 1; Coding = 2; Coding-synon = 3; Coding-nonsynon = 4;
mRNA-UTR = 5; Intron = 6; Splice-site = 7; Contig-reference = 8; Coding-exception = 9;
NearGene-3 = 13; NearGene-5 = 15; Codi
nq-nonsynonymous nonsense = 41; Codinq-nonsynonymous missense = 42; Codinq-nonsynonymous
frameshift = 44; Coding-nonsynonymous cds-indel = 45; UTR-3 = 53; UTR-5 = 55; Splice-3 = 73;
Splice-3 = 75.
##Filtering no filtering.
#Chr
        Source Type
                                        Pos_End Score
                                                        Strand Phase
                                                                        Attributes
                        Pos_Start
```

#### Base Space Example:

chr9 AB\_SOLiD Small Indel Tool insertion\_site 127466840 127466840 1.0000 ID=37171; ins\_len=15; clustered-indel-sizes=15; allele-callpos=127466840;reference=aggaaaa;allele-call=aggaaaa/TAGGGCC/ AGGGCCTCTAAAAAAGGAAAA; rightmost-allele-call-pos=127466840; rightmostreference=aggaaaa;rightmost-allele-call=aggaaaa/TAGGGCC/AGGGCCTCTAAAAAAAGGAAAA;contextpos=127466839; context-reference-seq=aaggaaaa; context-variant-seq=ATAGGGCC/ AAGGGCCTCTAAAAAAAGGAAAA/AAGCTTCTCTAAAAAAAGGAAAA/NO\_CALL/ GATCTAAAAAAAAAAAAAAAAAAAGGAAAA; reference-reads=13; context-variant-reads=1; context-variantnonPCRdup-reads=1;gap-total-reads=5;gap-nonred-reads=3;gap-nonPCRdupreads=5;tight\_chrom\_pos=none;loose\_chrom\_pos=127466838-127466846; coverage\_ratio=4.3333; zygosity=HEMIZYGOUS-REF-MULTI-INDEL-ALLELE; zygosityscore=1.0000;rg\_id\_nums=1\_1;bead\_ids=705\_312\_1080,585\_1149\_1247,486\_211\_1170,674\_1449\_36,145 3\_413\_82;overall\_qvs=100,78,82,22;no\_mismatches=3,4;read\_pos=24,29,27,26,39;from\_end\_pos=36, 31,33,34,21;strands=-;tags=R3,F3;indel\_sizes=21,15,-369;non\_indel\_no\_mismatches=0,1;unmatched-lengths=0;ave-unmatched=0.0000;anchor-match-



lengths=59;ave-anchor-

#### Color space example:

```
AB_SOLiD Small Indel Tool
                                                    4865598 4865605 1.0000
chr1
                                     deletion
ID=3743;del_len=8;clustered-indel-sizes=-8;allele-call-
pos=4865598;reference=CTCCACAGC;allele-call=CTCCACAGC/A;rightmost-allele-call-
pos=4865598; rightmost-reference=CTCCACAGC; rightmost-allele-call=CTCCACAGC/A; context-
pos=4865596; context-reference-seq=GGCTCCACAGCT; context-variant-seq=GGAT; reference-
reads=17; context-variant-reads=8; context-variant-nonPCRdup-reads=8; gap-total-reads=8; gap-
nonred-reads=7;gap-nonPCRdup-reads=8;tight_chrom_pos=4865598-
4865598;loose_chrom_pos=4865598-4865598;coverage_ratio=2.4286;zygosity=HEMIZYGOUS-
REF; zygosity-
score=0.9450;rg_id_nums=3_3,4_4,1_1,5_5,2_2;bead_ids=1043_1773_368,1779_878_646,979_899_1763
,1753_1103_184,1838_1695_1151,650_1885_304,1389_2001_1593,2219_1326_1756;overall_qvs=72,51,5
4,36;no_mismatches=4,5;read_pos=11,20,21,22,32;from_end_pos=39,30,29,28,18;strands=-
;tags=F3,R3;indel_sizes=-8;non_indel_no_mismatches=0,-1;unmatched-lengths=0;ave-
unmatched=0.0000; anchor-match-lengths=49,99; ave-anchor-
length=67.7500;read_seqs=T02000122011320020111101110
011112222000323100111220, T10102101132000122011320020111101110011112222000323, G00010210113200
012201132002011110111001111212200032,G00010210113200012201132002011110111001111222200032,T32
001021011320001220113200201111011100111122220003,T223020211002001021011320001220113200201111
01110011; base_gvs=; non_indel_seqs=G00100020131231013223231211310101000313020212312010,, T1113
122022,G20211023021003013123202010031022220210123101021322;non_indel_qvs=;rsID=34028970
```

# Small indel TXT and SQL formats

The GFF file serves as the primary output format for the small indel caller. However, there is the TXT file which is an alternative format of the data contained in the GFF file. The information contained here is generally only a subset of that of the GFF. The SQL file contains a CREATE TABLE command for MySQL, and the contents of the SQL file can be imported into MySQL.

Table 49 describes specifically what information the small indel TXT file contains.

Column name	Description	Example
chrom	Chromosome number of indel.	1
min-chrom-pos	Start position of the indel.	713662
max-chrom-pos	End position of the indel.	713663
called-range	Range of chromosome position range of the feature.	713661–713661
	Note: This ambiguity is resolved in the final indel call of the GFF.	
tight-range	Conservative estimate of chromosome position range of the feature.	713662–713667
loose-range	Estimate of the maximum chromosome position range of the feature.	713662–713667

 Table 49
 Small indel TXT file format description



#### Table 49 Small indel TXT file format description (continued)

Column name	Description	Example
num-pos-strand	Number of reads that were mapped to the positive strand.	3
num-neg-strand	Number of reads that were mapped to the negative strand.	0
num-r3-hits	Number of reads where the indel was found on the R3 or F5 tag.	0
num-f3-hits	Number of reads where the indel was found in the F3 tag.	3
num-frag-hits	Number of reads where the indel was found from a Fragment tag.	0
indel-type	The type of indel: INSERTION, DELETION, or COMBINATION. For COMBINATION, there were reads indicating both an insertion and a deletion, and no indel size was called.	DELETION
unique-indel-size	Called indel size.	-2
indel-size range	The indel sizes reported from each of the reads.	-2 to -2
num-uniq-align	Number of non-redundant alignments in the pileup.	3
num-tot-align	Total number of reads in the pileup.	3
average-read- position	Average read position where the gap occurred.	15.6667
ave-from-end-read- position	Average from end read position where the gap occurred.	34.3333
indel-read-pos-list	List of read positions where gap occurred.	20;17;10
dbsnp-indel	Legacy. Not used.	-
uw-hgsv-indel	Legacy. Not used.	-
read-lengths	Lengths of the reads (full read sequence, not the extended match length).	50;50;50
paired-distances	The clone sizes of each of the bead pairs (NIL for fragments).	1426;1370;1454
ave-pair-dist	Average of the paired distances.	1416.667
tags-R3-F3	Tags where the indel was found. Possible values are F3, R3, and FRAG.	F3;F3;F3
	Note: For PE data, R3 represents the F5 tag.	
chrom-pos-s	Chromosome positions of the indel tag's match location.	713641;713644;713651
strands	Strand for each bead id.	+;+;+
indel-sizes	List of indel sizes found for each read.	-2;-2;-2
nums-mismatches	List of number of mismatches for each read.	-1;-1;-1
ave-numb- mismatches	Average of the number of mismatches found in the indel tag.	-1
indel-lower-pos-s	List of lower ranges of the indel.	20;17;10
indel-upper-pos-s	List of upper ranges of the indel. The lower and upper ranges represent the ambiguity of the gap alignment, for example, $AT/$ in the context of $ATAT/AT$ .	25;22;15
run-names	Run names (one for each input file) for each read. For BAM files, the 1 in 50_1_r is the runIdNum in the ##@HD header line of the GFF file.	L1_1_50_1_r;L1_1_50_1_r ;L1_1_50_1_r

Column name	Description	Example
clone-ids	Bead IDs for each read.	984_536_1054;1431_2007_ 1567;116_364_1582
read-seqs	The read sequences for each read.	T302000;T120320;T232132
ref-allele	Reference allele.	acagagagaag
var-allele1	Most common variant allele (if it passes the color error correction, otherwise it is NULL).	aCAGAGAAG
other-var-alleles	Other alleles of the variant, if present.	-
var-counts1	Number of reads that have the most common indel allele.	3
other-var-counts	The list of the other allele counts.	NO_CALL
ungapped- unmatched-lengths	For a particular bead-id, the length of the read that was left unmatched (equal to the read length if no ungapped match was found or if the detail level is not set to 9).	50,50,50
ave-ungapped- unmatched	Average of the ungapped, unmatched lengths.	50
anchor-match- lengths	Anchor tags match length.	49,44,49
ave-anchor-length	The average of the anchor match lengths.	47.3333
coverage-ratios	Clipped normal coverage/number of non-redundant reads.	1.6667
zygosity-call	Experimental zygosity call.	HEMIZYGOUS
zygosity-p-value	Experimental zygosity score.	0.9656

Table 49 Small indel TXT file format description (continued)

#### Example TXT file content

The following is an example of small indel text file output. See Table 49 on page 227 for an explanation of the fields in this file.

1 3534096	3534096	3534095-	3534095	3534096-3	534098	3534096-
3534098 1	2	0	)	3 DE	LETION	-
1 -1 to -1	2	3	24.666	57 25.3333	30;22;2	2
\N \N 50	;50;50	$\setminus N$	\N	FRAG	G;FRAG;F	'RAG
3534065;-353411	9;-35341	19	+;-;-	-1;-1;-1		0;1;0
0.3333 30;20;2	0	32;22;22				
L1_1_50_1_r;L1_	1_50_1_r	;L1_1_50_	<u>1_</u> r			
894_1039_1506;1	267_2006	_1555;275	_1513_1	309		
Т33222211112120	13022022	112031113	3033230	3333000003	31;T021	1213300
000333303233033	11130211	220220310	1121111	2;T0211213	3000003	3330323
303311130211220	22031021	211112	АТАААТА	ATAATA	3	
50,50,50	50.0000	NIL,NIL,	NIL	NIL 7	.0000	
HEMIZYGOUS	1.0000					

Small indel ALIGNThe ALIGN file contains a text visual of each pileup that makes an indel call. Below is<br/>an example record from this file, with approximately 30 base pairs trimmed from each<br/>side for clarity.

> 64,chr1:16765166-16765167(),DELETION,-2,;allele-call-pos=1676 5166;allele-call=CT/;allele-pos=16765164;alleles=CACTCTGA/CACTG

A;allele-counts=REF,15		
CGAAGGGGTCAAAGGACACTCTGAGTTAGTG	16765118	16765208
3202000121002021112221221032113		
CACTCTGA	16765164	16765170
1122212		
T1001210020211121221032113	1122212/11212+	CACTCTGA/CACTGA
20001210020211121221032113	1122212/11212+	CACTCTGA/CACTGA
T3020001210020211121221032113	1122212/11212+	CACTCTGA/CACTGA
020000210020221121221032133	1122212/11212+	CACTCTGA/CACTGA
2020002210020221121222032113	1122212/11212+	CACTCTGA/CACTGA
2020000210020211121221032113	1122212/11212+	CACTCTGA/CACTGA
02020001210020211121221030113	1122212/11212+	CACTCTGA/CACTGA
02220001210020211121221032113	1122212/11212+	CACTCTGA/CACTGA
12020001210020211121221032113	1122212/11212+	CACTCTGA/CACTGA
12020001210020211121221032113	1122212/11212+	CACTCTGA/CACTGA
12020001210020211121221032113	1122212/11212+	CACTCTGA/CACTGA
1202000121002021112122103213T	1122212/11212+	CACTCTGA/CACTGA
1202000121002021112120T	1122212/11212+	CACTCTGA/CACTGA
1210000211121221032113	1122212/11212+	CACTCTGA/CACTGA
1210020211121222032113	1122212/11212+	CACTCTGA/CACTGA

Table 50 Small indel ALIGN file format

Line	Name	Format and description
1	Indel	> ID,chr#:start-end(),INDEL_TYPE,INDEL_SIZE. The rest of the line contains several tags that are in common with the GFF file.
2-5	Reference	REFERENCE_SEQUENCE START_POS END_POS
		Reference sequence in base and color space for entire pileup region, and region around the indel. The coordinates are given after the sequence.
6+	Reads	COLOR_SPACE_READ_REF_COLORS/READ_COLORS+_REF_BASES/ READ_BASES
		Contains all the reads of the pileup, extracted colors around the indel, and the corresponding bases. The sign after the extracted colors indicates color space compatibility (+) or incompatibility (-).

# UNGAPPED and PAS.SUM formats

**D and** ormats The UNGAPPED file contains more details of the reads counted towards the number of ungapped alignments used for the coverage ratio. For every pileup there are as many header lines as number of BAM files as input. The header line has this format:

# INDEL\_ID BAM\_CHROM\_NUM POS1 POS2

Each line after the header is a list of bead ids along with properties of that bead extracted from the BAM file.

The PAS.SUM file is an intermediate the pileup file extracted from the BAM file. The file starts with BAM header information from each of the inputs. Following that, each pileup has a separate line. An example and format are given in Table 51.



Name	Components	Example
Position, id, and ungapped	{CHROM_NUM}:{POS}- {POS2}:{INDEL_ID}:{NUM_UNGAPPED_READS}	7:315505-315512:347316:2
# reads	<pre>{NUM_NON_REDUNDANT}: {NUM_NON_OPT_DUPS} ( {NUM_TOTAL_READS})</pre>	10:9(13)
Indel size	{MIN_INDEL_SIZE}-{MAX_INDEL_SIZE}	2-2
Reference	{REF_POSITION}:{REF_SEQUENCE}	315455:ACGCAGGAGCA
Align 1	<pre>{BEAD_ID} {UNMATCHED- LENGTH}{ISDUP}_{READ_LENGTH}_{BAMID}_ {READGROUPID}, {PAS_ALIGNMENT} {BASE_QVS_TAG1} {BASE_QVS_TAG2} {MAP_QV_SCORE}</pre>	1524_856_1942 0NonDup_60_1_1,7 315519.57.2(12:12_13)[!TAACT]  7314783 1:(59.2.0):q255[!CCACG] AB@>:9 AA<9%%* 22
Align 2 and higher	[same as above]	1970_1821_1324 0Dup_60_1_1,7 317273 1:(59.0.0):q255[!CTGAG] 7 315522.57.0(12:12_12)[!GCAAC] >BA7?4> BB?9B; 100

Table 51 Column descriptions of the PAS SUM file

CHROM\_NUM is 1-based. Position 1 and position 2 are the approximate position of the indel from the alignments. The position is recalculated later from the actual alignments. The number of non-redundant reads is calculated by the module and is used for the coverage ratio calculation, while the number of non-optical duplicates is counted from the BAM 0x0400 flag. The individual alignments follow either TAG2|TAG1 for paired libraries, or just TAG1 for fragment ones. In each tag, the format for a gapped alignment is:

chromNum\_ReadPos.MatchLen.MisMatch(IndelPos:Start\_End)[!Base/Color Seq]

The format for an ungapped alignment is:

chromNum\_ReadPos.MatchLen.MisMatch(MatchLength.0.0)[!Base/Color Seq]

Base sequences start with !, while color start with the primer base.

The following is an example from base space indel calling using ECC mate-pair library.

7\_-317273.-1: (59.0.0):q255[!CTG...AAG] 7\_-315522.57.0(12:12\_12)[!GCA...CAC]

## Small indel detection algorithms

Detection of indels variants with a split-read technique is achieved by using LifeScope<sup>™</sup> Software's small indel caller on BAM files produced from long mate-pairs, mate-pair, fragment, and pair-end library types. The combination of multiple libraries of any combination of these types is also possible. For pair-end and long mate-pair libraries, the small indel module, by default, is able to determine sizes up to 500 for deletions and 20 for insertions. For all fragment libraries, the default size range is up to

19 for deletions and up to 4 for insertions. Also, 75mers from either pair-end and fragment libraries are capable of insertions up to at least size 29. Furthermore, the module allows for detection of more complex variants such as indel-SNP combinations.

The small indels module determines high-quality calls for insertions and deletions in two stages. In the first stage, the LifeScope<sup>™</sup> Software mapping module (see Chapter 8, "Run a Resequencing Mapping Analysis" on page 113) produces gapped alignments on a bead-by-bead basis and writes them to the resulting BAM file. For all libraries, this determination occurs during the mapping phase of this using a single tag approach. For paired tag libraries, this determination additionally occurs in the pairing phase using a paired tag approach. In the second stage, the indel caller takes these gap alignments, extracts them out of the BAM file, forms pileups, filters the pileups based on certain heuristics, determines zygosity, and annotates the indel sequences. This results in concisely annotated and highly accurate indel calls.

Figure 18 illustrates the F3 with the indel. The algorithm also determines indels in the R3/F5 tag.



Figure 18 Gap alignment detection using the paired reads from mate-pair and paired-end data

For paired tags, the algorithm surveys only those indels with one-end anchored (OEA) mate-pairs (see Figure 18). It does so by realigning the OEA pairs using an anchor tag (that is one tag which can be aligned to the genome by itself) and by performing a more aggressive alignment with the other tag in a several kbp window (depending on the orientation of the tags and on the minimum and maximum insert sizes set in mapping's pairing phase) around the anchored mate. A pair is considered OEA if one read of the tag fails to align or if its aligned length was not higher than a certain length threshold, specified by the parameter pairing.gap.min.non.matched.length. In addition, for anchor tags, only alignments above a minimum length are used (those alignments are close to full length). Finally, the anchor tag may only have a maximum number of anchor alignments (close to full length alignments) in the genome, which by default is set to 10 by the second value of pairing.gapped.max.hits.

Using the unanchored or non-fully extended tag, the module starts aligning both ends of the read localized by the other tag's match location. With the region of the genome determined by this match location and insert size distribution, the module makes a catalog of end locations by extending the read starting with a minimum value

#### Paired tag approach (paired libraries only)

(specified by pairing.gap.edge.length.insertions and pairing.gap.edge.length.deletions parameters rescue for insertions and deletions, respectively) until the alignment hits the maximum of number of mismatches allowed. These maximums are specified by these parameters:

- pairing.anchor.max.mismatches: the number of mismatches in both tags
- pairing.gap.max.mismatches.tag1: mismatches for the F3 tag
- pairing.gap.max.mismatches.tag2: mismatches for the R3/F5 tag

The default value for the single tag number of mismatches is 5 for read lengths 50 and higher (50 set by min.length.for.aggressive.gapped.mapping), and 2 for read lengths shorter than that. Then the single tag number of mismatches for paired-end libraries is 5 and 2 by default for the F3 and F5 tags, respectively.

With this catalog, the process attempts to join the ends of the read to find a single gapped alignment within a certain size range. The maximum gap size allowed depends on parameters specified in the mapping's pairing phase parameters. Furthermore, it identifies it as a gapped alignment if the above joining could be done with the fewest number of mismatches. Ambiguity of the location of this joining is common, mainly due to the presence of short tandem repeats. This is represented by the XA field in the BAM file. Later on downstream, the indel caller (as described later) using the consensus of reads, produces a left most, concise representation of the variant. Furthermore, gap alignments must also satisfy a minimum edge length which is the number of base pairs before the end of the read.

For determining small indels (deletions to size 11 and insertions to size 3), the algorithm disallows indels within 3 bases from either end of the read. The module identifies if it is able to piece together both ends, allowing only for a single gap of up to 4 base pairs inserted (present in read but not in reference), or up to 11 base pairs deleted. Larger indels up to size 500 are also found using different edge length criteria. These sizes and criteria are specified in several mapping parameters as shown in the Table 52. The values that effect gap alignments are the second through fourth ones, and are specified in the table as small indels, medium insertions, larger insertions, and larger deletions, respectively. s1 and s2 are specific tag lengths of the library used, for instance, a 75x35 PE library would have 75-35.

Larger indels are found for 60-60, 50-50, and 75-35 read length libraries, but not found for smaller 50-35, 50-25, 35-35, or 25-25 read length libraries.

 Table 52
 XML representation of gap size ranges

Parameter	Small indels	Medium insertions	Larger insertions	Larger deletions
pairing.gap.max.deletion.size pairing.gap.max.deletion.size.s1-s2	11	0	0	500
pairing.gap.max.insertion.size pairing.gap.max.insertion.size.s1-s2	4	15	21	0
<pre>pairing.gap.edge.length.deletions pairing.gap.edge.length.deletions.s1-s2</pre>	3	0	0	20
<pre>pairing.gap.edge.length.insertions pairing.gap.edge.length.insertions.s1-s2</pre>	3	14	14	0



#### Single tag approach (for both paired and fragment libraries)

1) Take reads that in the first stage of mapping do not align, or align only partially.



Figure 19 Global gap alignment detection using the single read technique used in all libraries

Using only a single tag, gap alignments are determined in the map phase of resequencing mapping. Although gap alignments can be found along with ungapped alignments in the first stage of mapping, the process here describes finding gap alignments only in the second stage of mapping. This process is illustrated in Figure 19 and Figure 19. As with mate-pair libraries, whole genome searches for gapped alignments is prohibitively expensive. Similar to the paired tag approach, a read is only considered for this process for both local and global alignment methods if the tag fails in the first stage of mapping to align or if its alignment length did not extend past a certain length threshold, specified by the gap aligner parameter mapping.gap.min.non.matched.length. Then each read is localized by performing a 20 base pair ungapped alignment allowing for 1 mismatch. The 20mer taken is from both the beginning and 30 bases pairs into the read. The anchor properties are specified by the second.map.scheme.unmapped.xx mapping parameters, and for 50mers and longer, the value is 20.1.0:30.

Two algorithms are available in the system, GLOBAL (default) and LOCAL, specified by second.map.gapped.algorithm. For GLOBAL (full read) gap alignments, each 20mer alignment then defines a search region of [A-40, A+80], where *A* is the position of the alignment. The downstream range (80) is settable using the parameter second.map.gapped.seed.window.right, and upstream (40), using second.map.gapped.seed.window.left. With this region, the same full read alignment strategy done with mate-pairs is performed: a catalog of partial begin and end read hits is formed, and an attempt is made to join them with a gap of some size. For LOCAL gap alignments, full read length matches are not required, but rather a gap is allowed during extension of the original anchor hit with a gap and mismatch penalty of the parameters second.map.gapped.penalty (default of 5) and second.map.mismatch.penalty (default of -2.0), respectively. Once the penalty threshold is met, the extension of the read terminates.



As in the paired approach, all gap alignments are restricted by both maximum size and minimum edge length requirements. By default, this approach finds deletions to size 19 and insertions to size 4. Larger insertions up to size 29 have been found effectively with 75mers using more restrictive parameters, which at the same time reduce sensitivity for smaller insertions and do not perform deletions. Insertions specifically at size 30 were set, but not evaluated. Both the default and larger insertion parameters are specified in the Table 53.

Parameter	Small indels (default)	Medium insertions (optional for 75mers)
second.map.gapped.deletion.tag1	19	0
<pre>second.map.gapped.insertion.tag1</pre>	4	30
<pre>second.map.gapped.insertion.tag2 second.map.gapped.edge.length.</pre>	12	0
second.map.gapped.edge.length.	12	18

 Table 53
 Various size and edge length requirements for the small indels and larger insertions

## Small Indel Calling Overview

The small indel caller uses alignments contained in the BAM file, which in the LifeScope<sup>TM</sup> Software mapper have two general types, ones that contain a single gap in the alignment (those that have an I or D in the CIGAR string) and ones that do not, which is referred to as ungapped alignments. The caller uses a variety of algorithms to produce the indel calls present in the GFF file. These algorithms are discussed in the sections below, and are summarized in this figure.



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#### Pileup handling

The gap alignments contained in the BAM files form the basis for calling indels. The gap alignments go through a series of processes before being reported in the final GFF output. Those alignments that have a minimum overall mapping quality (small.indel.min.mapping.quality, 8 by default) are extracted from the BAM file. The exception to meeting this quality threshold is the class of F3 gap alignments from pair-end libraries (see "Mapping quality adjustment (paired-end libraries only)" on page 238).

Properties of the gap alignment tag (small.indel.min.non.matched.length) and anchor tag (small.indel.min.anchor.mapping.quality and small.indel.min.map.length) are also assessed and are affected by small.indel.detail.level. Only the first 6 (detail level x 2) anchor reads are considered for the default setting of 3. Alternative alignments for the non-matched length filter are considered only for a detail level of 9. Alternative alignments are those that are found without a gap at the same chromosome location. Furthermore, anchor reads are present only in paired approaches, but alternative alignments could be present in either paired or single tag approaches.

Next the gap alignments are grouped together by genomic location to form pileups of reads. Because of the positional ambiguity of indels, pileups are formed by proximity; specifically, alignments that are within 5 base pairs between consecutive evidences are combined together. For pileups that have 6 or more non-redundant reads, the sixth and additional reads after that are only grouped together if the pileup is 2 or fewer base pairs from the last gap position. The default behavior is set by the parameter small.indel.consGroup (1 by default). A value of 9 makes every gapped alignment into a separate pileup.

Finally, the pileup is taken if the number of non-redundant reads is between small.indel.min.num.evid and small.indel.max.num.evid, inclusively. By default, pileups with two or more reads are taken at this stage. Non-redundant reads are those reads that have unique F3 and R3/F5 positions for paired tags, or unique F3 positions for single tag analysis. Mixed single and paired tag reads are considered unique from each other. With each pileup extracted from gap alignments, ungapped alignments that span the gap by at least 5 base pairs are counted, and are used in the normal vs. indel coverage ratio. The module stores the pileup information extracted from the BAM file into an intermediate PAS.SUM file for further analysis.





Figure 20 Small indel caller heuristics

The pileups go through additional filters, as illustrated in Figure 20, each with corresponding optional parameters in Table 47 on page 217. The best mapping quality cutoff (specified by small.indel.min.best.mapping.quality, with a default of 10) is the cutoff for the highest overall mapping quality from the pileup. For paired tags, this cutoff is the highest pairing quality value; for fragment, the highest mapping quality; and for mixed, the highest of either. Filtering for the maximum value allows lower quality reads to act as supportive evidence. Ambiguous indel size filtering requires that either 75% of the alignments have the same size or that they are able to form at least one valid clustered indel size. The latter method, in the case of multiple clustered sizes, allows including non-reference hemizygote indels. This method is described in more detail in "Indel size determination" on page 242.

If only two non-redundant reads are required to make an indel call, false positives can become prevalent at higher coverage levels (for instance, 100x). By using the number of ungapped alignments queried from the BAM file, the software can determine a coverage ratio of ungapped (with 5 base pairs clipped) coverage over a number of nonredundant indel supports. A high ratio is indicative of a false positive and are filtered out at values higher than the setting of small.indel.max.coverage.ratio (12 by default). Even in low coverage situations, there are regions of high coverage where this filter is active and further helps to reduce false positive indel calls. Finally, the pileup must also be comprised of reads where the majority of reads have gaps that are color space compatible. The parameter small.indel.colorspace.compatibility.level sets filtering based on this test, where a setting of 0 indicates no color space-based filtering and 1 filters out pileups were NO\_CALL is most prevalent (1 is the default for LifeScope<sup>™</sup> Software and recommended for max mapping). NO\_CALL for the allele occurs in reads where the gap is not color space compatible.

#### Mapping quality adjustment (paired-end libraries only)

Certain paired alignments may have poor overall mapping quality because of poor mapping of one tag. However the gap alignment of the better tag still constitutes strong evidence for indels and should be used for increased sensitivity. In order to effectively use these alignments in BAM files missing the single tag mapping quality (SM), the tool performs an adjustment to the mapping quality for the sole purpose of reducing the quality required to meet the minimum overall mapping quality (small.indel.min.mapping.quality) for certain otherwise high quality reads. This adjustment has no effect in either the best mapping quality filter or the reported output of quality scores in the GFF3 file. The QV adjustment follows this formula:

#### $QV_{adj} = elc \cdot EL - mmc \cdot MM + cc$

where *elc*, *mmc*, and *cc* are the coefficients for edge length, number of mismatches, and constant, respectively and EL and MM are the actual minimum edge match length and number of mismatches, respectively for that read. By default this is done only for the F3 tag of a pair-end library, with *elc* = 0.25, *mmc* = 0.50, and *cc* = 5. These settings are not readily available, as it may require editing the small indel XML file to add small.indel.mapping.quality.adjustment.libraries.tag1 (default value of PE) and small.indel.mapping.quality.adjustment.coefficients.tag1 (default of 0.25,0.50,5) parameters (editing the file is not recommended). This adjustment has no effect on the second (F5) tag.

# Color space considerations

When an indel occurs in a sequence, and that sequence is measured using color space, the color-space sequence not only has a gap of the same size of the indel, but also leaves a signature that can indicate if there is a measurement error within the gap. This is especially important in the case of insertions when you have a small number of evidences and there is a disagreement in the bases of the inserted sequence. With methods that directly measure bases, there would be no indication, based on the inserted sequence alone, on which inserted bases is more trusted.

In color space, this signature can be used to see if the color that spans the gap is compatible with the set of colors that go through the gap. For example, the alignment AACG/A--G would be 013/2-- in color space. Here the color 2 spans the gap (measuring both A and G), while 013 goes through the gap (measuring AACG). The color 2 is compatible with the sequence 013 because both would end with a G in base space. However, an alignment of 213/2-- would not be compatible because, using the same starting base A as the above example, 213 would measure AGTA. Because the rest of the color-space sequence beyond this would be aligned, 213/2-- would be indicative of a measurement error within the insertion if 213 is the sample, or if the sequence is the reference, the color 2 that spans the deletion. The alignment's color space compatibility can be calculated for any sequence using color space addition. This signature for color space compatibility is used to more accurately call the inserted sequence of the insertion, important if only a small number of reads are used to call an indel. Also an entire pileup is more likely to be a false positive if most of the reads indicate a gap that is not color space compatible.



# Allele calling and sequence context determination

For every gap alignment, the LifeScope<sup>™</sup> Software aligner reports the position of the gap and an ambiguity of that placement using the XA BAM field. The caller takes that information and the color-space reads to determine the base-space sequence for each of the reads, reporting the reference and all the sequences found. The module takes these calls further by taking the reference and the most common allele(s) present and then making a concise representation.

#### Context sequence extraction accounting for ambiguous placement

The small indel caller has three representations of the reference and variant sequences that contain the indel:

- Context sequences Represents the sequence context around the indel.
- Leftmost sequences Represents the sequence after trimming all common bases between the variant reads and the reference. Trimming goes from the right to the left first, and then left to the right.
- **Rightmost sequences** Same as leftmost, except that trimming goes from left to right first.

The context region is specified by taking the loose\_chrom\_pos (which is contained in the output GFF), expanding the start position by 1 and the end position by 2, and then trimming the first two positions and the last two positions, only if the bases at those particular positions are identical in all of the reads and the reference. In addition, if any of this trimming process produces no bases for the reference sequence, the position is represented as that of the base immediately before the insertion.

An example of this is the following indel call that has a loose\_chrom\_pos=94446948-94446955 and is annotated in dbSNP as rs58864345. The reference sequence at positions 94446948 to 94446956, and the corresponding reads is:

Ref	ACTTCTTCCC9444694894446956
	120220200
Read1	ACTTCTTCTTCCC
Read2	ACTTCTTCTTCCC
Read3	ACTTCTTCTTCCC
Read4	ACTTCGTCTTCCC

Because the beginning A and the ending CC are all the same, these would be trimmed off. This would result in the context sequence related tags being reported as:

context-pos=94446949;context-reference-seq=CTTCTTC;context-variant-seq=CTTCTTCTTC

The leftmost trimming of this result is represented here:

CTTCTTC Ref	erence @ 94446949
-------------	-------------------

CTTCTTCTTC Sequence found in reads

This results in the following positioning and allele call:

allele-call-pos=94446948;reference=-;allele-call=-/CTT

Since the reference here was entirely trimmed, the position reported, as per the GFF standard, is the one immediately before the CTT insertion. Finally, the rightmost trimming is represented here:

CTTCTTC	Reference	e @ 94	446949	
	0	c		

<u>CTTCTTC</u>TTC Sequence found in reads



and results in a different position:

rightmost-allele-call-pos=94446955;rightmost-reference=-;rightmost-allele-call=-/TTC

All three of these representations are present to preserve the CTT repeat context and have positioning compatible with both VCF (leftmost) and dbSNP (rightmost) positioning.

#### Making concise allele calls

The allele-call and rightmost-allele-call tags also represent information about the sequences detected. It always contains the most common indel variant sequence. Also, if a large enough number of ungapped alignments are present, it includes the sequence taken from the reference. Finally, if the second most common variant sequence has at least 75% of the reads as the most common, it too is included.

The following indel call from chromosome 2 at position 220,378,748 in hg18 illustrates the allele calling. The sequence context of this detected indel is:

and the read counts is reference-reads=18; context-variant-reads=4, 4, 1. From the read counts, both the reference (ATTT) and the two different variant sequences (ATA and TAT) are called. However, the variant with just one read is be called. All in all, the resultant allele call would be

allele-call-pos=220378748;reference=attt;allele-call=attt/ATA/TAT

#### VCF format

This section describes the differences between of indels in GFF format with indels in VCF format.

Insertion VCF record

The following is an example of a VCF record with an insertion:

20 3. C CTAG . PASS DP=100

This record contains an insertion since the reference base C is being replaced by C (the reference base) plus three insertion bases TAG. This example contains only two alleles so we have the two following segregating haplotypes:

Ref: a t C - - - g a // C is the reference base

: a t C T A G g a // following the C base is an insertion of 3 bases

The following shows how the LifeScope<sup>™</sup> Software small indels module represents the same insertion, in GFF3 format:

```
gff start/end position: 3/3
reference = -
variant = TAG
```

Deletion VCF record

The following is an example of a VCF record with a deletion:

20 2. TCG T . PASS DP=100



This record contains a deletion of two reference bases since the reference allele TCG is being replaced by just the T (the reference base). This example contains only two alleles so we have the two following segregating haplotypes:

Ref: a t C g a // C is the reference base

: a t - - a // following the C base is a deletion of 2 bases

The following shows how the small indels module represents the same deletion, in GFF3 format:

```
gff start/end position: 3/4
reference = CG
variant = -
```

Mixed VCF record for a microsatellite

The following is an example of a mixed VCF record:

```
20 2. TCGCG TCG, TCGCGCG . PASS DP=100
```

This example is a mixed type record containing a 2-base insertion and a 2-base deletion. This example contains three segregating alleles so we have the three following haplotypes:

Ref: a t c g c g - - a // C is the reference base
 : a t c g - - - a // following the C base is a deletion of 2 bases

: a t c g c g c g a // following the C base is a insertion of 2 bases

In all of these examples dashes have been added to make the haplotypes clearer, but of course the equivalence among bases is not provided by the VCF. Technically the following is an equivalent alignment:

Ref: atcg - - cga // C is the reference base

: a t c g - - - a // following the C base is a deletion of 2 bases

: a t c g c g c g a // following the C base is a insertion of 2 bases

The following discussion describes how the small indels module represents the same mixed record, in GFF3 format.

Leftmost representation of the indel actually represents the haplotypes as:

Ref: a t - - c g c g a // C is the reference base

: a t - - - c g a // following the C base is a deletion of 2 bases

: a t c g c g c g a // following the C base is a insertion of 2 bases

Given the above representation, the small indels module represents the deletion as:

```
gff start/end position: 3/4
reference=cg
variant= -
```

The insertion is represented as:

gff start/end position: 2
reference=variant=cg

When both the insertion and deletion occur at the same time, the small indels module represents them as:

gff start/end position: 3/4
reference = CG
variant = -/CGCG

**Note:** The indel caller may be missing some of the heterozygous indel sequences when there are multiple sizes present in a single indel call. In these cases, the allele-call is represented by the possibleOthers tag. This occurs when the second most common size does not have 75% of the reads as the most common indel size. The indel sequence is also missing with the third most common and rarer indel sizes, if present.

Indel size determination Indel sizes are determined by two methods. The first method is if the pileup has a single called indel size, where three-fourths or more of the alignments have a particular size. The GFF file reports this as ins\_len or del\_len for an insertion or deletion, respectively. The second method, that is performed on all pileups, is clustering based on the indel size. The list of indel sizes for a particular pileup is collected. Insertions and deletions, if they both occur, are split into two clusters. If both insertions and deletions do not occur, the starting point is a single cluster of all the sizes.

From those clusters that are present, the algorithm makes a decision whether or not to split that cluster based on the size differences contained in that cluster. For small indel clusters, those whose average size is less than 20, single base pair differences are split. For example, a size list of 1,1,1,2,2 are split into two clusters of 1,1,1 and 2,2. For the other clusters, they are split if the ratio of the largest size difference over the smallest indel size is greater than 1/4. For instance, a size list of -300,-340,-499,-500 results in two clusters of -300,-340 and -499,-500. If a cluster is not split by either of these, it still can be split if both the highest distance minus the average distance of the rest is greater than 6, and the highest distance is greater than 5/4 of the average distance of the rest.

Clusters are split recursively until any resulting cluster cannot be split further, and this process forms a list of clusters. Any cluster that has a single alignment is removed. For small indels (those less than 20 in size), the algorithm uses the single size that is present in that cluster. For a cluster of larger indels, the algorithm uses an average size found within that cluster. All cluster sizes are reported by the clustered-indel-sizes tag in the GFF. In addition, if there is not an indel size called with 3/4 agreement in size, then the cluster size(s), if at least one exists, is reported in the GFF as a comma-separated list after ins\_len, del\_len, or len. The first two are for when all the sizes are insertions or deletions, respectively, and the last (len) is used if there is both an insertion and deletion, where deletions indicated by a negative value.

Indels are also filtered based on whether these methods were successful in determining a consistent indel size or set of indel sizes, depending on the small.indel.indel.size.distribution.allowed setting. The default setting (can-cluster) requires that at least one size cluster is formed, and also allows for multiple clusters of different sizes. Overall, this method allows for more variety in the types of indels called. For sample long mate-pair and targeted resequencing paired-end datasets, this method does not significantly decrease concordance to dbSNP. A more restrictive setting of similar-size does not allow for this variety, by requiring that the pileup has at least 75% of the reads having the exact same size. A setting of similar-size-any-large-deletions allows for larger deletions to have

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different sizes. With similar-size-any-large-deletions, the caller includes additional indels that are discordant in size but have at least two reads with deletions of size 20 or higher. The setting any turns off filtering based on indel size, but this setting usually results in greatly increasing false indel calls.The coverage ratio (described in "Pileup handling" on page 236) also serves to call an indel hemizygous, specifically the type of zygosity detected here is one where the

Reference allele calling reference is present. Because the coverage ratio is the number of ungapped alignments over the number of the gapped ones, the ratio is an indicator of this localized hemizygosity. The length scale of this hemizygous call is that of the gap size range present in the input from the aligner, mostly in the small indel range. The size range is much smaller in length scale than the whole chromosome hemizygosity present in, for example, human males for chromosome X. The software calls the presence of reference alleles by utilizing a table of coverage ratios, by utilizing a table of coverage ratios located in a file located at <installdir>/etc/small-indels/zygositycalibration.conf. The table was derived by matching DH10B reads to an indel introduced reference to simulate the non-hemizygous state (indels of the same length on both alleles). For the hemizygous state, reads that occurred over a simulated indel had a 50% chance of being altered to contain that simulated indel. Because indel alignments are generally less sensitive, hemizygous situations occur frequently above coverage ratios values above 1. Different situations were simulated and are available using the parameter small.indel.zygosity.profile.name. The DH10B data comes from a 50mer long mate-pair sequencing run, using both tags for the paired approach and using only the F3 tag for a single tag approach.

**Zygosity Calling** Determination of the presence of the reference allele is analogous to heterozygous calling in SNPs where one allele is the reference. However, indels have greater complexity, because a single indel can have different inserted base pair sequences or it can have different gap sizes. For example, an indel can be the same with respect to size, but different with respect to the actual inserted sequence (see "Example 7, multiple inserted alleles" on page 246). Also, indels can also have different distinct sizes that are both not the reference (see "Indel size determination" on page 242).

In the end, three factors are used to call zygosity:

- Presence of the reference allele
- Presence of different instructed sequences
- Presence of multiple indel sizes

Non-diploid organisms or regions of higher than two in copy number can even have two or all three of the above factors. To take into consideration all the possible combinations of the above factors, the final zygosity call has values of HEMIZYGOUS-REF, HOMOZYGOUS-NON-REF, HEMIZYGOUS-REF-MULTI-INDEL-ALLELE, HETEROZYGOUS-NON-REF, MULTI-HEMIZYGOUS-REF, and HEMIZYGOUS-NON-REF.

**Note:** Having multiple different sizes, without having multiple different sequences, is not possible.

Sampling of gap alignments Sampling of gap alignments occurs when the number of gap alignments contained in a single pileup is greater than small.indel.num.aligns.per.pileup. A larger list than this is initially kept, but that size is limited to 30 times this value. Future gap alignments coming in order from the BAM file are ignored. From this larger list, each item has a probability of being included in the sample list of small.indel.num.aligns.per. pileup divided by the size of the larger list. This manifest itself in the context-variant-reads tag of the gff. For example,

context-variant-seq=GG,AT; context-variant-reads=15464,1497(899,87)

means that in the sample, 899 reads had GG in the actual sample, but based on the ratio of the sample size to the total size, an estimated 15,464 reads had that allele in the full sample.

**Examples** Below are several examples that illustrate the allele calling. The positions reported here are from chromosome 1 of the human hg18 reference.

#### Example 1, a simple insertion:

ins\_len=1;allele-call-pos=55076169;reference=-;allele-call=A/A;reference-reads=5;contextvariant-reads=5;coverage\_ratio=1.000;zygosity=HOMOZYGOUS-NON-REF;zygosity=score=0.1000

An insertion of A is reported here at 55,076,169 because it occurs between position 55,076,169 and 55,076,170. The allele call here is both alleles are variant. Even though there are as many reference reads as variants, it is called homozygous because the generally greater sensitivity of finding ungapped alignments.

```
> 4223,chr1:55076169-55076169(),INSERTION,1,;allele-call-pos=55076169;allele-call=/A;allele-
pos=55076169; alleles=/A; allele-counts=REF, 5
AATCTATACAGATCATTTCATCTTTTTCTGGCATTGAGT-TATATCTGTACTGGATACCTATGTTTAAGGCTATG 55076131 55076204
GT-TA
                                    55076168
                         Ref
                                                55076170
                             10 - 3
                         Reads GTATA
                              1333
 T03331122321300213220000221031301221333333221131210
               32200002210313012213333332211312102331023311003022T
               32200002210313012213333332211332102331023311003022T
                  T00221031301221333333221131210233102331100302032331
```

#### Example 2, a simple deletion:

del\_len=3;allele-call-pos=91763033;reference=AAA;allele-call=AAA/-;rightmost-allele-callpos=91763033;rightmost-allele-call=AAA/-;zygosity=HEMIZYGOUS-REF;

This example shows a deletion of AAA where position 91,763,033 would be the start of the deletion and 91,763,035 the end. It is a hemizygous reference call, so the reference is present in the allele call, and there is no difference in this case between the rightmost and leftmost representations.

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#### Example 3, an ambiguous insertion:

```
ins_len=3;allele-call-pos=2045476;reference=-;allele-call=GTA/GTA;rightmost-allele-call-
pos=2045478;rightmost-reference=-;rightmost-allele-call=AGT/AGT;context-pos=2045477;context-
reference-seq=gt;context-variant-seq=GTAGT
```

In this example there are two possibilities, -/GTA where the insertion is after position 2,045,476 or -/AGT, where the insertion is after 2,045,478. Following the left-most rule, the allele call and allele call position are reported as the lowest chromosome position. The full representation is reported here as gt/GTAGT.

```
> 167,chr1:2045476-2045476(),INSERTION,3,;allele-call-pos=2045476;allele-call=/GTA;allele-
pos=2045477;alleles=gt/GTAGT;allele-counts=REF,2
cacggcggtg---gttagggtcacggctgtag 2045467 2045495
1130330110---103200121130321132
    Ref tg---gtta 2045475 2045479
        10---103
    Reads tGGTAGTTA
        10132103
T3310110132103200121130321
        T1110132103200121100321132
```

#### Example 4, insertion of a short tandem repeat:

```
ins_len=3;allele-call-pos=5274096;reference=-;allele-call=-/TGT;
rightmost-allele-call-pos=5274100;rightmost-reference=-;rightmost-allele-call=-/GTT;
context-pos=5274097;context-reference-seq=tgtt;context-variant-seq=TGTTGTT
```

In this example there is a repeat in the sample that is not in the reference, so the full reference is TGT/TGTTGT, where TGT in the reference starts at position 5,274,097. The small indel caller trims this and then takes the position immediately before the insertion yields a position of 5,274,096 for the leftmost, and 5274100 for the rightmost representations.

```
> 532, chr1:5274096-5274096(), INSERTION, 3, ;allele-call-pos=5274096; allele-call=/TGT; allele-
pos=5274097; alleles=tgtt/TGTTGTT; allele-counts=REF, 6
gttcccatctcctaactggggctaattatcatccctc---tgtttgagtgtttcgaggatgaattgag
                                                                             5274060 5274124
1020013222023012100032303033213200222---110012211100232202312030122\\
                                    tc---tgtttg
                                                      5274095 5274101
                             Ref
                                    22---11001
                             Reads tCTGTTGTTTG
  T20132220230121000323030332132002221101100122111002
01322202301210003230303321320022211011001221110021T
     13222023012100032303033213200222110110012211100232T
T20230121000323030332132002221101100122111002322023
                T10032303033213200222110110012211100232202312030122
                T10032303033213200222110110012211100232202312030122
```

#### Example 5, deletion of short tandem repeats:

```
del_len=4;allele-call-pos=24115702;reference=agag;allele-call=-/-;rightmost-allele-call-
pos=24115707;rightmost-reference=gaga;rightmost-allele-call=-/-;context-
pos=24115700;context-reference-seq=acagagagagagac;context-variant-seq=aCAGAGAAC
```

The AG repeat occurs 4 times in the reference and only twice in the sample, or more concisely, AGAGAGAG/AGAG. Because it is a deletion, the left-most position of this repeat is reported here at position 24,115,702.

```
> 2130,chr1:24115702-24115705(),DELETION,-4,;allele-call-pos=24115702;allele-call=agag/;allele-
pos=24115700;alleles=acagagagagaac/aCAGAGAAC;allele-counts=REF,5
24115746
ref
                    acagagagagaac
                              24115700
                                       24115711
                    112222222201
                reads aC----AGAGAAC
 T2212303313231213112201203210011111----2222010201212012
    ТЗ313231213112201203210011211----2222010201212012000011
           T2210011211----222201020121201
```

#### Example 6, an insertion/SNP combination variant:

```
allele-call-pos=5658680;reference=a;allele-call=a/CT
```

At position 5,658,680 the reference has an A. The sample however has a CT, so this complex variant is simultaneously a SNP and an insertion.

#### Example 7, multiple inserted alleles

ins\_len=3;context-pos=60612131;context-reference-seq=A;contextvariant-seq=TAGA/TTGA/NO\_CALL;

This example has two main inserted alleles, TAG and TTG.

```
TGGGGGGATAAGGTGTTTATTAAGGATGACAGAAACCTCCTGAT---AGAGACAATATCATTCACCTTATAGATCCATCTCTG 60612088
TGAT---AG
                            Ref
                                             60612127
                                                           60612131
                                  1233---2
                                  12---332
                            Read1 TGATTTGAG
                                  12300122
                            Read3 TGATTAGAG
                                  12303222
 T10233020111003303020231211220010220212300122222110
 T10233020111003303020231211220010220212300122222110
  T12330201110033030202312112200102202123032222221103
              33030202312112200102202123032222221103332130211023T
                          T3010220212303222222110333
                          T30102202123032222221103332130211020333223201322222
                                Т0123001222221103332130211
```



# FAQ – Small indels

1

What size indels are detectable in LifeScope?

The size of the indels found in the caller is limited by the size of the gap found in the aligner which is dependent on the read lengths of the library. Indels, up to size 500 for deletions, and size 21 for insertions are found in 60-60, 50-50, and 75-35 read length libraries, and up to size 19 for deletions and size 4 for insertions in smaller 50-35, 50-25, 35-35, or 25-25 read length libraries. For long mate pair libraries, the practical limit for the insertion size, found in McKernan, et. al. (Genome Res. 2009. 19:1527-1541), was 20. More recent results show that for 75mer fragment or pair-end libraries, the limit is at least 29 (AGBT 2011), although non-default settings are needed to achieve this. (See Table 53 on page 235.)

#### Can the caller find SNP-Indel events?

Yes. These events, which are also referred to as replacements, are annotated by the allele-call tag in the GFF. For instance, allele-call=AT/G is an A/G SNP followed by a T deletion, which can equally be represented as an A deletion followed by a T/G SNP. Similarly, C/TGA is a C/T SNP followed by a GA insertion, which can also be represented as a TG insertion followed by a C/A SNP. There is no hard limit placed on the complexity of this variant (for example, double, triple, quad SNP, followed by an indel are all allowed), as long as the differences in the number of bases is within the gap size range contained in the BAM file, and the variable region is reported accurately by the aligner in the XA or XW BAM tag.

## 3

#### How can I speed up the small indel caller?

You can run the caller on different chromosomes at the same time making multiple INI files, one for every chromosome, and using the small.indel.genomic.region parameter to specify each specific chromosome.

## 4

#### Can I get the number of reference ad variant reads used for each indel call?

Yes. These are directly reported as reference-reads and context-variant-reads tags in the output GFF3 file. Actual bead ids and read sequences are also reported for the variant reads in the GFF3 file using the bead\_ids and read\_seqs tags. Bead ids for the reference, ungapped reads are reported in the \*.ungapped file.

Note that the LifeScope<sup>™</sup> Software aligner is generally less sensitive to gap alignments than to ungapped alignments, so two or three times more ungapped alignments are to be expected for hemizygous indels. Also, this calculated number of reference alignments, along with all the other allele counts may include redundant (PCR optical duplicates) alignments. The number of non-redundant reads per allele call is currently not reported.

### 5

# Can I perform gap alignments on all reads for greater sensitivity? Presently only those reads that do not match fully are considered for gap alignments.

The following parameters allow all alignments to be considered for gap alignments:

- For single tag: small.indel.frag.min.non.matched.length=0
- For paired tags: indel.min.non-matched.length=0

However, processing with these settings could be significantly slower.

# 6

#### Can I use a legacy PAS file as input to the small indel caller?

All inputs files used with the small indel caller should be BAM files. Legacy files such as PAS and MATES files are not supported as they may give unexpected results, such as missing indel IDs or less accurate zygosity calling.

# 7

#### Why is a duplicate insertion or deletion reported?

Occasionally an indel is reported twice in the output file. This situation occurs when an indel is detected separately in two different locations, from different sets of reads. After further processing, the two locations are resolved to the same location. These two entries represent a single insertion or deletion.

# 13

# Run a Large Indels Analysis

This chapter covers:

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## **Overview**

The large indel module identifies deviations in clone insert size. These deviations indicate intrachromosomal structural variations compared to a reference genome. Insertions and deletions (indels) up to 100 kbp are inferred by identifying positions in the genome in which the pairing distance between mapped mate-pairs deviates significantly from what is expected at the given level of clone coverage.

The module creates a look-up table of standard deviations at each level of clone coverage. The table produces an asymptotic curve in which the minimum size of detectable indels at a given level of significance drops rapidly as the clone coverage increases. The look-up table is used to determine the significance of the deviation in average insert size at each position in the genome.

Regions of the genome that are significantly deviated are selected as candidate indels. Hierarchical clustering is used to segregate the clones into groups in which the difference in the sizes of all clones in a group is less than a specified range. Clusters with too few clones, as specified by the user, are removed and the candidates are assessed to determine if a homozygous or heterozygous population of deviated insert sizes remains. All clones deviated by more than 100 kbp are discarded. Clones from various libraries with various insert sizes contribute to a single indel call by combining the probabilities associated with the clones from each library.

The large indel module only accepts paired data as input.

# Examples of running a large indel analysis

In a standard workflow The following standard workflows provide examples of running the large indels module:

- genomic.resequencing.lmp
- genomic.resequencing.pe

The following are example shell commands using reads and references from the LifeScope<sup>™</sup> Software repository.

```
lscope.sh shell -u username -w password
cd /projects
mk ecoli
cd ecoli
mk run1
cd run1
set workflow genomic.resequencing.pe
add xsq run0209a_50_PE.xsq
add xsq run0209b_50_PE.xsq
set reference hg19
# optionally change parameter defaults here
ls
run
```

See for Chapter 7, "Run a Standard Workflow Analysis" on page 97 for more information on running standard workflows. Chapter 6, "Run a Command Shell Analysis" on page 71 for information on shell commands and syntax.

**As a demo analysis** The optional examples download includes an example of how to run the large indel module by itself, as a single analysis that is not part of a standard workflow. The demo example is not recommended for general use.

See Appendix E, "Demo Analyses" on page 507 for information on running the large indel module as a standalone analysis.

## Large indel module parameters descriptions

In order to change a parameter value in your analysis, use the set param shell command to replace the line # *optionally change parameter defaults here* in the example commands in "In a standard workflow". For instance, to change the ploidy parameter, use this shell command:

```
set large.indel.ploidy 0 /tertiary/large.indel.ini
```



Table 54	Large indel	parameter	descriptions
----------	-------------	-----------	--------------

Parameter name	Default value	Description
library.type	matepair	Specifies the library type. Allowed values are:
		• matepair
		• pairedend
large.indel.bas.file	_	( <i>Optional</i> ) If you have a BAS file and specify it with this parameter, the file is not regenerated.
		The BAS file contains metadata information about the data source(s) and relevant mapping/pairing statistics. The BAS format is a pseudo-standard file format for storing BAM file metadata. For details see this site:
		ftp://ftp-trace.ncbi.nih.gov/1000genomes/ftp/pilot_data/README.bas
large.indel.max.clone.cov	1000	The maximum physical (clone) coverage for analysis. Loci with clone coverage above this threshold are not analyzed. Allowed values: Integers >= 3.
		<b>Note:</b> You can use this parameter in combination with the parameter large.indel.high.coverage to reduce false positives in high density genomes, for example, bacteria.
large.indel.min.coverage	3	The minimum physical (clone) coverage allowed. Loci with clone coverage below this threshold are not analyzed. Allowed values: Integers >= 1.
large.indel.ploidy	2	Determines zygosity and allele frequencies in eukaryotic genomes. Currently two scenarios are supported haploid (n=1) and diploid (n=2). Allowed values:
		• 1: Haploid.
		• <b>2</b> : Diploid.
large.indel.max.insert.size	100000	Maximum insert size, in base pairs.
large.indel.call.stringency	high	Specifies the large indel call stringency. Automates parameter adjustment to the desired stringency level.
		Allowed values:
		• <b>highest</b> : Recommend when a very low false positive tolerance is allowed.
		high: Increased filtering.
		• <b>medium</b> : Default values.
		low: Very aggressive.
		Lower settings results in more indel calls, but with more false positives. Higher settings result in fewer indel calls, but with fewer false positives.
large.indel.min.pairing.quality	25 (for MP) 10 (for PE)	Paired reads below this threshold are ignored. Allowed values: Integers 0–100.
large.indel.p.value	1e-10	P-value threshold, that is, the raw probability of committing a Type 1 error incorrectly identifying a large indel. Allowed values: Floats 0.0–1.0.



#### Table 54 Large indel parameter descriptions (continued)

Parameter name	Default value	Description
large.indel.high.coverage	0 (off)	Eliminate clone coverage weighting. This option significantly reduces the number of false positives when analyzing very high coverage genomic data. (Very high coverage genomic data is typically greater than 1000x read coverage, and is common for bacterial genomes.)
		Allowed values:
		• <b>0</b> : No effect on the module.
		• 1: Eliminate clone coverage weighting. The algorithm ignores coverage when calculating the structural significance of candidate indels.
Resource parameters		
memory.request	7gb	Memory request. Include the units gb in the setting.
java.heap.space	3000	Dynamic memory requirement.

# Large indel internal parameters

Table 55 describes parameters which we do not recommend changing.

Table 55 Large indel module internal parameters.

Parameter name	Default value	Description
large.indel.run	1	Whether or not to run the large indel module.
		• <b>0</b> : Do not run the large indel module.
		• 1: Run the large indel module during this analysis.
		The run parameter is set automatically by the shell during a standard workflow. Accept the default for most use.
large.indel.warnings	0	Whether or not to enable warning messages. Allowed values:
		• <b>0</b> : Do not enable warning messages.
		• 1: Enable warning messages.

# (Optional) Genomic annotation parameters

Table 56 describes the annotation parameters you can optionally add to your large indel analysis, to have annotation files generated. See Chapter 14, "Add Genomic Annotations to Analysis Results" on page 267 for information about annotations.

Table 56 Annotation parameters description

Parameter name	Default value	Description
Annotation general parameters		


#### Table 56 Annotation parameters description (continued)

Parameter name	Default value	Description
annotation.run	_	Specifies whether or not to add annotations. Allowed values:
		• <b>0</b> : Do not generate annotations.
		• 1: Generate annotations.
annotation.input.gff.file	_	The input variant GFF file, generated by the LifeScope <sup>™</sup> Software large indel analysis module (see "Annotation input files" on page 274).
		Expected file extensions: .gff, .gff3.
		Example: /data//HuRef_chr22/DB_out/HuRef_rmm13_SNP.gff3
		Must be a valid path, and the file must be readable. If this file is not found or is problematic, or if the parameter is missing, annotations output is not created.
Annotation source parameters	1	
annotation.indel.border.slack	5	An indel is considered to be matched to a dbSNP indel if the two overlap or if the distance between them is less than or equal to the value of annotation.indel.border.slack.
		Allowed values: Integers >= 0.
Annotation filtering parameters		
annotation.show.only.variants.in.exons	0	When set to 1, filters the annotated output: restricts the Filtered Variant annotation output file (see "Annotation output files" on page 282) to only the variants that overlap any exon.
		Allowed values:
		• <b>0</b> : Do not filter the output for variants that overlap any exon.
		• 1: Restrict output to variants that overlap any exon.
annotation.show.only.variants.in.genes	0	When set to 1, filters the annotated output: restricts the Filtered Variant annotation output file (see "Annotation output files" on page 282) to only the variants that overlap any gene, even if it does not overlap any exon.
		Allowed values:
		• <b>0</b> : Do not filter the output for variants that overlap any gene.
		• 1: Restrict output to variants that overlap any gene.
		<b>Note:</b> if the parameter annotation.show.only.variants.in.exons is set to 1, then the setting for the parameter annotation.show.only.variants.in.genes is ignored.
Resource parameters	1	
memory.request	22gb	Memory request. Include the units gb in the setting.
	-	For hg18 or hg19 sources, use memory.request=15gb.
	21500	Ear half or half courses use involves hear $c_{1000}$

If you are creating your own INI files, annotations parameters for this module are by convention defined in the file:

<project>/<analysis>/tertiary/annotation.large.indel.ini.



### Large indel output files

This section provides descriptions of the large.indel.gff file created by a large indels analysis run (see Table 57).

Table 57 large-indels.gff3 file format description

Column Title <sup>1</sup>	Description	Example				
Sequence ID	Chromosome name	chr11				
Source	Tool name	AB SOLID Large Indel Tool				
Туре	Indel type	insertion, deletion, and so forth				
Start Position	Estimated 5' breakpoint	1081331				
End Position	Estimated 3' breakpoint	1084700				
Score	Significance of the candidate Indel (p-value)	1e-10				
Strand	-	-				
Phase	-	-				
Attributes <sup>2</sup>						
dev	Indel size, measured in base pair	46				
avgDev <sup>3</sup>	Average deviation from the population average	-1.7198				
Zygosity <sup>4</sup>	Results from pair partitioning	Homozygous				
nRef <sup>5</sup>	Number of reference alleles	0				
nDev	Number of deviated alleles	5				
refDev <sup>5</sup>	Average deviation of the reference-allele pairs	0				
devDev <sup>3</sup>	Average deviation of the deviated-allele pairs	-3.567				
refVar <sup>5</sup>	Variance of the reference-allele pairs	0				
devVar	Variance of the deviated-allele pairs	0.8972				
beadIds <sup>6</sup>	Bead IDs providing support for the candidate indel	1806_975_1088,				
<sup>1</sup> genome.ucsc.edu/golder	Path/help/customTrack.html#GFF					
<sup>2</sup> Semicolon-separated list of large indel module-specific field names followed by an equal '=' sign, for example dev=46						
<sup>3</sup> Insertions have negative values						
<sup>4</sup> Either homozygous, hete	rozygous, or double					
<sup>5</sup> Value is always zero for	homozygous indels					

<sup>6</sup>Comma-separated list

Figure 21 shows an example of an output GFF file displayed in a spreadsheet.



	A	В	C	D	E	F	G	Н	1	J	
1	chr20	AB_SOLID Large Indel Tool	deletion	13022198	13022445	2.79E-11			dev=361;a	vgDev=0.60	5991;
2	chr20	AB_SOLID Large Indel Tool	deletion	13924362	13924978	1.88E-12			dev=271;a	vgDev=1.09	822; 2
3	chr20	AB_SOLID Large Indel Tool	deletion	14000747	14000930	2.29E-12			dev=193;a	vgDev=0.78	328; z
4	2010-2010 1										
5											
6	Attribute details										
7	dev=361										
8	avgDev=0.605991										
9	zygosity=HETEROZ	YGOUS									
10	nRef=55										
11	nDev=62										
12	refDev=-0.361414										
13	devDev=1.46417										
14	refVar=0.529432										
15	devVar=0.605724										
16	beadIds=466_1704_	1363,2199_554_1306									
17											

Figure 21 large-indels.gff3 file example

### Large indel analysis algorithm

Large indel detection is a tertiary module in LifeScope<sup>™</sup> Software. Data from the pairing phase of the mapping module serve as direct inputs for large indel discovery (see Figure 22 on page 256).

The large indel module uses pairing statistics are available in BAM headers of the input BAM files. If the pairing statistics are not available, the large indel module estimates the required parameters.

The large indel module processes inputs using an alignment window to hold and analyze sets of locus-spanning pairs. Regions with pairs that have significant insert size deviations are chosen as candidate indel sites, which are processed further to determine zygosity. The output of the analysis is one file in the GFF format. The GFF file contains information about the location, size, and significance of each large indel detected by the module.



Figure 22 Large indel analysis module

This paragraph refers to Figure 22. The large indel module accepts as input BAM files (blue) are generated during the secondary analysis module (mapping and pairing). Pairing statistics are included in the BAM headers. Pairing statistics provide more accuracy and can improve the final results of the large indel module. Final output consists of a GFF file (orange).

### Identify candidate indels

Pairing distances (sometimes called insert sizes) for each pair are assigned during the mapping and pairing module and subsequently used by the large indel module to determine indel candidacy.

**Note:** Insert sizes can be non-unique in the case of multiple feasible mapping/pairing combinations. When insert sizes are non-unique, the primary (optimal) pair is chosen for large indel analysis.



Clones that have been mapped and paired to a reference genome are classified as either concordant or discordant (see Figure 23 on page 257). Concordant pairs are those with insert sizes (sometimes called inter-read distances) that are not significantly deviated from the expected insert size of the library as a whole. Discordant pairs have insert sizes that deviate significantly from the expected value. Discordant pairs containing a putative deletion appear larger when mapped to the reference. Pairs containing a putative insertion appear smaller. Multiple discordant pairs in close proximity provide evidence for a candidate indel within the covered region.



Figure 23 Discordant clones used to identify candidate insertions or deletions

#### In Figure 23:

Pairs spanning a candidate indel (red) appear distorted when mapped to a reference genome. Insert sizes appear larger for deletions (left) and smaller for insertions (right).

The module moves across individual chromosomes in order of genomic position to generate an alignment window of overlapping locus-spanning pairs (see Figure 24 on page 258). When the window encounters the first read in a pair, the corresponding alignment is incorporated into the alignment window. Simultaneously, several moving statistics, including the number of locus-spanning pairs as well as their average insert size and variance, are updated. When the window encounters the second read in a pair, the corresponding alignment is dropped and the moving statistics are updated appropriately. A genomic region (sometimes called a window) is considered to contain a candidate indel when the average insert size of the set of locus-spanning clones is significantly deviated from the average insert size of the library as a whole.



Figure 24 Sets of locus-spanning pairs

The following sections refer to Figure 24.

#### Top panel

An alignment window, *W*, contains the set of overlapping locus-spanning pairs (black loops) at a particular genomic position *i* (arrow) of the reference genome (black line).

#### Middle panel

The window advances to the next position by incorporating the next alignment i+1. The corresponding alignment is the second in the pair, so the alignment i is dropped from the alignment window and the alignment statistics are updated accordingly. When the window encounters the first alignment, the corresponding pair is added to the window.



#### Lower panel

As this process proceeds along the chromosome, pairs are added or dropped from the window, moving statistics are continuously updated, and regions with significant insert size deviations are detected and analyzed.

Note: Insert size variations are exaggerated for illustrative purposes.

### Assign statistical significance to candidate indels

Regional insert size deviations can be calculated directly from the moving statistics associated with each clone window. The deviations that achieve statistical significance indicate relatively large structural variations compared to the reference genome (see Figure 25). Hypothesis testing determines the significance of deviations, where the null hypothesis asserts an insignificant difference between the local average insert size and the population average ( $H_0$ :  $x = \mu$ ). Candidate deviations are chosen where the probability of falsely rejecting the null hypothesis in favor of the alternative ( $H_a$ :  $x \mu$ ) falls below a user-defined confidence threshold.



**Figure 25** Hypothesis testing example The following paragraph refers to Figure 25.

The population average insert size  $\mu$  and standard deviation  $\sigma$  are calculated from the input BAM file. The alignment window contains a very small subset of pairs sorted by insert size (grey bars). Moving statistics including the number of locus-spanning pairs  $n_i$  as well as the sample average insert  $\overline{x_i}$  size are calculated from this subset of pairs at each genomic position *i*. The parameters are used to z-normalize insert sizes according

to  $z_i = \frac{|\bar{x}_i - \mu|}{\sigma}$ , which measures the absolute insert size deviation between the sample

and the population in units of standard deviation. The normalization step allows multiple libraries with variable insert sizes to be combined into one analysis. A candidate indel is considered significant if  $p(z_i|zn_i) < \alpha$  where probability values (p-values) are calculated according to the standard normal distribution and is a user-defined threshold parameter, where large.indel.p.value default is p=1e–10).

### **Determine zygosity**

After a candidate indel is detected and deemed significant, the alignment window is partitioned to remove erroneous pairs because of mapping/pairing artifacts and to further characterize indel alleles and zygosity (see Figure 26 on page 261). Locus-spanning pairs are partitioned to remove two groups because polyploidy is currently not supported.

Each partition represents a disjoint subset of pairs from the alignment window optimally grouped by insert size so that pairs with similar insert sizes are placed in the same partition. Partitions with only one pair (sometimes called outliers) are removed from consideration. The candidate indel is also removed if the average insert size for pairs in the remaining partition are not significantly deviated from the population average. Removing the candidate indel can occur when mapping or pairing errors are responsible for observed insert-size deviations.

The number of pairs per partition and various summary statistics are calculated for each partition to determine alleles, allele frequencies, and zygosity. Candidate regions with minor allele frequencies less than one-third are removed from consideration.

The large indel module uses a heuristic method to categorize each candidate indel. If both partitions contain pairs with insert sizes that are significantly deviated from the reference but not from each other, the region is designated HOMOZYGOUS. If pairs are significantly deviated from the reference and from each other, the region is designated DOUBLE, which indicates two indel alleles of different types or sizes. DOUBLE regions can be placed into one of several indel/indel categories, for example, insertion/deletion. If one pair is deviated from the reference and the other pair is not, the region is designated HETEROZYGOUS, which indicates the presence of indel and reference alleles.





Figure 26 Partitioning the alignment window to determine zygosity

#### Left panel

Heuristic methods are used to characterize indel alleles including homozygous reference alleles (lines), which are removed from consideration; insertions (hatched boxes), and deletions (broken lines).

#### Middle panel

Each partition contains pairs with characteristic insert size distributions. Summary statistics describing the distributions are used to determine indel and reference alleles (black and red bell curves, respectively).

#### **Right panel**

Based on the results of the analysis, each candidate indel is assigned an appropriate zygosity category.

Filter alignments and parameter optimization A user-defined pairing quality value (PQV) filtering threshold is set with the parameter large.indel.min.pairing.quality. The PQV threshold setting places constraints on which alignments are incorporated into the alignment window and subsequently used to determine large indel candidate regions. The default PQV threshold values are 25 for mate-pair data, and 10 for paired-end data. Adjusting the PQV threshold down (less than the default) to include lower-quality alignments generally improves sensitivity but increases the number of false positives. The opposite is true if you adjust the PQV threshold up (greater than the default). 13

	Adjusting the p-value and PQV thresholds together might be required to optimize the module for analysis of data that differ significantly from normal HuRef samples, for example, non-human or cancer genomes. However, the default settings provide a convenient starting point for this process.
	Additional optimization might be required for very high coverage samples. For example, alignments from high-density slides mapped to small prokaryotic genomes typically result in clone coverage values above 1000x. In the case of clone coverage values that are above 1000x, use the high-coverage flag (large.indel.high.coverage in LifeScope <sup>TM</sup> Software), which adjusts the p-value calculation to $p(z_i) < \alpha$ , eliminating the conditional coverage parameterization (weighting), and reducing the number of false positives that might otherwise result.
Input files for large indel analysis	The only acceptable inputs for large indel analysis are BAM files containing paired- end or mate-pair data. Other alignment types are not compatible with the large indel module.
	<b>Note:</b> LifeScope <sup>™</sup> Software does not support combining paired-end and mate-pair data into a single analysis.
Interpreting results from the large indel module	Large indel results are written to a GFF file, which can be uploaded directly into the UCSC Genome Browser ( <b>cbse.ucsc.edu/research/browser</b> ) or other similar visualization tools. See Table 57 on page 254 for a description of the GFF file format. See Figure 27 on page 263 and Figure 28 on page 264 for examples of GFF files generated by the large indel module





Figure 27 Identifying a 46bp insertion in MUC2

In Figure 27, paired-end data was analyzed using the SOLiD<sup>™</sup> mapping and pairing modules (against a human reference). The resulting BAM file was used for large indel analysis with default settings. The top panel uses the Integrated Genomics Viewer (IGV) to represent all BAM alignments covering the region chr11:1081331–1084700, which contains the partial coding region for the human mucin 2 precursor (MUC2) (bottom panel) as well as an indel previously identified by Levy et al., 2007 (not shown). Further information about IGV, including alignment color-encodings, is available at **broadinstitute.org**. The middle panel represents the subset of alignments used by the large indel module to identify a 46bp homozygous insertion breakpoint at the indicated position (dotted grey line). Forward and reverse strand reads are highlighted red and blue, respectively. The blue rectangle displays various alignment features, including a deviated insert size (97bp) that is significantly smaller than the average insert size of the population (170bp) providing evidence for an insertion at this site. The blue rectangle was generated by mousing over one of the reverse strand reads.



Figure 28 Identifying a 413bp deletion at IL2RA

In Figure 28, the top IGV panel represents long mate-pair alignments (mapped to HuRef) covering the region chr10:6135605-6139655, which contains an intron of human interleukin 2 receptor alpha (bottom panel) as well as several indels previously identified by Ahn et al., 2009; Bentley et al., 2008; Wang et al., 2008; Wheeler et al., 2008; and Levy et al., 2007 (not shown). Several alignment features are consistent with the presence of a deletion at this site, including six gapped alignments (dotted lines) flanking the putative deletion. The region between the six gapped alignments contains very few reads with high mapping quality, indicating sequence that is present in the reference but lacking in the sample. Surrounding the gapped alignments is the set of deviated pairs identified by the large indel module as supportive evidence for a 413bp homozygous deletion (middle panel). Reads with significantly deviated insert sizes, > 2000bp compared to the population average (1575bp), are highlighted (in pink).



### FAQ – Large indels

## 1

#### Why are there so many more deletions than insertions?

Insertion detection is limited by the average insert size of the library, typically around 1500 base pair for mate-pairs. The average insert size of paired-end libraries is much smaller (around 150 base pairs), and only insertions smaller than the insert size can be detected.

### 2

#### What is the resolution of large indel detection?

This depends on clone coverage, insert size variability, statistical threshold, and library type. For example, mate-pair data detects large insertions and deletions ranging from 30 base pair to 1.2 kbp and 86 bp to 100 kbp, respectively, in human hg18 (McKernan et al, 2009). For paired-end data, deletion sizes range from 100 base pairs to 2 kbp, and insertions are essentially undetectable.

3

Why are there so many large indels around 300 base pair and deletions around 6 kbp?

Alu elements (SINEs) and LINEs, respectively (McKernan et al, 2009).



#### How many large indels can I expect?

The quantity of large indels to expect depends on:

- Genome size
- Clone (read) coverage
- Average insert size
- Significance threshold
- Pairing quality value threshold
- Library type

**Note:** The Human hg18 has 4075 deletions and 1515 insertions.



### 5

#### Why do pooled samples take so long to analyze?

In areas of higher coverage, more significant candidate regions are found, causing more clustering. Analysis at CNVs can slow down as the coverage exceeds 2x.

### 6

#### How long does the module take to run and how much space is required?

Significant algorithmic improvements have cut processing time. Additional parameters and better implementation have reduced lag times associated with too much coverage and zygosity calculation (clustering).

- Non-parallelized human genome runs consisting of approximately 500 million reads typically take two to four hours to run, depending on platform and resource load.
- Running jobs in parallel at one job per chromosome, significantly reduces run time. Intermediate file generation is negligible for BAM file inputs.

# Add Genomic Annotations to Analysis Results

#### This chapter covers:

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### **Overview**

The annotations module is an optional post-processing step available with select  $LifeScope^{TM}$  Genomic Analysis Software analysis modules. The annotations processing makes a copy of the analysis module's GFF file and adds new attributes to the GFF entries in the copied file. The new attributes are taken from information in publicly available sources about the variants in the input, about features intersecting the variants in the input file, or about any biological function potentially changed by the variants.

Annotations processing does not modify the original LifeScope<sup>™</sup> Software analysis module's output file.

Several output files are potentially generated by the annotations processing, depending on the analysis modules involved. One annotations output file contains all of the content of the GFF file generated by the modules, with added attributes for annotations. Another annotations output file is a subset of the previous file, and it contains only those entries that pass filtering requirements that you specified. Annotation processing also produces output files containing only mutated genes, or only SNPs. Several statistics files are also produced.

Two sources of annotation are used:



- A UCSC or Ensembl GTF file, used to determine whether a variant overlaps a gene or exon.
- The National Center for Biotechnology Information (NCBI) dbSNP database, which contains information on SNPs and indels already found by other studies. We use the following fields of the of the dbSNP entries:
  - The reference SNP identifier (rsID)
  - The functional code
  - The corresponding locus ID (the gene)

If a conflict is found between the dbSNP data and the GTF data, both annotations are reported. The annotation module does not attempt to resolve these conflicts.

Annotation sources can be either the refGene GTF file with respect to the human hg18 reference build and the dbSNP 130 build or the human hg19 reference build and the dbSNP 132 build (both available with LifeScope<sup>™</sup> Software), or a source that you provide. Only GTF files and dbSNP data can be used as sources. The data from these three tables in dbSNP is required:

- SNPChrPosOnRef
- SNPContigLoc
- SNPContigLocusId

Annotations processing supports these annotation types:

- DNA features: genes and protein-coding features
- Verified variants in existing database

Post-processing types include:

- Annotate genomic variants with the annotation types that apply
- Filter based on annotation
- Report a list of gene and protein-coding annotations involved the variants
- Report statistics

The annotations module is available with the following modules:

- Resequencing:
  - SNPs
  - Small indels
  - Large indels
  - Human CNV
- Targeted resequencing:
  - SNPs
  - Small indels

**Input file handling** Annotations processing uses the GFF output of an analysis module, but does not modify the GFF file. Annotations are added to a copy of the GFF file.

# FiltersAnnotation processing optionally generates an output file containing only those<br/>entries that fulfill *all* the conditions you specify with annotation filtering options.<br/>Annotation filters are described in Table 58.



By default, the filtering options report all variants (regardless of whether or not they appear in dbSNP and whether or not they occur in exons or genes).

Table 58	Annotation	filtering	options
----------	------------	-----------	---------

Filtering option	Default
Report only variants in exons	Off
Report only variants in genes	Off
Report only variants that appear in dbSNP	Off
Report only variants that do not appear in dbSNP	Off

Table 59 shows the annotation filtering supported in select LifeScope<sup>™</sup> Software modules.

Annotation type	SNP	CNV	Small indel	Large indel
genelD	Yes	Yes	Yes	Yes
exonID	Yes	Yes	Yes	Yes
rsID	Yes	_	Yes	_
functionCode	Yes	—	_	—

 Table 59
 Support for annotation filtering in each variant type

#### Statistics

Table 60 lists the statistics included in the annotations output files and the LifeScope<sup>™</sup> Software modules that support these annotations. See Table 60 for information about SNPs transitions and transversions.

 Table 60
 Statistics generated in annotations output files

Analysis type	Statistics
All supported modules	Number of variants Number of variants per chromosome
(SNPs, CNVs, Indels)	Number of heterozygous variants Number of homozygous variants Number of heterozygous SNPs per chromosome Number of homozygous SNPs per chromosome
SNPs	Number of heterozygous SNPs that are transitions, transversions Number of homozygous SNPs that are transitions, transversions (compared to the reference)
Indels	Indel variant length distribution (negative for deletion, positive for insertion)
CNVs	Copy number distribution CNV length distribution

Analysis type	Statistics
Annotations from dbSNPs	Number of SNPs or indels in dbSNP Number of homozygous SNPs or indels in dbSNP Number of heterozygous SNPs or indels in dbSNP Overall dbSNP concordance (percentage of SNPs or indels in dbSNP) Heterozygous dbSNP concordance (the percentage of heterozygous SNPs or indels found in dbSNP) Homozygous dbSNP concordance (the percentage of homozygous SNPs or indels found in dbSNP)
Annotations from GTF file content	Number and percentage of variants in exons Number and percentage of heterozygous variants in exons Number and percentage of homozygous variants in exons Number and percentage of variants in genes Number and percentage of heterozygous variants in genes Number and percentage of homozygous variants in genes

 Table 60
 Statistics generated in annotations output files (continued)

Table 61 describes how transitions and transversions are determined based on the reference allele and the diBayes output genotype.

 Table 61
 Transitions and transversions

diBayes genotype	Reference allele   second allele	A	C	G	т	Other
А	А	0	TV	TS	TV	0
С	С	TV	0	TV	TS	0
G	G	TS	TV	0	TV	0
Т	Т	TV	TS	TV	0	0
М	AIC	TV	TV	TV	TV	TV
R	AlG	TS	TS	TS	TS	TS
W	AIT	TV	TV	ΤV	ΤV	TV
S	ClG	TV	TV	TV	TV	TV
Y	CIT	TS	TS	TS	TS	TS
K	TIG	TV	TV	TV	TV	TV
	Other	0	0	0	0	0



### Workflows

Annotation is integrated into the resequencing and targeted resequencing standard workflows. Table 62 shows the modules with integrated annotation support, in these workflows.

	, i i i i i i i i i i i i i i i i i i i		J		· I· ·
Workflow	Library type	SNP	CNV	Small indel	Large indel
Genomic resequenci	ng Fragment	Yes	Yes	Yes	_
Genomic resequenci	ng Mate-pair	Yes	Yes	Yes	Yes
Genomic resequenci	ng Paired-end	Yes	Yes	Yes	Yes
Targeted resequenci	ng Fragment	Yes	_	Yes	_
Targeted resequenci	ng Paired-end	Yes	_	Yes	_

Table 62 Standard workflows and analysis modules with integrated annotation support

Example shell	The following are example shell commands to run an analysis with annotations. This
commands	example uses reads and references from the LifeScope <sup><math>TM</math></sup> Software repository.

# log into the shell
lscope.sh shell -u <i>username -w password</i>
# cd to the projects repository
cd /projects
# make a project, and open it
mk ecoli
cd ecoli
# make an analysis, and open it
mk run1
cd run1
# define the analysis type
set workflow genomic.resequencing.frag
# define your input data
add xsq DH10B_ColorSpaceOnly_Unbarcoded_50_Frag.xsq
# specify the reference to be used
set reference hg18
<i># optionally change parameter defaults after this line or</i>
# set analysis-specific parameters
# list the configuration of the analysis
ls
# start the run
run
# list progress information about the run
ls
Because the set reference hg18 command uses a known assembly, reference-
related parameters including annotation parameters are set automatically.
To specify your own annotations files, use the set param command, as shown in these examples:

set annotation.dbsnp.file /path/to/dbsnp/file tertiary/INIfile
set annotation.gtf.file /path/to/gtf/file tertiary/INIfile

Place your set param commands after the set reference command (after the line # optionally change parameter defaults after this line). The INI file to use with the set param command depends on the type of LifeScope<sup>TM</sup> Software analysis you are running. Replace the string *INIfile* with one of these INI file names:

- annotation.cnv.ini
- annotation.dibayes.ini
- annotation.large.indel.ini
- annotation.small.indel.ini

### Annotation sources

Table 63 lists common annotation sources supported by LifeScope<sup>™</sup> Software. Files required to support certain annotations are available with LifeScope<sup>™</sup> Software. These files support the following:

- Annotations based on the hg18 and the dbSNP 130 build
- Annotations based on the hg19 and the dbSNP 132 build

Note: The dbSNP 132 file used is a VCF file containing 1000 Genome data.

Table 63 Annotation source	Table 63	Annotation	sources
----------------------------	----------	------------	---------

Type of annotation	Location and notes		
Variants in existing databases	NCBI Variation dbSNP resources: includes single nucleotide polymorphisms, microsatellites, and small-scale insertions and deletions. dbSNP contains population-specific frequency and genotype data, experimental conditions, molecular context, and mapping information for both neutral polymorphisms and clinical mutations. (This text is taken from NCBI website http:// www.ncbi.nlm.nih.gov/guide/variation.) http://www.ncbi.nlm.nih.gov/snp		
Genes and protein-coding features	RefGene data from UCSC:		
Note: Supported by LifeScope <sup>™</sup>	ftp://hgdownload.cse.ucsc.edu/goldenPath/hg18/database/refGene.txt.gz		
Software only if the data is downloaded in GTF format.	The European Bioinformatics Institute's Alternative Splicing Database Project: http://www.ebi.ac.uk/asd		

**Note:** While every effort is made to provide correct web addresses, Applied Biosystems is not responsible for the changes to these addresses, nor for the data provided by these locations.

### Use custom reference and custom dbSNP files

**Note:** LifeScope<sup>™</sup> Software supports only one format for GTF files. Public sources may have a variety of formats. To use custom GTF files, it may be necessary to download and modify these files into our supported GTF format. This section contains instructions to convert files into the supported GTF format

This section describes the steps required to use a custom reference with  $LifeScope^{TM}$ Software. These steps are an alternative to using the reference-related files available with  $LifeScope^{TM}$  Software.

Two main processes are required (listed here and described below):

- 1. Download and convert or reformat a GTF file. Choose one of the following:
  - Download and convert a UCSC GTF file.
  - Download and reformat an Ensembl GTF file.
- 2. Download and unzip dbSNP files.
- The UCSC GTF file Follow either the steps in this section or in the section "The Ensembl GTF file" on page 273.

After download, a conversion step is required to normalize the file by gene ID as required by LifeScope<sup>™</sup> Software. Genome annotations that are downloaded from the UCSC Genome Browser and converted by the annotation conversion script are optimal because they contain Human Genome Organization (HUGO)-style gene names. HUGO-style gene names allow interpretation when using a genome browser or reading reports.

The annotation conversion script refgene2gff.sh works with the most recent format of refGene.txt files (as of the time of this writing). Some assemblies such as the rat genome use an alternate format for the refGene.txt file. The refgene2gff.sh script does not convert these alternate formats.

Follow these steps to download and convert the UCSC GTF file:

- 1. Download the file from the following URL: ftp://hgdownload.cse.ucsc.edu/goldenPath/hg19/database/refGene.txt.gz
- **2.** Unzip the file with the following command: gunzip refGene.txt.gz
- **3.** Use this command to convert the file to GTF format: refgene2gff.sh -i refGene.txt -o refGene.gtf
- **4.** Specify the file, refGene.gtf, with the annotation.gtf.file parameter. (If you use the set reference *assembly* command and specify a known assembly, the annotation.gtf.file parameter is automatically set.)

The Ensembl GTF Follow either the steps in this section or in the section "The UCSC GTF file" on page 273.

> Ensembl GTF files have the following characteristics that apply to their use with LifeScope<sup>™</sup> Software.

- They are properly normalized by gene and transcript IDs.
- They use gene accession numbers instead of HUGO-style gene names.
- They use unprefixed sequence identifiers, such as 1, 2, 3, ....X, Y, MT.
- They are incompatible with genome reference FASTA files that have UCSC-style sequence IDs with the prefix "chr", for example, chr1, chr2, chr3, ....chrX, chrY, chrM.
- In order to be used as input for annotations with LifeScope<sup>™</sup> Software, they must be reformatted to use UCSC-style gene IDs.

Follow these steps to download and reformat the Ensembl GTF file:

file



- Download the file from the following URL: http://www.ensembl.org/
- Use this command to reformat the GTF file: reformat\_ensembl\_gtf.pl Homo\_sapiens.GR
- **3.** Specify the reformatted file with the annotation.gtf.file parameter. (If you use the set reference *assembly* command and specify a known assembly, the annotation.gtf.file parameter is automatically set.)

**The dbSNPs files** Annotations based on dbSNPs require the data from these three dbSNPs tables: SNPChrPosOnRef, SNPContigLoc, SNPContigLocusId. LifeScope<sup>™</sup> Software is verified to work with the format of these files as of February 2011. The dbSNP files follow this naming pattern:

b<dbSnp\_build\_no>\_<table\_name>\_<reference\_ncbi\_build\_no>.bcp.gz

For example, for build 131 and reference hg19 (NCBI build 37.1), the files required would be:

dbSNPs table	File name
SNPChrPosOnRef	b131_SNPChrPosOnRef_37_1.bcp.gz
SNPContigLoc	b131_SNPContigLoc_37_1.bcp.gz
SNPContigLocusId	b131_SNPContigLocusId_37_1.bcp.gz

Follow these steps to use hg19 dbSNPs files with LifeScope<sup>™</sup> Software annotations:

1. Download the dbSNP files from this site:

http://www.ncbi.nlm.nih.gov/snp

2. Specify the information from these files in your analysis with the annotation.dbsnp.file parameter.

(If you use the set reference *assembly* command and specify a known assembly in the reference repository, the annotation.dbsnp.file parameter is automatically set.)

**3.** Increase the memory and heap space for your analysis, if necessary. See "Set your memory requests" on page 277 for instructions.

"Annotation parameters" on page 277 provides more instructions on running annotations.

### Annotation input files

	This section describes the input files used during annotation processing.
GFFv3 variant file	This file is the output file of a previous analysis module such as SNPs, small indels, large indels, or CNV. The variant file must be in GFFv3 format.
GTF	The input genome annotation file is in GTF format. Refer to the following websites for information on the GTF file format:



#### genes.cse.wustl.edu/GTF2.html genome.ucsc.edu/FAQ/FAQformat.html#format4

The GTF file must match the genome reference to ensure that the analysis modules work correctly.

**IMPORTANT!** Make sure that the GTF file is for the same genome assembly as the genome FASTA reference file, and that matching sequence identifiers are used. Gene and transcript identifiers in the GTF file must be properly normalized. Identifier normalization is a known issue in GTF files from the UCSC Genome Browser, which is a popular source of GTF-formatted annotation. The UCSC GTF files always report the same value for gene and transcript IDs.

Use the LifeScope<sup>™</sup> Software scripts to transform a GTF file into the format required for use with LifeScope<sup>™</sup> Software modules. (See "The UCSC GTF file" and "The Ensembl GTF file" on page 273 for more information.)

Data downloaded from the publicly-available dbSNP database is included in the LifeScope<sup>™</sup> Software reference repository. This data is based on human reference builds hg18 dbSNP version 130 and hg19 dbSNP version 132.

LifeScope<sup>™</sup> Software automatically determines your dbSNP data file from the assembly name you specify, when you use the set workflow *assembly* command. For example:

set workflow hg18
set workflow hg19

You also have the option to specify a different dbSNP database (that you must supply) for your annotations. See "Add new reference files" on page 517 in Appendix G, The Reference Repository.

The following information is parsed from the dbSNP data:

- The refSNP identifier, rsID
- Chromosome number
- Chromosome start coordinate
- Chromosome end
- Whether or not the variant is a SNP
- Function code (consequence to transcript, if any)
- The gene name, locus\_symbol (if available)

#### hg19

dbSNP

The hg19 dbSNP file is dbSNP\_b132\_00-All.vcf.

#### hg18

Table 64 describes the NCBI hg18 files used for annotations. These are a set of three tab-separated files.

Table 64 dbSNP hg18 annotation files

File	Description	Fields used
b <dbsnp build="" number="">_ SNPChrPosOnRef_<reference build<br="">version&gt;.bcp</reference></dbsnp>	A 4-column, tab-separated file, corresponding to the table SNPChrPosOnRef	The first 3 columns: • The rsID • Chromosome • Position
b <dbsnp build="" number="">_ SNPContigLocusId_<reference build version&gt;.bcp</reference </dbsnp>	A 21-column, tab-separated file, corresponding to the table SNPContigLocusId	<ul> <li>snp_id: The rsID</li> <li>fxn_class: The function class code</li> <li>(<i>Optional</i>) locus_symbol: The gene name</li> </ul>
b <dbsnp build="" number="">_ SNPContigLoc_<reference build<br="">version&gt;.bcp</reference></dbsnp>	A 22-column, tab-separated file, corresponding to the table SNPContigLoc	<ul> <li>snp_id: The rsID</li> <li>asn_from, asn_to: Used to determine the feature length</li> <li>loc_type: Used to determine whether an entry is a SNP or an indel</li> </ul>

### Prepare to run annotation processing

Select the required	This section only applies to references <i>not</i> in the reference repository.		
input files	If you supply your own annotation source files, you must know the absolute path to those files.		
	<b>Note:</b> This section does not apply if you specify the human hg18 or hg19 assembly, because this option uses the annotation sources available with LifeScope <sup>™</sup> Software.		
	To include gene and exon annotations, you must know the absolute path to:		
	Your GTF file for gene and exon annotations		
	To include dbSNP annotations, you must know the absolute path to your dbSNP annotation file or files. Your dbSNP annotations might be in a single VCF file, or in a set of three BCP files:		
	<ul> <li>b<dbsnp_build_no>_SNPChrPos_<reference_ncbi_build_no>.bcp</reference_ncbi_build_no></dbsnp_build_no></li> </ul>		
	<ul> <li>b<dbsnp_build_no>_SNPContigLoc_<reference_ncbi_build_no>.bcp</reference_ncbi_build_no></dbsnp_build_no></li> </ul>		
	<ul> <li>b<dbsnp_build_no>_SNPContigLocusId_<reference_ncbi_build_no>.bcp</reference_ncbi_build_no></dbsnp_build_no></li> </ul>		
	You control which types of annotation are generated:		
	• To turn off gene and exon annotations, do not provide a GTF file.		
	<ul> <li>To turn off dbSNP annotations, do not provide the dbSNP files.</li> </ul>		
	<b>Note:</b> If any one of the three required dbSNPs files is missing from a set of three BCP files, no dbSNPs annotation is generated.		

# Set your memory requests

The memory requirements for dbSNPs annotations vary by reference build and dbSNPs build. Table 65 lists the hardware memory requirements for dbSNPs annotations for supported hg and dbSNPs builds.

hg18hg19dbSNP 13022GBNot supporteddbSNP 131Not supported22GBdbSNP 132Not supported22GB

 Table 65
 Hardware memory requirements for dbSNPs annotations

These parameters must be added to your module's properties file. Parameters in a module properties file affect all runs of that module, for all users.

These requirements are for full annotations (dbSNPs as well as exon-based annotations). The annotation module requires considerably less memory when run with exon annotation only (that is, with a GTF file but without a dbSNP file). However, the requirement increases linearly with the number of exons in the GTF file.

Complete the<br/>prerequisitesAdditional prerequisites specific to each type of analysis are described in the<br/>instructions for each analysis module.

### **Annotation parameters**

Table 66 describes the annotation parameters used with LifeScope<sup>™</sup> Software analyses.

For filtered output files, all annotation filter options that are turned on must be fulfilled by the variant entry, for the variant to appear in the output file.

Table 66 Annotation parameters description

Parameter name	Default value	Description	
Annotation source parameters			
annotation.dbsnp.annotate.snp	0	Whether or not to annotate variants with SNP entries from the dbSNP file. Allowed values:	
Nets and a substa CNDs and such		• <b>0</b> : Do not annotate variants with SNP entries from the dbSNP file.	
note: applies only to SNPS and small indels modules		• 1: Annotate variants with SNP entries from the dbSNP file.	
annotation.dbsnp.annotate.indel	0	Whether or not to annotate variants with indel entries from the dbSNP file. Allowed values:	
Nets and a substa CNDs and such		• <b>0</b> : Do not annotate variants with indel entries from the dbSNP file.	
note: applies only to SNPs and small indels modules		• 1: Annotate variants with indel entries from the dbSNP file.	
annotation.indel.border.slack	5	An indel is considered to be matched to a dbSNP indel if the two overlap or if the distance between them is less than or equal to the value of annotation.indel.border.slack.	
		Allowed values: Integers >= 0.	
Annotation filtering parameters			



#### Table 66 Annotation parameters description (continued)

Parameter name	Default value	Description	
annotation.show.only.variants.in.exons	0	When set to 1, filters the annotated output: restricts the Filtered Variant annotation output file (see "Annotation output files" on page 282) to only the variants that overlap any exon.	
		Allowed values:	
		• <b>0</b> : Do not filter the output for variants that overlap any exon.	
		• 1: Restrict output to variants that overlap any exon.	
annotation.show.only.variants.in.genes	0	When set to 1, filters the annotated output: restricts the Filtered Variant annotation output file (see "Annotation output files" on page 282) to only the variants that overlap any gene, even if it does not overlap any exon.	
		Allowed values:	
		• <b>0</b> : Do not filter the output for variants that overlap any gene.	
		• 1: Restrict output to variants that overlap any gene.	
		<b>Note:</b> if the parameter annotation.show.only.variants.in.exons is set to 1, then the setting for the parameter annotation.show.only.variants.in.genes is ignored.	
annotation.show.only.variants.in.dnsnp	0	When set to 1, filters the annotated output: restricts the Filtered Variant annotation output file (see "Annotation output files" on page 282) to only SNPs or small indels that exist in dbSNP.	
		Allowed values:	
		• <b>0</b> : Do not filter the output for variants that <i>appear</i> in dbSNP.	
<b>Note:</b> applies only to SNPs and small indels modules		<ul> <li>1: Restrict output to SNPs or small indels variants that appear in dbSNP.</li> </ul>	
annotation.show.only.variants.not.in. dnsnp	0	When set to 1, filters the annotated output: restricts the Filtered Variant annotation output file (see "Annotation output files" on page 282) to only SNPs or small indels that do not exist in dbSNP.	
		Allowed values:	
Note: applies only to SNDs and small		• <b>0</b> : Do not filter the output variants that <i>do not appear</i> in dbSNP.	
indels modules		<ul> <li>1: Restrict output to SNPs or small indels variants that <i>do not</i> appear in dbSNP.</li> </ul>	
Resource parameters			

memory.request	22gb	Memory request. Include the units gb in the setting.
java.heap.space	21500	Dynamic memory requirement.

Annotations parameters are by convention defined in INI files with the following names and location:

- <analysis>/tertiary/annotation.cnv.ini
- <analysis>/tertiary/annotation.dibayes.ini
- <analysis>/tertiary/annotation.large.indel.ini
- <analysis>/tertiary/annotation.small.indel.ini

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Table 67 describes annotations parameters that are automatically specified by LifeScope<sup>™</sup> Software when run as part of a standard workflow.

 Table 67
 Annotations internal parameter description

Parameter name	Default value	Description
annotation.run	1	Specifies whether or not to add annotations. Allowed values:
		• <b>0</b> : Do not generate annotations.
		• 1: Generate annotations during this analysis.
		The run parameter is set automatically by the shell during a standard workflow. Accept the default for most use.
annotation.dbsnp.file	_	The path to your input dbSNP file or files.
annotation.gtf.file	_	The path to your input genome annotation file.
annotation.output.file.name.stem	The base name of the input GFF	The common prefix for all output file names.
file (without directory path and without file extension).		Supplied by the system when run in a standard workflow.

### **Output annotation**

This section describes the new attributes that annotation adds to the GFF entries from the input file. The added information comes from publicly-available sources about variants in the input, features intersecting the variants in the input file, or biological function potentially changed by the variants.

Table 68 describes the new attributes added as output annotation.

 Table 68
 New attributes added as annotations

Attribute		Description
genelD		Adds a geneID attribute to applicable entries in the output GFF file.
		The following apply to the geneID attribute:
		The attribute is valid for all variants.
		• The attribute's value is the gene descriptor taken from the input GTF file.
		• If there are multiple overlapping genes, the attribute has multiple values (in a comma-separated list).
	Supported modules	Requirement to qualify for annotation
	SNPs Small indels Large indels CNVs	The variant must be within a gene (including introns and UTRs).



Table 68	New	attributes	added	as	annotations	(continued)
						·,

Attribute		Description
exonID		Adds exonID attributes to applicable entries in the output GFF file.
		The following apply to the exonID attribute:
		The attribute is valid for all variants.
		• The attribute's value is of the form < <i>genelD</i> >-< <i>Exon#</i> >, where <i>Exon#</i> is the number of the exon counted within the gene (not within each transcript) in the transcription direction.
		<ul> <li>If there are multiple overlapping exons, the attribute has multiple values (in a comma-separated list).</li> </ul>
		• If two exons start from the same position, the shorter one is listed first.
	Supported modules	Requirement to qualify for annotation
	SNPs Small indels Large indels CNVs	The variant must overlap an exon.
rsID		Adds rsID attributes to applicable entries in the output GFF file.
		The following apply to the rsID attribute:
		The attribute is valid for SNPs and Indels.
		<ul> <li>The attribute's value is the rsID value of the dbSNP entry.</li> </ul>
		The attribute cannot have multiple values.
	Supported modules	Requirement to qualify for annotation
	SNPs	The SNP must appear in dbSNP.
	Small indels	Indels must appear in dbSNP and also must be labeled as InDel in dbSNP.
functionCode		Based on functional codes from dbSNP, annotate SNP entries in the output GFF file.
		The following apply to the SNPType attribute:
		The attribute is valid for only for SNPs.
		• The attribute's value is a pair in which the first value is the locus ID and the second the function code from dbSNP. Function codes are listed in Table 69.
		• The attribute can have multiple values, one for each SNP type that applies to the variant.
	Supported modules	Requirement to qualify for annotation
	SNPs	Must be a SNP.
-	Small indels	Must be a small indel.

Table 69 describes function codes for refSNPs in gene features. This table is taken from the dbSNP website

:http://www.ncbi.nlm.nih.gov/bookshelf/br.fcgi?book=handbook&part=ch5



Functional class	Description	Database code
Locus region	Variation is within 2 kbp 5' or 500 bp 3' of a gene feature (on either strand), but the variation is not in the transcript for the gene. This class is indicated with an "L" in graphical summaries.	1
	<b>Note:</b> As of build 127, function code 1 has been modified into a <i>two-digit code</i> that more precisely indicates the location of a SNP. The two-digit code has function code 1 as the first digit, which keeps the meaning as described above, and 3 or 5 as the second digit, which indicates whether the SNP is 3' or 5' to the region of interest. See function codes 13 and 15 in this table.	
Coding	Variation is in the coding region of the gene. This class is assigned if the allele-specific class is unknown. This class is indicated with a 'C' in graphical summaries.	2
	Note: This code was <i>retired</i> as of dbSNP build 127.	
Coding-synon	The variation allele is synonymous with the contig codon in a gene. An allele receives this class when substitution and translation of the allele into the codon makes no change to the amino acid specified by the reference sequence. A variation is a synonymous substitution if all alleles are classified as contig reference or coding-synon. This class is indicated with a 'C' in graphical summaries.	3
Coding-nonsynon	The variation allele is nonsynonymous for the contig codon in a gene. An allele receives this class when substitution and translation of the allele into the codon changes the amino acid specified by the reference sequence. A variation is a nonsynonymous substitution if any alleles are classified as coding-nonsynon. This class is indicated with a "C" or "N" in graphical summaries.	4
	<b>Note:</b> As of build 128, function code 4 has been modified into a <i>two-digit code</i> that more precisely indicates the nonsynonymous nature of the SNP. The two-digit code has function code 4 as the first digit, which keeps its original meaning, and 1, 2, or 4 as the second digit, which indicates whether the SNP is nonsense, missense, or frameshift. See function codes 41, 42, and 44 in this table.	
mRNA-UTR	The variation is in the transcript of a gene but not in the coding region of the transcript. This class is indicated by a "T" in graphical summaries.	5
	<b>Note:</b> As of build 127, function code 5 has been modified into a <i>two-digit code</i> that more precisely indicates the location of a SNP. The two digit code has function code 5 as the first digit, which keeps its original meaning, and 3 or 5 as the second digit, which indicates whether the SNP is 3' or 5' to the region of interest. See function codes 53 and 55 in this table.	
Intron	The variation is in the intron of a gene but not in the first two or last two bases of the intron. This class is indicated by an "L" in graphical summaries.	6
Splice-site	The variation is in the first two or last two bases of the intron. This class is indicated by a "T" in graphical summaries.	7
	<b>Note:</b> As of build 127, function code 7 has been modified into a <i>two-digit code</i> that more precisely indicates the location of a SNP. The two-digit code has function codes 7 as the first digit, which keeps its original meaning, and 3 or 5 as the second digit, which indicates whether the SNP is 3' or 5' to the region of interest. See function codes 73 and 75 in this table.	
Contig-reference	The variation allele is identical to the contig nucleotide. Typically, one allele of a variation is the same as the reference genome. The letter used to indicate the variation is a "C" or "N", depending on the state of the alternative allele for the variation.	8



Functional class	Description	Database code
Coding-exception	The variation is in the coding region of a gene, but the precise location cannot be resolved because of an error in the alignment of the exon. The class is indicated by a "C" in graphical summaries.	9
NearGene-3	Function Code 13, where:	13
	1: locus region (see function code 1 in this table)	
	3: SNP is 3' to and 0.5 kbp away from gene	
NearGene-5	Function Code 15, where:	15
	1: locus region (see function code 1 in this table)	
	3: SNP is 5' to and 0.5 kbp away from gene	
Coding-	Function Code 41, where:	41
nonsynonymous	4: Coding-nonsynonymous (see function code 4 in this table)	
50130	1: Nonsense (changes to the Stop codon)	
Coding-	Function Code 42, where:	42
nonsynonymous	4: Coding-nonsynonymous (see function code 4 in this table)	
1113361136	2: missense (alters codon to make an altered amino acid in the protein product)	
Coding- nonsynonymous frameshift	Function Code 44, where:	44
	4: Coding-nonsynonymous (see function code 4 in this table)	
	4 (as the second digit): frameshift (alters codon to make an altered amino acid in protein product)	
UTR-3	Function code 53, where:	53
	5: UTR (untranslated region: see function code 5 in this table)	
	3: SNP located in the 3' untranslated region	
UTR-5	Function code 55, where:	55
	5: UTR (untranslated region: see function code 5 in this table)	
	5 (as the second digit): SNP located in the 5' untranslated region	
Splice-3	Function code 73, where:	73
	7: splice site (see function code 7 in this table)	
	3: 3' acceptor dinucleotide	
Splice-5	Function code 75, where:	75
	7: splice site (see function code 7 in this table)	
	5: 5' donor dinucleotide	

#### Table 69 Function codes for refSNPs in gene features (continued)

### Annotation output files

These types of output files are potentially generated by the annotations processing, depending on the LifeScope<sup>TM</sup> Software modules being run:

• Annotated Variants

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	Attribute/	Deceriation		
	Table 70         The Anno	tated Variant and Filtered Variant output file added fields		
	Table 70 describes of the Annotated V	the additional annotation attributes added to the last (ninth) column Variants and Filtered Variants files. Except for rsID, attributes can es listed as a comma-separated list		
	where <i><annotation.output.file.name.stem></annotation.output.file.name.stem></i> is a common prefix generated by the annotations module and used with all output files from the same run.			
	<annotation.output.file.name.stem>_annotated_filtered.gff</annotation.output.file.name.stem>			
	<pre><annotation.output.file.name.stem>_annotated.gff</annotation.output.file.name.stem></pre>			
	<ul> <li>In GFF v3 for</li> <li>Named according</li> </ul>	mat iding to these filename patterns:		
	Ine Annotated Va	mants and riftered variants output mes are:		
Variants output files	A Filtered Variant you specify with t	s output file contains only those entries that fulfill <i>all</i> the conditions he annotation UI options.		
	An Annotated Variants output file contains all the entries of the input GFF file.			
Annotated Variants and Filtered	The Annotated Va (see "Annotation i	riants and Filtered Variants output files are the input GFF files nput files" on page 274) with annotations added.		
	For filtered outpu fulfilled by the var	t files, all annotation-filter options that are turned on must be riant entry for the variant to appear in the output file.		
	The output file con chapter in the sect	ntent is typically labeled within each file, or as described in this ions concerning each annotation output file type.		
	Not all output file the type of analyst "About annotation information on an parameters" on pa	s are generated on each run. Which files are produced depends on is modules being run and on the annotation options you select. See and LifeScope <sup>™</sup> Software modules" on page 297 for more notations available with the different modules, and "Annotation age 277 for a description of annotation options.		
	<ul><li>diBayes tab-c</li><li>diBayes anno</li></ul>	lelimited output file otated tab-delimited output file		
	Mutated Gen	es		
	Variant Statis	tics		
	Filter Variant	s		

Attribute/ Metadata	Description		
Metadata keyword			
Annotation	A metadata line beginning with the word "Annotation" and followed by these tab-separated fields:		
	The date and the sources of annotation used		
	The path to the flat files		
	The dbSNP version, if available		
	The human reference build, if available		

Attribute/ Metadata	Description		
Filtering	A metadata line beginning with the word "Filtering" and followed by these tab-separated fields:		
	• The values "no filtering" or "show only"		
	<ul> <li>One or more of the values for "coding variants", "variants in genes", "variants in dbSNP", and "variants not in dbSNP"</li> </ul>		
Attributes			
genelD	Add a geneID attribute to applicable entries in the output GFF file.		
	The following apply to the geneID attribute:		
	• The attribute is valid for all variants.		
	• The attribute's value is the gene descriptor taken from the input GTI file.		
	If there are multiple overlapping genes, the attribute has multiple value (in a comma-separated list).		
exonID	Adds exonID attributes to applicable entries in the output GFF file.		
	The following apply to the exonID attribute:		
	• The attribute is valid for all variants.		
	• The attribute's value is of the form < <i>GeneID</i> >-< <i>Exon#</i> >, where <i>Exon#</i> is the number of the exon counted within the gene (not within each transcript) in the transcription direction.		
	• If there are multiple overlapping exons, the attribute has multiple values (in a comma-separated list).		
	• If two exons start from the same position, the shorter one is listed first		
rsID	Adds rsID attributes to applicable entries in the output GFF file.		
	The following apply to the rsID attribute:		
	The attribute is valid for SNPs and Indels.		
	• The attribute's value is the rsID value of the dbSNP entry.		
	The attribute cannot have multiple values.		
functionCode	Based on functional codes from dbSNP, annotate SNP entries in the output GFF file.		
	The following apply to the functionCode attribute:		
	• The attribute is valid for only for SNPs.		
	• The attribute's value is a pair in which the first value is the locus ID an the second the function code from dbSNP. Function codes are listed i Table 69.		
	The attribute can have multiple values, one for each SNP type that applie to the variant.		

 Table 70
 The Annotated Variant and Filtered Variant output file added fields (continued)

#### Variant Statistics 4 output file for SNPs i

A Variant Statistics file is a text file containing the statistics about the variants in the input GFF file. The file is named according to this filename pattern:

<annotation.output.file.name.stem>.stats



where <annotation.output.file.name.stem> is a common prefix generated by the annotations module and used with all output files from the same run.

The following is an example of the content of a Variant Statistics file for SNPs:

**************************************	* * * * * * * * * * * * * * T * * * * *	**************	**********	**********
Input File:/dat solid0064_20101 color_lmp_SNP.g Date:Apr 14, 20 Annotation file lifetech/hg18/d Annotation file LifeScope_resou	a/swdev/min: 210_MP_2x60 ff3 11 2:17:54 1 , dbSNP:/da bSNP/dbSNP_1 , Genes and rces//lifete	ita_test/ _T3_1/2.0/color_ PM ta/analysis/Life o130.tab Exons, GTF:/dat ech/hg18/GTF/ref	lmp/outputs Scope_reso a/analysis Gene.20090	s/dibayes/ urces/ / 513.gtf
* * * * * * * * * * * * * * * *	* * * * * * * * * * * *	* * * * * * * * * * * * * * * * *	* * * * * * * * * * *	* * * * * * * * * *
Statistics Over	view			
**********	* * * * * * * * * * * * *	* * * * * * * * * * * * * * * * *	* * * * * * * * * * * *	* * * * * * * * * * *
Basic Statis	tics			
Number of varia	nts			3072881
Number of heter	ozvgous var:	iants		1639649
Number of homoz	ygous varia	nts		1433232
Variant-Spec	ific Statis	tics (SNP)		
Number of trans	ition SNPs			2070418
Number of trans	ition heter	ozygous SNPs		1103589
Number of trans	ition homozy	ygous SNPs		966829
Number of trans	version SNP:	5		1002463
Number of trans	version hete	erozygous SNPs		536060
Number of trans	version homo	ozygous SNPs		466403
Transition:Tran	sversion rat	tio	2.065	: 1.000
dbSNP Annota	tion Statis	tics		
NT 1 C	CITE	' 11 CMD		0010001
Number of	SNP	s in dbSNP		2913081
Number of heter	ozygous SNP:	s in absNP		1506/40
Number of homoz	ygous SNP:	s in dbSNP		1406341
dbSNP concordan	ce			94.80%
dbSNP neterozyg	ous concorda	ance		91.898
absirp nomozygou	s concordano	ce		98.12%
Per-Chromoso	me Statistic	cs		
nVar. Number of	variants			
nHetVar. Number	of heteroz	voous variante		
nHomVar. Number	of homozyce	ous variants		
ivanoei	52 1101102399			
contig	nVar	nHetVar	nHomVar	



chr1	233502	125860	107642
chr2	247682	132717	114965
chr3	210137	115661	94476
chr4	225996	118326	107670
chr5	185137	102825	82312
chr6	191793	104138	87655
chr7	170500	94092	76408
chr8	160126	86553	73573
chr9	128205	70291	57914
chr10	157345	85408	71937
chr11	154271	80792	73479
chr12	139569	72915	66654
chr13	118842	60095	58747
chr14	93526	48742	44784
chr15	90141	46673	43468
chr16	95819	53400	42419
chr17	79447	44311	35136
chr18	85824	43117	42707
chr19	59142	31904	27238
chr20	65136	38340	26796
chr21	45656	26098	19558
chr22	36633	20355	16278
chrX	76085	18909	57176
chrY	22328	18127	4201
chrM	39	0	39

#### Variant Statistics output file for small indels

A Variant Statistics file is a text file containing the statistics about the variants in the input GFF file. The file is named according to this filename pattern:

<annotation.output.file.name.stem>\_stats
where <annotation.output.file.name.stem> is a common prefix
generated by the annotations module and used with all output files from the
same run.

The following is an example of the content of a Variant Statistics file for small indels:

***************************************	ł
LIFESCOPE REPORT	
***************************************	ł

Input File:/data/swdev/minita\_test/ solid0064\_20101210\_MP\_2x60\_T3\_1/2.0/color\_lmp/outputs/ small.indel/color\_lmp.gff3 Date:Apr 14, 2011 10:01:01 AM Annotation file, dbSNP:/data/analysis/LifeScope\_resources/ lifetech/hg18/dbSNP/dbSNP\_b130.tab Annotation file, Genes and Exons, GTF:/data/analysis/ LifeScope\_resources//lifetech/hg18/GTF/refGene.20090513.gtf

-- Basic Statistics -----

Number of variants	296	296183		
Number of heterozygo	ous variants 194	194752 100934		
Number of homozygous	s variants 100			
Variant-Specific	Statistics (InDel)			
Distribution of InDe	el length			
Length(bases)	Number			
(-Inf:-500]	0			
(-500:-490]	14			
(-490:-480]	8			
(-480:-470]	13			
(-470:-460]	21			
(-460:-450]	14			
(-450:-440]	18			
(-440:-430]	17			
(-430:-420]	18			
(-420:-410)	18			
$(-410 \cdot -400)$	15			
(-400:-390]	24			
$(-390 \cdot -380)$	21			
$(-380 \cdot -370)$	24			
(-370, -360)	30			
(-360, -350]	30			
(-500; -550]	53			
(-340, 220)				
(-340:-330]	113			
(-330:-320]	202			
(-320:-310]	181			
(-310:-300]	80			
(-300:-290]	43			
(-290:-280]	52			
(-280:-270]	42			
(-270:-260]	56			
(-260:-250]	49			
(-250:-240]	35			
(-240:-230]	39			
(-230:-220]	35			
(-220:-210]	33			
(-210:-200]	37			
(-200:-190]	34			
(-190:-180]	30			
(-180:-170]	40			
(-170:-160]	49			
(-160:-150]	42			
(-150:-140]	55			
(-140:-130]	78			
(-130:-120]	55			
(-120:-110]	68			
(-110:-100]	51			
(-100:-90]	60			
(-90:-80]	61			
(-80:-70]	94			
(-70:-60]	107			

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(-60:-50]	115
(-50:-40]	229
(-40:-30]	332
(-30:-20]	1064
(-20:-19]	220
(-19:-18]	392
(-18:-17)	268
(-17, -16]	568
$(-16 \cdot -15)$	436
$(-15 \cdot -14)$	624
$(-14 \cdot -13)$	541
$(-13 \cdot -12)$	1126
$(-12 \cdot -11)$	1009
(-11, -10)	1636
(-10, -9)	1000
(-10, -9]	1270
(-9; -0]	1566
(-8:-7)	2400
	5490
(-0:-5]	2420 16600
(-5:-4]	11020
(-4:-3]	11836
(-3:-2]	22210
(-2:-1)	//896
(-1:0]	
	65663
(1:2)	19317
(2:3)	14044
	25082
(4:5)	3/49
	2199
(6:7)	1011
(/:8]	1596
(8:9]	892
(9:10]	1169
	705
(11:12)	1289
(12:13]	803
(13:14]	1264
(14:15]	3547
(15:16]	192
(16:17]	164
(17:18]	123
(18:19]	141
(19:20]	268
(20:30]	530
(30:+Inf)	0

-- dbSNP Annotation Statistics -----

Number	of		InDels	in	dbSNP	246619
Number	of	heterozygous	InDels	in	dbSNP	150203
Number	of	homozygous	InDels	in	dbSNP	96105
dbSNP o	cond	cordance				83.27%
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	dbSNP heterozygous concordance dbSNP homozygous concordance								
	Per-Chro	omosome Statis	tics						
	nVar: Number of variants nHetVar: Number of heterozygous variants nHomVar: Number of homozygous variants								
	contig	nHomVar							
	chr1	22753	15323	7385					
	chr2	24215	16211	7975					
	chr3	20327	13613	6686					
	chr4	22322	14798	7485					
	chr5	18224	12212	5989					
	chr6	19046	12807	6213					
	chr7	16887	11032	5824					
	chr8	14530	9625	4889					
	chr9	11539	7926	3601					
	chr10	14864	9836	5000					
	chr11	14951	9502	5428					
	chr12	14000	9180	4793					
	chr13	11885	7685	4184					
	chr14	9439	6105	3323					
	chr15	8945	5912	3022					
	chr16	7705	5304	2382					
	chr17	7849	5323	2508					
	chr18	8605	5530	3066					
	chr19	5866	3825	2026					
	chr20	6107	4120	1977					
	chr21	5067	2977	2072					
	chr22	3587	2301	1280					
	chrX	6351	2650	3675					
	chrY	1118	954	151					
	chrM	1	1	0					
Variant Statistics output file for large	A Variant Statistics file is a text file containing the statistics about the variants in the input GFF file.								
indels	The file is named according to this filename pattern:								
	<annota< td=""><td>tion.output.f</td><td>ile.name.stem&gt;_</td><td>stats</td><td></td></annota<>	tion.output.f	ile.name.stem>_	stats					
	where <i><annotation.output.file.name.stem></annotation.output.file.name.stem></i> is a common prefix generated by the annotations module and used with all output files from the same run.								
	The following	is an example of t	ne content of a Varia	nt Statistics file for	large indels:				
	* * * * * * * * * * * *	* * * * * * * * * * * * *	* * * * * * * * * * * * * * *	* * * * * * * * * * * * * *	*******				
	LIFESCOPE H	REPORT							



```
Input File:/data/swdev/minita_test/
solid0064_20101210_MP_2x60_T3_1/2.0/color_lmp/outputs/
large.indel/large-indels.gff3
Date:Apr 14, 2011 10:00:05 AM
Annotation file, dbSNP:/data/analysis/LifeScope_resources/
lifetech/hg18/dbSNP/dbSNP_b130.tab
Annotation file, Genes and Exons, GTF:/data/analysis/
LifeScope_resources//lifetech/hg18/GTF/refGene.20090513.gtf
Statistics Overview
                   -- Basic Statistics ------
Number of variants
                                                   21
Number of heterozygous variants
                                                    2
Number of homozygous variants
                                                   19
-- Variant-Specific Statistics (InDel) ------
Distribution of InDel length
Length(bases)
                         Number
(-Inf:-500000]
                              0
(-500000:-100000]
                              0
(-100000:-50000]
                              0
(-50000:-20000]
                              0
(-20000:-10000]
                              0
(-10000:-5000]
                              0
(-5000:-1000]
                              0
(-1000:-500]
                              0
(-500:-200]
                              7
                              5
(-200:-100]
                              2
(-100:0]
(0:100]
                              2
                              4
(100:200]
(200:500)
                              1
                              0
(500:1000]
(1000:5000]
                              0
                              0
(5000:10000]
(10000:20000]
                              0
(20000:50000]
                              0
(50000:100000]
                              0
(100000:500000]
                              0
(500000:+Inf)
                              0
-- dbSNP Annotation Statistics ------
Number of
                    InDels in dbSNP
                                                    0
                                                    0
Number of heterozygous InDels in dbSNP
Number of homozygous InDels in dbSNP
                                                    0
dbSNP concordance
                                                 0.00%
dbSNP heterozygous concordance
                                                 0.00%
```

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	dbSNP homozygo		0.00%						
	Per-Chromosome Statistics								
	nVar: Number o	of var	iants						
	nHetVar: Numbe	er of	heterozy	gous variants	5				
	nHomVar: Numbe	er of	homozygo	ous variants					
	contig	nVa	r	nHetVar	nHomVar				
	chr2		1	0	1				
	chr9		1	0	1				
	chr10		7	2	5				
	chr18		4	0	4				
	chr19		2	0	2				
	chrY		4	0	4				
	chrM		2	0	2				
Variant Statistics output file for CNVs	A Variant Statistics file is a text file containing the statistics about the variants in the input GFF file.								
	The file is named according to this filename pattern:								
	<annotation.output.file.name.stem>_stats</annotation.output.file.name.stem>								
	where <annotation.output.file.name.stem> is a common prefix generated by the annotations module and used with all output files from the same run.</annotation.output.file.name.stem>								
	The following is an example of the content of a Variant Statistics file for CNVs:								
	****								
	LIFESCOPE REPORT								
	***************************************								
	Input File:/data/swdev/minita_test/ solid0064_20101210_MP_2x60_T3_1/2.0/color_lmp/outputs/cnv/ OutputCNVs.gff3								
	Date:Apr 14, 2011 9:58:12 AM								
	Annotation file, dbSNP:/data/analysis/LifeScope_resources/ lifetech/hq18/dbSNP/dbSNP b130.tab								
	Annotation file, Genes and Exons, GTF:/data/analvsis/								
	LifeScope_resources//lifetech/hg18/GTF/refGene.20090513.gtf								
	* * * * * * * * * * * * * * * *	* * * * * *	* * * * * * * *	* * * * * * * * * * * * * * *	****	* * * * * * * * * *			
	Statistics Ove	erview *****	******	* * * * * * * * * * * * * * *	*****	* * * * * * * * * *			
	Basic Stat	istics							
	Number of var	iants				120			
	Variant-Spe	ecific	Statist	cics (CNV)					
	Distribution of Copy_Number	of CNV	' copy nu Number	umber					



0	5
1	20
2	23
3	20
4	22
5	8
6	9
7	1
8	2
9	10
>9	0

Distribution of CNV length	
Length(bases)	Number
(0:1000]	0
(1000:2000]	0
(2000:5000]	0
(5000:10000]	0
(10000:20000]	41
(20000:50000]	54
(50000:100000]	11
(100000:500000]	10
(500000:+Inf)	4

-- Per-Chromosome Statistics -----

nVar: Number of variants

contig	nVar
chr1	7
chr2	5
chr3	11
chr4	4
chr5	7
chr6	11
chr7	4
chr8	5
chr9	3
chr10	3
chr11	8
chr12	4
chr13	3
chr14	2
chr15	4
chr16	2
chr17	4
chr19	4
chr20	1
chr22	2
chrX	17
chrY	9



#### Mutated Genes output file The Mutated Genes output file contains the list of genes whose coding regions were modified by at least one of the variants in the input file. The Mutated Genes file is a text file in tab-separated format. The file contains an entry per row, and one gene per entry. The first 12 columns are the same as the columns in the BED file (genes.bed). Additional columns are added by the annotations processing.

The file is named according to this filename pattern:

<annotation.output.file.name.stem>\_genes.tab

where *<annotation.output.file.name.stem>* is a common prefix generated by the annotations module and used with all output files from the same run.

Table 71 lists the information included in the Mutated Genes output file.

 Table 71
 Description of the Mutated Genes output file

Field number	Field name	Description
1	Chromosome/Contig	The contig of the gene.
2	Start	The left-most coordinate of the gene.
3	End	The right-most coordinate of the gene.
4	GenelD	The gene identifier as provided by the GTF file.
5	Score	A score between 0 and 1000. The score is a normalized value for the number of coding variants such that all numbers scale between 0 and 1000. The score is equal to the number of coding variants for this entry divided by the maximum number of coding variants for any gene.
6	Strand	The strand of the gene.
7	Start	The left-most coordinate of the gene.
8	End	The right-most coordinate of the gene.
9	Variant color	The color depends on the type of the variant that overlaps this gene:
		Red: SNP
		Green: Indel (small indels, large indels)
		Cyan: CNV
10	Number of exons	The number of exons of the gene.
11	Exon start position list	A comma-separated list of start-coordinates for exons in the gene. All coordinates are relative to the start position of the gene.
12	Exon length list	A comma-separated list of the lengths of the exons in the gene.
13	Total exon length	The sum the length of all exons.
14	Number of coding variants	The number of variants overlapping at least one exon.
15	Number of variants	The number of variants overlapping the gene.
16	Variant type	The type of the variant. (One type is listed per line.)
17	Variant start list	A comma-separated list for start coordinates of variants overlapping the gene. All coordinates are relative to the start position of the gene.
18	Variant length list	A comma-separated list of lengths of the variants overlapping the gene.

#### Potential uses of a Mutated Genes file

You can use a spreadsheet application or scripts to remove unwanted columns. First save a backup of your output file. The following are examples of uses for a Mutated Genes file:

- The first 12 columns provide information for visualization of modified genes and exons.
- Column 4, the GeneID, provides a list of modified genes. This column provides a gene list to focus on for enrichment analysis.
- A combination of the first 10 and the last 2 fields provides information useful for visualizing variants that overlap the gene.
- Fields 14 and 15 provide the number of variants that overlap exons and the number of variants that overlap the gene. With these columns you can filter by the number of variants overlapping the exons in the gene. With the GeneID in column 4, this information identifies genes to focus on for further analysis.

#### (Optional) View a Mutated Genes output file

Open the BED file in the genomic browser.

#### Example of a Mutated Genes output file

The following is a truncated example (3 lines) of a Mutated Genes output file:

Chromosome/Contig Start End GeneID Score Strand Start End Variant color Number of Exons Exon start position list Exon length list Total exon length Number of coding variants Number of variants Variant type Variant start list Variant length list chr1 752927 779603 LOC643837 0 + 752927 779603 255,0,0 0,1319,19970,24243,24987,25707,25893 7 92,102,153,184,96,132,784 1543 2 15 SNP 3041,5189,6765,12362,13482,14058,15505,17721,20240,21986,22925, 24198,24335,25649,26192 1,1,1,1,1,1,1,1,1,1,1,1,1,1,1,1 chr1 850984 869824 SAMD11 0 850984 869824 + 0,181,4414,5298,10031,13299,13534,15403,16395,1 255,0,0 14 6669,16818,17512,17957,18167 60,92,182,51,125,90,186,163,116 ,79,500,125,111,674 2554 1 3 SNP 1003,6463,18566 1,1,1 chr1 945366 981355 AGRN 0 945366 981355 + 360, 2078, 15154, 20542, 21050, 21355, 21833, 23116, 23415, 2 255,0,0 3700,23986,24211,25038,25236,25610,25841,26037,26274,26697,2720 4,27450,27653,27889,28744,29113,29443,29780,30110,30304,30603,3 1130,31330,31605,33630,34325,34701 251,262,48,216,225,225,20 7,219,195,201,149,106,117,165,144,125,106,339,138,128,115,120,3 54, 193, 216, 230, 135, 97, 165, 112, 117, 193, 88, 225, 104, 1289 7319 16631,17955,18722,18991,31697,32807,33000, SNP 1 8 34914 1,1,1,1,1,1,1,1

diBayes tabdelimited output file

The first eight columns of this output file are the same as the input GFF file. The ninth column in the input file is replaced by individual columns for each attribute, as shown in Table 72. This output file is generated only on SNPs module runs.

This is a tab-separated text file. The file is named according to this filename pattern:

# 14

#### <annotation.output.file.name.stem>.tab

where <annotation.output.file.name.stem> is a common prefix generated by the annotations module and used with all output files from the same run.

Table 72 Attributes in the ninth column

Attribute	Description
Seqid	The string ID of the sequence to which the start and end coordinates refer.
Source	The source of the data.
Туре	Sequence ontology derived type for this variation. For diBayes, this is always SNP.
Start	Start position of the SNP.
End	End position of the SNP.
Score	Calculated p-value of the SNP.
Strand	Not used.
Phase	Not used.
Genotype	Genotype in the form of IUB codes for bases observed in all the reads. The base of the reference sequence at the current position.
Coverage	The number of the reads that cover the current position.
refAlleleCounts	The number of reads of the reference allele at the current position.
refAlleleStarts	The number of different start positions of reads having the reference allele at the current position.
refAlleleMeanQV	The mean of quality values of all reference allele reads at the current position.
novelAlleleCount	The number of reads of the most abundant non-reference allele at the current position.
novelAlleleStarts	The number of different start positions of reads having the most abundant non-reference allele at the current position.
novelAlleleMeanQV	The mean of quality values of all novel allele reads.
mostAbundantAlleleDiColor2	The most abundant allele in the reads (not necessarily the reference allele) in dicolor encoding (for example, 00, 01, 32, 33), of 16 possible dicolors.
secondAbundantAlleleDiColor3	The second most abundant allele in the reads.
Het	Heterozygosity flag. Allowed values: 0,1
	• 0: Homozygous SNP
	• 1: Heterozygous SNP

The following is a truncated example of diBayes tab-delimited output file content:

Seqid Source Type Start End Score Strand Phase genotype reference coverage refAlleleCounts refAlleleStarts refAlleleMeanQV novelAlleleCounts novelAlleleStarts novelAlleleMeanQV mostAbundantAlleleDiColor2 secondAbundantAlleleDiColor3 het chr1 SOLiD\_diBayes SNP 10007 10007 0.806886 G . Α 1 16 2 2 25 21 21 3 1 0 chr1 SOLiD\_diBayes SNP 224472 224472 0.879526 С т . . 3 1 1 26 2 2 24 10 10 0 chr1 SOLiD\_diBayes SNP 553488 553488 0.0625 С т . . 3 0 0 0 2 2 25 20 20 0



chr1 SOLiD_diB	ayes SNP	556955	556955 0.	.095335	•	Y	Т
10 7 6 14	2 2	26	13 31	1			
chr1 SOLiD_di	ayes SNP	558326	558326 0.	.093729	•	R	А
8 6 6 26	2 2	22	03 21	1			
chr1 SOLiD_di	ayes SNP	558554	558554 0.	.0761	•	Y	Т
14 11 9 20	3 3	22	03 21	1			

#### diBayes annotated tab-delimited output file

This output file is the same as the diBayes tab-delimited output file (see "diBayes tabdelimited output file" on page 294), with four additional columns added. As with the diBayes tab-delimited output file, the first eight columns of this output file are the same as the input file. The ninth column in the input file is replaced by individual columns for each attribute, as shown in Table 73. This output file is generated only on SNPs module runs.

This is a tab-separated text file. The file is named according to this filename pattern:

<annotation.output.file.name.stem>\_annotated.tab

where *<annotation.output.file.name.stem>* is a common prefix generated by the annotations module and used with all output files from the same run.

Table 73 Additional columns in the diBayes annotated tab-delimited output file

Field	Description
genelD	The gene id as provided by the GTF file.
exonID	The exon id formed by concatenating the GeneID and the exon index number in the list of exons of the gene sorted in transcription order. Transcription order ties are broken listing by the shortest exon first ( <geneid>-<exon #="">).</exon></geneid>
rsID	The dbSNP id of the SNP.
functionCode	The dbSNP functional code.

The following is a truncated example of diBayes annotated tab-delimited output file content:

Seqid	Sourc	е Т	ype	Stai	st Er	ıd	Score	Strar	nd Pl	hase g	genot	zype	
refere	ence	cove	erage	refA	Allel	leCo	ounts r	efAll	leleSt	tarts			
refAll	LeleMe	anQV	nove	elAll	LeleC	Cour	nts nov	elAll	leleSt	tarts			
novelA	Allele	Mean	iQV ma	ostAk	ounda	ant <i>I</i>	AlleleD	iCold	or2				
second	lAbund	antA	llele	eDiCo	olor3	8 he	et gene	ID ex	onID	rsID	func	tion	.Code
chr1	SOLiD	_diE	Bayes	SNP	1000	)7	10007	0.80	06886			А	G
3	1	1	16	2	2	25	21	21	0	WASH	5P		
chr1	SOLiD	_diE	Bayes	SNP	2244	ł72	224472	0.87	79526			С	т
3	1	1	26	2	2	24	10	10	0				
chr1	SOLiD	_diE	Bayes	SNP	5534	188	553488	0.06	525			С	т
3	0	0	0	2	2	25	20	20	0				
chr1	SOLiD	_diE	Bayes	SNP	5569	955	556955	0.09	95335			Y	т
10	7	6	14	2	2	26	13	31	1	9326	622		
chr1	SOLiD	_diE	Bayes	SNP	5583	326	558326	0.09	93729			R	A
8	6	6	26	2	2	22	03	21	1	2153	587		
chr1	SOLiD	_diE	Bayes	SNP	5585	554	558554	0.07	761			Y	т
14	11	9	20	3	3	22	03	21	1	8179	256		

chr1	SOLiD_	_di	Bayes	SNP	658	055	658055	0.06	25	•	т	С
3	0	0	0	2	2	22	20	20	0			

diBayes filtered annotated tabdelimited output file This file is the same as the diBayes annotated tab-delimited output file, except that it contains only those entries that fulfill all the conditions set by the your annotation filtering settings (described in "Annotation parameters" on page 277).

This is a tab-separated text file. The file is named according to this filename pattern:

<annotation.output.file.name.stem>\_annotated\_filtered.tab

where <annotation.output.file.name.stem> is a common prefix generated by the annotations module and used with all output files from the same run.

This output file is generated only on SNPs module runs.

The following is a truncated example of diBayes filtered annotated tab-delimited output file content:

Segid Source Type Start End Score Strand Phase genotype reference coverage refAlleleCounts refAlleleStarts refAlleleMeanQV novelAlleleCounts novelAlleleStarts novelAlleleMeanQV mostAbundantAlleleDiColor2 secondAbundantAlleleDiColor3 het geneID exonID rsID functionCode chr1 SOLiD\_diBayes SNP 556955 556955 0.095335 Т Y • 6 13 9326622 10 7 14 2 2 26 31 1 chr1 SOLiD diBayes SNP 558326 558326 0.093729 . R Α 8 6 6 26 2 2 22 03 21 1 2153587 chr1 SOLiD\_diBayes SNP 558554 558554 0.0761 т Υ . 14 11 9 20 3 3 22 03 21 1 8179256 chr1 SOLiD\_diBayes SNP 708249 708249 0.003906 G Α . 0 3 23 30 10900602 5 0 0 3 30 0 chr1 SOLiD diBayes SNP 708418 708418 0.0625 . С т 2 Ο 0 0 2 15 13 13 Ο 10751453 2 chr1 SOLiD diBayes SNP 710103 710103 0.003906 С Т 0 17 01 3121393 3 0 0 3 3 01 0 chr1 SOLiD diBaves SNP 713754 713754 0.003906 . С G 0 0 0 3 3 21 33 33 Ω 2977670

## About annotations and LifeScope<sup>™</sup> Software modules

This section describes which annotation functionality is available with the various LifeScope<sup>™</sup> Software modules.

Table 74 shows which LifeScope<sup>™</sup> Software modules support the annotation attributes.

Label	SNPs	CNV	Small indel	Large indel	
genelD, exonID	Yes	Yes	Yes	Yes	
rsID - SNPs	Yes	_	_	—	
rsID – Indels	—	—	Yes	_	

Table 74 Support for annotation labels

Label	SNPs	CNV	Small indel	Large indel
functionCode	Yes	_	Yes	—

Table 75 lists which LifeScope<sup>™</sup> Software modules support annotation filtering.

Table 75 Support for the annotation filtering options

Label	SNPs	CNV	Small indel	Large indel
Only in exons	Yes	Yes	Yes	Yes
Only in genes	Yes	Yes	Yes	Yes
Only not in dbSNP	Yes	_	Yes	_
Only in dbSNP	Yes	_	Yes	_

Table 76 lists which LifeScope<sup>™</sup> Software modules support for the various annotation statistics. Heterozygous and homozygous statistics are included for applicable variants.

 Table 76
 Support for annotation statistics

Label	SNPs	CNV	Small indel	Large indel
Number of variants (total and per chromosome)	Yes	Yes	Yes	Yes
Transitions and transversions	Yes	_	_	_
Variant length distribution	-	Yes	Yes	Yes
Copy number distribution	-	Yes	_	_
dbSNP concordance	Yes	_	Yes	_
Overlapping exons (number and percent)	Yes	Yes	Yes	Yes
Overlapping genes (number and percent)	Yes	Yes	Yes	Yes

## **FAQ** - Annotations

### 1

#### How are errors with input files handled?

If an entry in the GTF file is malformed then the entry would be skipped. If there is a problem with the file then the entire file would be skipped. If no file is provided or is skipped then the GTF annotation is turned off. Similarly, if one of the dbSNP files is missing then the dbSNP annotation is turned off. The \*.stats files may still show zeros in the dbSNP fields if the dbSNP annotation is turned on.



#### Can I add annotations to my GFF file from an earlier version of the software?

Yes. Run the annotations as a standalone module. Set the parameter to point to your existing GFF file.

## 3

2

Why is the function code missing locus information in hg19 while the information is in hg18? Can I do anything to get the locus information?

While doing annotation with hg19 we use the VCF file, which does not have locus information yet. You can use the dbSNP b132 BCP files to get the locus information, but please note that the indels are not completely annotated in that file. Also, using the BCP files requires more memory than the VCF file.



# PART IV Targeted Resequencing

# Run Targeted Resequencing and Enrichment Analyses

This chapter covers:

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Enrichment statistics input files	307
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Targeted resequencing read selection algorithm	313
Run other targeted resequencing modules	314

## **Overview**

#### Targeted Targeted resequencing is a set of analyses designed for target-enriched sequencing data. Given a set of coordinates representing a region of interest in a genome, a library resequencing may be enriched for reads within those regions prior to sequencing. The workflow for analysis of target enriched data is nearly identical to a standard resequencing workflow (see Chapter 8, "Run a Resequencing Mapping Analysis" on page 113). In fact, adequate results can be obtained using the standard workflow. The targeted resequencing application consists of many of the same modules as standard resequencing analysis. However, two additional modules are enabled by default for targeted resequencing analysis, and several modules have specific parameters set that are optimized for analysis of targeted resequencing data. The SAET module is enabled by default for enriched runs, as the smaller target size makes spectral error correction viable, even if it may not be practical for unenriched sequencing runs on a large reference sequence. Targeted resequencing mapping produces a filtered BAM file for variant calling. The variant calling modules diBayes and small indel are run on the filtered data sets instead of the entire read population, and limit SNP and indel calls to the target region. The input list of regions of interest (ROI) is provided by the user. Supported ROI input files include third-party files of baits, probes, or tiles. The input ROI files can be in BED format or in a text file as a list of "chr:start-end" values. The targeted resequencing mapping module sorts the input ROI file by chromosome and start position, and makes the resulting file available as output. Figure 29 describes the targeted resequencing analyses.



#### Figure 29 Components of targeted resequencing analyses



The targeted resequencing analyses support data from these library types:

- Fragment
- Paired-End
- Mate-Pair
- Mate-pair libraries are accepted, but pairing information is not considered for mate-pair libraries.
- Libraries generated with up to 96 barcodes Individual sequencing and enrichment reports are provided for each barcoded library.

Targeted resequencing output files are compatible with third-party genome browsers, including the IGV browser.

The targeted resequencing mapping module provides analysis of sequencing runs that have been run through targeted enrichment.

Enrichment statistics

Targeted resequencing support optional enrichment statistics, which provide a means to assess enrichment platform performance by:

- Looking at variations in coverage, both across all targets and on a per-target basis.
- Addressing the uniformity and completeness of coverage within the target region.
- Calculating the degree of enrichment.

# Examples of running the enrichment module

In a standard workflow The following standard workflows provide examples of running the enrichment module:

- targeted.resequencing.frag
- targeted.resequencing.pe



The following are example shell commands using reads and references from the LifeScope<sup>™</sup> Software repository.

```
lscope.sh shell -u username -w password
# cd to the projects repository
cd /projects
# make a project, and open it
mk ecoli
cd ecoli
# make an analysis, and open it
mk run1
cd run1
# define the analysis type
set workflow targeted.resequencing.frag
# define the input data
add xsg DH10B ColorSpaceOnly Unbarcoded 50 Frag.xsg
# specify the reference for the analysis
set reference hg19
# optionally change parameter defaults here
set analysis.regions.file /data/results/readsdata/dh10b.bed
# list the configuration of the analysis
ls
# start the run
run
# list progress information about the run
15
```

The command to set the analysis.regions.file does not include the optional INI file argument. As written, the command sets the analysis.regions.file parameter in both the secondary and tertiary global.ini files. In this way the parameter is available to all analysis modules in the workflow. If the parameter is instead set in only one INI file, for example the enrichment.ini file, then the regions of interest file is available to the enrichment module, but not available to the SAET module and other modules in the workflow.

See for Chapter 7, "Run a Standard Workflow Analysis" on page 97 for more information on running standard workflows. Chapter 6, "Run a Command Shell Analysis" on page 71 for information on shell commands and syntax.

**As a demo analysis** The optional examples download includes an example of how to run the enrichment module by itself, as a single analysis that is not part of a standard workflow. The demo example is not recommended for general use.

See Appendix E, "Demo Analyses" on page 507 for information on running the enrichment module as a standalone analysis.

## **Enrichment parameters**

Table 77 describes the parameters used with the enrichment statistics module.

In order to change a parameter value in your analysis, use the set param shell command to replace the line # *optionally change parameter defaults here* in the example commands in "In a standard workflow". For instance, to turn off the summary report, use this shell command:



#### set enrichment.summary.report 0 /tertiary/enrichment.ini

#### Table 77 Enrichment statistics parameters

Parameter name	Default value	Description				
Optional module parameters	Optional module parameters					
enrichment.run	1	Whether or not to run the enrichment module. Allowed values:				
		• <b>0</b> : Do not run the enrichment module.				
		• 1: Run the enrichment module during this analysis.				
		The run parameter is set automatically by the shell during a standard workflow. Accept the default for most use.				
enrichment.extend.bases	0	The number of bases on either side of the target region in which alignments may be captured. The target sequence is extended by this number of bp on either side. Used by both target capture and target coverage statistics.				
		Example: enrichment.extend.bases=0				
		Allowed values: Integers >= 0.				
enrichment.minimum.mapping. score	8	Minimum mapping quality value (MAPQ) allowed for aligned reads. Reads below this threshold are not used.				
		Allowed values: Integers >= 0–80.				
enrichment.minimum.target. overlap	0.0001	The fraction of an alignment that must be overlapped by a target in order to be classified as on target. "0.50" is interpreted as 50%, and implies that the midpoint of the read must fall within target region.				
		Allowed values: 0.0001 < X <= 1.0				
		Example: enrichment.minimum.target.overlap=0.50				
enrichment.minimum.target. overlap.reverse	1.0	The fraction of a Reverse Read (F5) alignment which must be overlapped by the target in order to be classified as on-target. To require 100% alignment, use 1.0 (the default).				
		Example: enrichment.minimum.target.overlap.reverse=.9				
		Allowed values: 0.0001 < X <= 1.0				
Required output parameters		I				
enrichment.summary.report	true	Whether or not to create the summary statistics file. Allowed values:				
		• false, 0: Do not create the summary statistics.				
		• true, 1: Create the summary statistics file.				
enrichment.target.coverage. stats	true	Whether or not to output per-target coverage statistics (min, max, mean) in tabular format. Allowed values:				
		• false, 0: Do not create per-target coverage statistics.				
		• true, 1: Create per-target coverage statistics.				
enrichment.target.coverage. frequency	false	Whether or not to create the per-target coverage frequency histogram. Allowed values:				
		• false, 0: Do not create the per-target coverage frequency histogram.				
		• true, 1: Create the per-target coverage frequency histogram.				

1	5
	$\mathbf{U}$

Table 77	Enrichment	statistics	parameters	(continued)
----------	------------	------------	------------	-------------

Parameter name	Default value	Description
enrichment.target.coverage. bedgraph	false	Whether or not to create a BEDGRAPH format coverage file for on-target reads. Allowed values: true, false, 1, 0
		• false, 0: Do not create this file.
		• <b>true</b> , <b>1</b> : Create a BEDGRAPH format coverage file for on-target reads.
enrichment.genome.coverage. frequency	false	Whether or not to create a per-chromosome coverage frequency histogram. Allowed values:
		• false, 0: Do not create this histogram.
		• true, 1: Output a per-chromosome coverage frequency histogram.
Resource parameters		
memory.request	15GB	Memory requested per run. Must be at least 13GB to run enrichment statistics on the largest human chromosome (chr1).
		Specified in GB, MB, or KB.
		Example usage: memory.request=15GB
java.heap.space	1536	Dynamic memory requirement.

# **Enrichment statistics input files**

This section describes the input files required for by the enrichment statistics module. Parameters mentioned in this section are described in Table 77 on page 306.

Table 78 lists the two input files for the enrichment statistics module.

Table 78 Enrichment Statistics required input files

regions file

File	File format	Associated parameter	Description
target regions	BED, GFF v3, or text	enrichment.target.file	A list of target regions, either as regions of interest (ROI) or as target lists provided by vendors of enrichment platforms (such as baits, probes, or tiles files).
aligned reads	BAM	enrichment.input.file	Alignments generated by the sequencing run after being matched to the entire reference genome.

The input target The target regions file may be either in BED format or a list of chr:start-end elements. An example of partial content for a target region file is shown in Figure 30.

chr13	31787000	31788297
chr13	31788404	31788957
chr13	31789616	31789695
chr13	31790390	31790601
chr13	31790852	31791495
chr13	31791644	31791981

Figure 30 Example content for an input target region file

The following information applies to the input target region file:

5

	<ul> <li>Target region coordinates must be specified using the same reference used in mapping.</li> </ul>
	• Target regions should be sorted by chromosome/position. If not sorted, they will be sorted as the first step in the enrichment statistics module.
	• After the first 3 required fields per line, any additional fields are ignored.
	• The first field, "chrom", must have identifiers which match the sequence names used in the BAM file. For example, for the hg18 reference:
	– Use "chr1", not "1".
	<ul> <li>Use "chrX", not "23" or "chr23".</li> </ul>
	• This file is in standard BED format, using zero-based indexing and a half-open interval (meaning the specified range is up to, but not including, the last position).
	The input can optionally be a list of target regions in the "chr:start-end" format. This format uses one-based, closed intervals to represent genomic regions. An example of this format is:
	chr1:1234-4321
	<b>Note:</b> The input target region file and the chr:start-end list use different indexing and different interval formats.
The input aligned reads file	This file is a BAM-formatted file of aligned reads generated by the sequencing run after being matched to the entire reference genome.
	The following information applies to the input aligned reads file:
	• The BAM file must have a header listing sequence identifiers.
	The header is mandatory in the SAM specification for the binary BAM files.
	• These sequence identifiers must be consistent with those used in the target list.
	• The BAM file must be sorted by coordinate. The SO annotation in the header is not examined. All alignments within a chromosome must be contiguous. For sequential records, start coordinates within a chromosome must be equal or increasing.
	• The BAM file may be from a fragment, paired-end, or mate-pair library.
	All reads are treated independently. No special handing is performed based on library type.
	• Reads from paired libraries tend to have improved mapping scores based on unique paired hits, and are more likely to pass uniqueness filtering than unpaired reads.

# **Enrichment statistics output files**

This section describes the enrichment statistics module's output files. The output files are used in downstream analysis with LifeScope<sup>™</sup> Software and are also compatible with external viewers. Genomic reads and annotations in GFFv3, BED, BEDGRAPH, and BAM format can be displayed with third-party graphical genome browsers. Targeted resequencing reports and tabular data can be viewed and graphed in third-party spreadsheet applications.

Table 79 lists the output files for the enrichment statistics module. These files' purpose and content are described in the following pages. The enrichment statistics parameters mentioned in this section are described in Table 77 on page 306.

#### Table 79 Enrichment Statistics output files

File	File format	Optional	Controlling parameter (if optional)
Output target file	BED	No	_
Output alignment file	BAM	No	_
Target coverage BEDGRAPH file	BEDGRAPH	Yes	enrichment.target.coverage.bedgraph
Genome coverage frequency file	Text	Yes	enrichment.genome.coverage.frequency
Target coverage frequency file	Text	Yes	enrichment.target.coverage.frequency
Target statistics report file	Text	Yes	enrichment.target.coverage.statistics
Enrichment statistics reports	Text	No	_
			·

The output target file	This file is a BED file containing the sorted target list for use by later modules.					
The output alignment file	This file is a filtered BAM file which includes only those alignments that overlap target regions. The file is a filtered version of the input file and retains the input file's original characteristics. The BAI index file is also created.					
	The defau	It file names for the BAM and index file are:				
	<inp< th=""><th>out BAM file name&gt;.ontarget.<extend_bases>.bam</extend_bases></th></inp<>	out BAM file name>.ontarget. <extend_bases>.bam</extend_bases>				
	<inp< th=""><th>out BAM file name&gt;.ontarget.<extend_bases>.bam.bai</extend_bases></th></inp<>	out BAM file name>.ontarget. <extend_bases>.bam.bai</extend_bases>				
	where < <i>inj</i> extension) enrichme	<pre>put BAM file name&gt; is the name of the input file (without path or file name , and <extend_bases> is optional and taken from the .nt.extend.bases parameter, if that parameter is non-zero.</extend_bases></pre>				
The target coverage BEDGRAPH file	This targe the depth is generate parameter	t coverage BEDGRAPH file is a text file in BEDGRAPH format and describes of coverage within targets by on-target alignments. The target coverage file ed, as determined by the enrichment.target.coverage.bedgraph :				
	By default the target coverage file is created with the name \${enrichment.output.prefix}_target_coverage.bedgraph.					
	Example c	content for a target coverage BEDGRAPH file is shown below:				
	track type=bedGraph name="On-Target Coverage" descriptio of Coverage of On-Target Reads"					
	chr1	2974349 2974349 1				
	chr1	2974350 2974351 2				
	chr1	2974352 2974352 5				
	chr1	2974353 2974353 18				
	chr1	2974354 2974354 40				

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#### Table 80 describes the file content. These are tab-separated text fields.

#### Table 80 Description of the target coverage BEDGRAPH file fields

Field name	Example content	Description
Track annotation headers	track type=bedGraph name="Coverage"	Annotations used for UCSC browser display purposes.
Contig name	chr1	Contig name of the feature. This value comes from the sequence name (SN) in the BAM header. However, in order to be viewed in the UCSC browser, these values must be of the form chr1, chr2, etc.
Feature start	148	Start location of the feature. Zero-based, inclusive.
Feature end	150	End location of the feature. Zero-based, exclusive (half-open ranges).
Coverage	1	Depth of coverage over the given (start-end) range.

# The genomeThe genome coveragecoverage frequencycoverage depth forfilegenome coverage frequency

The genome coverage frequency file is a text file which reports frequency of the coverage depth for each reference sequence contig listed in the BAM file header. The genome coverage frequency file is optionally generated, as determined by the enrichment.genome.coverage.frequency parameter.

By default the genome coverage frequency file is named
\${enrichment.output.prefix}\_genome\_coverage.frequency.tab.

Example content for a genome coverage frequency file is shown below:

track type=bedGraph name="On-Target Coverage" description="Depth of Coverage of On-Target Reads"

chr1	0	233314335	247249719	0.943638
chr1	1	10264341	247249719	0.0415141
chr1	2	2195560	247249719	0.00887993
chr1	3	581693	247249719	0.00235265
chr1	4	216736	247249719	0.000876587
chr1	5	114047	247249719	0.000461262

Table 81 describes the content of the genome coverage frequency file.

 Table 81
 Description of the genome coverage frequency file fields

Column	Field name	Example content	Туре	Description
1	Contig name	chr1	Text	Contig name of the feature. This value comes from the sequence name (SN) in the BAM header.
2	Coverage	0	Text, Integer	Coverage depth value.
3	Coverage count	211562282	Text, Integer	Number of positions with the given depth of coverage.
4	Total contig size	247249719	Text, Integer	Total number of base pairs in the contig.
5	Frequency of depth in contig	0.855662	Text, Float	Fraction of contig covered at given depth. Count / Total = Frequency



The target<br/>coverage frequency<br/>fileThe target coverage frequency file is a text file which reports the frequency of coverage<br/>depth for each target. The target coverage frequency file is optionally generated, as<br/>determined by the enrichment.target.coverage.frequency<br/>parameter.

By default the target coverage frequency file name ends in \*\_target\_coverage.frequency.tab.

Example content for a target coverage frequency file is shown below:

chr1_2974357_2974879	33	4	522	0.00766284
chr1_2974357_2974879	34	3	522	0.00574713
chr1_2974357_2974879	35	2	522	0.00383142
chr1_2974357_2974879	36	2	522	0.00383142
chr1_2974357_2974879	37	4	522	0.00766284
chr1_2974357_2974879	38	5	522	0.00957854

Table 82 describes the file content. The target coverage frequency fields are the same as for the genome coverage frequency file, except that location are given as target ranges.

Table 82 Description of the target coverage frequency file fields

Column	Field name	Example content	Туре	Description
1	Target range	chr1_2974357_2 974879	Text	One of the input target ranges. The contig name and the start and end positions from a single input target file line are concatenated.
2	Coverage	33	Text, Int	Coverage depth value.
3	Coverage count	4	Text, Int	Number of positions with the given depth of coverage.
4	Total target size	522	Text, Int	Total number of base pairs in the target.
5	Frequency of depth in target	0.00766284	Text, Float	Fraction of target covered at given depth. Count / Total = Frequency

# The target statistics report file

The target statistics report file is a text file which reports per-target coverage and enrichment details in tab-delimited format, with one line per target. The target statistics report file is optionally generated, as determined by the enrichment.target.coverage.statistics parameter.

By default the target statistics report file name ends in \*\_target\_statistics.txt.

Example content for a target statistics report file is shown below:

Target Name	Target	Size	Minimum	Coverage	9	Maximum	
Coverage	Average	Coverage	e	Total BI	P Coverag	le	
chr1_2974357_29	74879	522	33	2334	605.515	316079	522
chr1_2974634_29	75198	564	0	892	210.183	118543	547
chr1_2975115_29	75633	518	22	892	295.367	153000	518
chr1_2975421_29	75989	568	0	571	110.783	62925	218
chr1_3092354_30	92941	587	347	1875	991.16	581811	587
chr1_3150338_31	50930	592	75	1301	546.51	323534	592



chr1_3291283_3291753	470	468	4436	1705.47	801571	470
chr1_3302721_3303234	513	611	3545	1804.12	925514	513
chr1_3309173_3309758	585	314	2267	940.32	550087	585
chr1_3311082_3311483	401	120	5775	2960.54	1187176	401

Table 83 describes the content of a target statistics report file.

 Table 83
 Description of the target statistics report file fields

Column	Field name	Example content	Туре	Description
1	Target name (target range)	chr1_2974357_2 974879	Text	One of the input target ranges. The contig name and the start and end positions from a single input target file line are concatenated with underscores.
2	Target Size	522	Text, Integer	Total size of this target range, in base pairs.
5	Min	33	Text, Integer	Minimum fold coverage of position in target.
6	Max	2334	Text, Integer	Maximum fold coverage of position in target.
7	Avg	605.515	Text, Number	Average fold coverage of all positions in target.
8	Total BP Coverage	316079	Text, Number	Enrichment fold of the target range.
8	BP Covered at least 1X	522	Text, Number	Bases covered at least 1X.

# The enrichment statistics report

These reports provide data on the targeted regions in your analysis. The percent of coverage data is for the regions of interest, not for the whole genome reference. The end of the report summaries the number of target bases covered at various depths. The file name pattern is \*enrichment\_report.stats. Example content is given below.

Run Reads On Percent On Reads Off Percent Off Enrichment Fold ./outputs/enrichment/output.bam 69260675 88.6216% 8892585 11.3784% 1338.33

Reads In Targets: 69260675 88.6216% Reads Off Targets: 8892585 11.3784% Ratio of Percent on/off Target: 7.78859 Total Target BP: 2039803 Total Genome Size: 3080436051 Ratio of target to genome: 0.00066218 Enrichment fold relative to target size: 1338.33

# Targets Not Covered Target Bases Not Covered Percent of Target Bases Not Covered Percent of Target Covered >= 1X Percent of Target Covered >= 5X Percent of Target Covered >= 10X Percent of Target Covered >= 20X Average Depth of Target Coverage

1 12061 0.59% 99.41% 98.98% 98.63% 98.22% 2351.24

Number of target regions with no coverage: 1 Percent of target bp not covered: 0.59% (12061 bp) Percent of target bp covered at >= 1X: 99.41% Percent of target bp covered at >= 5X: 98.98%



Percent of target bp covered at >= 10X: 98.63% Percent of target bp covered at >= 20X: 98.22% Maximum Depth of Coverage within targets: 49453 Average Depth of Coverage within targets: 2351.24

## Targeted resequencing read selection algorithm

Reads are classified as "on-target" by a set of user-definable criteria. First, all reads are filtered for a minimum mapping quality. Reads which align to multiple places in the genome are assigned a lower mapping quality. Reads with very low mapping qualities may be false positive hits arising from repetitive sequence in the reference genome.

The default target definition criteria used in targeted resequencing mapping are:

- Reads must have a minimum mapping quality of 8. Recommended values are between 0 (in which case there may be a population of repetitive reads which inflate enrichment statistics and may contribute to false positives in variant calling), and 20 (which ensures only uniquely mapping reads are used, but results in a loss of sensitivity in regions with low mappability).
- Alignments may be marked as duplicate by the alignment algorithm. These reads are typically excluded from variant calling, but are counted for purposes of assessing enrichment.
- Only primary alignments are considered. Alignments marked as secondary (alternative placements) are excluded. Note that the mapping quality threshold may excluded reads with multiple secondary hits.
- Target regions are considered as provided. Reads aligning adjacent to targets may be included in the analysis by specifying the number of bases flanking the target regions that each target should be extended by.
- Mapping to target region is defined as having at least one base overlapping the target region.
- For paired-end runs, by default the reverse read must align 100%, as these are often shorter and less accurate. The default 100% alignment requires that all bases align to coordinates that are fully contained by the target region. For this test, trimmed bases are not considered. The overlap percentage required is controlled by the enrichment.minimum.target.overlap.reverse parameter.

Both alignment thresholds are user-adjustable, down to  $1 \times 10^{-4}$  (meaning at least one base, for reads up to 1000bp) or up to 1.0, meaning the read and target must completely overlap.

While only one list of target regions can be provided by the user, multiple target region sets may be analyzed by re-running the enrichment analysis, and specifying a different output folder or prefix for each target list. Similarly, to assess near-target alignments, reanalyze while specifying a larger number of bases by which to extend the target.

The targeted resequencing module by default runs the SAET utility prior to mapping.



# Run other targeted resequencing modules

The instructions for targeted resequencing modules are the same as for the resequencing tools, except that the Targeted modules require an additional input file, the regions of interest (ROI) file. You must provide the ROI file, in any of the following supported formats:

- a bait, probe, tile, or other file in BED format
- a text file as a list of "chr:start-end" values

To run resequencing pairing, find SNPs, and find small indels analyses, complete the instructions in the following chapters and also specify your the regions of interest file:

- Chapter 8, "FAQ Pairing" on page 155
- Chapter 9, "Run an SNPs Analysis" on page 159
- Chapter 12, "Run a Small Indels Analysis" on page 215

The differences when these analyses are run for targeted resequencing, compared to standard resequencing, are:

- The find SNPs and find small indels analyses are both run with an input file containing only on-target reads. Additionally, find SNPs takes the processed target list as input, and restricts SNP calls to events occurring within the target regions.
- SNPs analysis is also passed the dibayes.het.skip.high.coverage=0 parameter, to avoid excluding regions of very high coverage, which are common in enriched regions.
- The SAET module is enabled by default.

**Note:** If you turn off SAET, comment out the analysis.input.readset.file line, if it appears in your mapping INI file:

analysis.input.readset.file = \${analysis.output.dir}/saet/\*.rrs

# 16

# Run the SOLiD<sup>™</sup> Accuracy Enhancement Tool

This chapter covers:

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## **Overview**

The SOLiD<sup>TM</sup> Accuracy Enhancement Tool (SAET) improves color call accuracy prior to mapping. This method implements a proprietary spectral alignment error correction algorithm that uses quality values and properties of color space. Significantly improving its performance over conventional error correction methods, the use of SAET produces more accurate mapping with both an increased number of reads with zero miscalls and an increased number of mapped reads. The SNP calling on error-corrected data results in an increase in true positive calls and a decrease in the number of false positive calls. Using the *de novo* assembly on error-corrected reads results in an increase that is as much as three to five times higher in average contig length.

SAET performance was tested on various datasets, including a large spectrum of genome sizes and complexities, coverages, read lengths, and experiment types. SAET shows similar performance on datasets sequenced from enriched regions or genomes of 1 kbp to 3 Gbp, with coverage from 10 to 4000x and a read length of 25 to 75 bp.

On average SAET requires up to 5 GB RAM for each 1 Gbp of sequenced region. SAET can be applied for diploid organisms. SAET should not be used for rare variant detection datasets where the frequency of rare variant may be less that twice that of the color calling error rate.

# SAET usage guidelines

SAET is recommended for use with targeted resequencing and is on by default in targeted resequencing standard workflows. SAET has not been validated with other experiment types and may have insufficient memory to run. If you do choose to run SAET with other experiment types, you can do the following:

• Run SAET before you run the resequencing or whole transcriptome mapping and pairing tools.

- Use the SAET output with resequencing and whole transcriptome mapping and pairing.
- Use updated quality values file for SNP calling.
- Avoid the generation of the updated quality values, if the downstream analysis modules in your workflow do not use quality values. Your SAET runtime is reduced approximately by half if you do not update quality values.

# Examples of how to run SAET

Workflows using	The following standard workflows provide examples of running SAET:				
SAET	<ul> <li>targeted.resequencing.frag</li> </ul>				
	targeted.resequencing.pe				
	The optional examples download includes an example of how to run SAET by itself, as an individual analysis that is not part of a standard workflow.				
	See the following sections for information on these methods:				
	<ul> <li>As part of a workflow: Chapter 7, "Run a Standard Workflow Analysis" on page 97</li> </ul>				
	• As an individual analysis: Appendix E, "Demo Analyses" on page 507				
SAET example in a workflow	The targeted resequencing Fragment standard workflow gives an example of running SAET on with targeted resequencing analyses. The analyses are:				
	• SAET				
	• Mapping				
	• Enrichment				
	• SNPs				
	Small indels				
	Annotations				
	SAET error corrects reads prior to mapping. This correction significantly improves mapping quality and the quality of variant calls.				
	<b>Note:</b> If you turn off SAET, make sure your mapping INI file does not contain the line:				

analysis.input.readset.file = \${analysis.output.dir}/saet/\*.rrs

Comment out this line if it appears in your mapping INI file.

## **SAET** input files

By default, SAET takes an input file of  $SOLiD^{TM}$  system reads in XSQ format, and an expected length of the sequenced region, such as the:

- Genome size for whole genome sequencing
- Size of enriched region in targeting resequencing
- Size of transcribed region in whole transcriptome sequencing



The required input file is in XSQ format, which is typically generated by 5500 Series  $SOLiD^{TM}$  Sequencer. The file must contain reads in color-space that require correction. SAET constructs a spectrum of all *k*-mers from the set of all reads. Then each read is corrected individually if necessary. (The value *k* is the seed length. The default can be overridden with the parameter saet.seed.)

In targeted resequencing, the SAET module accepts as an input the BED, GFF, or TXT file with segments of target regions. In this case the refLength parameter is computed from the targeted regions information.

# **SAET** parameters

This section describes the parameters that control the runtime behavior of the SAET module. In order to change a parameter value, use the set param shell command. For example, to turn off quupdate, change the value of the parameter saet.quupdate with this shell command:

set saet.qvupdate 0 secondary/saet.ini

Table 84 SAET parameter descriptions

Parameter name	Default value	Description
Required parameters		·
saet.refLength	1000000	Expected length of sequenced (or enriched) DNA region. For example, 4,600,000 for the E.Coli 4.6 MB genome or 30,000,000 for the entire Human Transcriptome.
		Allowed values: Integers >= 200.
Parameters required f	for targeted resequencing	
saet.on.target.ratio	0.5	The expected ratio of reads that come from the targeted region. Allowed values: Floats 0.0–1.0.
Optional parameters	·	
saet.qvupdate	1	Whether or not to update the quality value of modified calls. Allowed values:
		• <b>0</b> : Do not update the quality value of modified calls. This option is approximately twice as fast.
		• 1: Update the quality value of modified calls. Required for SNPs.
saet.qvhigh	25	Correction is applied to calls with a quality value below the value of this parameter. Allowed values: Integers >= 1.
saet.seed	0	Size of $k$ -mer (>=5) used in spectrum construction and error correction. (If set to 0, then the value is computed internally.)
		Allowed values: Integers 0, 5-28.
saet.suppvotes	3 for targeted reseq; 2 for other analyses	The minimum number of $k$ -mer votes required to make a correction. Allowed values: Integers >= 1.
		For datasets forming coverage less than 50x, the recommended value is 1 or 2. For high coverage data, the value can be increased up to k-1. Increasing this value results in a lower number of over-corrections in repetitive regions and a lower number of corrections over all. The maximum recommended value is 7.



#### Table 84 SAET parameter descriptions

Parameter name	Default value	Description
saet.trustfreq	0	The lowest multiplicity of the seed to be included in the spectrum. (If set to 0, then the value is computed internally.) Allowed values: Integers >= 0.
saet.localrounds	0	Maximum number of allowed corrections per read. (If set to 0, then the value is set to [readLength/8]). Reduce if over-corrections are observed, or increase if under-corrections are observed. Allowed values: Integers 0–9.
saet.globalrounds	1 for targeted reseq and reseq of large genomes; 2 for general reseq of small genome	The error correction step is repeated the provided number of times. Reduce if over-corrections are observed, or increase if under-corrections are observed. Allowed values: 1, 2, or 3.
saet.trustprefix	0	Position in the read at which the error rate inflates, for instance, 35-40 for 50bp long reads. If set to 0, then the value used is $0.8$ *readLength. Allowed values: Integers >=k, where k is the size of the k-mer (either set by saet.seed or computed internally).
saet.nosampling	0	Disables random sampling in spectrum building. If set to 0, then for large datasets (coverage > 300x), a subset of reads is used in spectrum building. Allowed values:
		• <b>0</b> : Do not disable random sampling in spectrum building.
		• 1: Disables random sampling in spectrum building.
Resource parameters	1	
processors.per.node	8	Number of parallel threads used to run SAET on a node.
saet.minreads.per. node	1000000	If SAET is run on multiple nodes, then each node receives at least this amount of reads to process.
memory.request	15GB	Maximum amount of memory available on the node.
		<b>IMPORTANT!</b> SAET Accuracy Enhancement Tool requires 20GB to run on large genomes, such as experiments sequencing whole human genomes. With the minimum memory (15MB), SAET may not run on whole large genomes, and in this case the original reads are used for mapping.
		For applications such as targeted resequencing, SAET runs with the minimum memory.
		If SAET does not have enough memory, it issues a warning and allows mapping to continue with the original uncorrected reads.

#### Explanation of parameters

The processing time and the level of error correction aggressiveness depend mainly on the values defined for saet.localrounds and saet.globalrounds. The saet.localrounds parameter allows you to correct up to saet.localrounds errors in a read by using a precomputed spectrum. The saet.globalrounds parameter recomputes the spectrum after each global round, and allows you to correct up to saet.localrounds errors in a read based on the precomputed spectrum.

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SAET is designed to reduce the error rate in the reads generated by the SOLiD<sup>™</sup> System. Reducing the error rate increases the number of mapped reads for resequencing analyses, which can for example lead to an increase in true positive SNPcalls and a decrease in false positive SNP-calls. (If the SAET is used for SNP calls, the saet.qvupdate parameter must be set to 1.) Use the saet.qvhigh parameter to restrict corrections to positions that have a quality value below the saet.qvhigh threshold.

The optional parameters available in SAET allow you to modify the algorithm's behavior. For example, you might find that the globally computed cutoff for frequency of trusted seeds does not meet your purpose. For example, the cutoff might be too low, causing too many junk seeds to be considered correct. However, if the cutoff is too high, then many correct but low-frequency seeds are inappropriately filtered out. You can change the saet.trustfreq parameter to overwrite the estimated frequency cutoff.

If you trust the quality of your reads more or less than the value specified in the saet.trustprefix parameter, then make the corresponding changes to the value. If SAET makes many corruptions in the regions of reads with highly packed errors, you can increase saet.suppvotes to improve the tendency to correct only isolated errors. SAET provides options for reading and writing spectrum files. The options enable you to build spectrum from better-quality reads and use the spectrum to correct lower-quality reads. You can also build a spectrum from a reference, or build a spectrum from multiple files, such as data from multi-run experiments.

# SAET internal parameters

Table 85 lists SAET parameters that we do not recommend changing.

saet.run	1	Whether or not to run the SAET module. Allowed values:
		• <b>0</b> : Do not run SAET.
		• 1: Run SAET during this analysis.
		The run parameter is set automatically by the shell during a standard workflow. Accept the default for most use.
saet.outspecbin	0	Whether or not to output a spectrum in binary format in the file \${task.out.dir}/analysis.input.readset.file.spect.bin
		Allowed values:
		• <b>0</b> : Do not output the spectrum.
		• 1: Output the spectrum.
saet.inspecbin	-	Uses pre-generated file(s) with spectrum (in binary format) for error correction. Use a comma to separate multiple files.
saet.maxtrim	0	Trims erroneous tails of reads up to first trusted seed or up to maxtrim. If the remaining part of a read is shorter than seed size + 2, then the read is discarded. Do not use this option together with the seat.qvupdate parameter.
		If this parameter is set to 0, then no trimming is done.
		Allowed values: Integers >= 0.

 Table 85
 SAET internal parameters



Table 85 SAET internal parameters (continued)

saet.trimqv	_	Trims erroneous tails of reads up to first trusted seed or up to a position with quality value higher than trimqv. If the remaining part of a read is shorter than seed size + 2, then the read is discarded. Do not use this option together with the seat.qvupdate parameter.
		If this parameter is set to 0, then no trimming is done.
		Allowed values: Integers >= 0.
saet.log	saet.log	Log file name.
saet.progress.file	.progress	Progress file name.
saet.progress.start	0	Progress start point. Allowed values: Integers 0–100.
saet.progress.end	100	Progress end point. Allowed values: Integers 0-100.
maximum.workers	4	Maximum number of nodes that can be used by SAET.

# SAET output files

SAET writes the corrected reads and quality values into new XSQ files.

The SAET output files are used as input to mapping analyses.

# **Algorithm description**

#### SAET SAET implements a modified version of the spectral alignment error correction algorithm proposed by Pevzner et al., 2001. SAET extends the original algorithm by implementation taking quality values and properties of color-space into account, significantly improving the performance of the original method. The method corrects reads given a set of k-mers (seeds) spectrum. Given a set of reads (R), and a frequency threshold (m), the spectrum is defined as the set of all *k*-mers that appear at least *m* times in reads from *R*. The set of such trusted *k*-mers approximates the set of all *k*-mers in the genome. A read is defined as error-free if all of its *k*-mers are trusted. If not all *k*-mers are trusted, SAET attempts to make a read error-free by mutating a few colors in the read. SAET only considers substitution mutations since there no insertions or deletions in SOLiD<sup>™</sup> system reads. When making error corrections in the read, SAET gives priority to positions with missing calls, followed by positions that have lower-quality values. In addition, SAET avoids correcting two adjacent colors with color-call quality values above a certain threshold, in order to preserve any SNP evidence. SAET computes the *k*-mer size and the multiplicity of trusted *k*-mers, based on the estimated reference length and number of reads in the dataset. Using default parameters, the color call changes introduced by SAET are expected to be 99.99% accurate. SAET supports multi-core runs. SAET provides a mechanism for multi-node runs by creating sub-spectra from subsets of reads. SAET merges the sub-spectra into the final spectrum that can be used for correction of any subset of reads. Phases The SAET algorithm has three steps as described below.



#### Spectrum build

In this step, a temporary file with a spectrum constructed from all of the reads is generated in the output directory. This process is very fast and does not require large RAM resources However, SAET requires a large amount of disk space (~1 GB for each 1Mbp of genome). SAET creates temporary swap files when memory usage approaches 2 GB.

#### Error correction

The error correction step takes each read in the input file and attempts to correct it if any of its *k*-mers are not present in the spectrum. This step is the most time-consuming. LifeScope<sup>TM</sup> Software runs SAET on multiple cores to improve the error correction speed.

#### Update of quality values for corrected calls

The quality value of each corrected position is replaced with a quality value derived from spectral probability. This step is time-consuming and can be disabled if the update of quality values is not required. (To disable, set saet.gvupdate to 0.)

**SAET run times** The SAET processing time depends on the input size and number of global and local rounds. With a large number of global and local rounds, an expected throughput is 1 Gbp per hour on a single core. The amount of used RAM is approximately 5 GB for 1 Gbp of genome size.



# PART V Whole Transcriptome Analyses
# 17

# Run a Whole Transcriptome Mapping Analysis

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## **Overview**

High-throughput sequencing of the transcriptome using the SOLiD<sup>™</sup> System enables genome-wide expression profiling with high sensitivity and a wider dynamic range than microarray technology. The whole transcriptome library preparation also preserves the strandedness of the RNA transcripts. Preserving the strandedness simplifies data analysis, allows determination of the directionality of transcription and gene orientation, and facilitates detection of opposing and overlapping transcripts.

You can use the 5500 Series SOLiD<sup>™</sup> Sequencer to sequence RNA prepared with RNA-Seq sample preparation kits. RNA sequencing produces high-depth, short-read sequencing data that can be used to measure RNA expression. Like microarray analysis, RNA-Seq measures expression intensity across many genomic features. Unlike microarray analysis, RNA-Seq can be used to identify novel transcriptome features in a sample.

Like other SOLiD<sup>TM</sup> System applications, RNA-Seq produces short reads in the form of XSQ files. Whole Transcriptome Analysis (WTA) may be applied to these reads to produce alignment, coverage, count, and splice-finding results. WTA is a reference-based analytical method, which means a reference genome is required as input. Much WTA functionality also requires gene annotation associated with the reference genome sequence.

WTA analysis can be divided into secondary and tertiary analysis. Secondary analysis produces the aligned reads using sequences and quality values as input. Tertiary analyses produce biologically meaningful results (gene expression level, feature discovery, etc.) from aligned reads. This chapter describes how WTA is used to align reads to a reference genome, the methods used by WTA, and the configurable parameters of WTA.

The 5500 Series SOLiD<sup>™</sup> Sequencer System produces RNA-Seq reads in both singleread and paired-end configurations. While there is overlap in the analytical techniques, the analysis of single-read and paired-end data is performed with two distinct pipelines. The single-read pipeline consists of alignment, counting, and coverage steps. The paired-end pipeline extends single-read analysis with an alternative alignment step and an additional junction finding step.

The aligned reads produced by WTA can be used with the following tertiary modules:

- **Count annotations –** Computes the number of reads that align within genomic features.
- Coverage Computes the read coverage at each genomic position.
- **Splice finder** Identifies splice junctions of various types, including fusion junctions from paired-end reads.

The 5500 Series SOLiD<sup>™</sup> Sequencer<sup>™</sup> System produces RNA-Seq reads in both singleread and paired-end configurations. While there is overlap in the analytical techniques, the analysis of single-read and paired-end data is performed with two distinct pipelines. The single-read pipeline consists of alignment, counting, and coverage steps. The paired-end pipeline extends single-read analysis with an alternative alignment step and an additional splice finding step.



Figure 31 WTA single-read and paired-end internal workflows

## Examples of running the WT mapping module

In a standard The following standard workflows provide examples of running the WT mapping module: workflow whole.transcriptome.frag whole.transcriptome.pe The following are example shell commands using reads and references from the LifeScope<sup>™</sup> Software repository. This example turns off tertiary modules in the workflow. # log into the shell lscope.sh shell -u username -w password # cd to the projects repository cd /projects # make a project, and open it mk ecoli cd ecoli # make an analysis, and open it mk run1 cd run1 # define the analysis type set workflow genomic.resequencing.pe # define the input add xsg run0209a\_50\_PE.xsg add xsg run0209b\_50\_PE.xsg # specify the reference to be used set reference hg19 # these commands turn off the tertiary modules set coverage.run 0 tertiary/coverage.ini set splice.finder.run 0 tertiary/splice.finder.ini set wtcounts.run 0 tertiary/wtcounts.ini # optionally change mapping parameter defaults after this line # list the configuration of the analysis 15 # start the run run # list progress information about the run 15 In order to change a mapping parameter value in your analysis, use the set param shell command after the line # optionally change mapping parameter

defaults after this line. For instance, to turn on the mapping.in.base parameter, use this shell command:

set mapping.in.base 1 secondary/wt.pe.map.ini

This command shows how to turn on the mapping.in.base parameter for the whole.transcriptome.frag workflow:

set mapping.in.base 1 secondary/wt.frag.map.ini

See Chapter 7, "Run a Standard Workflow Analysis" on page 97 for more information on running standard workflows.



# **As a demo analysis** The optional examples download includes an example of how to run the whole transcriptome analyses outside of a standard workflow. A standard workflow is recommended.

See Appendix E, "Demo Analyses" on page 507 for more information on running a demo analysis.

# Input files

**Reads input files** The WT mapping module accepts as input one or more input XSQ files. The input reads can have different read lengths but they must be of the same library type (fragment or paired-end). For example, paired-end read-sets of length 50x25 and 60x25 can be processed together, but a paired-end read-set cannot be processed with a fragment read-set.

#### **RNA-Seq reads**

The RNA-Seq reads used in WTA are different from the genomic reads. For RNA-Seq reads:

- Only transcribed sequences are measured by the system.
- Genome coverage is non-uniform due to variation in transcriptional intensity.
- A large subset of reads originate from uninteresting sequences such as ribosomal RNA (rRNA).
- A subset of the reads originates from splice junctions and do not align contiguously on the genome.

#### Reads input specification

Plan your analysis input carefully. The way you define your reads input affects the behavior of your analysis. The following factors control how your input data is analyzed:

- Index (barcode) IDs Using an index ID restricts your input to the reads data of one or more indices.
- **Grouping of reads** Each group of reads is analyses together as one specimen. The output data for a group is combined into one set of results files.
- Multiple sample runs Unrelated reads can be processed together in one run of LifeScope<sup>™</sup> Software, but analyses separately as separate input data.

See "Define input data" on page 85 for more information on designing how to add input data to your analysis.

Mapping one tag of paired data is not supported.

#### Legacy data

If the data you want to process with LifeScope<sup>™</sup> Software is in CSFASTA and QUAL files, these files must be converted to the XSQ file format, through one of the following methods:

• The LifeScope<sup>™</sup> Software UI handles converting these files to the XSQ format before mapping.



	• If you are using the LifeScope <sup>™</sup> Software command shell, you must first convert the CSFASTA and QUAL files to the XSQ format. See Appendix C, "XSQ Tools" on page 479 for information on the standalone XSQ converter.
Reference input files	WTA uses several files defining reference information: filter-reference, genome- reference and annotation files. The filter-reference file is a FASTA-formatted file defining uninteresting reference sequences such as Ribosomal RNA, tRNA, or vector sequences. To increase the speed of analysis, reads that align to sequences in the filter reference are excluded from downstream alignment steps. Many of the entries in the filter reference are species-specific. The filter reference file must be appropriate for the species being studied.
	The principal output of alignment analysis is a set of alignments between the reads and the genome-reference in BAM format. The genome-reference file is a FASTA- formatted file defining sequences in the reference genome.
	The annotation file is a GTF-formatted file defining known genes, transcripts, and exons in the genome reference. See "Annotations input files".
	Junction-reference and exon-reference files are generated internally by the software. These files are entirely derived from the GTF and genome-reference files. A user is not required to provide these references. The junction-reference file is a FASTA-formatted file with an entry for every potential intragenic splice-junction defined in the annotation file. The entry contains the position of the junction and the associated flanking sequence. The exon-reference is a FASTA-formatted file with an entry for every exon defined in the annotation file.
Annotations input files	While genomic resequencing relies only on alignment to a reference sequence, WTA also makes use of gene annotations. Gene annotations define the exons, genes, and transcripts used to improve alignment.
	WTA modules use genome annotation files in GTF format. The following sites explain the GTF format:
	genes.cse.wustl.edu/GTF2.html genome.ucsc.edu/FAQ/FAQformat.html#format4
	The GTF file must match the genome reference to ensure that the WTA modules work correctly.
	<b>IMPORTANT!</b> Make sure the GTF file is for the same genome assembly as the FASTA file, and that matching sequence identifiers are used. Gene and transcript identifiers in the GTF file must be properly normalized. Identifier normalization is a known issue in GTF files from the UCSC Genome Browser, which is a popular source of GTF-formatted annotation. The UCSC GTF files always report the same value for gene and transcript IDs.
	LifeScope <sup>™</sup> Software includes a script (refgene2gff.sh) that produces a compliant GTF file from the UCSC Genome browser database.

#### UCSC genome annotations

Format UCSC Genome Browser Database annotations for WT analyses

The UCSC Genome Browser Database has genome annotations available for many assemblies at

hgdownload.cse.ucsc.edu/goldenPath/

The GTF-formatted annotations available for download are not properly normalized by gene ID. The required content is present for each assembly in the file export of the refGene database table database/refGene/txt/gz.

For example, annotation for human genome build 18 is available at:

hgdownload.cse.ucsc.edu/goldenPath/hg18/database/refGene.txt.gz

**Note:** The GrefGene.txt.gz file is not in GTF format. You must convert the annotation before using it in WTA.

Convert the refGene.txt.gz file

Run the script bin/refgene2gff.sh to convert the refGene.txt.gz file:

```
gunzip refGene.txt.gz
refgene2gff.sh -i refGene.txt -o refGene.gff
```

Genome annotations that are downloaded from the UCSC Genome Browser and converted by the annotation conversion script are preferable because they contain Human Genome Organization (HUGO)-style gene names. HUGO-style gene names allow interpretation when using a genome browser or reading reports.

The annotation conversion script works with the latest format of refGene.txt files. Assemblies, such as the rat genome, use an alternative format for the refGene.txt file. The refgene2gff.sh script does not convert alternative formats.

#### **Ensembl GTF files**

Format Ensembl GTF files for WTA pipelines

The Ensembl website **ensembl.org**/ has GTF-formatted genome annotations available for many popular assemblies. Unlike the GTF files directly downloadable from UCSC (see "UCSC genome annotations" on page 330), Ensembl GTF files are properly normalized by gene and transcript IDs. Visit the following site to download an Ensembl GTF file:

http://www.ensembl.org/

Ensembl GTF files use gene accession numbers instead of HUGO-style gene names. Ensembl GTF files also use unprefixed sequence identifiers, such as 1,2,3....X,Y,MT. The Ensembl GTF files are incompatible with genome reference FASTA files that have UCSC-style sequence IDs with the prefix "chr", for example, chr1, chr2, chr3....chrX, chrY, chrM.

Reformat the Ensembl GTF file

To reformat the Ensembl GTF file to use UCSC-style gene IDs, run this script:

reformat\_ensembl\_gtf.pl Homo\_sapiens.GR



## Stages of mapping

A WT mapping run consists of the following phases:

- Scatter The entire set of reads is divided into small non-overlapping subsets and defines one mapping job per subset.
- **Mapping** A single computer processes a mapping job, which takes reads as input and produces aligned reads as output.
- **Gather** The results of multiple mapping jobs are combined to produce final mapping result.



Figure 32 WT mapping phases

The Mapping phase alone computes the alignment records that compose the results. This phase can be broken down into several steps:

- **Filter mapping** Reads that align to the filter reference are counted and excluded from further analysis.
- Genome mapping Reads are aligned to the genome reference.
- Junction mapping Reads are aligned to the junction reference (F3 reads only).
- Exon mapping Reads are aligned to the exon reference (F5 reads only).
- Alignment merge Results from genome and junction or exon mapping are combined to produce a single set of alignments for each read type (F3 or F5).
- Exon table rescue Supplements the set of alignments by conducting a more sensitive search for a alignments of a read (target) near the alignments found for the other member of the pair (anchor). These regions are selected using the gene structure of the genome. (Paired-end only).
- **Pairing** Selection of most likely alignments based on alignment quality and the distance between alignments of each member of a pair. (Paired-end only).
- **BAM file generation** The process producing results in BAM format. For colorspace data, the base translation step is also included. Due to the scattering step, the BAM files at each compute node are a sub-set of the final BAM file and are referred to as baby BAMs.

BAM generation is composed of several substeps.

- Ma2BAM, Pa2BAM A BAM file (lacking actual sequences or quality values) is written. Mapping quality values are computed (MAPQ)
- **Refcor** Reference-assisted color-to-base translation. Sequences and quality values are added to the BAM records.
- **Sorting –** Sorting the BAM file by genome coordinate.



Figure 33 The phases of mapping WT fragment data





Figure 34 The phases of mapping WT paired-end data

**Scatter** The scatter stage distributes reads to the available compute nodes, in order to gain the efficiency benefits of parallel computing.

The mapping module can accept multiple read-sets, which are allowed to have different read lengths. The input set of read-sets from multiple XSQ files is first split based on read length. Fragment read-sets are partitioned into separate categories based on these read length categories:

- $0 \leq \text{Length} < 25$
- $25 \leq \text{Length} < 35$
- $35 \leq \text{Length} < 50$
- $50 \leq \text{Length} < 75$
- $75 \leq \text{Length}$
- Trimmed reads, with variable read lengths

The reads from each category are scattered into multiple jobs based on the parameters fragmap.minreads.per.node and fragmap.number.of.nodes. These parameters are described in the section "Mapping performance" on page 342.

# Single-read<br/>mappingWT Single Read mapping produces a set of aligned reads. Stated briefly, the method<br/>performs the following steps (see Figure 33 on page 332).• Remove reads that align to the filter reference.

- Map reads to genome and junctions.
- Merge results.
- Convert to BAM format.

Alignment finding uses a proprietary algorithm (see Chapter 8, "Run a Resequencing Mapping Analysis" on page 113). The filter mapping step identifies and counts the reads that align to the filter reference. The counts are written to the filter report. Reads that align to entries in the filter reference are omitted from further analysis and do not appear in the set of aligned reads produced by the module.

The genome mapping step finds alignments between the reads and the genome reference. Likewise the junction mapping step finds alignments with the junction reference. Alignments of a read to a genome and to junctions can produce multiple similar alignments at the same location. When multiple similar alignments occur, the alignment with the highest alignment score replaces all others. If there is a tie, the genomic alignment is used. The alignment score is calculated as follows:

*Score=len-nm*×1(1+*mp*)-*jp* 

where:

- *len* = number of colors in the alignment
- *nm* = number of color mismatches
- *mp* = mapping mismatch penalty
- *jp* = penalty for alignment to a junction

The parameters to control mapping algorithm are in Table 88 on page 341. For details of the mapping algorithm, see "Mapping schemes" on page 345.

# **Filter mapping** The reads are mapped to the filter reference in the filter mapping step. Reads that align to the filter reference are called filtered reads. The result of this mapping step is used to populate the filter report. In the single-read pipeline, filtered reads are populated in a separate BAM file. Filtered reads are omitted from the BAM file. A filter reference for use with human reads is available with LifeScope<sup>™</sup> Software. This reference contains:

- SOLiD<sup>™</sup> adapter sequences
- Human ribosomal RNAs (rRNAs)
- Human transfer RNAs (tRNAs)
- Other human sequences
- Single-base-repeat sequences, including Poly-A, Poly-T, and more

Complete the following steps to construct a filter reference for another species.

- 1. Copy the adapter and single-base-repeat sequences to a new file.
- **2.** Append FASTA-formatted species-specific ribosomal, transfer RNA, and other sequences to the filter reference.

1	7
	/

Junction mapping	Reads are mapped to the flanking sequence of junctions defined by the genome annotation in the junction mapping step. Junctions are inferred from the exon, gene, and transcript definitions in the genome annotation. Junction sequences are produced for all splices between pairs of exons within a gene. Junctions present in annotated
	are called putative junctions. Junctions not present in annotated transcripts are called putative junctions.

**Exon mapping** In the exon mapping step, paired-end reads are mapped to the set of exon sequences defined by the genome annotation. This step maps the shorter F5 reads in the paired-end pipeline. By default, more mismatches are allowed in exon mapping of F5 reads in the paired-end pipeline. Single-read mapping skips this stage.

Table 86 describes a human exon sequence database and a human junctions database.

 Table 86
 Comparison of alternative human references

	Reference	Number of references	Bp size	Mappability	Novelty	Junctions
	Genome	25	3 billion	OK	OK	_
F3	Junctions	2,012,075	201,207,500	OK	_	OK
	Refseq	32,699	99,02,749	OK	_	Partial
	Genome	25	3 billion	_	OK	_
F5	Exons	216,884	99,026,749	OK	_	_
	Refseq	32,699	99,026,749	OK	_	Partial

In a paired-end alignment, the F3 and F5 paired-ends are initially processed through independent mapping paths. Similar to single-read-mapping, the F3 reads are mapped to filter, genome, and junction sequences. The F5 reads are also mapped to filter, genome, and exon sequences. Reads that map to filter sequences are discarded and do not proceed further in the pipeline.

The F3 genomic and junction mapping results are merged into a non-redundant set of alignments. The F5 genomic and exon mappings are merged into a non-redundant set of alignments. At this point in the analysis, the output consists of a set of independently processed F3 and F5 genomic alignments.

# **Rescue method** You can optionally use the rescue method to find additional alignments. Rescue is an alignment method that is applied to read pairs that have at least one alignment, but no pair of alignments occurring within an expected range (see Figure 35 on page 336). The expected range is set to 100,000 bases by default.



Figure 35 Annotation-aided rescue for WT

The next section refers to Figure 35.

#### Row A

For alignments that fall on an exon of a gene, and do not have a mate alignment above a certain score threshold, a special exon rescue is performed. Rescue region (rr) is defined as a rescue distance that is downstream of the alignment. The rescue distance is determined by user-defined thresholds. The rescue distance starts from the left-most position of the alignment for overlapping mates. The rescue region also includes certain position range in downstream exons of the same gene. The approach defined in the previous sentences helps rescue mates on different exons.

#### Row B

A rescue is still performed on downstream exons if a read is mapped on the intron of a given gene.

#### Row C

No special exon rescue is performed if an alignment falls on an intergenic region. Instead, regular rescue within the defined pairing distance is performed.

Alignments that lack a sibling read aligned nearby are called anchors. If anchors are found, the rescue tool conducts a more sensitive search for the sibling near the anchor, within a limited region of the genome. In the WT paired-end pipeline, the search is limited to anchors that occur within the introns or exons of annotated genes, with an allowance for a few overhanging bases.

Rescue is performed only within a set of expected rescue distances determined by a gene's exon structure and the insert size distribution. Rescue distances are a function of the transcript rescue distance:

*transcript rescue distance = mean insert size + 3 standard deviations* 

This distance is longer than the majority (99.7%) of inserts. The formula above describes the rescue distance without taking into account the presence of introns. Because inserts are very likely to contain introns, a rescue distance is calculated for each potential splice configuration of the gene. For each configuration, the rescue distance is the transcript rescue distance + length of introns within the relevant transcribed sequence interval. Alternatively, rescue can be performed on all exons to improve sensitivity, but with a substantial increase in both false positives and run time. Rescue can be applied using F3 anchors only, F5 anchors only, or both.

Regardless of the rescue steps employed, the rescue results supplement the alignments detected in individual mapping steps. The end result of rescue is a set of F3 and F5 alignments in the same format as that produced by mapping.

Pairing reads In the pairing step, pairs of reads are evaluated, assigned a mapping quality value, and written to a BAM file. The pairing range is set to 100,000 so that reads in adjacent exons are tagged as proper pairs. Unlike the pairing of genomic resequencing data, the mapping quality of read pairs is a function of genome annotation. Alignment pairs that do not occur within the same gene are penalized. For each pair, the alignment with the highest quality value is designated as the primary alignment. In the case of multiple highest quality, a single-pair alignment is selected randomly. Figure 36 shows an example of pairing range calculation with junction alignments.



Figure 36 WT annotation-aided alignment Pairing Quality Value (PQV)

See Table 87 for details about rows A to F in Figure 36.

 Table 87
 Annotation-aided alignment PQV description

Row(s)	Description	PVQ penalized?
А	Mates fall on the same exon.	No
В	One mate falls on an exon and the other falls on an intron.	Yes
C and D	Mates fall on separate exons of the same gene.	No
E	Mates fall on exons of different genes.	Yes
F	Spliced alignment where one mate partially falls on a known gene and the other falls on the exon of the same gene.	No



# BAM file generation

#### Ma2BAM, Pa2BAM

Ma2BAM and pa2BAM are format converters that change match files to a pro-BAM format. These pro-BAM files do not contain sequence or quality information but contain relevant BAM meta data. The pro-BAM files are sorted by incoming bead-id order. Relevant meta data from XSQ files is captured in pro-BAM files at this stage. The parameters bamgen.primary.output.filter.type, bamgen.mqv.threshold, and create.unmapped.bam.files control the functionality of the ma2BAM and pa2BAM utilities. The pro-BAM files are generated on the scratch folder.

Ma2BAM takes output of mapping (MA files) and converts them to pro-BAM format. The pa2BAM takes output of pairing (PA files) and converts them to pro-BAM format. During BAM generation, the mapping and pairing quality values are also computed.

For reads with multiple ungapped alignments, the read with the highest mapping or pairing quality value is chosen as the primary alignment for the read, and is reported to the BAM file. In cases where there are multiple alignments with the same quality value, the primary alignment is chosen at random from among the alignments with the same quality value. The details of calculating mapping and pairing quality values are described below.

#### Mapping and pairing quality values

This section describes mapping quality value (MQV) and pairing quality value (PQV) scores. Quality values are Phred-scaled quality scores. For any given alignment, an MQV is a measure of the confidence of a read aligning to the particular location, given all possible alignments for the read. A PQV represents the confidence in both alignments of a pair, and is a combined quality score for both reads. Quality values are within the range 0–100:

- 0 The highest probability of error
- 100 The lowest probability of error

The following factors increase confidence and increase a read's quality value:

- Longer alignment
- Fewer possible alignments
- Fewer mismatches within the alignment

These factors reduce confidence and reduce a read's quality value:

- Shorter alignment
- More possible alignments
- More mismatches within the alignment

The pairing algorithm reports multiple sets of possible alignments for any given pair of reads (F3/R3 tags for a mate-pair run and F3/F5-P2 tags for a paired-end run). The pairing quality algorithm uses a Bayesian approach to calculate the quality of a given alignment for a pair of reads and the alignment with the highest pairing quality value (PQV) is chosen as the primary alignment for the pair of reads. The PQVs represent the Phred-scaled quality score, and they are useful for downstream variant detection modules such as diBayes, small indels, large indels, and CNV.

es of mapping

In order to be consistent with the Phred quality score (-10\*log10[prob(error)]) used widely in literature, the quality is computed as the negative log odds of misaligning the read (or pairs of reads, for PQV). The resulting quality values are normalized by the maximum possible value to ensure that the quality values are within the range 0–100.

#### Gapped alignments

The pairing algorithm searches for gapped alignments (indels) when one of the tags (F3/R3/F5-P2/F5-BC) maps to the reference genome and the other tag does not map to the genome within the insert-size range, or it maps within the insert-range but the alignment is short. If both an ungapped and a gapped alignment are found for a given read, then, due to the low prior probability of 10^-4 assigned to the gapped alignments, the PQV for gapped alignments is zero.



Figure 37 Example of a gapped alignment and a partial alignment

In calculating the PQV for gapped alignments, the alternative hypothesis tested is the probability of finding the partial ungapped alignments. The read with the gapped alignment is treated as two partial reads on either side of the indel start point. The partial read with the greater length is used as the partial alignment length for the alternate hypothesis.

#### Refcor

Reference Assisted Color-to-Base Translation (also known as refcor) combines information from color calls, optional ECC extra primer round calls, and the reference sequence, to enhance the quality of base-space reads by detecting and fixing color errors that convert into multiple base-space errors.

Refcor accepts a set of pro-BAM files, which contain alignments information without base or color sequence, and generates a corresponding set of BAM files containing alignments with base-space and color-space calls (on the scratch folder). These BAM files are sorted by bead-ID and contain dummy base calls and QVs, but are otherwise compatible with the BAM specification. The refcor module produces bead-ID-sorted BAM files that have valid calls and QVs.

For each alignment, the module constructs the primer-transition graph with vertices corresponding to all possible k-mers (k=4 for 2BE+4BE, k=2 for 2BE) in each read position. Two adjacent vertices are connected with an edge if they overlap in k-1 bases. A score corresponding to the (1-Pe), where *Pe* is the probability of being erroneous, is assigned to each of the vertices. Scores are derived from quality values corresponding to enumerated calls. When making a base call in a position, the current algorithm considers the cumulative probability of all sequences that have called base in that

position. The evidence from the reference (base) is assigned to the vertex with *k*-mer ending in corresponding base only if the color transition between current and following reference bases matches the color call in a read. Such assignment reduces the reference over-correction in SNP positions. In order to reduce reference over-correction in positions with color errors, we assign to reference base a very low weight corresponding to QV of 8. The weight can be reduced to QV of 0, completely reducing the reference guiding in the color-to-base translation. This results in a smaller skew between reference to non-reference allele ratio. Increasing the weight results in false positive variant calls reduction. This process restores the phase shift caused by color errors or completely eliminates the errors.

The parameters bamgen.refcor.addcs, refcor.reference.weight, and refcor.base.filter.qv control the refcor program. The defaults provided are validated for a wide range of data.

- If the XSQ file contains base-space data only, the XSQ base call and quality value are transferred to BAM file. In this case, no color calls are present in BAM file irrespective of the value of bamgen.refcor.addcs.
- If the XSQ file contains 2BE color-space data only, refcor uses aligned 2BE and reference to generate base translation for the aligned part of the read. The base sequence and quality value are generated using a proprietary algorithm. For reads with no alignment, bases are determined by naively translating color to base. For these unaligned reads, the QV for the base at position *k* is the minimum color QV for all positions  $\leq k$ .
- If the XSQ file contains ECC data, then reference with 2BE and 4BE color are used to generate the new base sequence. The base sequence and quality value are generated using a proprietary algorithm. For reads with no alignment, the original base-space data from ECC is preserved.

#### Sort BAM

The output of refcor is a set of BAM files in bead-id sorted order. These files are sorted in this step using Picard sort into coordinate sorted order before merge. These BAM files are referred to as mini-BAM files and are stored in the analysis temporary directory.

#### Merge BAM

This phase accepts a set of mini-BAM files and outputs a set of BAM files. These BAM files are stored in the analysis output directory (which is typically a network-attached storage device).

One BAM file is generated per read-set, which is defined as a barcode index in one XSQ file. For a 96-barcoded dataset in a single XSQ file, either 96 or 97 BAM files are created. The 97th BAM contains reads that have the default barcode index.

### Single-read mapping parameters

Table 88 describes the algorithm parameters that you can configure for single-read whole transcriptome mapping.

This table lists mapping, resource, parallelization, and BAM-generation parameters. WT tertiary parameters are listed in their chapters:



- Chapter 18, Run a WT Coverage Analysis
- Chapter 19, Run a WT Count Known Genes and Exons Analysis
- Chapter 20, Run a WT Splice Finder Analysis

#### Table 88 Single-read WTA mapping parameters

Parameter	Default	Description
Mapping parameters		
enable.filter.mapping	1	Enables filter mapping.
enable.junction.mapping	1	Enables junction mapping.
mapping.in.base	false	Whether to map in base space (true) or color space (false). Set to true if input data has base space available.
		Allowed values:
		• <b>false</b> : Map in color space.
		true: Map in base space.
		If only color space is available, then the module fails when base space mapping is turned on.
wt.merge.min.junction.overhang	8	Alignments to junctions are not reported if they have fewer than this number of bases unaligned on either side of the splice. Allowed values: Integers 0–100.
Optional resource parameters		
java.heap.space	1500	Dynamic memory requirement.
mapping.memory	15gb	The total memory, in gigabytes, that is available for map reads, per compute node. Include the units gb in the setting.
		This number is set to 15 GB because the minimum hardware requirement for memory is 16 GB.
		<b>Note:</b> Typically, the job scheduler will allocate on nodes that have the requested memory available. For mixed hardware, it is suggested that different queues are created based on the memory available. Mapping jobs should be launched on high memory systems. For human references (hg18, hg19) and a 25.2 scheme, 45 GB is recommended for fastest performance. The smallest recommended RAM for human reference is 19 GB.
wt.filter.mapping.memory	14gb	Include the units gb in the setting.
wt.genomic.mapping.memory	14gb	Include the units gb in the setting.
wt.junction.mapping.memory	14gb	Include the units gb in the setting.
mapping.schema.file	-	The mapping schemas file.
Optional parallelization parameters		
fragmap.number.of.nodes	4	The number of compute nodes available for this analysis.
fragmap.max.number.of.jobs	100	The maximum number of nodes allowed for mapping.
fragmap.minreads.per.node	4000000	If the total number of fragments exceeds this amount, then the analysis is distributed across the available nodes.
		Allowed values: Integers 1-125000000 (125 Million)



#### Table 88 Single-read WTA mapping parameters (continued)

Parameter	Default	Description
fragmap.maxreads.per.node	15000000	The maximum number of fragments that can be processed on a single node.
		Allowed values: Integers 1–150000000 (150 Million)
processors.per.node	8	The number of processors per node use for mapping. The reads are divided into the number of chunks specified for this parameter.
wt.filter.mapping.np.per.node	8	The number of processors per node use for filter mapping.
wt.genomic.mapping.np.per.node	8	The number of processors per node use for genomic mapping.
wt.junction.mapping.np.per.node	8	The number of processors per node use for junction mapping.
Optional BAM generation parameters	;	
bamgen.refcor.addcs	1	Whether or not to add the color sequence to BAM records. If the XSQ file does not contain color space, this parameter is ignored, and the resulting BAM file output does not contain color space. See "Refcor" on page 339 for information about refcor. Allowed values: 0,1.
create.unmapped.bam.files	FALSE	<ul> <li>If set to TRUE, creates an additional BAM output file containing unmapped reads. Allowed values:</li> <li>FALSE: Do not create the unmapped reads output file.</li> <li>TRUE: Create a BAM output file containing the unmapped reads.</li> </ul>
bamgen.mqv.threshold	0	Provides control over the contents written to the output BAM file depending on the quality value of the alignment. To preserve only high quality alignments, set this value to a positive integer. Allowed values: Integers 0–255.
refcor.base.filter.qv	10	Bases with a quality value below the value of this parameter provided are replaced with 'N'. Allowed values: Integers 0–255.
refcor.reference.weight	8	Reference weight. This parameter is used during base translation. In the read reconstruction process, multiple signals are combined to generate the final base call. This parameter adds weight (in terms of Phred score) to the signals which are compatible with reference. Color combinations that result in a variant are considered compatible with reference. Additional weight helps to eliminate base errors caused by color error(s) during base translation. Allowed values: Integers 0–100.

#### Mapping performance

Mapping speed depends on hardware properties as well as on the scattering logic. Mapping runs are split into multiple jobs for processing efficiency. The following parameters affect how a mapping run is split:

- **fragmap.number.of.nodes** Specifies the number of jobs that are created, only if a split is necessary. The size of each split job is approximately the total number of reads divided by the number of jobs.
- **fragmap.max.number.of.jobs** Determines the maximum number of jobs that can be launched per analysis. Setting this parameter to a very large value can cause the scheduler to behave incorrectly.



- **fragmap.minreads.per.node** Determines the threshold of beads beyond which splitting occurs.
- **fragmap.maxreads.per.node** Determines the largest number of beads that can be mapped per node. The default is based on the minimum scratch space requirements. If you scratch space is smaller or larger, this number can be made smaller or larger respectively.
- **processors.per.node** This is the number of cores available for mapping analysis on each node.
- **mapping.memory** For human mapping and 25.2.0 scheme, there are 3 schema lines. Simultaneous handling of schema lines reduces I/O and therefore improves mapping speed.

Memory requirements for mapping jobs are the following:

- Reference: ~6 GB
- 1 schema line: ~13 GB
- 1 schema line and reference: ~19 GB
- 2 schema lines and reference: ~32 GB
- 3 schema lines and reference: ~45 GB

Below 19 GB, the reference is split up and I/O increases significantly. Pre-built hash tables are not used for memory less than 24 GB. LifeScope<sup>™</sup> Software supports the reuse of hash tables for human genome reference files (hg18 and hg19) and the default mapping schemes 35.2, 25.2, and 20.1. Pre-calculated color-space are included in the reference directory available with LifeScope<sup>™</sup> Software. If system RAM is 24 GB or higher, the hash tables are loaded into memory. Depending on the pipeline, this hash table reuse typically saves 2–3 hours in analysis processing time.

In general, set the mapping.memory parameter to be 1 GB less than the total system memory (RAM) available.

Mapping only scales when the number of reads that are mapped together is greater than 100 M. Splitting such that the total number of fragments per node is significantly less than 100 M is not recommended. The splitting can be explained using the following pseudo-code.

```
if (num.reads <= fragmap.minreads.per.node ) { num.jobs = 1}
reads.per.job = num.reads / fragmap.number.of.nodes
if (reads.per.job > fragmap.maxreads.per.node) {
    reads.per.job = fragmap.maxreads.per.node}
num.jobs = celing (num.reads / reads.per.job)
if (num.jobs > fragmap.max.number.of.jobs ) {
    throw exception; }
```

In above pseudo-code, the input read-sets (possibly from multiple XSQ files) have num.reads that fall in one read length category. Num.jobs is the total number of jobs launched for mapping analysis of num.reads beads.

In general, the mapping parameters allow mixing and matching single-anchor and multiple-anchor schemes, and also mixing and matching different seed lengths and different numbers of mismatches. For example, an unmapped scheme of 25.2.0, 35.2.0:15:25, 20.1.0, with a repetitive scheme of 35.2.0:30, 45.2.0, 60.2.0, is supported, but not recommended. The impact on performance can be significant.



# WT mapping<br/>internalTable 89 describes the single-read algorithm parameters that we do not recommend<br/>changing.parametersChanging.

 Table 89
 Single-read
 WTA internal parameters

Parameter	Value	Description
wt.frag.map.run	1	Enables fragment mapping.
		The run parameter is set automatically by the shell during a standard workflow. Accept the default for most use.
wtexonsequenceextractor.secondary.run	1	Enables exon extraction.
		The run parameter is set automatically by the shell during a standard workflow. Accept the default for most use.
wtsplicejunctionextractor.run	1	Enables splice junction extraction, which is required for junction mapping.
		The run parameter is set automatically by the shell during a standard workflow. Accept the default for most use.
Algorithm parameters		
mapping.use.iub.reference	FALSE	Whether or not to support reference sequences that contain IUB codes. Allowed values:
		• FALSE: Do not support IUB codes in the reference file.
		• <b>TRUE</b> : Mapping allows matching to either alleles of bi- allelic IUB codes in the reference file.
bamgen.primary.output.filter.type	primary_ only	<ul> <li>Filters the content written to the output BAM file depending on whether the alignment is gapped or not. Note that unmapped reads are considered primary. Allowed values:</li> <li>no_filtering: Report all alignments.</li> </ul>
		<ul> <li>primary_only: Report all primary alignments (both gapped and ungapped).</li> </ul>
wt.filter.map.max.hits	10	The maximum number of alignments for a read, during filter mapping. Allowed values: Integers 1–100000.
wt.genomic.map.max.hits	10	The maximum number of alignments for a read, during genomic mapping. Allowed values: Integers 1–100000.
wt.junction.map.max.hits	10	The maximum number of alignments for a read, during junction mapping. Allowed values: Integers 1–100000.
junction.reference.file	_	The junction reference file. Must be a FA or FASTA file.
wt.mask.positions	-	A string of 0s and 1s that show positions to be masked or not. If blank (default), no masking is performed. There is no penalty in mapping towards positions that are masked.
wt.map.mismatch.penalty	-2.0	Specifies a negative scoring penalty for mismatch that is used in local alignment mode. Setting this value to the negative of a number greater than the read lengths has the effect of disallowing mismatches. Allowed values: Floats.

Mapping scheme parameters



Parameter Value Description map.scheme.unmapped.1 0.0.0 Mapping scheme used for reads with lengths less than 25. map.scheme.unmapped.25 25.2.0 Mapping scheme used for reads with lengths 25-34. 25.2.0:10 Mapping scheme used for reads with lengths 35-49. map.scheme.unmapped.35 25.2.0:20 Mapping scheme used for reads with lengths 51-74. map.scheme.unmapped.50 map.scheme.unmapped.75 25.2.0:20 Mapping scheme used for reads with lengths of 75 and greater. 25.2.0:20 map.scheme.unmapped.variable Mapping scheme used for trimmed reads.

#### Table 89 Single-read WTA internal parameters (continued)

#### Mapping schemes

The mapping scheme parameters, listed in Table 89, determine the behavior of the mapping module. The LifeScope<sup>™</sup> Software mapping module supports two types of mapping, classic mapping and seed-and-extend mapping.

- **Classic mapping** With this approach the seed length matches the full read length, and the seed anchors at the beginning of the read. For example, classic mapping for 50-bp reads uses a scheme such as 50.6.0. (In this example, 50, the seed length, matches the read length, and 0, the seed start position, starts at the beginning of the read.)
- Seed-and-extend The initial alignment step that locates short matches between a read and the reference sequence. The seed specifies the length of the attempted match. This approach is also called local alignment.

#### **Classic mapping**

To map a 50 bp read using classic mapping, a typical scheme to use is 50.6.0. This allows for up to 6 mismatches. Because the seed length equals the length of the read, there is no extension.

#### Seed-and-extend

With the seed-and-extend approach, a mapping scheme parameter value such as 25.2.0 is interpreted as follows:

- **First value –** The seed length (25).
- Second value The quantity of allowed mismatches in the seed (2).
- **Third value** The start site of the seed within the read (0).



allowed in the seed

Figure 38 Seeds for local alignments



For example, with a 25.2.0 seed extend scheme, the mapping module examines the first 25 bases of a 50 bp read, and if that portion aligns to the reference with two or fewer mismatches, then the mapping module attempts seed extension. Seed extension begins at the anchor alignment and extends along the read, to find the best ungapped local alignment with a score *S*. The alignment length can be shorter than the read length.

For information on sliding periodic seeds, refer to the publication at this site:

http://bioinformatics.oxfordjournals.org/content/25/19/2514.full

## **Paired-end parameters**

Table 90 describes the algorithm parameters that you can configure for paired-end whole transcriptome mapping.

 Table 90
 Paired-end WTA mapping parameters

Parameter	Default	Description			
Mapping parameters					
enable.exon.mapping	1	Enables exon mapping.			
enable.filter.mapping	1	Enables filter mapping.			
enable.junction.mapping	1	Enables junction mapping.			
wt.f3.rescue.enabled	true	Whether or not to attempt rescue of F3 reads.			
wt.f5.rescue.enabled	true	Whether or not to attempt rescue of F5 reads.			
std.insert.size	60	The standard deviation of the insert size. Allowed values: Integers 1–100000.			
mapping.in.base	FALSE	Whether to map in base space (TRUE) or color space (FALSE). Set to TRUE if input data has base space available.			
		Allowed values:			
		• FALSE: Map in color space.			
		TRUE: Map in base space.			
		If only color space is available, then the module fails when base space mapping is turned on.			
wt.merge.min.junction.overhang	8	Alignments to junctions are not reported if they have fewer than this number of bases unaligned on either side of the splice. Allowed values: Integers 0–100.			
Splice junction extractor parameters	1	1			
read.length	75	Read length. Allowed values: Integers 25–100.			
Rescue parameters	Rescue parameters				
avg.insert.size	120	The average the insert size. Used to compute the rescue distance: rescueDistance = avgInsertSize + (3 * stdInsertSize) Allowed values: Integers 1–100000.			
wt.f3.rescue.max.mismatches.allowed	8	The maximum number of mismatches allowed in a rescued F3 alignment. Allowed values: Integers 0–15.			



Table 90	Paired-end	WTA	mapping	parameters	(continued)
	i un cu chu		mapping	parameters	(contaca)

Parameter	Default	Description
wt.f5.rescue.max.mismatches.allowed	6	The maximum number of mismatches allowed in a rescued F5 alignment. Allowed values: Integers 0–10.
Optional resource parameters		
java.heap.space	1500	Dynamic memory requirement.
mapping.memory	15gb	The total memory, in gigabytes, that is available for map reads, per compute node. This number is set to 15 GB because the minimum hardware requirement for memory is 16 GB.
		<b>Note:</b> Typically, the job scheduler will allocate on nodes that have the requested memory available. For mixed hardware, it is suggested that different queues are created based on the memory available. Mapping jobs should be launched on high memory systems. For human references (hg18, hg19) and a 25.2 scheme, 45 GB is recommended for fastest performance. The smallest recommended RAM for human reference is 19 GB.
Optional parallelization parameters		
fragmap.number.of.nodes	4	The number of compute nodes available for this analysis.
fragmap.max.number.of.jobs	100	The maximum number of nodes allowed for mapping.
fragmap.minreads.per.node	4000000	If the total number of fragments exceeds this amount, then the analysis is distributed across the available nodes.
		Allowed values: Integers 1-125000000 (125 Million).
fragmap.maxreads.per.node	150 M	The maximum number of fragments that can be processed on a single node.
		Allowed values: Integers 1-150000000 (150 Million).
processors.per.node	8	The number of processors per node use for mapping. The reads are divided into the number of chunks specified for this

Table 91 describes WTA paired-end algorithm parameters that we do not recommend changing.

 Table 91
 Paired-end WTA internal parameters

Parameter	Value	Description
wt.pe.map.run	1	Enables paired-end mapping.
		The run parameter is set automatically by the shell during a standard workflow. Accept the default for most use.
Splice junction parameters		
wt.f3.splext.output.reference	-	The name of the reference file output by the splice junction extractor stage.
wt.gtf.file	—	The GTF file used to create the genemodel.
		The default is \${annotation.gtf.file}.
Splice junction extractor parameters	1	-

parameter.



#### Table 91 Paired-end WTA internal parameters (continued)

Parameter	Value	Description
wtspljunctionextractor.run	1	Enables splice junction extraction, which is required for junction mapping.
		The run parameter is set automatically by the shell during a standard workflow. Accept the default for most use.
Exon extractor parameters		
wtexonsequenceextractor.secondary.run	1	Enables exon extraction, which is required for F5 pairing.
		The run parameter is set automatically by the shell during a standard workflow. Accept the default for most use.
wt.f5.exseqext.output.reference		The name of the reference file output by the exon extractor stage.
Algorithm parameters		
mapping.use.iub.reference	FALSE	Whether or not to support reference sequences that contain IUB codes. Allowed values:
		• <b>FALSE</b> : Do not support IUB codes in the reference file.
		• <b>TRUE</b> : Mapping allows matching to either alleles of bi- allelic IUB codes in the reference file.
wt.map.mismatch.penalty	-2.0	Specifies a negative scoring penalty for mismatch that is used in local alignment mode. Setting this value to the negative of a number greater than the read lengths has the effect of disallowing mismatches. Allowed values: Floats.
wt.f3.filter.map.max.hits	10	The maximum number of alignments for an F3 read, during filter mapping. Allowed values: Integers 1–100000.
wt.f3.genome.map.max.hits	10	The maximum number of alignments for an F3 read, during genomic mapping. Allowed values: Integers 1–100000.
wt.f3.junction.map.max.hits	10	The maximum number of alignments for an F3 read, during junction mapping. Allowed values: Integers 1–100000.
wt.f3.map.mask.positions	-	A string of 0s and 1s that show F3 positions to be masked or not. If blank (default), no masking is performed. There is no penalty in mapping towards positions that are masked.
wt.f3.mapping.schema.file	_	Mapping schema file for F3 reads.
wt.f5.filter.map.max.hits	10	The maximum number of alignments for an F5 read, during filter mapping. Allowed values: Integers 1–100000.
wt.f5.genome.map.max.hits	10	The maximum number of alignments for an F5 read, during genomic mapping. Allowed values: Integers 1–100000.
wt.f5.exon.map.max.hits	10	The maximum number of alignments for an F5 read, during exon mapping. Allowed values: Integers 1–100000.

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	/

Table 91 Paired-end WIA internal parameters (cont	inued)
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Parameter	Value	Description	
wt.f5.map.mask.positions	-	A string of 0s and 1s that show F5 positions to be masked or not. If blank (default), no masking is performed. There is no penalty in mapping towards positions that are masked.	
wt.f5.mapping.schema.file	_	Mapping schema file for F5 reads.	
wt.map.mismatch.penalty	-2.0	Specifies a negative scoring penalty for mismatches. Used in local alignment mode. Setting this value to the negative of a number greater than the read lengths has the effect of disallowing mismatches. Allowed values: Floats.	
exon.reference.file	exons.fasta	Path to the exon reference file. Must be a FA or FASTA file.	
junction.reference.file	_	Path to the junction reference file. Must be a FA or FASTA file.	
Pairing parameters			
pair.uniqueness.threshold	10.0	This value defines the clear zone. This value is considered only when pairing.output.uniqueness.type is set to -1. If the score of the best alignment is this threshold more than the score of the second best alignment, the best alignment is considered uniquely placed. A threshold of zero means that this uniqueness is strictly adhered to, regardless of the alignment's relative scores.	
		Allowed values: Floats 0.0–100.0.	
pairing.anchor.max.mismatches	4	Total mismatches for both reads of a pair.	
		Allowed values: Integers 0–100.	
pairing.gapped.max.hits	100:0	For gapped and ungapped alignment determination during rescue, this value determines the maximum number of rescues attempted for each read. If a read has more than this number of hits, rescue is not performed on this bead. The first value is for ungapped rescue, and second, gapped.	
pairing.mismatch.penalty	-2.0	A single penalty applied to alignments, to compare the significance between alignments. Helps evaluate the alignments. Allowed values: Floats –100 to 0.	
pairing.output.uniqueness.type	-1	Whether to output non-unique hits. Allowed values:	
		<ul> <li>-1: A clear zone determines uniqueness. One or two best pairs are output, depending on whether it is unique.</li> </ul>	
		• <b>0</b> : All good pairs are output.	
		• 1: Only unique good pairs are output.	
		Allowed values: Integers –1, 1.	
Rescue parameters			
wt.rescue.input.generation.exon.fuzzy. border.width	10	The amount of slack allowed around exon borders, for doing rescue with fuzzy exon borders. Allowed values: Integers 0–2147483647.	



#### Table 91 Paired-end WTA internal parameters (continued)

Parameter	Value	Description
wt.rescue.input.generation.min.alignment. distance.for.rescue	100000	When rescuing reads that have already been mapped to the reference, this value is the minimum distance between the anchor alignment and any of the alignments of the rescued read downstream of the anchor alignment. Allowed values: Integers 0–2147483647.
wt.rescue.input.generation.rescue.anchor. alignments.not.overlapping.exons	1	<ul> <li>Enables the additional rescue downstream and upstream of alignments that do not overlap exons but do overlap genes.</li> <li>Allowed values:</li> <li>0: No change in behavior.</li> <li>1: Enables the rescue described above.</li> </ul>
wt.rescue.input.generation.rescue.fuzzy. exon.borders	1	<ul> <li>Enables allowing some slack around exons borders.</li> <li>Allowed values:</li> <li>0: No change in behavior.</li> <li>1: Enables slack around exons borders.</li> </ul>
wt.rescue.input.generation.rescue.only.for. the.best.anchor.alignment	0	<ul> <li>Limits rescue to only for the best anchor alignment.</li> <li>Allowed values:</li> <li>0: Rescue for all anchor alignments.</li> <li>1: Rescue only for the best anchor alignment.</li> </ul>
wt.rescue.input.generation.rescue.only. unaligned.reads	0	<ul> <li>Rescue ignores target reads that already have an alignment.</li> <li>Allowed values:</li> <li>0: Rescue all the target reads.</li> <li>1: Rescue ignores target reads that already have an alignment.</li> </ul>
wt.rescue.input.generation.rescue.only.within. rescue.distance	1	<ul> <li>Rescue only within rescue distance. Rescue only within the rescue region starting from the exon border, to a limit inside the exon. The limit is computed based on the insert size and the location of the anchor alignment inside the previous exon.</li> <li>Allowed values:</li> <li>0: Rescue on the entire exon.</li> <li>1: Rescue only within rescue distance.</li> </ul>
wt.rescue.valid.adjacent.mismatches.count. as.one	0	<ul> <li>Two adjacent mismatches that can account for a SNP count only as one mismatch.</li> <li>0: No change in behavior.</li> <li>1: Two adjacent mismatches that can account for a SNP count only as one mismatch.</li> </ul>
wt.rescue.input.generation.rescue.short. range	1	<ul> <li>Limits rescue to all the target reads in the vicinity of the anchor read. Allowed values:</li> <li>0: No change in behavior.</li> <li>1: Limits rescue to all the target reads in the vicinity of the anchor read.</li> </ul>
wt.rescue.input.generation.zLimit	100	The Z limit. Allowed values: Integers 0–2147483647.

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	-

Parameter	Value	Description
wt.rescue.standalone.rescue.executable	rescue	The name of the executable file used for rescue.
Mapping scheme parameters		
wt.f3.mapping.scheme.unmapped.1	0.0.0	Mapping scheme for F3 reads with lengths less than 25.
wt.f3.mapping.scheme.unmapped.25	25.2.0	Mapping scheme for F3 reads with lengths 25–34.
wt.f3.mapping.scheme.unmapped.35	25.2.0:10	Mapping scheme for F3 reads with lengths 35–49.
wt.f3.mapping.scheme.unmapped.50	25.2.0:20	Mapping scheme for F3 reads with lengths 51–74.
wt.f3.mapping.scheme.unmapped.75	25.2.0:20	Mapping scheme or F3 reads with lengths of 75 and greater.
wt.f3.mapping.scheme.unmapped.variable	25.2.0:20	Mapping scheme for F3 reads with variable lengths (trimmed reads).
wt.f5.mapping.scheme.unmapped.1	0.0.0	Mapping scheme for F5 reads with lengths less than 25.
wt.f5.mapping.scheme.unmapped.25	25.2.0	Mapping scheme for F5 reads with lengths 25–34.
wt.f5.mapping.scheme.unmapped.35	25.2.0:10	Mapping scheme for F5 reads with lengths 35–49.
wt.f5.mapping.scheme.unmapped.50	25.2.0:20	Mapping scheme for F5 reads with lengths 51–74.
wt.f5.mapping.scheme.unmapped.75	25.2.0:20	Mapping scheme for F5 reads with lengths of 75 and greater.
wt.f5.mapping.scheme.unmapped.variable	25.2.0:20	Mapping scheme for F5 reads with variable lengths (trimmed reads).
wt.f5.exon.mapping.scheme.unmapped.1	0.0.0	Mapping scheme for F5 reads with lengths less than 25.
wt.f5.exon.mapping.scheme.unmapped.25	25.3.0	Mapping scheme for F5 reads with lengths 25–34.
wt.f5.exon.mapping.scheme.unmapped.35	25.3.0:10	Mapping scheme for F5 reads with lengths 35–49.
wt.f5.exon.mapping.scheme.unmapped.50	25.3.0:20	Mapping scheme for F5 reads with lengths 51–74.
wt.f5.exon.mapping.scheme.unmapped.75	25.3.0:20	Mapping scheme for F5 reads with lengths of 75 and greater.
wt.f5.exon.mapping.scheme.unmapped. variable	25.3.0:20	Mapping scheme for F5 reads with variable lengths (trimmed reads).

Table 91 Paired-end WTA internal parameters (continued)

# Mapping output files

**Overview** 

The mapping module generates a BAM file containing alignments in coordinate order. For information about the BAM file, see Appendix B, "File Format Descriptions" on page 455.

The number of BAM files generated depends on the input XSQ files. One BAM file is generated per read-set (a read-set is defined as data from one barcode, in one XSQ file). If the input data is an XSQ file containing 96 barcodes, then the mapping module generates at least 96 output BAM files. Beads that are unclassified in any barcode are output into a separate additional BAM file. If create.unmapped.bam.files is set to TRUE, then an additional 96 output BAM files corresponding to the unmapped reads are also generated.

The filenames for the output BAM files are created using information from the input XSQ file, including: file base name, file id, index name, and index id. The BAM files are named according to the following patterns:

- Non-indexed BAM files: xsqname-fileID-1.bam
- Indexed BAM files: xsqname-fileID-idx\_bcIndex-bcID.bam

The fields in the filenames are:

- **xsqname** The XSQ file base name, without the .xsq extension.
- fileID The file ID for internal XSQ file tracking.
- **bcIndex** The barcode index.
- **bcID** The barcode identifier for internal barcode tracking.

The directory structure for mapping output is as follows:

The directory name <mappingIniFileBasename> is the basename of the mapping INI file. The defaults are wt.frag.map and wt.pe.map.

The string *<sampleName*\*> is determined by the sample description for the particular read-set.

The mapping output files are used by the BAMStats mapping statistics module and by the whole transcriptome tertiary analysis modules: WT counts, WT coverage, and WT splice junctions.

This section describes two types of differences seen in WT BAM files, compared to other LifeScope<sup>™</sup> Software BAM files.

#### Single-read optional fields

A BAM file produced by the WT paired-end pipeline is identical to that produced by the resequencing pipeline. However, the BAM format from the WT single-read pipeline differs from BAM files produced elsewhere in LifeScope<sup>™</sup> Software. The single-read pipeline produces separate BAM files for filtered, unmapped, and mapped reads.

**Note:** The WTA single-read pipeline produces a BAM file that uses the optional fields described in Table 92.

Optional field	Description
IH:i:	Number of stored alignments containing the current query.
HI:i:	Query hit index.
NH:i:	Number of reported alignments containing the current query.
CS:z:	Color read sequence.
CQ:z:	Color quality sequence.
CC:z:	Reference name of the next hit.

Table 92 WT single-read pipeline BAM file optional fields

**BAM** file

differences

Optional field	Description
CP:z:	Coordinate of the next hit.
AS:i:	locationAlignment Score generated by the aligner.
XN:i:	Alignment score of the best non-primary alignment for query in the current record.
XF:z:	T for true or F for false. Set to "T" if this read is filtered.
XJ:z:	"K" for Known Junction or "P" for Putative Junction.

 Table 92
 WT single-read pipeline
 BAM file optional fields (continued)

#### **Clipped records**

By design, BAM records produced in WT modules may have soft-clips in the CIGAR string. For records that have soft-clips at the beginning, the correct start position of reported sequence is interpreted as the start position minus the soft-clipped length. However, BAM-consuming external tools that are not BAM-specification complaint may present soft-clipped records as shifted or missed alignments.

#### WT filtering stats

An example whole transcriptome human filter reference is provided under the <*examplesdir*>/demos/WholeTranscriptome/references folder in the optional LifeScope<sup>TM</sup> Software examples distribution. This example reference fasta includes contigs from barcode primers, human ribosomal RNAs, tRNAs, and other known targets for filtering. The filter reference may be expanded or removed with more csfasta reference records at the discretion of the user.

The stats for filtered reads are generated after a LifeScope<sup>™</sup> Software run. First two lines of the stats file states the number of reads processed, and the number of reads mapped to the filtered reference followed by its percentage. Each following line reports a contig name and the count of reads that aligned to that contig. Paired-end filtering stats may be found in the Intermediate folder. The following is a truncated example of filtering stats output:

```
countOfReadsProcessed: 175,839,941
countOfReadsMapping:
                        18,783,220 (10.7%)
....
Barcode-047-3-end reverse
                            83
Barcode-048-3-end reverse
                            2,350
gi|124517659|ref|NR_003286.1| Homo sapiens 18S ribosomal RNA
(LOC100008588)
                 5,175,899
gi|142372596|ref|NR_003285.2| Homo sapiens 5.8S ribosomal RNA
                 78,936
(LOC100008587)
gi|124517661|ref|NR 003287.1| Homo sapiens 28S ribosomal RNA
(LOC100008589)
                 7,616,212
chr6.trna95-AlaAGC (58249908-58249836) Ala (AGC) 73 bp
                                                           Sc:
42.26
       5
chr6.trna25-AlaAGC (26859897-26859969)
                                         Ala (AGC) 73 bp
                                                           Sc:
46.89
       2
chr6.trna94-AlaAGC (58250620-58250548)
                                         Ala (AGC) 73 bp
                                                           Sc:
54.62
        4
chr6.trna160-AlaAGC (26881822-26881750)
                                         Ala (AGC) 73 bp
                                                           Sc:
54.69
       35
```



# **Mapping statistics**

Mapping statistics occur after mapping as an optional post-processing step named BAMStats. BAMStats accepts the output of the mapping step and generates statistics files to provide an in-depth understanding of the experimental data and to better detect the presence of anomalies. LifeScope<sup>™</sup> Software shell users can display the mapping statistics output data as a chart with a spreadsheet program or other third-party program. A subset of the output from mapping can be visualized in the LifeScope<sup>™</sup> Software UI as a series of line and bar charts, if the analysis is run in the projects repository.

Mapping statisticsTable 93 lists the parameters which control the BAMStats output.parameters

Table 93 BAMStats parameter description

Parameter name	Default value	Description
Mandatory parameters		
bamstats.run	1	Whether or not to run the BAMStats module. Allowed values:
		• <b>0</b> : Do not run the BAMStats module. BAMStats statistics are not generated.
		• 1: Run the BAMStats module during this analysis.
		The run parameter is set automatically by the shell during a standard workflow. Accept the default for most use.
Optional parameters		
bamstats.maximum.coverage	10000	Defines the maximum coverage allowed for locations in the reference. Locations with coverage more than the maximum coverage value are ignored during coverage calculations. Allowed values: Integers 1–10000.
bamstats.maximum.isize	100000	The maximum inset size for LMP and PE libraries. Reads with an insert size more then the specified value are ignored for Insert Range Report calculation.
		Allowed values: Integers 1–100000.
bamstats.wig.primary.only	1	Use only primary alignments for coverage in WIG file format. Allowed values:
		• <b>0</b> : Do not restrict coverage in WIG file format to only primary alignments.
		<ul> <li>1: Restrict coverage in WIG file format to only primary alignments.</li> </ul>
bamstats.combined.report.both. strands	0	Whether or not to combine data from both strands for coverage in WIG format. Allowed values: 0,1.
bamstats.wig.binsize	100	The bin size for coverage in WIG file format.
		Allowed values: Integers 1–100000.
bamstats.bin.isize	100	The bin size for insert range distribution.
		Allowed values: Integers 1–100000.



 Table 93
 BAMStats parameter description (continued)

Parameter name	Default value	Description
bamstats.wig.combined.report.both .strands	0	Whether to combine data from both the strands for coverage in WIG format. Allowed values:
		• <b>0</b> : Do not combine data.
		<ul> <li>1: Combine data from both the strands for coverage in WIG format.</li> </ul>
Optional resource parameters		
wall.time	120	Total time for the process to complete.
java.heap.space	13000	Dynamic memory requirement.
number.of.nodes	4	The number of compute nodes available for this analysis.
memory.request	14gb	Memory request. Include the units gb in the setting.

Table 94 lists BAMStats parameters that we do not recommend changing.

Table 94	BAMStats	internal	parameter	description
----------	----------	----------	-----------	-------------

Parameter name	Default value	Description
bamstats.group.stats	1	Enable combined statistics covering all read-sets in the group or sample. Allowed values:
		• <b>0</b> : Only statistics for BAM files are generated.
		• 1: Also generate combined statistics for the group or sample.
bamstats.maximum.mismatches	100	The maximum mismatches allowed in the alignments. Any alignment with more then the specified number of mismatches is ignored while generating reports related to number of mismatches. Allowed values: Integers 0–100.
bamstats.maximum.baseqv	100	Max base quality values. Any base with base quality value more then the specified value is ignored while generating reports. Allowed values: Integers 0–100.
bamstats.maximum.mappingqv	255	Maximum mapping quality value. Any alignment with mapping quality value more than the specified value is ignored. Allowed values: Integers 0–255.
bamstats.wig.minimum.mappingqv	2	Defines the minimum mapping quality allowed for coverage in WIG format. Any alignment with a mapping quality value less than this value is ignored. Allowed values: Integers 0–100.
bamstats.enable.probe.position	0	Enable probe and position error reports. Allowed values:
		• <b>0</b> : Do not generate probe and position error reports. Off is recommended for WT.
		• 1: Also generate probe and position error reports.

#### Summary of mapping statistics output

#### File formats

For every input BAM file, a set of statistics files are generated. These files are in CHT, CSV, TXT, and WIG formats. Each CHT file corresponds to one displayed chart. A CHT file specifies the type of chart, the displayed range of each axis, and the data points, without using external references.



The CHT file format is an internal file format based on the CSV file format, with addition header information included. CHT header information is the following:

```
# name:
# type: scatter2d | pie | vbar | line
# title:
# xaxisname:
# yaxisname:
# yrange: <min>:<tickinterval>:<max>
# yrange: <min>:<tickinterval>:<max>
XAXISNAME, SERIES1NAME, SERIES2NAME, ...
x1, y1.1, y1.2, ...
x2, y2.1, y2.2, ...
x3, y3.1, y3.2, ...
```

The wiggle format (.wig) is a public format typically used for coverage. Visit their site for more information:

hgdownload.cse.ucsc.edu/goldenPath/help/wiggle.html

A genome browser such as the Integrative Genomics Viewer (IGV) can be used to visualize the coverage. For information is available from their site:

```
www.broadinstitute.org/igv/
```

For a collection of input BAM files that belong to a sample, a set of cumulative statistics files are generated. The cumulative statistics files are also in CHT, CSV, TXT, and WIG formats. The cumulative statistics can be visualized in the LifeScope<sup>™</sup> Software UI.

#### Summary information

A tab-separated summary file (*BAMfilename*-summary.tbl) is generated that summarizes key mapping quality statistics per input BAM file. This file contains one row for each input BAM file in the sample. The summary file is displayed in the LifeScope<sup>™</sup> Software UI. See "Summary file" on page 361.

#### **Directory structure**

The output of the BAMStats module has the following directory structure:

bamstats/

```
<sampleName>/*.cht, *.tbl
<sampleName>/<bam>/.*cht
<sampleName>/<bam>/Misc/*.csv, *.txt, *.wig
<sampleName>/Misc/*.csv, *.txt
```

The position error files and probe errors files are created in the BAM file directories.

#### **Overview**

The summary report contains a snapshot of statistics in all BAM files present in this sample. This report is displayed in the LifeScope<sup>™</sup> Software UI.

<sampleName1>/BAMfilename-summary.tbl

Following directories and reports contain the cumulative statistics from all BAM files that belong to this sample. The Misc folder is not displayed in UI.

<sampleName1>/\*.cht

```
<sampleName1>/Misc/*.csv, *.wig, *.txt
```

Following statistics are generated per BAM file in the mapping directory. These reports are not displayed in the UI.

```
<sampleName1>/<BAMfilename>/.*cht
<sampleName1>/<BAMfilename>/Misc/*.csv, *.txt, *.wig
```

In a sample, some of the BAM files possibly represent unhealthy DNA or RNA, causing the cumulative statistics to look poor. The summary tbl file is the unified location for examining the quality of data of all BAM files in the sample.

If the data for a particular BAM file is not as expected, look at the directory-level reports for that BAM file, for details.

## Mapping statistics output files

This section describes the mapping statistics files generated by the BAMStats module. When statistics reports are separated by tag type, the report's file name includes the tag in the file name. The tags used in file names are:

Library type	Tag
Fragment	F3
Mate-pair	F3
	R3
Paired-end	F3
	F5-P2

Table 95 Tags in mapping statistics output file names

The output files generated by BAMStats include the name of the BAM file. The file name pattern is:

BAMFileName-fileID-barcodeID.StatisticsReportName.tag.extension

This string is referred to by *prefix* in Table 96, the mapping statistics output table. In the description of mapping statistics output file names in Table 96, the string *tag* is used to refer to the tags in Table 95.

Table 96 Mapping statistics output files

Report	Description, axis information, filename		
Alignment Length Distribution			
	A bar plot giving the distribution of alignment lengths found in various tags used in alignment. The report is separated by tag type to provide visibility into the accuracy of individual tags.		
	Only the primary alignment for each bead is considered in calculating the distribution.		
	Y axis: Frequency X axis: Alignment length (from 0 to the maximum read length)		
	Output file name: prefix.Alignment.Length.Distribution.tag.cht		
Alignment Length Distribution for Unique Alignments			



#### Table 96 Mapping statistics output files (continued)

Report	Description, axis information, filename
Th	e same report as the Alignment Length Distribution, except that only tags that have a primary gnment are considered in calculating the distribution.
By Ali Ho dis	default BAM files contain primary alignments only. In this case, the reports gnmentLengthDistribution and AlignmentLengthDistribution for Unique Alignments are identical. wever, users can also print all alignments or secondary alignments in BAM file. In this case, the two stributions are different.
No	te: The title Unique Alignments is interpreted to mean primary alignments, for this report.
Y a X a	axis: Frequency axis: Alignment length (from 0 to the maximum read length)
Ou	tput file name: <i>prefix</i> .Alignment.Length.Distribution.Unique. <i>tag</i> .cht
Base Mismatch Dis	tribution
A t Th	par plot giving the distribution of total number of mismatches found in various tags used in alignment. e bins are summed over all alignment lengths.
On	ly the primary alignment for each bead is considered in calculating the distribution.
Y a X a	axis: Frequency axis: Alignment length (from 0 to the maximum mismatches allowed)
Ou	tput file name: <i>prefix</i> .Mismatch.Distribution. <i>tag.</i> cht
Base Mismatch Dis	tribution for Unique Alignments
Th coi	e same report as the Mismatch Distribution, except that only tags that have a unique alignment are nsidered in calculating the distribution.
No	te: The title Unique Alignments is interpreted to mean primary alignments, for this report.
Y a X a	exis: Frequency exis: Alignment length (from 0 to the maximum mismatches allowed)
Ou	tput file name: <i>prefix</i> .Mismatch.Distribution.Unique. <i>tag.</i> cht
Distribution of Alig	nment Length and Number of Mismatches in Tags
A r in v dis mi	report providing a simultaneous picture of the alignment length and number of mismatches distributions various tags used in alignment. Only primary alignments for each tag are considered in calculating this tribution. The bins range from 0 to the maximum read length, and from 0 to the maximum number of smatches allowed.
Thi	is report is located in the ${\tt Misc}$ folder, and is not displayed in the UI.
Z a Y a X a	axis: Frequency axis: Alignment length (from 0 to the maximum mismatches allowed) axis: Number of mismatches (from 0 to the maximum mismatches allowed)
Ou	tput file name: <i>prefix</i> .AlignmentLength.Mismatch. <i>tag</i> .csv
Base QV Distributio	n
A t cor ind	par plot providing a distribution of base quality values generated using the reference assisted error rrection/base conversion algorithm. The base QV distributions are separated for each tag as quality of lividual tags could be very different.
Y a X a	axis: Frequency axis: Base QV (from 0 to the maximum QV)
Ou	tput file name: <i>prefix</i> .BaseQV. <i>tag</i> .cht
Base QVs by Position	on



Report	Description, axis information, filename
	A report providing a distribution of base quality values by individual base positions. This report identifies if certain base positions (particularly towards the end of the read) have poor base quality values. All the tags (F3/R3/F5-P2) used in the alignment are combined into a single bin.
	Y axis: Base QV X axis: Base position (from 0 to read length)
	Output file name: <i>prefix</i> .BaseQV.by.Position.csv
Distribution d	of Mismatches by Base QV
	A 3D surface plot providing a distribution of errors (mismatches to reference) by base quality values bins. This report measures whether the base QVs generated are well calibrated to the probability of error in that particular base position.
	Y axis: Error rate X axis: Base QV (from 0 to maximum QV)
	Output file name: <i>prefix</i> .Mismatches.By.BaseQV. <i>tag</i> .cht
Distribution of	of Mismatches by Position
	A bar plot providing a distribution of errors (mismatches to reference) by position within the read. Only the primary alignments for each bead are used to generate this distribution.
	Y axis: Frequency X axis: Position (from 0 to maximum read length)
	Output file name: <i>prefix</i> .Mismatches.By.Position. <i>tag</i> .cht
Distribution b	y Mapping QVs by Tag
	A bar plot providing a distribution of mapping quality values for individual tags (F3/R3/F5-P2). Only the primary alignment for each bead is used in calculating this distribution.
	Y axis: Frequency X axis: Position (from 0 to maximum mapping QV)
	Output file name: <i>prefix</i> .MappingQV. <i>tag</i> .cht
Distribution L	y Pairing QVs
	A bar plot providing a distribution of pairing quality values for individual tags (F3/R3/F5-P2). Only the primary alignment for each bead is used in calculating this distribution.
	Y axis: Frequency X axis: Pairing QV (from 0 to maximum pairing QV)
	Output file name: <i>prefix</i> .PairingQVs. <i>tag</i> .cht
Coverage Re	port
	A line plot providing a distribution of coverage obtained after mapping/pairing. Only the primary alignment for each bead is used in this calculation.
	Y axis: Number of bases X axis: Coverage (from 0 to number of reads)
	Output file name: <i>prefix</i> .Coverage. <i>tag</i> .cht
Coverage Re	port by Chromosome (Contig) and Base Windows

#### Table 96 Mapping statistics output files (continued)



#### Table 96 Mapping statistics output files (continued)

Report	Description, axis information, filename
	A line plot providing a distribution of coverage within each reference window. The coverage is calculated within each window along a reference chromosome. The window size can theoretically be anywhere from a single base to the contig length. However the calculations of base level coverage are computationally expensive and less interpretable as the window size increases. Only the primary alignment for each bead is used in this calculation.
	Y axis: Coverage (from 0 to number of reads) X axis: Contig and window number
	Output file name: <i>prefix</i> .Coverage.By.Chromosome.contig <i>nn.tag</i> .cht
Coverage by S	trand
	A line plot providing the distribution of coverage within each reference window separated by reference strand (+/-). Only the primary alignment for each bead is used in this calculation.
	Y axis: Number of bases X axis: Coverage (from 0 to number of reads)
	Output file name: <i>prefix</i> .Coverage.By.Strand. <i>tag</i> .cht
Coverage files	
	Coverage reports in wiggle format. For each genome position, reports the number of reads that cover (map to or span) the position. Because reporting coverage for each position results in very large files, coverage is reported for bins, with each bin spanning a user-defined number of bases. For each bin, the mean coverage of all the positions in that bin is reported. The parameter <code>bamstats.wig.binsize</code> controls the size of the bins in this file.
	By default one coverage file is generated, per chromosome, per strand. If the parameter bamstats.combined.report.both.strands is set to true, then one file per chromosome is generated, combining coverage from both strands into one file per chromosome.
	Each coverage file includes a header as the first line. The header lines follow this pattern:
	track type=wiggle_0 name=< <i>chrname&gt;</i> description=< <i>coverage from positive/negative/both strand&gt;</i> visibility=full color=0,0,255 fixedStep chrom=< <i>chrname&gt;</i> start=< <i>startpos&gt;</i> step=< <i>binsize&gt;</i> span=< <i>binsize&gt;</i>
	Output file name: coverage_chrnn.POS.wig, coverage_chrnn.NEG.wig
Insert Range D	Distribution
	A line plot providing the distribution of insert sizes for paired data (PE and LMP library types).
	Y axis: Frequency X axis: Insert Range bins
	Output file name: <i>prefix</i> .Insert.Range.Distribution.cht
Distribution of	Read Pair Types
	For paired data, this report calculates the distribution of read pair types (AAA, AAB, C**, etc.).
	Y axis: Frequency X axis: Read pair type
	Output file name: <i>prefix</i> .ReadPair.Type.cht
Pairing Statist	lics
1	7
---	---

Report	Description, axis information, filename		
	A pie chart showing the percentages of the following F3, R3 (or F5) combinations, for paired data,:		
	Mapped, Mapped		
	Mapped, Unmapped		
	Unmapped, Mapped		
	Unmapped, Unmapped		
	Mapped, Missing		
	Missing, Mapped		
	Unmapped, Missing		
	Missing, Unmapped		
	Output file name: <i>prefix</i> .Pairing.Stats.cht		
Unique Start	Position		
	A text report with the following statistics about the start position on the genome, based only on primary alignments. The report contains:		
	1. The number of starting points in uniquely placed tags:		
	Reports the positions in the reference with at least one uniquely placed alignment starting at that position. Unique here does not mean primary.		
	Also reports the percentage of this number within the total number of positions in the reference.		
	2. The average number of uniquely mapped reads per starting point:		
	Reports the total number of unique alignments divided by the number of starting points in uniquely placed tags.		
	3. An estimated number of starting points for all mapped tags:		
	Reports the total number of primary alignments divided by the number of starting points in uniquely placed tags.		
	Output file name: <i>prefix</i> .Unique.Start.Positions.txt		

#### Table 96 Mapping statistics output files (continued)

# Mapping statistics This section provides example output of some mapping statistics output files. example output Cummer: file

#### Summary file

The summary file provides statistics for each BAM file in the sample. This list describes labels used in the summary file:

- **NumUnFilteredBeads** The total number of beads, before any filtering on the instrument. This value is also the fragment count in the input XSQ file.
- NumFilteredBeads The number of beads that pass filtering on the instrument.
- NumMapped The number of reads with primary alignment.
- % filtered that mapped The number of primary reads divided by the number of beads passing instrument filtering (NumMapped / NumFilteredBeads).
- % total that mapped The number of primary reads divided by the number of total beads (NumMapped / NumUnFilteredBeads).

The number of beads filtered out in the instrument is found by subtracting the number of beads that pass filtering from the total number of beads:

NumUnFilteredBeads - NumFilteredBeads

The following is example contents for a summary file:

```
#title: BAMStats Summary
```

```
BamFileName,IsColorInBam,IsBaseInXSQ,IsECC,LibraryType,ReadLeng
th,PredictedInsertSize,NumFilteredBeads,NumUnFilteredBeads,Tag1
-NumMapped,Tag1- % total Mapped,Tag1- % filtered mapped,Tag2-
NumMapped,Tag2- % total Mapped,Tag2- % filtered mapped,(Tag1-
AlignmentLength;Min;Max;Avg;Median;StdDev),(Tag1-
NumMismatches;Min;Max;Avg;Median;StdDev),(Tag1-
MappingQV;Min;Max;Avg;Median;StdDev),(Tag1-
BaseQV;Min;Max;Avg;Median;StdDev),(Tag2-
AlignmentLength;Min;Max;Avg;Median;StdDev),(Tag2-
NumMismatches;Min;Max;Avg;Median;StdDev),(Tag2-
BaseQV;Min;Max;Avg;Median;StdDev),(Tag2-
MappingQV;Min;Max;Avg;Median;StdDev),(Tag2-
StdDev),Min;Max;Avg;Median;StdDev),(Tag2-
BaseQV;Min;Max;Avg;Median;StdDev),(Tag2-
BaseQV;Min;Max;Avg;Median;StdDev),(Tag2-
BaseQV;Min;Max;Avg;Median;StdDev),(Tag2-
BaseQV;Min;Max;Avg;Median;StdDev),(Tag2-
BaseQV;Min;Max;Avg;Median;StdDev),(Coverage;Min;Max;Avg;Median;
StdDev)
```

#### **Unique Start Position**

The following is an example of the output of a Unique Start Position report: # Starting Points within Placed Tags Number of Starting Points in Uniquely placed tag 109,068,047 (2.005740% of reference) Average Number of Uniquely Mapped reads per Start Point 1.992770 Estimate number of starting point for all mapped tags 152,843,627 (2.810765% of reference)

#### Coverage files

Example contents for the file coverage\_chr1\_positive.wig are:

```
browser position chr1:1-20000000
browser hide all
browser pack refGene encodeRegions
# minimumMapq=25, minimumCoverage=1,
alignmentFilteringMode=PRIMARY, filterOrpha-nedMates=false,
track name="BAM Coverage positive strand" description="BAM
Coverage positive strand" visibility="full color 0,0,255"
priority=10 yLineMark=0 type=wiggle_0 yLineOnOff=on
variableStep chrom=chr1 span=1
336171
336181
336191
336201
336211
336221
336231
```



Example contents for the file coverage\_chr1\_negative.wig are:

	browser position chr1:1-200000000
	browser hide all
	browser pack refGene encodeRegions
	<pre># minimumMapq=25, minimumCoverage=1, alignmentFilteringMode=PRIMARY, filterOrpha-nedMates=false,</pre>
	<pre>track name="BAM Coverage negative strand" description="BAM Coverage negative strand" visibility="full color 0,0,255" priority=10 yLineMark=0 type=wiggle_0 yLineOnOff=on</pre>
	variableStep chrom=chr1 span=1
	57432
	57442
	57452
	57463
	57473
	57483
	57493
	57503
	57513
	57523
	57533
S	section describes requirements to generate mapping statistics outside of the

Run BAMStats standalone

This section describes requirements to generate mapping statistics outside of the context of the mapping module. These requirements are:

• **Directory structure** – The input directory for BAMStats module must mimic the output directory structure of mapping:

```
Input.BAMStat.dir/
    <sampleName1>/*.bam, *.bai
    <sampleName2>/*.bam, *.bai
```

• **Index files** – For every BAM file, there must be a corresponding BAI-formatted indexed file. BAI files can be generated using the following command:

samtools index <bamfile>

This command generates index sorted alignment for fast random access. The index file <bamfile>.bai is created. For details, please refer to the samtools site:

http://samtools.sourceforge.net/samtools.shtml



• Write permission – The user must also have write permission for the input directory, because the BAMStats module generates position and probe error files in that directory.

# FAQ – Whole transcriptome

#### Why is SNPs analysis not included in the whole transcriptome pipeline?

A SNPs analysis is not offered for WTA pipelines. The reverse transcription process that is part of the WT library prep intrinsically adds a percentage of sequencing errors. In addition, alternative splicing and other alignment complexities around exon junctions may cause local misalignments, which can lead to false positive SNP calls. Using that data in a SNPs analysis may cause a high number of false positives.

## 2

1

#### What reads scenario achieves the best accuracy?

The best performance is achieved by considering 2BE and 4BE reads, their alignment, and reference. Typically, base-space reads generated by ECC have lower mapping throughput. Base-space reads generated from combined 2BE and 4BE calls and no reference have lower accuracy, compared to ECC calls.

## 3

# In the mapping statistics genome coverage calculation report, how are Ns in the reference counted?

Ns in the reference are counted as missing coverage. For example, in the human genome, about 7% of the reference sequence consists of Ns. This means that the coverage calculation will never be reported as higher than 93%. If the frequency distribution says 7.49% of the genome is uncovered, this means that about 0.34% of the non-N reference sequence does not have reads mapped to it, and that also the 7.15% of the genome that consists of N does not have reads mapped to it.

## 4

What should I do to ensure balanced allele ratios at heterozygous positions?



The base-translation algorithm, refcor, uses instrument color-space data (ECC or non-ECC) and the reference sequence to produce the most likely base calls, along with a Phred-scale quality value. The degree to which the reference influences the base call is controlled by the refcor.reference.weight parameter. A position is converted to reference only if the refcor.reference.weight setting is higher than the difference between the quality values of a correct color call and of an adjacent erroneous color call that supports reference.

The refcor.reference.weight default setting, 8, is chosen to be low enough to ensure balanced allele ratios, and also be large enough both to distinguish correct from incorrect color calls and to maximize base accuracy.

In the event an incorrect reference base call is made, the disagreement between evidence sources causes that call's QV to be low. Incorrect calls can be filtered out based on QV threshold, and this filtering improves allele ratios.

Another way of filtering low quality base-calls is to adjust the parameter refcor.base.filter.qv, that converts all base calls with a quality value less than refcor.base.filter.qv into Ns. This filtering has the effect of setting QV to zero in the calls with low confidence. To decrease the number of N base calls, reduce this threshold (for example, from the default of 10 to 5).

**Note:** Setting the refcor.reference.weight parameter to zero completely eliminates the reference as a guide for base-translation (completely eliminates reference bias), but also considerably reduces the quality of base-translation.

#### How is the uniqueness of a pair determined?

A pair of reads is unique when there is exactly one good AAA pair. With the advent of local alignment, the likelihood of finding only one good pair is decreased. As a result, a different heuristic is used to determine "uniqueness".

Consider an alignment of length *L* with *M* mismatches. If the local score is defined as L+(m-1)M, where m<0 is the mismatch penalty, then the score of each good pair candidate is the sum of its two constituent local scores.

A pair of tags is considered unique if it has only one good pair, or if the score of the best pair is at least X greater than the score of the second best pair. X is specified by the pair.uniqueness.threshold key, and has a default value of 10.0. See Table 91 on page 347 and Table 90 on page 346 for a description of this and all pairing parameters.

#### What do the 3-letter pair classifications mean?

Table 97 lists a summary of genomic code classifications for mate-pair pairing runs. See Table 98 on page 367 for a summary of genomic code classifications for paired-end runs.

Class name	Strand	Orientation	Insert size	Preference
AAA	Same	R3 to F3	Normal	1
AAB	Same	R3 to F3	< min	2
AAC	Same	R3 to F3	> max	2
ABA	Same	F3 to R3	Normal	2
ABB	Same	F3 to R3	< min	3
ABC	Same	F3 to R3	> max	3
BAA	Different	Outward	Normal	3
BAB	Different	Outward	< min	4
BAC	Different	Outward	> max	4
BBA	Different	Inward	Normal	3
BBB	Different	Inward	< min	4
BBC	Different	Inward	> max	4

 Table 97
 Genomic code classifications, for mate-pair pairing

The first letter determines whether the two tags match the same strand:

- A Match the expected strand.
- **B** Match different strands.

The third letter determines if the distance between two hit positions is within the correct range:

- **A** Within the correct range.
- B The distance is less than the insert size measurement minimum (denoted by "< min" in the table above).</li>
- C The distance is greater than the insert size measurement maximum (denoted by "> max" in the table above).

The default minimum insert size is 0. For paired-end and long mate pair workflows, the default maximum insert sizes are 2000 and 20,000, respectively. An insert size measurement automatically determines the actual values using the defaults as the largest interval, and the defaults are used if the insert size measurement fails.

The middle letter has different meanings, depending on whether the first letter is A or B. If the first letter is A, the second letter indicates if the order of the two tags is correct:

- A The order is correct, and R3 is upstream of F3.
- **B** The order is incorrect, and F3 is upstream of R3.

When the two tags are on different strands, the second letter indicates whether the two tags are pointing toward or away from each other

- A Pointing away from each other.
- **B** Pointing toward each other.

In addition, C\*\* classification is assigned to a pair of reads that have one hit each, but are on different chromosomes. If both F3 and R3 tags are present, but only one has a unique hit while the other one is unmapped, the pair is labeled D\*\*. Finally, if one tag from a pair has a unique hit but the other one is missing, as opposed to unmapped, then the pair is classified as E\*\*.

17

The Preference value indicates likelihood of the variation can occur in nature. Larger values indicate the variation is less likely to occur in nature.

Table 98 shows the classifications for paired-end runs. Summary statistics of these categories are calculated in the optional BAMStats Read pair Type Statistics Report (.ReadPair.Type.cht).

Class name	Strand	Orientation	Insert size	Preference
AAA	Different	Inward	Normal	1
AAB	Different	Inward	< min	2
AAC	Different	Inward	> max	2
ABA	Different	Outward	Normal	2
ABB	Different	Outward	< min	3
ABC	Different	Outward	> max	3
BAA	Same	F3 to F5	Normal	3
BAB	Same	F3 to F5	< min	4
BAC	Same	F3 to F5	> max	4
BBA	Same	F5 to F3	Normal	3
BBB	Same	F5 to F3	< min	4
BBC	Same	F5 to F3	> max	4

 Table 98 Genomic code classifications, for paired-end pairing





# Run a WT Coverage Analysis

This chapter covers:

Overview	369
WT coverage input files	369
WT coverage parameters	370
Coverage output files	371

## **Overview**

The whole transcriptome coverage module calculates read coverage per position.

# WT coverage input files

The WT coverage module takes as input one or more BAM files containing mapped data. The following describe the alignment input accepted. This module:

- Accepts only fragment data for RNA libraries.
- Accepts multiple BAM files as input.
- Accepts BAM files with different read lengths.
- Accepts both color space and base-space files, and a combination of color and base-space data.

#### BAM file metadata

• Table 99 lists BAM file metadata used by the WT coverage module. Table 99 BAM file meta data required by the WT coverage module

Field name	Field in BAM header	Requirement
Application Type	The AT field in the @CO line for every @RG line. (Every @RG read group line has one @CO comment line.)	@RG lines must be present. Every @CO line must have an AT field, and the AT field must be set to the string "Small RNA".
Sequence Name	The SN field in each @SQ header line.	Only contigs with an @SQ line with an SN field are analyzed. If a contig does not have an @SQ line in the header, the alignments in that contig are not counted.
Sort Order	The SO field in the HD header line.	The SO field must be present and must be set to the string "Coordinate".

# WT coverage parameters

Table 100 lists the WT coverage parameters.

Table 100 WT coverage parameter description

Parameter name	Default value	Description	
coverage.run	1	Whether or not to run the WT coverage module. Allowed values:	
		• <b>0</b> : Do not run the coverage module.	
		• 1: Run the coverage module during this analysis.	
		The run parameter is set automatically by the shell during a standard workflow. Accept the default for most use.	
RNA.bam.file	—	A comma-separated list of paths to the input BAM files. Relative and absolute paths are accepted.	
		Generated automatically by LifeScope <sup>™</sup> Software during a workflow. Use this parameter only when running this module standalone.	
Optional parameters	Optional parameters		
RNA.coverage.min.quality	2	( <i>Optional</i> ) The mapping quality value threshold for selecting the alignments. Alignments with a mapping quality value lower than this threshold are not written to output. Allowed values: Integers 0–100.	
RNA.coverage.min.value	0	( <i>Optional</i> ) The mapping quality threshold value for selecting alignments from the input BAM. In order to be included in the output, a position's coverage must be greater than or equal to this value. Allowed values: Integers >= 0.	



 Table 100
 WT coverage parameter description (continued)

Parameter name	Default value	Description
RNA.coverage.per.chromosome	true	( <i>Optional</i> ) Whether coverage output is generated as one file per every chromosome per strand, or as a single file with coverage of all chromosomes per strand. Allowed values:
		• <b>false</b> : Generate a single file with coverage of all chromosomes per strand.
		• <b>true</b> : Generate one file per every chromosome per strand.
RNA.coverage.primary.only	true	( <i>Optional</i> ) Whether only primary alignments or all the alignments from the input BAM file are used as input. Allowed values:
		• false: Consider all alignments.
		<ul> <li>true: Consider only primary alignments (both gapped and ungapped).</li> </ul>
Resource parameters		
memory.request	4gb	Memory request. Include the units gb in the setting.
java.heap.space	4000	Dynamic memory requirement.

## **Coverage output files**

The WT coverage module's output files are in the wiggle (\*.wig) file format. Wiggle files are visualized in genomics browsers such as the Integrative Genomics Viewer (IGV). You can download the IGV browser from the Broad Institute website:

#### www.broadinstitute.org/igv

The wiggle file specification is available from this site:

#### http://genome.ucsc.edu/goldenPath/help/wiggle.html

If the parameter RNA.coverage.per.chromosome is set to false, the WT coverage module generates two output files with coverage of all chromosomes per strand:

- coverage\_positive.wig
- coverage\_negative.wig

If the parameter RNA.coverage.per.chromosome is set to true, the WT coverage module generates one coverage file per chromosome per strand:

- coverage\_chr\*\*\_positive.wig
- coverage\_chr\*\*\_negative.wig



**Chapter 18** Run a WT Coverage Analysis *Coverage output files* 



# Run a WT Count Known Genes and Exons Analysis

This chapter contains:

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## **Overview**

The module performs counting of features provided in a GTF file. It counts:

- Exons
- Genes

The module allows filters, generates RPKM, and annotates how many exons are expressed in a gene. This tool annotates only the Exon features in a GTF file

**IMPORTANT!** Alignments must come from the same strand as the feature to contribute to the count. A non-gapped tag contributes to a feature's count if it overlaps the feature and has no more than three bases outside the feature. A gapped tag contributes to a features count if one of its match regions terminates at a feature boundary.

# Input files

Use the WT counts module to count the number of reads that align within genomic features. The WT counts module takes the following as input:

- A set of read alignments (BAM file)
- A set of user-defined (or default) stringency parameters
- A set of genome annotations (GTF file)

Paired-end and fragment libraries may be combined in the BAM input. Unmapped reads in the BAM input are not used.

# WT counts parameter description

Table 101 describes parameters that you can configure for the WT counts module.

Table 101 Counts module algorithm parameters

Parameter	Default	Description
Algorithm parameters	1	
wt.gtf.file	\${annotation.gtf.file}	The GTf file used to create the genemodel.
wt.count.features.gene.count	1	Generate whole transcriptome gene count. Allowed values:
		• <b>0</b> : Do not generate whole transcriptome gene count.
		• 1: Generate whole transcriptome gene count.
wt.count.features.primary	1	Whole transcriptome counts primary. Allowed values:
		• <b>0</b> : Count all alignments.
		• 1: Count only primary alignments. Recommended.
wt.count.features.min.mapq	10	The minimum alignment quality for an alignment to be counted. Allowed values: Integers 0–255.
wt.count.features.overflow.limit	3	Whole transcriptome overflow limit. An alignment may align outside the boundaries of a given exon feature (called overflowing). If an alignment overflows the feature more than the value specified in this parameter, then that read is not counted towards that feature.
		Allowed values: Integers 0–100.
Optional resource parameters	1	·
java.heap.space	8000	Dynamic memory requirement.
memory.request	8gb	Memory request. Include the units gb in the setting.

#### Table 102 lists WT counts parameters that we do not recommend changing.

Table 102WT Counts internal parameters

Parameter name	Description
wt.exon.reference	The exon FASTA reference file. Default is \${analysis.output.dir}/ wtexonsequenceextractor/exons.fasta.

Table 103 lists parameters that you do not specify when you run the module in a standard workflow.

Table 103 WI count parameters supplied by LifeScope <sup>TH</sup> Softwa
--

Parameter name	Description
wt.counts.run	Enable the WT counts module. Allowed values:
	• <b>0</b> : Do not run the module.
	• 1: Run the WT counts module. Default.
wt.exon.reference	The exon FASTA reference file. Default is \${analysis.output.dir}/wtexonsequenceextractor/exons.fasta.
	This file is defined by the wt.f5.exseqext.output.reference parameter in the WT exon sequence extractor.
wt.gtf.file	The GTf file used to create the genemodel. Default: \${annotation.gtf.file}.

# WT counts algorithm description

Filters

Stringency parameters govern the set of alignments in a BAM file. These alignments are considered in counting. Reads that do not meet these stringency parameters are filtered out and do not contribute to the result.

These parameters govern stringency:

- Filter alignment mode (wt.count.features.primary)
- Minimum mapping quality (wt.count.features.min.mapq)

The filter alignment mode governs how multi-mapping reads are handled. The option settings for filter alignment mode are all and primary (see Table 104). Both options are supported for single-read and for paired-end data.

 Table 104
 Alignment filter modes

Mode	Description
all	All alignments are considered.
primary	Only primary alignments are considered.

If a minimum mapping quality is selected, only alignments with the minimum quality or greater are considered. Likewise, if a minimum score is selected, only alignments having at least the minimum score are considered.

Filtering by unique or top criteria is included mainly for legacy reasons. We recommend that you use only default setting of mapping quality to filter reads.

Half (0.5) a count is assigned for each exon of a read that spans an exon-exon junction.

Mapping quality incorporates an assessment of score and uniqueness. To filter using only mapping quality specify a minimum mapping quality, and set the alignment filter mode to all, and specify a minimum mapping quality.

In previous releases of LifeScope<sup>™</sup> Software, filtering single-read alignments using the unique alignment filter mode was recommended. When unique filtering is enabled, only alignments that satisfy the following criteria pass the filter:

- The total number of alignments for the read is less than the maximum allowed to be reported in mapping (the Z parameter).
- The alignment must be the single best alignment of the set of alignments for the read.
- The score of the alignment must be sufficiently better than that of any suboptimal hits. If no suboptimal hits are present, a suboptimal hit is assumed to exist with a score one less than the minimum possible score for the anchor size and mismatch level used in mapping. The difference between the score of the alignment score and the best suboptimal alignment must be greater than or equal to the clear zone.

Each feature defined in the genome annotation is assigned a tag count. A tag count is the total number of alignments that pass the stringency filters and are consistent with the annotation. The criteria for counting an alignment toward a feature are:

- The alignment must overlap the feature's numeric coordinates (contig, start, end).
- The alignment strand orientation must be consistent with the feature strand orientation.

**Note:** F3 reads align to the sense genomic strand; F5 reads align to the antisense genomic strand.

- If the alignment does not span an intron, the alignment must include no more than three bases outside the feature.
- If the alignment spans an intron, the position of the exon-intron boundary must match in the alignment and the feature.

The read count for a given gene is defined as number of reads that map to the maximal region (features) of the merged gene. Maximal features are calculated from the transcripts provided in the GTF file by using the gene model. Reads that map to overlapping exons or features of a given gene are not counted multiple times.

The following figure describes the component flow during exon counting.



**RPKM** The RPKM of exons (the Reads Per Kilobase of exon model per Million mapped reads), which is a normalized measure of expression, is also reported. This metric is calculated with the formula:

 $RPKM = 10^9 \times \frac{ExonReadCount}{TotalReadCount \times ExonLength}$ 

Counts

The number of total uniquely mapped reads on exons is used as the normalization factor. The RPKM value is also reported by the WT splice finder module (see "Splice finder parameters" on page 380).

# **Output files**

This section describes the output of the WT counts module.

**GTF output** The output format of the WT counts module follows the same GTF file format as the supplied genome annotation. The score field of the GTF file contains the tag count. RPKM is reported in the attributes field.

The following is example GTF output.

```
# filterOrphanedMates=false, minMapq=10, minScore=0, scoreClearZone=5,
alignmentFilteringMode=PRIMARY
#number_of_mapped_reads=190375154
                                        . gene_id "WASH5P"; transcript_id
chr1 BioScope exon 4225 4692
                                  86 -
"NR_024540"; RPKM "0.97";
                                            gene id "WASH5P"; transcript id
chr1 BioScope exon 4833
                            4901
                                  5
                                      _
                                         .
"NR_024540"; RPKM "0.38";
chr1 BioScope exon 5659
                           5810 18 - .
                                            gene_id "WASH5P"; transcript_id
"NR_024540"; RPKM "0.62";
```

Tab output

The following fields are in the gene\_counts.tab output file:

- Gene name
- Chr
- Start
- End
- Strand
- ReadCount
- Length (the maximal gene region length)
- RPKM
- Number of features (exons)
- Number of expressed features (>10 reads)
- Transcript names

Every gene is reported whether it is expressed or not.



1

# FAQ – WT counts

Does the whole transcriptome module count invalid reads?

A: No. If a read is invalid (that is, considered invalid by Picard software), that read is not included in the count of mapped reads.



# Run a WT Splice Finder Analysis

This chapter contains:

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## **Overview**

Transcripts of eukaryotic genomes are transcribed from exons which are usually separated by one or more introns. The spliceosome machinery removes the introns from pre-messenger-RNAs to generate mature messenger-RNAs. The removed introns are called splice junctions. Alignments from an RNA-Seq experiment may map on these splice junctions, meaning the alignment has two parts that are separated by thousands of bases on the genome. The splice finder software is designed to discover these splice junctions efficiently and annotate the type and evidence for each discovered junction.

A fusion junction is a section of transcribed RNA that maps to an exon from one gene followed by an exon from another gene. It can occur as the result of a translocation, deletion, or chromosomal inversion. A splice junction excludes exon-to-exon boundaries that arise from alternative splicing for a gene.

The module takes as input:

- A BAM file of either paired-end or fragment libraries reads sequenced from transcribed RNA
- A list of exons. Each exon includes its color sequence and the reverse of that sequence.
- A GTF file
- A FASTA file of the reference genome

Unmapped reads in the BAM input are not used. Partially mapped reads are used to find gene fusions.



# Splice finder parameters

#### Table 105 lists the parameters for the splice finder module.

Table 105 Splice finder parameter description

Parameter name	Default	Description
	value	Description
Required parameters	1	
wt.splice.finder.first.read.max.read.length	50	F3 read length. This value is auto-populated if possible:
		Allowed values: Integers 25–150.
wt.splice.finder.second.read.max.read.length	25	F5 read length. This value is auto-populated if possible:
		Allowed values: Integers 25–150.
Optional parameters		
wt.splice.finder.min.exon.length	25	All exons shorter than this are removed:
		Allowed values: Integers 0–500.
Optional single read junction evidence collection		
wt.splice.finder.single.read	1	Enable single-read fusion finding. Allowed values:
		• <b>0</b> : Do not run single-read junction finding.
		• 1: Enable single-read junction finding.
wt.splice.finder.single.read.min.overlap	10	The minimum number of contiguous color positions that must align in a read-exon map.
		Allowed values: Integers 1–100.
wt.splice.finder.single.read.max.mismatches	2	The maximum number of mismatches allowed in a read- exon map.
		Allowed values: Integers 0–25.
wt.splice.finder.single.read.clip.size	2	Progressive unit size for clipping at the end of read.
		Allowed values: Integers 0–25.
wt.splice.finder.single.read.clip.total	10	Total size for clipping at the end of read.
		Allowed values: Integers 0–25.
wt.splice.finder.single.read.ReportMultihit	0	How to count single reads that align to multiple splice junctions. Allowed values:
		• <b>0</b> : Do not count them as evidence to any of the junctions.
		• 1: Count them as evidence to all of the junctions.
		• <b>2</b> : Count them as evidence to the first of the junctions.
		Option 0 provides uniqueness.
wt.splice.finder.single.read.remap	0	Single read remap. 0=false, 1=true. Allowed values:
		• <b>0</b> : SASR registers the evidence, but does not remap.
		<ul> <li>1: SASR remaps reads that have already been mapped to a junction, and registers evidence for that remap.</li> </ul>
wt.splice.finder.single.read.clip.5.prime	1	Single read clip 5 prime. Allowed values:
		• <b>0</b> : SASR clips only the 3' end.
		• 1: SASR clips both the 5' and the 3' end of the read.



 Table 105
 Splice finder parameter description (continued)

Parameter name	Default value	Description
wt.splice.finder.single.read.min.read.length	37	SASR considers shorter reads (in number of colors) to be inadmissible as evidence.
		Allowed values: Integers 0–100.
Optional paired read junction evidence collection		
wt.splice.finder.paired.read	1	Enable paired-end fusion finding. Allowed values:
		• <b>0</b> : Do not run paired-end junction finding.
		• 1: Enable paired-end junction finding.
wt.splice.finder.paired.read.min.mapq	10	A read-pair that has mapping quality lower than this is inadmissible as evidence.
		Allowed values: Integers 0–225.
wt.splice.finder.paired.read.avg.insert.size	120	Average insert size of the paired-end library.
		Allowed values: Integers 0–1000.
wt.splice.finder.paired.read.std.insert.size	60	Standard deviation of the insert size for the paired-end library.
		Allowed values: Integers 0–1000.

#### Combined caller for junction

The parameters below allow splice finder to combine evidence from single-reads and from paired-reads to call junctions, alternative splices, and fusions, respectively. In each case, an event (for example, a fusion) is reported if SRE>x and PRE>y and ((SRE+PRE)>z or MAX(SRE,PRE) > w), where

SRE = unique single-read evidence,

PRE = unique paired-read evidence, and

x, y, z, and w are specified by the parameters below.

wt.splice.finder.junction.split.read.min.evidence	1	x in SRE>x
		Minimum single read evidence for junction.
		Allowed values: Integers 0–50.
wt.splice.finder.junction.paired.end.min.evidence	1	y in PRE>y
		Minimum paired-end read evidence for junction.
		Allowed values: Integers 0–50.
wt.splice.finder.junction.combined.min.evidence	2	z in (SRE+PRE)>z
		Minimum combined evidence for junction.
		Allowed values: Integers 0–50.
Combined caller for (same gene) alternative splicing		
wt.splice.finder.alt.splicing.split.read.min.evidence	1	X
		Minimum single read evidence for alternative splicing.
		Allowed values: Integers 0–50.
wt.splice.finder.alt.splicing.paired.end.min.evidence	1	У
		Minimum paired-end evidence for alternative splicing.
		Allowed values: Integers 0–50.



#### Table 105 Splice finder parameter description (continued)

Parameter name	Default value	Description	
wt.splice.finder.alt.splicing.combined.min.evidence	2	Z	
		Minimum combined evidence for alternative splicing.	
		Allowed values: Integers 0–50.	
Combined caller for gene fusion			
wt.splice.finder.fusion.split.read.min.evidence	2	x	
		Minimum split read evidence for junction.	
		Allowed values: Integers 0–50.	
wt.splice.finder.fusion.paired.end.min.evidence	2	У	
		Minimum paired-end evidence for junction.	
		Allowed values: Integers 0–50.	
wt.splice.finder.fusion.combined.evidence	4	Z	
		Minimum combined evidence for junction.	
		Allowed values: Integers 0–50.	
Other options	1		
wt.splice.finder.output.format	3	Output format parameter. Allowed values:	
		1: Tabular output format	
		• 2: BED output format	
		• 3: BED, tabular, and SEQ output formats	
		• 4: All formats, including Circos.	
		With option 3, both tabular, BED, and also additional SEQ files are created separately. SEQ files contain 50 base pairs of sequences from each end of a junction exon and are useful for validation.	

#### Table 107 lists parameters that we do not recommend changing.

 Table 106
 Splice finder internal parameters

Parameter name	Default value	Description
wt.splice.finder.base.space	0	Call junctions in base space. Allowed values:
		• <b>0</b> : Call junctions in color space.
		• 1: Call junctions in base space.
Optional resource parameters		
java.heap.space	14000	Dynamic memory requirement.
memory.request	15gb	Memory request. Include the units gb in the setting.



Table 107 lists parameters that you do not specify when you run the module in a standard workflow.

Table 107	Splice	finder	parameters	supplied	by	LifeScope™	Software
-----------	--------	--------	------------	----------	----	------------	----------

Parameter name	Description
splice.finder.run	Enable the splice finder module. Allowed values:
	• <b>0</b> : Do not run the splice finder module.
	• 1: Run the splice finder module. Default.
wt.exon.reference	The exon FASTA reference file. Default is \${analysis.output.dir}/wtexonsequenceextractor/exons.fasta.
	This file is defined by the wt.f5.exseqext.output.reference parameter in the WT exon sequence extractor.
wt.gtf.file	The GTf file used to create the genemodel. Default: \${annotation.gtf.file}.

# **BAM** file metadata

Table 108 lists BAM file metadata used by the splice finder module. This information is taken from the @CO lines that apply to the first read group (@RG) line. See "XSQ metadata in BAM headers" on page 462 for information on LifeScope<sup>™</sup> Software's use of @RG and @CO fields in BAM headers.

 Table 108
 BAM file meta data

രCO field	Behavior
LT	If LT is fragment, the following parameters are set: wt.splice.finder.single.read is set to 1, and wt.splice.finder.paired.read is set to 0.
IA	Parsed for the average insert size. Overrides the setting of the wt.splice.finder.paired.read.avg.insert.size parameter.
IS	Parsed for the standard deviation of the insert size. Overrides the setting of the wt.splice.finder.paired.read.std.insert.size parameter.
BX	Determines if base space is present for the first tag.
BY	Determines if base space is present for the second tag.
TX	Gives the number of color calls in first tag (max read length). Overrides the wt.splice.finder.first.read.max.read.length parameter setting.
UX	Gives the number of color calls in second tag (max read length). Overrides the wt.splice.finder.second.read.max.read.length parameter setting.
ECC run	If both tags are in base space, sets wt.splice.finder.base.space to 1.
	If this field is set to 0 in the BAM header, then the value of wt.splice.finder.base.space is used.



# SASR splice finder

SASR splice finder	1. Read in and process the exon list.							
module summary	<b>2.</b> For each admissible read:							
	<b>a</b> . Let <i>L</i> be the set of exons that map to the prefix of the read.							
	<b>b.</b> Let $R$ be the set of exons that map to the suffix of the read.							
	<b>c.</b> For each exon $e$ in $L$ , and each exon $f$ in $R$ ,							
	if $overlap(e, r) + overlap(f, r) + 1 = length(r)$ , then							
	register read $r$ as evidence for junction <i>e-f</i> .							
	<b>3.</b> Output the list of junctions and evidence.							
SASR splice finder module evidence evaluator	After the single-read and paired-end splice finders fully process the BAM file, the evidence evaluator evaluates the existing evidence and makes junction calls based user-defined thresholds.							
	Pairs mapped less than a confidence threshold (set with the parameter wt.splice.finder.paired.read.min.mapq) are not used. By default, reads with an original length of less than 37 are not considered for splicing evidence. The parameter wt.splice.finder.single.read.min.read.length changes the default. Spliced reads are used as evidence unless remap is enabled (with the parameter wt.splice.finder.single.read.remap). When remap is enabled, the spliced read is remapped and then used as evidence if it falls on a junction.							
	The default thresholds require at least one single-read and one paired-end evidence to call a junction. The calling of special junctions, such as alternative splicing and fusion, is described in the following three paragraphs.							
	The evidence evaluator also counts the number of unique pieces of evidence. In the case of paired-end evidence, this count is the number of unique mate-alignment start positions, and in the case of single-read evidence, it is the number of unique overlap lengths for the "left" exon.							
	The candidate's data structure stores the evidence for junctions as properties of its exons. Each exon points to other exons for which there is evidence of a junction, where the current exon is a source, and the pointed-to exon is the destination. The data structure of the candidate corresponds to a directed exon evidence graph (shown in Figure 39 on page 385.							
	<ul> <li>In a junction evidence graph, nodes are exons, and each directed edge corresponds to junction evidence from a source exon to a destination exon. Each edge has two values:</li> <li>Unique SASR evidence</li> <li>Unique PR evidence</li> </ul>							





Figure 39 A directed exon evidence graph

Referring to Figure 39, the black arrows correspond to normal junctions, blue arrows correspond to alternative splice junctions, and red arrows correspond to fusions.

Most junctions found are regular junctions, meaning junctions between exons of the same gene in the RefSeq expected order. However, more interesting special junctions are also detected during this evaluation step. Alternatively, spliced junctions are defined as multiple junctions from a given exon to two other exons of the same gene within the given sample. Fusion junctions are defined as junctions between exons of different genes.

LifeScope<sup>™</sup> Software includes user-defined parameters that allow the software to call Call junctions and junctions and fusions with fragment (F3 only) datasets. However, LifeScope™ fusions with single Software's fusion caller is designed to work with paired-end reads. Calling fusions with only fragment reads likely results in a large number of false positives. We observed that on the same dataset, LifeScope<sup>™</sup> Software calls ten times or more fusions using only single-read evidence compared with calls using both paired-end and single-read evidence. We suggest rigorous post-filtering and sorting by total unique single-read evidence for validating those fusion calls.

> For detecting known same-gene junctions we observed that using only single-reads calls 80% of the junctions called by paired-end reads (see Figure 40). However, the remaining 20% usually contains lowly-expressed or harder to detect junctions, and we observed a need to double the total number of reads in order to achieve the same sensitivity. As a result, detection of exon junctions with fragment reads is not recommended due to lower specificity and sensitivity.

read only

Figure 40 compares single-read (SR) vs. paired-end (SR+PE) junction calling sensitivity. The left two bars show SR and SR+PE number of junction calls when using UHR barcode 1. The middle two bars are with UHR barcode 2. The right two bars show the increase in number of calls when the two barcodes are merged and calls are repeated with effectively double the amount of reads from the same library. For both SR and SR+PE, two unique evidences are required to call a junction. For SR+PE, at least one SR and at least one PE evidence is required as well.



Figure 40 Comparison of single-read (SR) vs. paired-end (SR+PE) junction calling sensitivity with two barcodes

# **Output files**

The following types of output are generated, depending on the parameter wt.splice.finder.output.format:

- Tabular (a tab-delimited junction database) files with the following contents:
  - All detected junctions
  - Junctions interpreted as arising from alternate gene splicing
  - Junctions interpreted as arising from inter-gene fusion
- BED
- SEQ
- Circos
- Statistics files, which contain the following parts:
  - Paired-end splice finder statistics
  - SASR splice finder statistics
  - Evidence matrix

These files are described in the remainder of this section.



# Tabular junctionThree output files are generated with the .tab extension for regular junctions,<br/>alternatively-spliced junctions, and fusion junctions. Table 109 describes the fields for<br/>these files.

Table 109 J	lunction	tab-delimited	output	format
-------------	----------	---------------	--------	--------

Field name	Description
Exon-1	The gene id followed by the exon order on the gene.
Exon-1-reference	The reference FASTA file.
Exon-1-strand	± strand.
Exon-1-start	The start position.
Exon-1-end	The end position.
Exon-2	The gene id followed by the exon order on the gene.
Exon-2-reference	The reference FASTA file.
Exon-2-strand	± strand.
Exon-2-start	The start position.
Exon-2-end	The end position.
Exon-1-size	The size of the first exon.
Exon-2-size	The size of the second exon.
Exon-1-readcount	The number of reads that map on the first exon.
Exon-2-readcount	The number of reads that map on the second exon.
Exon-1-F3-RPKM	The Reads Per kbp Per Million Reads (RPKM) exon-1 from F3.
Exon-2-F3-RPKM	The RPKM reads (RPKM) exon-2 from F3.
Exon-distance	Distance between two exons. The exon-distance is not applicable if it is on a different chromosome.
Total-PR-evidence	The total paired-end evidence for the junction.
Unique-PR-evidence	The unique paired-end evidence for the junction.
Total-SR-evidence	The total single-read evidence for the junction.
Unique-SR-evidence	The unique single-read evidence for this junction.
JCV	A junction confidence value.
Known	"K" if a junction is known, and "P" if a junction is putative.
E1-all-genes	The list of all genes to which exon-1 was mapped.
E2-all-genes	The list of all genes to which exon-1 was mapped.
Other	Other information provided about the junction.
	For alternatively spliced junctions, this field reports the length of the alternative exon.

The columns that are most relevant to the splice finder are Unique-PR-evidence and Unique-SR-evidence. Also of interest are the two metrics RPKM and JCV.

The purpose of the RPKM (Reads Per Kilobase of exon sequence, per Million reads) metric is to provide a normalized exon expression level. This metric is calculated with the formula:



 $RPKM = 10^9 \times \frac{ExonReadCount}{TotalReadCount \times ExonLength}$ The RPKM value is also reported by the WT counts module (see "WT counts parameter description" on page 374).

The purpose of the Junction Confidence Value (JCV) metric is to aid in the detection of false positives and in other decisions that depend on a confidence level. Depending on the coverage of the sequencing experiment and exons being tested, the algorithms might generate results that require different user-defined thresholds to detect the most likely fusions. The major contributor of false positives possibly is highly-expressed genes, which have a considerable random chance of encountering a fusion event due to sequencing errors and mapping to homologous exons. A statistical confidence metric (see below) was developed to detect such false positives and assign a confidence level to the evidenced junction.

$$JCV_{j_{x-y}} = \sum_{i=1}^{n} PQV_{i} - 10\log_{10} (EEM_{j_{x-y}})$$

**Equation 1. Junction Confidence Value (JCV)** 

$$EEM_{j_{x-y}} = \frac{RC_x}{\frac{l_x}{\mu_T + 3 \times \sigma_T}} \times \frac{RC_y}{\frac{l_y}{\mu_T + 3 \times \sigma_T}}$$

**Equation 2. Error expectation metric (EEM)** 

where  $PQV_i$  is the Phred-scale pairing quality value for the *i*'th unique paired read evidence for a candidate junction  $j_{x-y}$  and *x* and *y* are the junction exons. For each unique single read evidence, the  $PQV_i$  is set to 10. If there are multiple alignments *for* a given unique start point, take the PQV of the first such alignment. *RC* is the absolute proper mapped read count for the corresponding exon and is the length of the exon;  $\mu_T$ and  $\sigma_T$  are the mean and standard deviation of the insert size for the current experiment, *T*.

Error expectation metric (EEM) is used to quantify highly expressed junctions. This metric is hard to calculate due to genome complexity and homology of exons. Our estimation considers the number of reads mapped to the exons, the length of exons, and a conservative insert range.

After the equation is calculated, a JCV that is larger than 100 is set to 100, and those smaller then 0 are set to 0. If a JCV approaches 100, it is more likely to be a real junction.

**Junction examples** For a junction detected between exon-1 of size 5,000 and exon-2 of size 400, with mean insert 100 and standard deviation 33.3 bp, assume in case-1, there were three unique junction evidences with PQV 30, 20 and 40 respectively and in case-2 only a single unique evidence with PQV 20. There were 900 properly mapped alignments to exon-1 and 100 such alignments for exon-2. In case-3 has 3 unique evidences similar to case-1, but the exons are assumed to have 100 times more coverage each. The junction confidence value for a junction between these exons would be:

$$30 + 20 + 40 - 10 \times \log_{10} \left( \frac{900}{5000} \times \frac{100}{400} \times 200^2 \right) = 90 - 10 \times \log_{10} (1800) \approx 50 \text{ to } 60$$



Equation 3. Case-1: 3 unique junction evidence between 9x and 12x exons

$$20 - 10 \times \log_{10} \left( \frac{900}{5000} \times \frac{100}{400} \times 200^2 \right) = 20 - 10 \times \log_{10} (1800) \approx 0$$

Equation 4. Case-2: 1 unique junction evidence between 9x and 12x exons

$$30+20+40-10 \times \log_{10}\left(\frac{90000}{5000} \times \frac{10000}{400} \times 200^{2}\right) = 90-10 \times \log_{10}(18,000,00) \approx 10 \text{ to } 20$$

#### Equation 5. Case-3: 3 unique junction evidence between 900x and 1200x exons

A simplified table of outputs is shown in Table 110.

Table 110 Simplified junction tab output

E1	E1- reference	s	E1-start/ E1-end	E2	E2- reference	s	E2-Start/ E2-End	Unique-PR	Unique-SR
AGRN-1	chr1	+	886872/ 886993	KLHL17-4	chr1	+	887069/ 887290	3	2
KLHL17-8	chr1	+	888580/ 888747	KLHL17-9	chr1	+	889163/ 889251	1	1
AGRN-11	chr1	+	969352/ 969500	AGRN-12	chr1	+	969577/ 969682	2	1
AGRN-14	chr1	+	970602/ 970766	AGRN-15	chr1	+	970976/ 971119	2	1
AGRN-17	chr1	+	971403/ 971508	AGRN-15	chr1	+	971640/ 971978	6	2
AGRN-20	chr1	+	972570/ 972697	AGRN-21	chr1	+	972816/ 972930	2	3
AGRN-21	chr1	+	972816/ 972930	AGRN-22	chr1	+	973019/ 973138	2	1
AGRN-26	chr1	+	974809/ 975038	AGRN-27	chr1	+	975146/ 975280	3	0
AGRN-27	chr1	+	975146/ 975280	AGRN-28	chr1	+	975476/ 975572	3	1
PUSL1-4	chr1	+	1234697/ 1234846	PUSL1-5	chr1	+	1234924/ 1235094	2	0
PUSL1-6	chr1	+	1235877/ 1235931	PUSL1-7	chr1	+	1236152/ 1236314	2	0
VWA1-0	chr1	+	1362170/ 1362727	VWA1-3	chr1	+	1364324/ 136600	4	0



E1	E1- reference	s	E1-start/ E1-end	E2	E2- reference	s	E2-Start/ E2-End	Unique-PR	Unique-SR
MIB2-3	chr1	+	1548632/ 1548942	MIB2-4	chr1	+	1549017/ 1549188	5	1
MIB2-4	chr1	+	1549017/ 1549188	MIB2-5	chr1	+	1550038/ 1550144	1	1
MIB2-6	chr1	+	1550234/ 1550428	MIB2-7	chr1	+	1550529/ 1550671	2	1
MIB2-7	chr1	+	1550529/ 1550671	MIB2-8	chr1	+	1550789/ 1550896	2	1
MIB2-9	chr1	+	1551893/ 1551997	MIB2-10	chr1	+	1552080/ 1552242	2	2
MIB2-10	chr1	+	1552080/ 1552242	MIB2-11	chr1	+	1552317/ 1552450	3	0
MIB2-11	chr1	+	1552317/ 1552450	MIB2-12	chr1	+	1552539/ 1552687	2	0
MIB2-14	chr1	+	1553262/ 1553422	MIB2-15	chr1	+	1553516/ 1553642	3	1

 Table 110 Simplified junction tab output (continued)

#### Browser Extensible Display (BED) output

The BED format was developed to extend the UCSC Genome Browser with userdefined tracks. BED is used to visualize the splice and fusion junctions in the UCSC Genome browser and in the IGV browser (see Figure 41 on page 391). For general documentation about the BED format, including information about all of the BED fields, go to:

#### genome.ucsc.edu/FAQ/FAQformat.html

For information about visualization software, see "Pairing information in a BAM file" on page 466.

Each line in the track defines a junction where chromStart is the smaller of the coordinates of and chromEnd is the greater.

There are two blocks because a junction typically contains two exons. BlockSizes are the lengths of the exons. The block starts the beginning of the exons. When fusions on different strands or chromosomes, two lines are added to the output, with each line representing one chromosome. Different colors are used to color-code different types:

Figure 41 on page 391 shows the Upstream Hypersensitive Region (UHR) gene region displayed with the Integrative Genomics Viewer (IGV) for positions 3,530,193 to 3,548,355 of Human Chr-1. The following sections describe the tracks in Figure 41.

#### WIG (x2) tracks

The top two tracks show the genomic coverage using the negative strand and positive strand generated by the Bam2Wig tool (Max: 100 coverage).



#### **BAM** track

The middle track shows the alignments from the BAM file. For display purposes, reads are filtered with MAPQ threshold of 45 (a stringent filter). Bases with quality value 5 to 20 are shaded.

#### **BED track**

The fifth track shows the junctions detected by the splice finder (BED file). As shown in the figure, all junctions detected are "known" and so are shaded in green.



Figure 41 BED-IGV extended track example

SEQ	The optional SEQ output files are in FASTA-format and list regular junctions, alternatively-spliced junctions, and fusion junctions. The entries are 100bp from each end of putative junctions. If any of the exons is less than 100bp in size, the total exon sequence is added.						
Circos	The Circos-ready format is generated when the wt.splice.finder.output.format parameter is set to 4. Gene fusion calls are output in Circos-compatible format, to enable easy visualization with Circos. Follow these steps to use this file with Circos:						
	1. Copy the folder splice-finder output directory circus-fusion and paste it under the Circos installation directory.						
	<b>2.</b> By default, only the human karyotype file is distributed. If you are working with another species, you must replace the karyotype file.						
	<b>3.</b> Run this command to start Circos:						
	<pre>bin/Circos -conf circus-fusion/circos.conf</pre>						
Output format values	The value of the wt.splice.finder.output.format parameter determines which file formats are included in the splice-finder output, as shown in the following list:						
	<ul> <li>wt.splice.finder.output.format=1</li> </ul>						

- <bam\_prefix>.alternative\_splicing.tab
- <bam\_prefix>.fusion.tab
- <bam\_prefix>.junctions.tab
- <bam\_prefix>.stats
- wt.splice.finder.output.format=2
  - <bam\_prefix>.alternative\_splicing.tab
  - <bam\_prefix>.fusion.tab
  - <bam\_prefix>.junctions.bed
  - <bam\_prefix>.junctions.tab
  - <bam\_prefix>.stats
- wt.splice.finder.output.format=3
  - <bam\_prefix>.alternative\_splicing.seq
  - <bam\_prefix>.alternative\_splicing.tab
  - <bam\_prefix>.fusion.seq
  - <bam\_prefix>.fusion.tab
  - <bam\_prefix>.junctions.bed
  - <bam\_prefix>.junctions.seq
  - <bam\_prefix>.junctions.tab
  - <bam\_prefix>.stats
- wt.splice.finder.output.format=4
  - <bam\_prefix>.alternative\_splicing.seq
  - <bam\_prefix>.alternative\_splicing.tab
  - <bam\_prefix>.fusion.seq
  - <bam\_prefix>.fusion.tab
  - <bam\_prefix>.junctions.bed
  - <bam\_prefix>.junctions.seq
  - <bam\_prefix>.junctions.tab
  - <bam\_prefix>.stats
  - circos-fusion/fusion.genes
  - circos-fusion/fusion.links
  - circos-fusion/circos.conf
  - circos-fusion/ideogram.conf
  - circos-fusion/ticks.conf
  - circos-fusion/karyotype.human.txt

# PART VI Small RNA Analyses

# 21

# Run a Small RNA Mapping Analysis

This chapter covers:

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# **Overview**

The LifeScope<sup>™</sup> Software small RNA analysis modules are used to analyze high throughput small RNA sequencing raw data. These modules encapsulate a workflow to map reads to the target genome, to detect small RNAs in the genome, and to determine the expression of the detected small RNAs.

The small RNA mapping module maps small RNA (also known as microRNA or miRNA) reads using the mapping program mapReads. This module maps the small RNA reads to three different references in three steps:

- In the first step, the given set of reads are mapped to filter sequences in order to eliminate the reads generated from irrelevant sources (such as tRNA, adaptor sequences, or others).
- In the second step, the remaining reads from the first step are mapped to the list of known small RNA precursor sequences (miRBase annotations). This list is downloaded from Sanger's website. This lists for human hg18 and hg19 references are available in the LifeScope<sup>™</sup> Software reference repository.
- In the third step, the unmapped reads from the second step are mapped to the genome reference sequence, in order to find novel small RNAs in the sample.

The mapped reads outputs from the second and third steps are merged, and the merged file is the primary output of the small RNA mapping module.

The major components of the small RNA mapping module are:

- 1. Filter reference mapping
- 2. miRBase reference mapping
- **3.** Genome reference mapping
- 4. Merge of mapped reads output (MA files)
- 5. BAM generation (including ma2BAM, refcor, and sort steps) and BAM merge

Parameters are provided to skip any of the three mapping steps.

Usual read lengths from small RNA sequencing are 35–50 bases, and the length of the miRNA fragments is only 18–28 bp. The reads contain both the miRNA sequence and a P2 adapter sequence at the end. The mapReads phase does an extra step of extension onto adaptor, to correctly remove the adaptor sequence from the read and to report only small RNA sequences in the BAM file.

After the miRBase mapping step, the mapping output file (a MA file) has alignments with respect to the smaller miRBase reference locations. Before merging the MA files, the program converts these miRBase alignments to genomic reference locations, using the input GFF file (described in "Small RNA mapping input files" on page 397).

# Example of running the small RNA mapping module

The small.rna standard workflow provides an example of how to run the small RNA mapping module. Here are example LifeScope<sup>™</sup> Software shell commands for this workflow, which includes small RNA coverage and small RNA counts:

```
# log into the shell
lscope.sh shell -u username -w password
# cd to the projects repository
cd projects
# make a project, and open it
mk mirna
cd mirna
# make an analysis, and open it
mk run1
cd run1
# define the analysis type
set workflow small.rna
# define the input
add xsg rna.xsg
add xsg rna2.xsg
# specify the reference to be used
set reference hg18
# optionally change parameter defaults after this line
# list the analysis configuration
ls
# run the analysis
run
# display the run progress
ls
# exit the command shell
exit
```
Dummy XSQ file names are used in this example.

In order to change a parameter value in your analysis, use the set param shell command after the line # *optionally change parameter defaults after this line* in the example commands. For instance, to change the create.unmapped.bam.files parameter, use this shell command:

set create.unmapped.bam.files 0 /secondary/smallrnamapping.ini

You must know which INI file to use for set param command. The ls command, executed in an analysis virtual directory after the workflow has been defined, lists the INI files used in the analysis.

### Small RNA mapping input files

The small RNA mapping module requires the following input:

- One or more XSQ files containing sequencing data.
- A Genome Coordinates GFF file containing precursor sequences with genomic locations. This file is available the following link:

#### http://www.mirbase.org/ftp

- A single multi-fasta reference file for the filter mapping step.
- A single multi-fasta reference file for the genome mapping step.

The mapping modules accepts as input one or more input XSQ files. The input reads can have different read lengths but they must be of the same library type.

#### **RNA-Seq reads**

The RNA-Seq reads used in the small RNA modules are different from the genomic reads. For RNA-Seq reads:

- Only transcribed sequences are measured by the system.
- Genome coverage is non-uniform due to variation in transcriptional intensity.
- A large subset of reads originate from uninteresting sequences such as ribosomal RNA (rRNA).

#### Reads input specification

Plan your analysis input carefully. The way you define your reads input affects the behavior of your analysis. The following factors control how your input data is analyzed:

- Index (barcode) IDs Using an index ID restricts your input to the reads data of one or more indices.
- **Grouping of reads** Each group of reads is analyses together as one specimen. The output data for a group is combined into one set of results files.
- Multiple sample runs Unrelated reads can be processed together in one run of LifeScope<sup>™</sup> Software, while analyzed separately as separate input data.

See "Define input data" on page 85 for more information on designing how to add input data to your analysis.

### Legacy data

If the data you want to process with LifeScope<sup>™</sup> Software is in CSFASTA and QUAL files, these files must be converted to the XSQ file format, through one of the following methods:

- The LifeScope<sup>™</sup> Software UI handles converting these files to the XSQ format before mapping.
- If you are using the LifeScope<sup>™</sup> Software command shell, you must first convert the CSFASTA and QUAL files to the XSQ format. See Appendix C, "XSQ Tools" on page 479 for information on the standalone XSQ converter.

## Stages of mapping

The mapping algorithm involves following stages:

- **Scatter** The process of distributing reads to the available compute nodes, to achieve parallelization.
- Mapping Each tag is aligned to the reference.
- **BAM file generation** The process of converting the output of mapping and pairing to BAM format. For color-space data, the base translation step is also included. Due to the scattering step, the BAM files at each compute node are a sub-set of the final BAM file and are referred to as baby BAMs.

BAM generation involves these steps:

- Ma2BAM A conversion step, from MA format to BAM format, for fragment data only.
- Refcor Reference-assisted color-to-base translation.
- Sort BAM Sorting the BAM file from bead-id order to coordinate order.
- Merge BAM Merging the mini-BAM files created in the sort phase into the final output BAM files.

Figure 42 shows the phases involved in small RNA mapping.



Figure 42 The phases within the small RNA mapping module



**Scatter** The scatter stage distributes reads to the available compute nodes, in order to gain the efficiency benefits of parallel computing.

The mapping module can accept multiple read-sets, which are allowed to have different read lengths. The input set of read-sets from multiple XSQ files is first split based on read length. Fragment read-sets are partitioned into separate categories based on these read length categories:

- $0 \leq \text{Length} < 25$
- $25 \leq \text{Length} < 35$
- $35 \leq \text{Length} < 50$
- $50 \leq \text{Length} < 60$
- $60 \leq \text{Length} < 75$
- $75 \leq \text{Length}$
- Trimmed reads, with variable read lengths

The reads from each category are scattered into multiple jobs based on the parameters fragmap.minreads.per.node and fragmap.number.of.nodes. These parameters are described in the performance section.

Filter map The reads are mapped to the filter reference in the filter mapping step. Reads that align to the filter reference are skipped from mapping to miRBase and genome references. The filtered reads are populated in separate BAM file inside the outputs/filtered directory. A filter reference for use with human reads is available with LifeScope<sup>™</sup> Software. This reference contains:

- SOLiD<sup>™</sup> adapter sequences
- Human ribosomal RNAs (rRNAs)
- Human transfer RNAs (tRNAs)
- Other human sequences
- Single-base-repeat sequences, including Poly-A, Poly-T, and more

Complete the following steps to construct a filter reference for another species.

- 1. Copy the adapter and single-base-repeat sequences to a new file.
- **2.** Append FASTA-formatted species-specific ribosomal, transfer RNA, and other sequences to the filter reference.
- (Optional) Add the new filter reference file to the reference repository.
  - If you added the assembly for this species to the repository, copy the new filter file to the assembly directory. See "Add new reference files" on page 517. The filter reference is then set automatically when you use the follow shell command: set reference assembly\_name.
  - If you do not have the assembly for this species in the reference repository, use the analysis.filter.reference parameter to include the new filter reference in your analysis. For example:

set analysis.filter.reference filterfile secondary/global.ini

An absolute path is recommended for filterfile. A relative path is interpreted from the Linux directory in which the shell was started (when the lscope.sh shell command was given). (The secondary/global.ini INI file is the correct file for the analysis.filter.reference parameter.)



miRBase map	This phase maps to known miRBase sequences. In a pre-process step miRBase database is converted from GFF to FASTA format to create the miRBase reference. Then the map module is invoked with the unmapped reads from the filter map step and the miRBase reference. The outputs of this step are the miRBase match file (or files) and unmapped reads in XSQ format.			
Genome map	The unmapped reads from the miRBase map step are mapped to the genome reference in the genome mapping step to produce genome match files.			
miRBase2Genome	The output of the miRBase mapping step has alignments with respect to the smaller miRBase reference locations. This phase converts alignments from the miRBase coordinate system to the genome coordinate system required with LifeScope <sup>™</sup> Software BAM files.			
BAM file	Ma2BAM			
generation	Ma2BAM is a format converter that changes match files to a pro-BAM format. These pro-BAM files do not contain sequence or quality information but contain relevant BAM meta data. The pro-BAM files are sorted by incoming bead-id order. Relevant meta data from XSQ files is captured in pro-BAM files at this stage. The parameters smallRNA.bam.primary.align.only, smallrna.bam.mqv.threshold, and smallrna.create.unmapped.bam.files control the functionality of the ma2BAM utility. The pro-BAM files are generated on the scratch folder.			
	Ma2BAM takes output of mapping (MA files) and converts them to pro-BAM format. During BAM generation, mapping quality values are also computed.			
	For reads with multiple ungapped alignments, the read with the highest mapping quality value is chosen as the primary alignment for the read, and is reported to the BAM file. In cases where there are multiple alignments with the same quality value, the primary alignment is chosen at random from among the alignments with the same quality value. The details of calculating mapping quality values are described below.			
	Mapping quality values			
	Quality values are Phred-scaled quality scores. For any given alignment, an MQV is a measure of the confidence of a read aligning to the particular location, given all possible alignments for the read. Quality values are in the range 0–100:			
	<ul> <li>0 – The highest probability of error</li> <li>100 – The lowest probability of error</li> </ul>			
	The following factors increase confidence and increase a read's quality value:			
	<ul><li>Longer alignment</li><li>Fewer possible alignments</li><li>Fewer mismatches within the alignment</li></ul>			
	These factors reduce confidence and reduce a read's quality value:			
	Shorter alignment			
	More possible alignments			
	More mismatches within the alignment			



In order to be consistent with the Phred quality score (-10\*log10[prob(error)]) used widely in literature, the quality is computed as the negative log odds of misaligning the read. The resulting quality values are normalized by the maximum possible value to ensure that the quality values are within the range 0–100.

#### Refcor

Reference Assisted Color-to-Base Translation (also known as refcor) combines information from color calls, optional ECC extra primer round calls, and the reference sequence, to enhance the quality of base-space reads by detecting and fixing color errors that convert into multiple base-space errors.

Refcor accepts a set of pro-BAM files, which contain alignments information without base or color sequence, and generates a corresponding set of BAM files containing alignments with base-space and color-space calls (on the scratch folder). These BAM files are sorted by bead-ID and contain dummy base calls and QVs, but are otherwise compatible with the BAM specification. The refcor module produces bead-ID-sorted BAM files that have valid calls and QVs.

For each read, the module constructs the primer-transition graph with vertices corresponding to all possible *k*-mers (*k*=4 for 2BE+4BE, *k*=2 for 2BE) in each read position. Two adjacent vertices are connected with an edge if they overlap in *k*–1 bases. A score corresponding to the (1-Pe), where Pe is the probability of being erroneous, is assigned to each of the vertices. Scores are derived from quality values corresponding to enumerated calls. When making a base call in a position, the current algorithm considers the cumulative probability of all sequences that have called base in that position. The evidence from the reference (base) is assigned to the vertex with *k*-mer ending in corresponding base only if the color transition between current and following reference bases matches the color call in a read. Such assignment reduces the reference over-correction in SNP positions. In order to reduce reference overcorrection in positions with color errors, we assign to reference base a very low weight corresponding to QV of 8. The weight can be reduced to QV of 0, completely reducing the reference guiding in the color-to-base translation. This results in a smaller skew between reference to non-reference allele ratio. Increasing the weight results in false positive variant calls reduction. This process restores the phase shift caused by color errors or completely eliminates the errors.

The parameters bamgen.refcor.softclip, smallRNA.bamgen.refcor.addcs, refcor.reference.weight, and refcor.base.filter.qv control the refcor program. The defaults provided are validated for a wide range of data.

- If the XSQ file contains base-space data only, the XSQ base call and quality value are transferred to BAM file. In this case, no color calls are present in BAM file irrespective of the value of bamgen.refcor.addcs.
- If the XSQ file contains 2BE color-space data only, refcor uses aligned 2BE and reference to do base translation for the aligned part of the read. The base sequence and quality value are generated using a proprietary algorithm. For reads with no alignment, bases are determined by naively translating color to base. For these unaligned reads, the QV for the base at position *k* is the minimum color QV for all positions  $\leq k$ .
- If the XSQ file contains ECC data, then reference with 2BE and 4BE color are used to generate the new base sequence. The base sequence and quality value are generated using a proprietary algorithm. For reads with no alignment, the original base-space data from ECC is preserved.



#### Sort BAM

The output of refcor is a set of BAM files in bead-id sorted order. These files are sorted in this step using Picard sort into coordinate sorted order before merge. These BAM files are referred to as mini-BAM files and are stored in the analysis temporary directory.

#### Merge BAM

This phase accepts a set of mini-BAM files and outputs a set of BAM files. These BAM files are stored in the analysis output directory (which is typically a network-attached storage device).

One BAM file is generated per read-set, which is defined as a barcode index in one XSQ file. For a 96-barcoded dataset in a single XSQ file, either 96 or 97 BAM files are created. The 97th BAM contains reads that have the default barcode index.

## Small RNA mapping parameters

Table 111 lists small RNA mapping parameters.

Table 111 Small RNA mapping parameters

Parameter name	Default value	Description
Mandatory parameter		
smallrnamapping.run	1	Whether or not to run the small RNA mapping module. Allowed values:
		• <b>0</b> : Do not run the small RNA mapping module.
		• 1: Run the small RNA mapping module during this analysis.
		The run parameter is set automatically by the shell during a standard workflow. Accept the default for most use.
Optional parallelization parameter	ers	
fragmap.number.of.nodes	4	The number of compute nodes available for this analysis.
fragmap.max.number.of.jobs	100	The maximum number of jobs allowed for mapping.
		Allowed values: Integers 24–1000.
mapping.np.per.node	8	The number of processors per node use for mapping. The reads are divided into the number of chunks specified for this parameter.
fragmap.minreads.per.node	400000	If the total number of fragments exceeds this amount, then the analysis is distributed across the available nodes.
		Allowed values: Integers 1–125000000 (125 Million).
fragmap.maxreads.per.node	15000000	The maximum number of fragments that can be processed on a single node.
		Allowed values: Integers 1–150000000 (150 Million).

**Optional resource parameters** 

Parameter name	Default value	Description
wall.time	120	Total time for the process to complete.
java.heap.space	1500	Dynamic memory requirement.
mapping.memory	15gb	The total memory, in gigabytes, that is available for map reads, per compute node. Include the units gb in the setting.
		This number is set to 15 GB because the minimum hardware requirement for memory is 16 GB.
		<b>Note:</b> Typically, the job scheduler allocates on nodes that have the requested memory available. For mixed hardware, it is suggested that different queues are created based on the memory available. Mapping jobs should be launched on high memory systems. For human references (hg18, hg19) and a 25.2 scheme, 45 GB is recommended for fastest performance. The smallest recommended RAM for human reference is 19 GB.
Optional mapping parameters		
smallRNA.filter.mapping	true	Whether to turn on or off the filter mapping step. Allowed values:
		• <b>false</b> : Turn off the filter mapping step.
		• <b>true</b> : Turn on the filter mapping step.
smallRNA.genome.mapping	true	( <i>Optional</i> ) Whether to turn on or off the genome mapping step. Allowed values:
		• <b>false</b> : Turn off the genome mapping step.
		• <b>true</b> : Turn on the genome mapping step.
mapping.in.base	false	Whether or not to include base-space data in the mapping. Allowed values:
		• <b>false</b> : Do not include base-space data in mapping.
		• <b>true</b> : Include base-space data in mapping.
smallRNA.mirBase.mapping	true	( <i>Optional</i> ) Whether or not to turn on the small RNA miRBase mapping step. Allowed values:
		• <b>false</b> : Turn off the small RNA miRBase mapping step.
		• <b>true</b> : Turn on the small RNA miRBase mapping step.
Optional BAM file generation param	eters	
bamgen.refcor.addcs	true	Whether or not to add the color sequence to BAM records. If the XSQ file does not contain color space, this parameter is ignored, and the resulting BAM file output does not contain color space. See "Refcor" on page 401 for information about refcor. Allowed values:
		• <b>false</b> : Do not add color sequence to the output.
		• <b>true</b> : Add color sequence, if color space is present.

Table 111	Small RNA	mapping	parameters	(continued)
	SHIULT IN A	mapping	purumeters	(continucu)

#### Table 111 Small RNA mapping parameters (continued)

Parameter name	Default value	Description
refcor.reference.weight	8	Reference weight. This parameter is used during base translation. In the read reconstruction process, multiple signals are combined to generate the final base call. This parameter adds weight (in terms of Phred score) to the signals which are compatible with reference. Color combinations that result in a variant are considered compatible with reference. Additional weight helps to eliminate base errors caused by color error(s) during base translation. Allowed values: Integers 0–100.
refcor.base.filter.qv	10	Bases with a quality value below the value of this parameter provided are replaced with 'N'. Allowed values: Integers 0–255.
bamgen.mqv.threshold	0	Provides control over the contents written to the output BAM file depending on the quality value of the alignment. To preserve only high quality alignments, set this value to a positive integer.
		Allowed values: Integers 0–100.
create.unmapped.bam.files	false	If set to TRUE, creates an additional BAM output file containing unmapped reads. Allowed values:
		• <b>false</b> : Do not create the unmapped reads output file.
		<ul> <li>true: Create a BAM output file containing the unmapped reads.</li> </ul>

### Mapping performance

Mapping speed depends on hardware properties as well as on the scattering logic. Mapping runs are split into multiple jobs for processing efficiency. The following parameters affect how a mapping run is split:

- **fragmap.number.of.nodes** Specifies the number of jobs that are created, only if a split is necessary. The size of each split job is approximately the total number of reads divided by the number of jobs.
- **fragmap.max.number.of.jobs** Determines the maximum number of jobs that can be launched per analysis. Setting this parameter to a very large value can cause the scheduler to behave incorrectly.
- **fragmap.minreads.per.node** Determines the threshold of beads beyond which splitting occurs.
- **fragmap.maxreads.per.node** Determines the largest number of beads that can be mapped per node. The default is based on the minimum scratch space requirements. If you scratch space is smaller or larger, this number can be made smaller or larger respectively.
- **processors.per.node** This is the number of cores available for mapping analysis on each node.
- **mapping.memory** For human mapping and 25.2.0 scheme, there are 3 schema lines. Simultaneous handling of schema lines reduces I/O and therefore improves mapping speed.

Memory requirements for mapping jobs are the following:

- Reference: ~6 GB
- 1 schema line: ~13 GB
- 1 schema line and reference: ~19 GB
- 2 schema lines and reference: ~32 GB
- 3 schema lines and reference: ~45 GB

Below 19 GB, the reference is split up and I/O increases significantly. Pre-built hash tables are not used for memory less than 24 GB. LifeScope<sup>™</sup> Software supports the reuse of hash tables for human genome reference files (hg18 and hg19) and the default mapping schemes 35.2, 25.2, and 20.1. Pre-calculated color-space hash tables are included in the reference directory available with LifeScope<sup>™</sup> Software. If system RAM is 24 GB or higher, the hash tables are loaded into memory. Depending on the pipeline, this hash table reuse typically saves 2–3 hours in analysis processing time.

In general, set the mapping.memory parameter to be 1 GB less than the total system memory (RAM) available.

Mapping only scales when the number of reads that are mapped together is greater than 100 M. Splitting such that the total number of fragments per node is significantly less than 100 M is not recommended. The splitting can be explained using the following pseudo-code.

```
if (num.reads <= fragmap.minreads.per.node ) { num.jobs = 1}
reads.per.job = num.reads / fragmap.number.of.nodes
if (reads.per.job > fragmap.maxreads.per.node) {
    reads.per.job = fragmap.maxreads.per.node}
num.jobs = celing (num.reads / reads.per.job)
if (num.jobs > fragmap.max.number.of.jobs ) {
    throw exception; }
```

In above pseudo-code, the input read-sets (possibly from multiple XSQ files) have num.reads that fall in one read length category. Num.jobs is the total number of jobs launched for mapping analysis of num.reads beads.

In general, the mapping parameters allow mixing and matching single-anchor and multiple-anchor schemes, and also mixing and matching different seed lengths and different numbers of mismatches. For example, an unmapped scheme of 25.2.0, 35.2.0:15:25, 20.1.0, with a repetitive scheme of 35.2.0:30, 45.2.0, 60.2.0, is supported, but not recommended. The impact on performance can be significant.

# Internal Table 112 lists small RNA mapping parameters that we do not recommend changing. parameters

Table 112 Small RNA internal mapping parameters

Parameter name	Default value	Description
Mapping parameters	1	
smallRNA.adaptor.sequence	CGCCTTGGCCGTACAGCAG	( <i>Optional</i> ) The small RNA adaptor sequence. Used by mapreads in the extension step to identify the adaptor in the read.
smallRNA.filter.mapping.max.hits	10	The maximum number of hits for a read in the filter mapping step. Allowed values: Integers.

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Table 112	Small	RNA	internal	mapping	parameters	Icontinued
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Parameter name	Default value	Description
smallRNA.filter.mapping.mismatch .penalty	-2.0	Specifies a negative scoring penalty for mismatch that is used in local alignment mode. Setting this value to a negative number greater than the read lengths has the effect of disallowing mismatches. Allowed values: Floats.
smallRNA.filter.mapping.scheme	25.3.0	The mapping scheme for the filter mapping step. The values are: the size of the seed, number of mismatches allowed in the seed, and the start position of the seed in the read. See "Mapping schemes" on page 407 for information about mapping scheme notation.
smallRNA.genome.mapping.max. hits	30	( <i>Optional</i> ) The maximum number of hits allowed for a read in the genome mapping step. Allowed values: Integers <= 0.
smallRNA.genome.mapping. mismatch.penalty	-2.0	Specifies a negative scoring penalty for a mismatch. Used in local alignment mode. Allowed values: Floats <del>-&lt;= 0.0</del> .
		If the absolute value of this penalty is set to greater than the read length, effectively no mismatches are allowed on this pass.
smallRNA.genome.mapping. scheme	20.1.0	( <i>Optional</i> ) The mapping scheme for the filter mapping step. The values are: the size of the seed, number of mismatches allowed in the seed, and the start position of the seed in the read. See "Mapping schemes" on page 407 for information about mapping scheme notation.
smallRNA.mapping.fragment.tag	F3	Small RNA mapping fragment tag name.
smallRNA.mapping.schemas.file	-	Location of the schemas file for small RNA mapping.
smallRNA.mirBase.mapping.max. hits	30	( <i>Optional</i> ) The maximum number of hits per read in the miRBase mapping step.
smallRNA.mirBase.mapping. mismatch.penalty	-2.0	Specifies a negative scoring penalty for a mismatch. Used in local alignment mode. Allowed values: Floats <del>-&lt;= 0.0</del> .
		If the absolute value of this penalty is set to greater than the read length, effectively no mismatches are allowed on this pass.
smallRNA.mirBase.mapping. scheme	18.2.0	( <i>Optional</i> ) Mapping scheme for the miRBase mapping step. The values are: the size of the seed, number of mismatches allowed in the seed, and the start position of the seed in the read.
smallRNA.valid.adjacent. mismatch	false	( <i>Optional</i> ) Indicates how to count adjacent mismatches. Allowed values:
		• <b>false</b> : Treat two valid adjacent mismatches as two.
		• <b>true</b> : Treat two valid adjacent mismatches as one.
		Required if smallRNA.mirBase.mapping is set to true.

Parameter name	Default value	Description
task.scratch.dir.delete	true	Whether to keep the scratch files generated during small RNA. Allowed values:
		• <b>false</b> : Do not keep the scratch files.
		• <b>true</b> : Keep the scratch files.
analysis.clean.temp.files	1	Whether or not to keep temporary files generated during small RNA mapping. Allowed values are:
		• <b>0</b> : Do not keep temporary files.
		• 1: Keep the temporary files.
BAM file generation parameters		
bamgen.primary.align.only	true	Whether only primary alignments or all the alignments are reported in the BAM files. Allowed values:
		• false: Report all alignments.
		<ul> <li>true: Report only primary alignments (both gapped and ungapped).</li> </ul>
bamgen.refcor.softclip	0	Whether to try partial recovering of hard-clipped regions. For cases when the unaligned portion of a read needs to be presented in the BAM record, we use soft clipping in the CIGAR string. If a base-space sequence generated by ECC decoder is available, we use that sequence to represent corresponding part of the read. If the ECC decoded base-space is not available, then we naively translate using 2BE and/or +4BE evidence (if available).
		The soft clipped part of the read can be permanently clipped if either of these conditions is true:
		• The ECC base QV is less than 4.
		• The ECC base and 2+4 naive propagation have less than 50% similarity.
		Allowed values:
		• <b>false</b> : Do not try partial recovery.
		• true: Attempt partial recovery.

 Table 112
 Small RNA internal mapping parameters (continued)

Mapping schemes The mapping scheme parameters listed in Table 110 on page 385 determine the behavior of the mapping module. The LifeScope<sup>™</sup> Software mapping module is based on seed-and-extend mapping, where the initial alignment step locates short matches between a read and the reference sequence. The seed specifies the length of the attempted match. With the seed-and-extend approach, a mapping scheme parameter value such as 25.2.0 is interpreted as follows:

- First value The seed length (25).
- Second value The quantity of allowed mismatches in the seed (2).
- **Third value** The start site of the seed within the read (0).



Figure 43 Seeds for local alignments

For example, with a 25.2.0 seed extend scheme, the mapping module examines the first 25 bases of a 50 bp read, and if that portion aligns to the reference with two or fewer mismatches, then the mapping module attempts seed extension. Typically, because small RNA reads are short, the seed lengths also are short, for example 18 bp for miRBase and 20 bp for genomic mapping. The extension step involves identification of the adaptor sequence. For a hit of length 35, once a hit is found, say of length 18, the extension part is evaluated for best match for adaptor starting at 19 and ending at 35 or adaptor starting at 20 and ending at 35 and so forth.

### Small RNA mapping output files

The small RNA mapping module generates two BAM files containing alignments in coordinate order:

- A filtered BAM file, from the filter mapping step.
- A mapped BAM file with genomic locations. This content is merged output from the miRBase mapping and genomic mapping steps.

For information about the BAM file format, see Appendix B, "File Format Descriptions" on page 455.

The number of BAM files generated depends on the input XSQ files. One BAM file is generated per read-set (a read-set is defined as data from one barcode, in one XSQ file). So if the input data is an XSQ file containing 96 barcodes, then the mapping module generates at least 96 output BAM files. Beads that are unclassified in any barcode are output into a separate additional BAM file. If the parameter

smallRNA.create.unmapped.bam.files is set to TRUE, then an additional 96 or 97 output BAM files corresponding to the unmapped reads are also generated.

The filenames for the output BAM files are created using information from the input XSQ file, including: file base name, file id, index name, and index id. The BAM files are named according to the following patterns:

- Non-indexed BAM files: *xsqname-fileID-1.bam*
- Indexed BAM files: xsqname-fileID-idx\_bcIndex-bcID.bam

The fields in the filenames are:

- **xsqname** The XSQ file base name, without the .xsq extension.
- fileID The file ID for internal XSQ file tracking.
- **bcIndex** The barcode index.



• **bcID** – The barcode identifier for internal barcode tracking.

The directory structure for mapping output is as follows:

The directory name <*mappingIniFileBasename*> is the basename of the mapping INI file. The default is smallrnamapping.

The string *<sampleName*\*> is determined by the sample description for the particular read-set.

The small RNA mapping output files are used by the BAMStats mapping statistics module and by the small RNA tertiary analysis modules.

### **Mapping statistics**

Mapping statistics occur after mapping as an optional post-processing step named BAMStats. BAMStats accepts the output of the mapping step and generates statistics files to provide an in-depth understanding of the experimental data and to better detect the presence of anomalies. LifeScope<sup>™</sup> Software shell users can display the mapping statistics output data as a chart with a spreadsheet program or other third-party program. A subset of the output from mapping can be visualized in the LifeScope<sup>™</sup> Software UI as a series of line and bar charts, if the analysis is run in the projects repository.

Mapping statisticsTable 113 lists the parameters which control the BAMStats output.parameters

Table 113 BAMStats parameter description

Parameter name	Default value	Description
Mandatory parameters		
bamstats.run	1	<ul> <li>Whether or not to run the BAMStats module. Allowed values:</li> <li>0: Do not run the BAMStats module. BAMStats statistics are not generated.</li> <li>1: Run the BAMStats module during this analysis. The run parameter is set automatically by the shell during a</li> </ul>
Optional parameters		standard workflow. Accept the default for most use.
bamstats.maximum.coverage	10000	Defines the maximum coverage allowed for locations in the reference. Locations with coverage more than the maximum coverage value are ignored during coverage calculations. Allowed values: Integers 1–10000.



#### Table 113 BAMStats parameter description (continued)

Parameter name	Default value	Description
bamstats.maximum.isize	100000	The maximum inset size for LMP and PE libraries. Reads with an insert size more then the specified value are ignored for Insert Range Report calculation.
		Allowed values: Integers 1–100000.
bamstats.wig.primary.only	1	Use only primary alignments for coverage in WIG file format. Allowed values:
		• <b>0</b> : Do not restrict coverage in WIG file format to only primary alignments.
		<ul> <li>1: Restrict coverage in WIG file format to only primary alignments.</li> </ul>
bamstats.combined.report.both. strands	0	Whether or not to combine data from both strands for coverage in WIG format. Allowed values: 0,1.
bamstats.wig.binsize	100	The bin size for coverage in WIG file format.
		Allowed values: Integers 1–100000.
bamstats.bin.isize	100	The bin size for insert range distribution.
		Allowed values: Integers 1–100000.
bamstats.wig.combined.report.both .strands	0	Whether to combine data from both the strands for coverage in WIG format. Allowed values:
		• <b>0</b> : Do not combine data.
		<ul> <li>1: Combine data from both the strands for coverage in WIG format.</li> </ul>
Resource parameters		
wall.time	120	Total time for the process to complete.
java.heap.space	13000	Dynamic memory requirement.
number.of.nodes	4	The number of compute nodes available for this analysis.
memory.request	14gb	Memory request. Include the units gb in the setting.

#### Table 114 lists BAMStats parameters that we do not recommend changing.

Table 114 BAMStats internal parameter description

Parameter name	Default value	Description
bamstats.group.stats	1	Enable combined statistics covering all read-sets in the group or sample. Allowed values:
		• <b>0</b> : Only statistics for BAM files are generated.
		• 1: Also generate combined statistics for the group or sample.
bamstats.maximum.mismatches	100	The maximum mismatches allowed in the alignments. Any alignment with more then the specified number of mismatches is ignored while generating reports related to number of mismatches. Allowed values: Integers 0–100.

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	1	1
Parameter name	Default value	Description
bamstats.maximum.baseqv	100	Max base quality values. Any base with base quality value more then the specified value is ignored while generating reports. Allowed values: Integers 0–100.
bamstats.maximum.mappingqv	255	Maximum mapping quality value. Any alignment with mapping quality value more than the specified value is ignored. Allowed values: Integers 0–100.
bamstats.wig.minimum.mappingqv	2	Defines the minimum mapping quality allowed for coverage in WIG format. Any alignment with a mapping quality value less than this value is ignored. Allowed values: Integers 0–100.
bamstats.enable.probe.position	0	Enable probe and position error reports. Allowed values:
		• <b>0</b> : Do not generate probe and position error reports. Off is recommended for small RNA.
		• 1: Also generate probe and position error reports.

 Table 114
 BAMStats internal parameter description (continued)

Summary of mapping statistics File formats For every input BAM file, a set of statistics files are generated. These files are in CHT, CSV, TXT, and WIG formats. Each CHT file corresponds to one displayed chart. A CHT file specifies the type of chart, the displayed range of each axis, and the data points, without using external references.

The CHT file format is an internal file format based on the CSV file format, with addition header information included. CHT header information is the following:

```
# name:
# type: scatter2d | pie | vbar | line
# title:
# xaxisname:
# yaxisname:
# yrange: <min>:<tickinterval>:<max>
# yrange: <min>:<tickinterval>:<max>
XAXISNAME, SERIES1NAME, SERIES2NAME, ...
x1, y1.1, y1.2, ...
x2, y2.1, y2.2, ...
x3, y3.1, y3.2, ...
```

The wiggle format (.wig) is a public format typically used for coverage. Visit their site for more information:

hgdownload.cse.ucsc.edu/goldenPath/help/wiggle.html

A genome browser such as the Integrative Genomics Viewer (IGV) can be used to visualize the coverage. For information is available from their site:

www.broadinstitute.org/igv/

For a collection of input BAM files that belong to a sample, a set of cumulative statistics files are generated. The cumulative statistics files are also in CHT, CSV, TXT, and WIG formats. The cumulative statistics can be visualized in the LifeScope<sup>™</sup> Software UI.

### **Directory structure**

The output of the BAMStats module has the following directory structure:

```
bamstats/
```

```
<sampleName1>/*.cht, *.tbl
<sampleName1>/<bam1>/.*cht
<sampleName1>/<bam1>/misc/*.csv, *.txt, *.wig
```

#### **Overview**

BAMStats generates a tab-separated summary file (*BAMfilename*-summary.tbl) that summarizes key mapping quality statistics per input BAM file. The summary report contains a snapshot of statistics in all BAM files present in this sample. This file contains one row for each input BAM file in the sample. The summary file is displayed in the LifeScope<sup>™</sup> Software UI.

<sampleName1>/BAMfilename-summary.tbl

Following directories and reports contain the cumulative statistics from all BAM files that belong to this sample. The Misc folder is not displayed in UI.

```
<sampleName1>/*.cht
<sampleName1>/Misc/*.csv, *.wig, *.txt
```

Following statistics are generated per BAM file in the mapping directory. These reports are not displayed in the UI.

```
<sampleName1>/<BAMfilename>/.*cht
<sampleName1>/<BAMfilename>/Misc/*.csv, *.txt, *.wig
```

In a sample, some of the BAM files possibly represent unhealthy DNA or RNA, causing the cumulative statistics to look poor. The summary tbl file is the unified location for examining the quality of data of all BAM files in the sample.

If the data for a particular BAM file is not as expected, look at the directory-level reports for that BAM file, for details.

Mapping statisticsThis section describes the mapping statistics files generated by the BAMStats module.output filesThe output files generated by BAMStats include the name of the BAM file. The file<br/>name pattern is:

BAMFileName-fileID-barcodeID.StatisticsReportName.tag.extension

This string, without the extension, is referred to by *prefix* in Table 115, the mapping statistics output table. For small RNA reports, *tag* is F3,

 Table 115
 Mapping statistics output files

Report	Description, axis information, filename
Alignment Len	gth Distribution
	A bar plot giving the distribution of alignment lengths found in various tags used in alignment. The report is separated by tag type to provide visibility into the accuracy of individual tags.
	Only the primary alignment for each bead is considered in calculating the distribution.
	Y axis: Frequency X axis: Alignment length (from 0 to the maximum read length)
	Output file name: <i>prefix</i> .Alignment.Length.Distribution. <i>tag</i> .cht



Table 115	Mapping	statistics	output	files	(continued)
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Report	Description, axis information, filename
Alignment Le	ngth Distribution for Unique Alignments
	The same report as the Alignment Length Distribution, except that only tags that have a primary alignment are considered in calculating the distribution.
	By default BAM files contain primary alignments only. In this case, the reports AlignmentLengthDistribution and AlignmentLengthDistribution for Unique Alignments are identical. However, users can also print all alignments or secondary alignments in BAM file. In this case, the two distributions are different.
	Note: The title Unique Alignments is interpreted to mean primary alignments, for this report.
	Y axis: Frequency X axis: Alignment length (from 0 to the maximum read length)
	Output file name: <i>prefix</i> .Alignment.Length.Distribution.Unique. <i>tag</i> .cht
Base Mismate	ch Distribution
	A bar plot giving the distribution of total number of mismatches found in various tags used in alignment The bins are summed over all alignment lengths.
	Only the primary alignment for each bead is considered in calculating the distribution.
	Y axis: Frequency X axis: Number of mismatches (from 0 to the maximum mismatches allowed)
	Output file name: <i>prefix</i> .BaseMismatch.Distribution. <i>tag</i> .cht
Base Mismate	h Distribution for Unique Alignments
	The same report as the Base Mismatch Distribution, except that only tags that have a primary alignmen are considered in calculating the distribution.
	Note: The title Unique Alignments is interpreted to mean primary alignments, for this report.
	Y axis: Frequency X axis: Number of mismatches (from 0 to the maximum mismatches allowed)
	Output file name: <i>prefix</i> .BaseMismatch.Distribution.Unique. <i>tag</i> .cht
Distribution o	f Alignment Length and Number of Mismatches in Tags
	A report providing a simultaneous picture of the alignment length and number of mismatches distributions in various tags used in alignment. Only primary alignments for each tag are considered in calculating this distribution. The bins range from 0 to the maximum read length, and from 0 to the maximum number of mismatches allowed.
	Note: This report is located in the Misc folder, and is not displayed in the UI.
	Y axis: Alignment length (from 0 to the max read length) X axis: Number of mismatches (from 0 to the maximum mismatches allowed)
	Output file name: <i>prefix</i> .AlignmentLength.Mismatch. <i>tag</i> .csv
Base QV Disti	ibution
	A bar plot providing a distribution of base quality values generated using the reference assisted error correction/base conversion algorithm. The base QV distributions are separated for each tag as quality of individual tags could be very different.
	Y axis: Frequency X axis: Base QV (from 0 to the maximum QV)
	Output file name: prefix BaseQV cht



### Table 115 Mapping statistics output files (continued)

Report	Description, axis information, filename
Base QVs by	Position
	A bar plot providing a distribution of base quality values by individual base positions. This report identifies if certain base positions (particularly towards the end of the read) have poor base quality values. All the tags (F3/R3/F5-P2) used in the alignment are combined into a single bin.
	Y axis: Base QV X axis: Base position (from 0 to read length)
	Output file name: <i>prefix</i> .BaseQV.by.Position.csv
Distribution of	of Mismatches by Base QV
	A 3D surface plot providing a distribution of errors (mismatches to reference) by base quality values bins. This report measures whether the base QVs generated are well calibrated to the probability of error in that particular base position.
	Y axis: Error rate X axis: Base QV (from 0 to maximum QV)
	Output file name: <i>prefix</i> .Mismatches.By.BaseQV. <i>tag</i> .cht
Distribution d	f Mismatches by Position
	A bar plot providing a distribution of errors (mismatches to reference) by position within the read. Only the primary alignments for each bead are used to generate this distribution.
	Y axis: Frequency X axis: Position (from 0 to maximum read length)
	Output file name: <i>prefix</i> .Mismatches.By.Position.BaseSpace. <i>tag</i> .cht
Distribution L	y Mapping QVs by Tag
	A bar plot providing a distribution of mapping quality values for individual tags (F3/R3/F5-P2). Only the primary alignment for each bead is used in calculating this distribution.
	Y axis: Frequency X axis: Position (from 0 to maximum mapping QV)
	Output file name: <i>prefix</i> .MappingQV. <i>tag</i> .cht
Coverage Re	port
	A line plot providing a distribution of coverage obtained after mapping/pairing. Only the primary alignment for each bead is used in this calculation.
	Y axis: Frequency X axis: Coverage (from 0 to number of reads)
	Output file name: <i>prefix</i> .Coverage. <i>tag</i> .cht
Coverage Re	port by Chromosome (Contig) and Base Windows
	A line plot providing a distribution of coverage within each reference window. The coverage is calculated within each window along a reference chromosome. The window size can theoretically be anywhere from a single base to the contig length. However the calculations of base level coverage are computationally expensive and less interpretable as the window size increases. Only the primary alignment for each bead is used in this calculation.
	Y axis: Frequency X axis: Coverage (from 0 to number of reads)
	Output file name: <i>prefix</i> .Coverage.By.Chromosome.contig <i>nn</i> .cht
Coverage by	Strand



Report	Description, axis information, filename
	A line plot providing the distribution of coverage within each reference window separated by reference strand (+/-). Only the primary alignment for each bead is used in this calculation.
	Y axis: Frequency X axis: Coverage (from 0 to number of reads)
	Output file name: <i>prefix</i> .Coverage.By.Strand. <i>tag</i> .cht
Coverage files	
	Coverage reports in wiggle format. For each genome position, reports the number of reads that cover (map to or span) the position. Because reporting coverage for each position results in very large files, coverage is reported for bins, with each bin spanning a user-defined number of bases. For each bin, the mean coverage of all the positions in that bin is reported. The parameter <code>bamstats.wig.binsize</code> controls the size of the bins in this file.
	By default one coverage file is generated, per chromosome, per strand. If the parameter bamstats.combined.report.both.strands is set to true, then one file per chromosome is generated, combining coverage from both strands into one file per chromosome.
	Each coverage file includes a header as the first line. The header lines follow this pattern:
	track type=wiggle_0 name=< <i>chrname&gt;</i> description=< <i>coverage from positive/negative/both strand&gt;</i> visibility=full color=0,0,255 fixedStep chrom=< <i>chrname&gt;</i> start=< <i>startpos&gt;</i> step=< <i>binsize&gt;</i> span=< <i>binsize&gt;</i>
	Output file name: coverage_chrnn.POS.wig, coverage_chrnn.NEG.wig

#### Table 115 Mapping statistics output files (continued)

 Mapping statistics
 This section provides example output of some mapping statistics output files.

 example output
 Summary file

#### Summary file

The summary file provides statistics for each BAM file in the sample. This list describes labels used in the summary file:

- **NumUnFilteredBeads** The total number of beads, before any filtering on the instrument. This value is also the fragment count in the input XSQ file.
- NumFilteredBeads The number of beads that pass filtering on the instrument.
- NumMapped The number of reads with primary alignment.
- % filtered that mapped The number of primary reads divided by the number of beads passing instrument filtering (NumMapped / NumFilteredBeads).
- % total that mapped The number of primary reads divided by the number of total beads (NumMapped / NumUnFilteredBeads).

The number of beads filtered out in the instrument is found by subtracting the number of beads that pass filtering from the total number of beads:

NumUnFilteredBeads - NumFilteredBeads

The following are example contents for a summary file:

```
#title: BAMStats Summary
BamFileName,IsColorInBam,IsBaseInXSQ,IsECC,LibraryType,ReadLeng
th,PredictedInsertSize,NumFilteredBeads,NumUnFilteredBeads,Tag1
-NumMapped,Tag1- % total Mapped,Tag1- % filtered mapped,Tag2-
NumMapped,Tag2- % total Mapped,Tag2- % filtered mapped,(Tag1-
AlignmentLength;Min;Max;Avg;Median;StdDev),(Tag1-
NumMismatches;Min;Max;Avg;Median;StdDev),(Tag1-
```



```
MappingQV;Min;Max;Avg;Median;StdDev),(Tag1-
BaseQV;Min;Max;Avg;Median;StdDev),(Tag2-
AlignmentLength;Min;Max;Avg;Median;StdDev),(Tag2-
NumMismatches;Min;Max;Avg;Median;StdDev),(Tag2-
MappingQV;Min;Max;Avg;Median;StdDev),(Tag2-
BaseQV;Min;Max;Avg;Median;StdDev),(Coverage;Min;Max;Avg;Median;
StdDev)
hg19_Simulated_GR_CS_BC16_PE50X25_320Mil-1-Idx_BC15-
15.bam,Y,N,N,PE,50x25,200,18000000,18000000,16785960,93.26,93.2
6,14720544,81.78,81.78,(F3;25;50;46.19;50;6.20),(F3;0;25;1.64;1
;2.03),(F3;0;60;34.37;43;22.61),(F3;1;41;34.66;41;11.02),(F5-
P2;25;25;25.00;25;0.00),(F5-P2;0;16;0.61;0;0.91),(F5-
P2;0;60;33.35;41;23.21),(F5-
P2;1;41;36.15;41;9.56),(0;9921;0.34;0;7.11)
```

### **Unique Start Position**

The following is an example of the output of a Unique Start Position report: # Starting Points within Placed Tags Number of Starting Points in Uniquely placed tag 109,068,047 (2.005740% of reference) Average Number of Uniquely Mapped reads per Start Point 1.992770 Estimate number of starting point for all mapped tags 152,843,627 (2.810765% of reference)

### Coverage files

Example contents for the file coverage\_chr1\_positive.wig are:

```
browser position chr1:1-20000000
browser hide all
browser pack refGene encodeRegions
# minimumMapg=25, minimumCoverage=1,
alignmentFilteringMode=PRIMARY, filterOrpha-nedMates=false,
track name="BAM Coverage positive strand" description="BAM
Coverage positive strand" visibility="full color 0,0,255"
priority=10 yLineMark=0 type=wiggle_0 yLineOnOff=on
variableStep chrom=chr1 span=1
336171
336181
336191
336201
336211
336221
336231
336241
336251
336261
```



336271

Example contents for the file coverage\_chr1\_negative.wig are:

```
browser position chr1:1-20000000
                          browser hide all
                          browser pack refGene encodeRegions
                           # minimumMapg=25, minimumCoverage=1,
                          alignmentFilteringMode=PRIMARY, filterOrpha-nedMates=false,
                           track name="BAM Coverage negative strand" description="BAM
                          Coverage negative strand" visibility="full color 0,0,255"
                          priority=10 yLineMark=0 type=wiggle_0 yLineOnOff=on
                          variableStep chrom=chr1 span=1
                          57432
                          57442
                          57452
                          57463
                          57473
                          57483
                          57493
                          57503
                          57513
                          57523
                          57533
                          •••
Run BAMStats
                      This section describes requirements to generate mapping statistics outside of the
                      context of the mapping module. These requirements are:
                        • Directory structure – The input directory for BAMStats module must mimic the
                          output directory structure of mapping:
                                Input.BAMStat.dir/
                                    <sampleName1>/*.bam, *.bai
                                    <sampleName2>/*.bam, *.bai
                        • Index files – For every BAM file, there must be a corresponding BAI-formatted
                          indexed file. BAI files can be generated using the following command:
                               samtools index <bamfile>
                          This command generates index sorted alignment for fast random access. The
                          index file < bamfile>.bai is created. For details, please refer to the samtools
                          site:
                          http://samtools.sourceforge.net/samtools.shtml
                        • Write permission – The user must also have write permission for the input
                          directory, because the BAMStats module generates position and probe error files
                          in that directory.
```

standalone



1

# FAQ - Small RNA mapping

#### How does the local alignment approach affect mapping?

Using the local alignment approach removes the constraint of a whole-read alignment, and mapping rate is significantly increased. It is not uncommon for a poor dataset with a 30% mapping rate at 50\_6, to reach more than a 60% mapping rate with the mapping algorithm used in LifeScope<sup>TM</sup> Software.

While the majority of alignments are full length, some alignments can vary in length. If a read has many errors towards the end, the final alignment will not include these positions if a shorter alignment receives a better overall score.

#### How do I increase mapping rate?

Mapping rate increases with each additional round of mapping. However, the gain in rate comes at the cost of increased runtime and disk requirement. Testing has found sets of mapping schemes that strike a good balance between mapping rate and runtime for 25-, 35,- and 50-bp reads. These schemes are shipped as LifeScope<sup>™</sup> Software defaults and should work well for most purposes.

For 50-bp reads, two mapping parameters keys define how many rounds of mapping are run, and what happens in each round. The default values of the keys are shown below, and their meanings are further explained in Table 9 on page 76.

Note: Repetitive schemes are empty by default.

- mapping.scheme.unmapped.50 = 25.2.0,25.2.15
- mapping.scheme.repetitive.50 = 38.3.0,25.2.0

The repetitive scheme does not affect the number of mapped reads, and setting it only increases the chance of finding better alignments for repetitive reads. If your main goal is to improve throughput, first try to add more iterations at the unmapped scheme. For example, add another step of 25.2.25 at the end of the unmapped scheme above.

### 3

#### What is a good seed for my application?

Consider the following factors when picking seed parameters:

- Mapping is slower when more mismatches are allowed in the seed. However, allowing more mismatches improves the mapping rate.
- Shorter seeds have higher mapping rates, but also lead to more spurious alignments.



• Color-calls at the beginning of the read are more reliable than those at the end. Therefore, you have the option to anchor the seed near the front of the read. However, applications such as transcriptome sequencing are scenarios where it is an advantage to anchor the seed near the end of the read.

For 50-bp reads, the default setting of 25.2.0 for the first round, followed by 25.2.15 in the second round, delivers good results in most cases. For a single round, 30.3.0 is recommended.

# 4

#### What reads scenario achieves the best accuracy?

The best performance is achieved by considering 2BE and 4BE reads, their alignment, and reference. Typically, base-space reads generated by ECC have lower mapping throughput. Base-space reads generated from combined 2BE and 4BE calls and no reference have lower accuracy, compared to ECC calls.

### 5

#### How do I interpret a mapping quality value?

First of all, it is worth noting that the mapping quality value (QV) is not like the pvalue in BLAST. In BLAST, the p-value is used to estimate the likelihood that the aligned sequences are related. In general, reads come from the same source and it is known that the two are related.

The purpose of mapping QV is to estimate the probability that the read originates from the mapped genomic location. The estimate is determined mainly by the difference in significance between the best- and second-best hits.

The numeric interpretation of mapping QV is the same as the base call QV. A mapping QV of 10 means that there is a 90% chance that the alignment is correct. A mapping QV of 20 means that there is a 99% chance that the alignment is correct.

### 6

#### How much RAM do I need to analyze human samples?

For optimal performance, 24 GB of RAM per cluster node is recommended for human samples. If the cluster node has less than 24 GB of available RAM, mapreads can split the genome into smaller segments, and align to each segment sequentially. Splitting the genome into smaller segments is implemented inside mapreads and is completely transparent to the user.

Mapping with less than the recommended amount of RAM slows down performance. If the cluster node has 16 GB of RAM, mapping human samples will be roughly 30% slower compared to a machine with 24 GB of RAM.

7

# In the mapping statistics genome coverage calculation report, how are Ns in the reference counted?

Ns in the reference are counted as missing coverage. For example, in the human genome, about 7% of the reference sequence consists of Ns. This means that the coverage calculation will never be reported as higher than 93%. If the frequency distribution says 7.49% of the genome is uncovered, this means that about 0.34% of the non-N reference sequence does not have reads mapped to it, and that also the 7.15% of the genome that consists of N does not have reads mapped to it.

#### What should I do to ensure balanced allele ratios at heterozygous positions?

The base-translation algorithm, refcor, uses instrument color-space data (ECC or non-ECC) and the reference sequence to produce the most likely base calls, along with a Phred-scale quality value. The degree to which the reference influences the base call is controlled by the refcor.reference.weight parameter. A position is converted to reference only if the refcor.reference.weight setting is higher than the difference between the quality values of a correct color call and of an adjacent erroneous color call that supports reference.

The refcor.reference.weight default setting, 8, is chosen to be low enough to ensure balanced allele ratios, and also be large enough both to distinguish correct from incorrect color calls and to maximize base accuracy.

In the event an incorrect reference base call is made, the disagreement between evidence sources causes that call's QV to be low. Incorrect calls can be filtered out based on QV threshold, and this filtering improves allele ratios.

Another way of filtering low quality base-calls is to adjust the parameter refcor.base.filter.qv, that converts all base calls with a quality value less than refcor.base.filter.qv into Ns. This filtering has the effect of setting QV to zero in the calls with low confidence. To decrease the number of N base calls, reduce this threshold (for example, from the default of 10 to 5).

**Note:** Setting the refcor.reference.weight parameter to zero completely eliminates the reference as a guide for base-translation (completely eliminates reference bias), but also considerably reduces the quality of base-translation.

#### Can I run BAMStats (mapping statistics) on filtered BAM files?

Yes. You can run BAMStats on the filtered BAM files using the command shell. This scenario is not supported in the LifeScope<sup>™</sup> Software UI.

These instructions describe creating your own version of the small RNA workflow, and modifying your version to include BAMStats on the filtered BAM file.



- 1. Follow the instructions in "Create a new workflow" on page 106 to create a workflow based on the small.rna standard workflow.
- 2. Create a new INI file named filtermapping.bamstats.ini, with these contents:

import global.ini

##Run parameter

bamstats.run=1

##Bin size for insert range distribution.
#bamstats.bin.isize=100

##The input directory for BamStats. There should be one directory per sample containg the BAM files for that sample. bamstats.input.dir=\${analysis.output.dir}/bam/filtered

##Maximum Coverage of a locations in the reference. Locations
with coverage more than the maximum coverage values will be
ignored during coverage calculations.
#bamstats.maximum.coverage=10000

##Maximum inset size for LMP and PE libraries. Reads with insert size more then the specified value will be ignored for Insert Range Report calculation. #bamstats.maximum.isize=100000

##The path to the output directory where BamStats will write its
.cht files.
#bamstats.output.dir= \${task.output.dir}

##Bin size for Coverage in WIG file format
#bamstats.wig.binsize=100

##Whether to combine data from both the strands for coverage in WIG format #bamstats.wig.combined.report.both.strands=0

##Use only primary alignments for Coverage in WIG file format. #bamstats.wig.primary.only=1

analysis.genome.reference = \${analysis.filter.reference}

**3.** Edit the file secondary/secondary.pln in your workflow to add the filtermapping.bamstats step. The contents of the revised PLN file are:

smallrnamapping.ini
bamstats.ini < smallrnamapping.ini</pre>

This PLN file runs the BAMStats module on filtered BAM files also as part of workflow.





# Run a Small RNA Coverage Analysis

This chapter covers:

Overview	423
Example of running the small RNA coverage module	423
Small RNA coverage input files	424
Small RNA coverage parameters	424
Small RNA coverage output files	425

### **Overview**

The small RNA coverage module calculates read coverage per position.

# Example of running the small RNA coverage module

The small.rna standard workflow provides an example of how to run the small RNA coverage module. Here are example LifeScope<sup>™</sup> Software shell commands for this workflow, which includes small RNA mapping and small RNA counts:

```
lscope.sh shell -u username -w password
cd projects
mk mirna
cd mirna
mk run1
cd run1
set workflow small.rna
add xsq rna.xsq
set reference hg18
# optionally change parameter defaults here
run
ls
exit
```

In order to change a parameter value in your analysis, use the set param shell command after the line # *optionally change parameter defaults here* in the example commands. For instance, to change the RNA.coverage.min.value parameter, use this shell command:

set RNA.coverage.min.value 0 /secondary/srCoverage.ini



# Small RNA coverage input files

The small RNA coverage module takes as input one or more BAM files containing mapped data. The following describe the alignment input accepted. This module:

- Accepts only fragment data for RNA libraries.
- Accepts multiple BAM files as input.
- Accepts BAM files with different read lengths.
- Accepts both color space and base-space files, and a combination of color and base-space data.

## Small RNA coverage parameters

Table 116 lists the small RNA coverage parameters.

Parameter name	Default value	Description
Optional parameters		
RNA.coverage.min.quality	2	The mapping quality value threshold for selecting the alignments. Alignments with a mapping quality value lower than this threshold are not written to output. Allowed values: Integers 0–100.
RNA.coverage.min.value	0	The mapping quality threshold value for selecting alignments from the input BAM. In order to be included in the output, a position's coverage must be greater than or equal to this value. Allowed values: Integers >= 0.
RNA.coverage.per.chromosome	true	Whether coverage output is generated as one file per every chromosome per strand, or as a single file with coverage of all chromosomes per strand. Allowed values:
		• <b>false</b> : Generate a single file with coverage of all chromosomes per strand.
		• <b>true</b> : Generate one file per every chromosome per strand.
RNA.coverage.primary.only	true	Whether only primary alignments or all the alignments from the input BAM file are used as input. Allowed values:
		false: Consider all alignments.
		<ul> <li>true: Consider only primary alignments (both gapped and ungapped).</li> </ul>
Resource parameters		
memory.request	4gb	Memory request. Include the units gb in the setting.
java.heap.space	4000	Dynamic memory requirement.

Table 116 Small RNA coverage parameter description



# Table 117 lists the small RNA coverage parameters that we do not recommend changing.

Table 117	Small RNA	coverage	internal	parameter	description
	JINULL INIA	coverage	micrinat	purumeter	ucscription

Parameter name	Default value	Description
coverage.run	1	Whether or not to run the small RNA coverage module. Allowed values:
		• 0: Do not run the coverage module.
		• 1: Run the coverage module during this analysis.
		The run parameter is set automatically by the shell during a standard workflow. Accept the default for most use.
analysis.input.bams	_	A comma-separated list of paths to the input BAM files. Relative and absolute paths are accepted.
		Generated automatically by LifeScope <sup>™</sup> Software during a workflow.

### Small RNA coverage output files

The small RNA coverage module's output files are in the wiggle (\*.wig) file format. Wiggle files are visualized in genomics browsers such as the Integrative Genomics Viewer (IGV). You can download the IGV browser from the Broad Institute website:

#### www.broadinstitute.org/igv

The wiggle file specification is available from this site:

#### http://genome.ucsc.edu/goldenPath/help/wiggle.html

If the parameter RNA.coverage.per.chromosome is set to false, the small RNA coverage module generates two output files with coverage of all chromosomes per strand:

- coverage\_positive.wig
- coverage\_negative.wig

If the parameter RNA.coverage.per.chromosome is set to true, the small RNA coverage module generates one coverage file per chromosome per strand:

- coverage\_chr\*\*\_positive.wig
- coverage\_chr\*\*\_negative.wig





# Run a Small RNA Counts Analysis

This chapter covers:

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Small RNA counts	428
Small RNA counts input files	428
Small RNA counts parameters	429
Small RNA counts output files	431

### **Overview**

This module generates tag counts for precursor and mature miRNA sequences. Run this analysis module to generate miRNA expression profiles.

**IMPORTANT!** Alignments must come from the same strand as the feature to contribute to the count. A non-gapped tag contributes to a feature's count if it overlaps the feature and has no more than three bases outside the feature. A gapped tag contributes to a feature's count if one of its match regions terminates at a feature boundary.

## Example of running the small RNA count module

The small.rna standard workflow provides an example of how to run the small RNA counts module. Here are example LifeScope<sup>™</sup> Software shell commands for this workflow, which includes small RNA mapping and small RNA coverage:

# log into the shell
lscope.sh shell -u username -w password
# cd to the projects repository
cd /projects
# create a project and open it
mk mirna
cd mirna
# create an analysis and open it



```
mk run1
cd run1
# define the analysis type
set workflow small.rna
# define the input data
add xsg rna.xsg
add xsg rna2.xsg
# specify the reference
set reference hg18
# optionally change parameter defaults here
# list the analysis configuration
ls
# start the analysis
run
# list progress information about the run
ls
# exit the command shell
exit
```

Dummy XSQ file name are given in the example. In order to change a parameter value in your analysis, use the set param shell command after the line # *optionally* change parameter defaults here in the example commands. For instance, to change the parameter SmallRNA.counts.min.quality, use this shell command:

set SmallRNA.counts.min.quality 0 /secondary/srCount.ini

You must know which INI file to use for the set param command. The ls command, executed in an analysis virtual directory after the workflow has been defined, lists the INI files used in the analysis.

### Small RNA counts

The LifeScope<sup>™</sup> Software small RNA counts module calculates counts per precursor or mature sequence, where counts are the number of reads mapped. This module is similar to the WT counts module, which computes counts per exon.

This module requires an annotation features file, which contains either precursor sequences information in a GFF file, or mature sequences information in a text file. The counts module uses the annotation features file and alignments in the form of one or more BAM files, and computes the number of alignments that match every feature. Non-primary and low quality alignments can be filtered by setting parameters appropriately.

## Small RNA counts input files

The small RNA mapping module generates two BAM files containing alignments in coordinate order. These two BAM files are input to the small RNA counts module for separate counts processing:

• The filtered BAM file, from the filter mapping step. The small RNA counts module uses this file to provide counts for the number of reads mapped to the sequences in the filter reference. See "Filtered BAM file parameters" on page 430 and "Filtered counts output" on page 433.



	• One or more mapped BAM files with genomic locations. The module uses these files as input with the mapped BAM files:
	<ul> <li>A Genome Coordinates GFF file containing precursor sequences with genomic locations.</li> </ul>
	<ul> <li>(Optional) A file containing mature form sequences with locations relative to precursor sequences. If the mature sequences file is not provided as input, the module computes counts per precursor sequence.</li> </ul>
	These files are described in the following sections.
Alignments files	The following describe the alignment input accepted by the small RNA counts module. This module:
	Accepts only fragment data for RNA libraries.
	Accepts multiple BAM files as input.
	<ul> <li>Accepts BAM files with different read lengths.</li> </ul>
	• Accepts both color space and base space files, and a combination of color-space and base-space data.
Precursor sequences	This file is a Genome Coordinates GFF file containing precursor sequences with genomic locations. You must use the same file that was used during the small RNA mapping step.
	This file is included in the LifeScope <sup>™</sup> Software reference repository and is also available the following link:
	http://www.mirbase.org/ftp
Mature form sequences	The mature form sequences file provides locations relative to precursor sequences. This tab-separated file is included in the LifeScope <sup>™</sup> Software reference repository and is also available the following link:
	ftp://mirbase.org/pub/mirbase/CURRENT/miRNA.xls
	If you download a new file, you must save the file locally in tab-delimited format. Only this specific mature forms file is supported.
	If the mature sequences file is not provided as input, the module computes counts per precursor sequence.

# Small RNA counts parameters

Mapped file	Table 118 lists the small RNA counts parameters used to process the mapped BAM file.
parameters	

Table 118 Small RNA counts parameter description

Parameter name	Default value	Description
SmallRNA.counts.min.quality	2	The mapping quality value threshold for selecting the alignments. Alignments with a mapping quality value lower than this threshold are not written to output. Allowed values: Integers 0–100.



Table 118	Small RNA	counts	parameter	description	(continued)
	Sindle Internet	counts	purumeter	acocription	(continucu)

Parameter name	Default value	Description
SmallRNA.counts.overflow.limit	3	The maximum allowed offset on either side of an alignment or feature, when looking for overlap between an alignment and a feature. If this limit is set to 0, then the start and end coordinates of the alignment record and feature must match exactly. Allowed values: Integers 0–20.
SmallRNA.counts.per.feature	true	Controls the aggregated count output. Allowed values:
		• <b>false</b> : The aggregated count of all the reads mapped to a feature are output on the same line.
		• <b>true</b> : Reads of different starts and ends are counted separately for every feature and are output on separate lines.
		The primary key in the output GTF file is: {Feature, start, end}.
SmallRNA.counts.primary.only	true	Whether only primary alignments or all the alignments from the input BAM file are used as input. Allowed values:
		• false: Consider all alignments.
		<ul> <li>true: Consider only primary alignments (both gapped and ungapped).</li> </ul>
Resource parameters		
memory.request	4gb	Memory request. Include the units gb in the setting.
java.heap.space	3000	Dynamic memory requirement.

Table 119 lists the small RNA counts parameters that we do not recommend changing.

Table 119	Small RNA	counts	internal	parameter	description
-----------	-----------	--------	----------	-----------	-------------

Parameter name	Default value	Description
counts.run	1	Whether or not to run the small RNA counts module. Allowed values:
		• 0: Do not run the small RNA counts module.
		• 1: Run the small RNA counts module during this analysis.
		The run parameter is set automatically by the shell during a standard workflow. Accept the default for most use.
SmallRNA.counts.mismatches	3	The maximum number of mismatches used when computing statistics. Allowed values: Integers 0–10.
SmallRNA.matureForms.file	—	Path to the tab-separated file with mature form sequences in every precursor sequence. Relative and absolute paths are supported.
SmallRNA.precursor.gff.file	—	Path to the input GFF file with genomic coordinates of the precursor sequences. Relative and absolute paths are supported.

### Filtered BAM file Tab parameters

Table 120 lists the small RNA counts parameters related to the filtered BAM file.



	1	
Parameter name	Default value	Description
filtermapping.counts.run	1	Whether or not to run the small RNA counts on the filtered BAM file. Allowed values:
		• 0: Do not run small RNA counts on the filtered BAM file.
		• 1: Run small RNA counts on the filtered BAM file.
		The run parameter is set automatically by the shell during a standard workflow.
analysis.filter.reference	_	The path to the filter reference file. This parameter is set automatically by the shell when you use the set reference <i>assembly</i> command.
SmallRNA.counts.primary.only	true	Whether only primary alignments or all the alignments from the input BAM file are used as input. Allowed values:
		• false: Consider all alignments.
		<ul> <li>true: Consider only primary alignments (both gapped and ungapped).</li> </ul>

Table 120 Small RNA filter counts parameter description

### Small RNA counts output files

Mapped outputThis section describes the output files the small RNA counts module creates from the<br/>mapped BAM input.

#### GTF files

The small RNA counts module creates a GTF file for precursor counts and one for mature counts (if mature parameter is provided). Each input GFF file contains the counts per feature in column 5. Table 121 describes the counts file fields. The names of these files are:

- precursor\_counts.gtf
- matureForms\_counts.gtf

The following is example content from a counts.gtf file:

```
chr1 . mirBase 3044539 3044599 0 - . Precursor_ID
"hsa-mir-4251"; RPM "0.00";
```

	Table 121	Small RNA	counts GTF	output file	format	description
--	-----------	-----------	------------	-------------	--------	-------------

Column name	Description	Examples
seqname	The name of the sequence. Must be a chromosome or a scaffold.	chr1
source	The program that generated this feature.	-
feature	The name of this type of feature.	*miRNA



Column name	Description	Examples
start	The starting position of the feature in the sequence. The first base is numbered 1.	3044539
end	The ending position of the feature (inclusive).	3044599
score	The tag count for the feature.	0
strand	<ul> <li>The strand containing the feature. Values:</li> <li>'+': The feature is on the 3' strand.</li> <li>'-': The feature is on the 5' strand.</li> <li>'.': Unknown or not required.</li> </ul>	-
frame	(Not applicable to miRNAs.)	-
attributes	A list of attributes delimited by semi-colons. Each attribute is a type/value pair, with type and value separated by a single space character. Type is a string matching the pattern [A-Za-z1-9_]+. Value is a number or a double-quoted string. Every attribute must end with a semi-colon.	ACC="MI000000": ID "hsa-mir-4251";

#### Table 121 Small RNA counts GTF output file format description (continued)

#### contig\_counts.txt file

This section describes the output file, contig\_counts.txt, generated from the mapped BAM file. This output file is a tab-separated file listing the following for each the sequence id in the filter reference:

- The sequence id
- The number of reads on the positive strand that map to that sequence
- The number of reads on the negative strand that map to that sequence
- The size of the reference sequence

The following is an example of a few lines from the mapped contig\_counts.txt file:

ID	Counts_	Pos_St	rand	Counts_Neg_Strand	d RefSeq_size
chr20	2932	1	630	025520	
chr21	1312	0	483	129895	
chr22	159	2	513	304566	

#### Mismatches stats files

The small RNA module generates two stats files based on the mapped BAM input:

- precursor\_mismatch\_stats.txt
- matureForms\_mismatch\_stats.txt

These files list the sequences from either the input precursor and or the mature forms file, and provide the number of alignments that map to each sequence with 0, 1, 2, and 3 mismatches. The total number of alignments that map to the sequence is also provided. The following is an example of a few lines from a mismatch\_stats.txt file:

	0MM	1MM	2MM	3MM	Total
--	-----	-----	-----	-----	-------


hsa-let-7a	48( 64.00%)	15( 20.00%)	3( 4.00%)	5( 6.67%)	75(100.00%)
hsa-let-7a*	1( 16.67%)	0( 0.00%)	4(66.67%)	0( 0.00%)	6(100.00%)
hsa-let-7a-2*	0( 0.00%)	0( 0.00%)	0( 0.00%)	0( 0.00%)	0( -%)
hsa-let-7b	17( 33.33%)	7(13.73%)	16( 31.37%)	7( 13.73%)	51(100.00%)
hsa-let-7b*	1( 33.33%)	0( 0.00%)	1( 33.33%)	1(33.33%)	3(100.00%)

# Filtered countsThis section describes the output file, contig\_counts.txt, generated from the<br/>filtered BAM file. This output file is tab-separated and lists the following for each the<br/>sequence id in the filter reference:

- The sequence id
- The number of reads on the positive strand that map to that sequence
- The number of reads on the negative strand that map to that sequence
- The size of the reference sequence

The following is an example of a few lines from the filtered contig\_counts.txt file:

ID Counts_Pos_Strand Counts_Neg_Strand RefS	eq_size
3000072055953=Tigger4a#DNA/MER2_type 0 0 236	
chr16.trna19-GlyGCC 69 0 71	
chr17.trna14-ThrCGT 2 0 72	
3000072056044=tRNA-Tyr-TAC#tRNA 0 0 76	
chr2.trna27-GlyCCC 128 0 71	



**Chapter 23** Run a Small RNA Counts Analysis *Small RNA counts output files* 

# PART VII Additional Analyses



# Run a ChIP-Seq Mapping Analysis

This chapter covers:

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Mapping algorithm	440
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## **Overview**

LifeScope<sup>™</sup> Genomic Analysis Software provides the ability to map data and create an output file type compatible with a variety of third-party Chromatin Immunoprecipitation Sequencing (ChIP-Seq) data analysis tools. The ChIP-Seq application has publicly available analysis software that can be used withLifeScope<sup>™</sup> Software output.

The ChIP assay is a method for analyzing epigenetic modifications and genomic DNA sequences bound to specific regulatory proteins. ChIP-Seq is a combined assay and sequencing technique for identifying and characterizing elements in protein-DNA interactions. It typically examines transcription factors (TF) bound to DNA and finds DNA sequence motifs common to binding sites.

Using the MAGnify<sup>TM</sup> ChIP-Seq kit with the SOLiD<sup>TM</sup> sequencing system enables you to generate sequence read data from a ChIP-Seq experimental approach. LifeScope<sup>TM</sup> Software gives you the option to map the read data.

## **Run ChIP-Seq mapping**

ChIP-Seq analysis is available as workflow in the LifeScope<sup>™</sup> Software command shell.

Run in the lscope<br/>command shellThis section describes how to run the fragment ChIP-Seq mapping workflow in the<br/>LifeScope™ Software command shell.



In the LifeScope<sup>™</sup> Software command shell, running the ChIP-Seq workflow requires the following steps:

- Create a LifeScope<sup>TM</sup> Software command shell project and analysis.
- Identify your input data (your reads).
- Identify the reference genome.
- Select the ChIP-Seq the workflow as the analysis to be executed on your data.
- Issue the run command to start your analysis.

#### Example steps to run ChIP-Seq mapping as standard workflow

This section shows the steps to run the ChIP-Seq workflow in the LifeScope<sup>™</sup> Software command shell. A description of each step is given in Table 122.

```
# log into the shell
lscope.sh shell -u username -w password
# import your data into the reads repository
cd /reads
import /data/xsg/xsg1.xsg
# cd to the projects repository
cd /projects
# make a project, and open it
cd /projects
mk chip_seq
cd chip_seq
# make a analysis, and open it
mk run1
cd run1
# define the analysis type (the ChIP-Seq workflow)
set workflow chip.seq
# define the input
add xsg xsq1.xsg lung
# specify the reference to be used
set reference /data/results/references/DH10B.fasta
# list the analysis configuration
ls
# run the analysis
run
# display progress information on your run
ls
```

Dummy XSQ file names are used in the example.

#### Information about your workflow job

Output of the 1s command displays the analysis' parameters, INI files, and run completion percentage or completion status.

Table 122 explains the purpose of the steps shown above.

 Table 122
 Description of sample shell commands to run the ChIP-Seq workflow

Command	Purpose	
lscope.sh shell -u <i>username</i>	Log into the LifeScope <sup>™</sup> Software command shell.	



	•		
Command		Purpose	
cd /reads import /data/xsq/DH10B_ColorSpaceOnly_ Unbarcoded_50_Frag.xsq		Go into the reads repository and import an XSQ file (the input data file). This command in effect brings the reads file into the shell reads repository.	
		Once in the reads repository, the reads file is available as input to your analysis runs.	
cd /projects mk chip_seq cd chip_seq		In the projects repository, create a project named ecoli, and in that project create an analysis named run1. Open the run1 analysis.	
mk run1		Note: A project name cannot include a dash character ("-").	
ca runi		<b>Note:</b> The open analysis (cd run1, in this case) command is important. The data, reference, and other configuration commands which follow apply only to the current analysis.	
set workflow chip.	seq	Define the workflow used with this analysis. The ChIP-Seq workflow includes mapping, mapping statistics, and duplicate bead finding.	
add xsq DH10B_Co ed_50_Frag.xsq	lorSpaceOnly_Unbarcod	Define the XSQ file containing the input reads data for this analysis.	
<pre>set reference /data/results/results/ references/DH10B.fasta</pre>		Define the reference file for this analysis.	
set saet.run 0 secondary/saet.ini		Turns off the SAET module. SAET IS NOT recommended for ChIP-Seq.	
ls		Display the configuration of your analysis, including input reads, reference files, workflow, and select parameter settings.	
run		Start the analysis run.	
ls		Display status information about the analysis run.	
	See Chapter 7, "Run a Standar on running workflows, includ	d Workflow Analysis" on page 97 for more information ing the ChIP-Seq workflow.	
Do not rename	Do not delete or rename any c	of the following directories or files for your workflow:	
	• The analysis.pln file		
	The secondary director	У	
	<ul> <li>The tertiary directory</li> <li>The second area with fill</li> </ul>		
	• The secondary.pin me		
Turn off SAET	We recommend not running S SAET in a workflow, make su	EAT with the ChIP-Seq workflow. When you turn off re your mapping INI file does not contain the line:	
	analysis.input.reads	et.file = \${analysis.output.dir}/saet/*.rrs	
	Comment out this line if it app	pears in your mapping INI file.	

Table 122 Description of sample shell commands to run the ChIP-Seq workflow (continued)



## Run ChIP-Seq as an individual analysis

The optional examples download includes an example of how to run the ChIP-Seq module by itself, as an individual analysis that is not part of a standard workflow.

See Appendix E, "Demo Analyses" on page 507 for information on the examples.

## Mapping algorithm

The ChIP-Seq module uses the resequencing mapping and pairing algorithm. See the following sections for more information:

- "Stages of mapping" on page 115
- "Mapping algorithm" on page 126
- "Mapping statistics" on page 141
- "Mapping output files" on page 140

## Use results files

After the mapping steps are complete, the resulting BAM file can be used with compatible third-party commercial and academic ChIP-Seq analysis software tools (see Figure 44 on page 441).

As of this writing, you can download a BAM-to-BED format converter from thirdparty tools sites, for example:

code.google.com/p/bedtools

Chapter 24 Run a ChIP-Seq Mapping Analysis Use results files





Figure 44 Compatible ChIP-Seq analysis software tools





# Run a MethylMiner<sup>™</sup> Mapping Analysis

This chapter covers:

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## **Overview**

DNA methylation is an epigenetic modification crucial for organism development and normal gene regulation. Life Technologies have introduced a versatile methyl-CpG binding protein-based system, the MethylMiner<sup>™</sup> Kit, for the enrichment of methylated sequences from genomic DNA, that, with the use of SOLiD<sup>™</sup> System sequencing, allows for focused evaluation of methylation patterns in genome-wide studies. The enriched reads can also be bisulfite-converted, which would additionally allow determination of methylation status of individual cytosines.

SOLiD<sup>™</sup> System sequencing can also, of course, be used to study methylation in unenriched, whole-genome bisulfite-converted read data.

The purpose of LifeScope<sup>TM</sup> Genomic Analysis Software MethylMiner<sup>TM</sup> module is to provide a data analysis pipeline for mapping and analyzing MethylMiner<sup>TM</sup> enriched and unenriched fractions of genomic DNA as well as bisulfite-converted and unconverted reads sequenced on the SOLiD<sup>TM</sup> System.

The MethylMiner<sup>™</sup> analysis module functionality currently includes:

- Mapping of unconverted and bisulfite-converted reads.
- Mapping statistics and statistics on read coverage and depth.
- Mapped reads output in BAM-format files.
- Visualization of mapped reads on publicly available genome browsers.



Figure 45 shows the MethylMiner<sup>™</sup> analysis workflow with SOLiD<sup>™</sup> software and third-party browsers.



Figure 45 Methylation data analysis module overview

## Run MethylMiner<sup>™</sup> mapping

Unconverted MethylMiner<sup>™</sup> reads must be mapped to a regular (unconverted) reference genome sequence. Bisulfite-converted reads must be mapped to a pair of appropriately converted reference sequences (forward and reverse conversions), as recommended below.

Follow these recommendations for MethylMiner<sup>™</sup> mapping:

- MethylMiner<sup>™</sup> supports fragment and paired-end libraries. Do not use this module with mate-pair libraries.
- For mapping bisulfite reads, the converted reference sequence pairs below are recommended. Use one of the following:
  - Pair 1:

Reference with all non-CpG C's converted to T's

Reference with all non-CpG G's converted to A's

– Pair 2:

Reference with all C's converted to T's

Reference with all G's converted to A's

MethylMiner<sup>™</sup> analysis is available as two standard workflows in the LifeScope<sup>™</sup> Software command shell:

- For fragment data: methyl.miner.frag
- For paired-end data: methyl.miner.pe



Run in the lscope<br/>command shellThis section describes how to run the fragment MethylMiner<sup>TM</sup> mapping workflow in<br/>the LifeScope<br/>TM Software command shell.

In the LifeScope<sup>™</sup> Software command shell, running the MethylMiner<sup>™</sup> workflow requires the following steps:

- Create a LifeScope<sup>™</sup> Software command shell project and analysis.
- Identify your input data (your reads).
- Identify the reference genome.
- Specify the MethylMiner<sup>™</sup> fragment workflow as the analysis type to be executed on your data.
- Issue the run command to start your analysis.

#### Example steps to run the MethylMiner<sup>™</sup> fragment mapping standard workflow

This section shows the steps to run the MethylMiner<sup>™</sup> fragment mapping workflow in the LifeScope<sup>™</sup> Software command shell. A description of each step is given in Table 123.

lscope.sh shell -u username -w password cd /reads import /data/xsq/DH10B\_ColorSpaceOnly\_Unbarcoded\_50\_Frag.xsq cd /projects mk methyl cd methyl mk run1 cd run1 set workflow methyl.miner.frag add xsq DH10B\_ColorSpaceOnly\_Unbarcoded\_50\_Frag.xsq lung set reference /data/results/references/DH10B.fasta set saet.run 0 secondary/saet.ini ls run ls

Dummy XSQ file names are used in the example.

Information about your workflow job

Output of the 1s command displays the analysis' parameters, INI files, and run completion percentage or completion status.

Table 123 explains the purpose of the steps shown above.

Table 123 Description of sample shell commands to run the MethylMiner<sup>™</sup> workflow

Command	Purpose
lscope.sh shell -u username	Log into the LifeScope <sup>™</sup> Software command shell.
cd /reads import /data/xsq/DH10B_ColorSpaceOnly_ Unbarcoded_50_Frag.xsq	Go into the reads repository and import an XSQ file (the input data file). This command in effect brings the reads file into the shell reads repository.
	Once in the reads repository, the reads file is available as input to your analysis runs.



Command	Purpose
cd /projects mk methyl cd methyl	In the projects repository, create a project named ecoli, and in that project create an analysis named run1. Open the run1 analysis.
mk run1 cd run1	<b>Note:</b> The open analysis (cd run1, in this case) command is important. The data, reference, and other configuration commands which follow apply only to the current analysis.
set workflow methyl.miner.frag	Define the workflow used with this analysis. The MethylMiner <sup>™</sup> workflow includes SAET, mapping, mapping statistics, and duplicate bead finding.
add xsq DH10B_ColorSpaceOnly_Unbarcod ed_50_Frag.xsq	Define the XSQ file containing the input reads data for this analysis.
set reference /data/results/results/ references/DH10B.fasta	Define the reference file for this analysis.
<pre>set saet.run 0 secondary/saet.ini</pre>	Turns off the SAET module. SAET IS NOT recommended for the MethylMiner <sup>™</sup> workflow.
ls	Display the configuration of your analysis, including input reads, reference files, workflow, and select parameter settings.
run	Start the analysis run.
ls	Display status information about the analysis run.

Table 123 Description of sample shell commands to run the MethylMiner<sup>™</sup> workflow (continued)

See Chapter 7, "Run a Standard Workflow Analysis" on page 97 for more information on running workflows, including the MethylMiner<sup>™</sup> workflows.

**Do not rename** When using a standard workflow, do not delete or rename any of the following directories or files:

- The analysis.pln file
- The secondary directory
- The tertiary directory
- The secondary.pln file

Visualize MethylMiner <sup>™</sup> results	After the mapping steps are complete, the resulting BAM file with mapped reads can be visualized with compatible third-party commercial software tools and publicly-available genome browsers (see Figure 45 on page 444).
	To visualize the mapped reads, launch your UCSC genome browser or IGV browser to import and view the mapped read BAM file generated by MethylMiner <sup>™</sup> mapping.

## Run MethylMiner<sup>™</sup> as an individual analysis

The optional examples download includes an example of how to run the MethylMiner<sup>™</sup> module by itself, as an individual analysis that is not part of a standard workflow.



See Appendix E, Demo Analyses for information on the examples.

## MethylMiner<sup>™</sup> mapping output files

Mapped reads are output in BAM files.

The output mapping statistics reports generated by MethylMiner<sup>™</sup> mapping include:

- The number and percent of all reads mapped and of reads mapped uniquely, in the file mapping-stats.txt and also in files ending in F3.stats and F5-P2.stats.
- Genome coverage tables for at least read depths 0X to 50X, in output files ending in coverage-histogram-F3.txt and coverage-histogram-F5-P2.txt.

## Mapping algorithm

The MethylMiner<sup>™</sup> module uses the resequencing mapping and pairing algorithm. See the following sections for more information:

- "Stages of mapping" on page 115
- "Mapping algorithm" on page 126
- "Mapping statistics" on page 141
- "Mapping output files" on page 140

## Further analysis of MethylMiner<sup>™</sup> mapping results

Besides viewing mapped reads in genome browsers, you can further analyze mapped reads outside of LifeScope<sup>TM</sup> Software with software available in the SOLiD<sup>TM</sup> development community or with other third-party tools.

MethylMiner<sup>™</sup> unconverted mapped reads can be processed with peak-finding programs like MACS to identify genome regions of significant methylation.

Similarly, MethylMiner<sup>™</sup> bisulfite-converted mapped reads can be processed with peak-finding programs to identify regions of significant methylation. These reads can also be processed at nucleotide resolution to report the methylation status of individual C bases, for bases covered at sufficient read depth.



# PART VIII Appendices

# FAQ

# A

This appendix covers:

- FAQ lists for LifeScope<sup>TM</sup> Genomic Analysis Software modules ...... 451

## FAQ lists for LifeScope<sup>™</sup> Genomic Analysis Software modules

Frequently-asked questions for modules are available in the analysis module chapters. See the following sections for module-specific FAQ lists:

- "FAQ Mapping"
- "FAQ Pairing"
- "FAQ Human CNVs"
- "FAQ SNPs"
- "FAQ Small indels"
- "FAQ Large indels"
- "FAQ Whole transcriptome"

## General FAQ for LifeScope<sup>™</sup> Software

#### 1

#### What is the performance increase vs. RAM increase?

For the human genome, 24GB of RAM per node provides approximately 30% to 40% performance improvement over systems that have 16GB of RAM per node.

## 2

#### Does LifeScope<sup>™</sup> Software support hyper-threading?

Yes, some LifeScope<sup>™</sup> Software components can take advantage of hyper-threading if it is present.



## Can I configure LifeScope<sup>™</sup> Software to submit jobs to the queues that are already set up on my cluster?

You can use pre-existing queues. During installation you will be asked which existing queues you would like to use with LifeScope<sup>™</sup> Software.

## 4

#### What is the expected time for my run to complete?

This depends on many factors, like the number of beads, the reference, the hardware of the cluster in use, the traffic condition if it's a shared cluster, the speed of storage system, etc.

In a typical dedicated three to five nodes cluster, with 8 cores per node and 24GB memory per node, for one (1) full slide in a mate-pair run, it takes about two days to do the mapping, pairing, and generate BAM files for 1 billion reads. Most analyses can be done within one or two days.

## 5

#### How do I know if my run is successfully complete?

Use the LifeScope<sup>™</sup> Software command shell 1s command in your analysis, or the 1scope.sh status command at the Linux prompt.

See "Review job status" on page 93 and "Review logs" on page 94.

### 6

#### What should I do if my run never seems to complete?

Verify that your run partition is not full and that the queueing system is working correctly. If your run directory's partition is 100% full, LifeScope<sup>™</sup> Software may report that your job has started, but LifeScope<sup>™</sup> Software is not able to process the job.

#### 7

#### Do you have example data to test my install with?

Ask your LifeScope  $^{\text{TM}}$  Software administrator if the optional Examples directory was installed.

#### 8

Can the "scratch" directory be on a shared NFS storage?



Scratch space can be a local disk or shared storage mounted via NFS, but there will be high I/O activity on the scratch disk. Ensure the storage has high throughput and will not degrade the overall performance of the system. LifeScope<sup>™</sup> Software is not tested in an environment that is configured with scratch space that is mounted via NFS.

#### Can multiple users try to install LifeScope<sup>™</sup> Software in different locations?

No. There is no need to install multiple copies. You just need to set the environment to point to LifeScope<sup>™</sup> Software. See "Set the LifeScope<sup>™</sup> Software environment" on page 46.

## 10

9

Can I continue to run my BioScope<sup>™</sup> Software after I install LifeScope<sup>™</sup> Software?

Yes. You must follow the instructions in "BioScope<sup>™</sup> Software users' PATH variable" on page 46.



Appendix A FAQ General FAQ for LifeScope<sup>™</sup> Software

# **File Format Descriptions**

This appendix covers:

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## Introduction

This appendix describes file formats used with LifeScope<sup>™</sup> Software.

Before reading the section about the SOLiD<sup>TM</sup> System BAM file contents, you should be familiar with the general SAM specification and with the SAM specification field definitions.

Before SOLiD<sup>TM</sup> 4.0, the primary SOLiD<sup>TM</sup> read and alignment format was based on the public GFF specification. The specification defined the location of an aligned read on a reference sequence and provided for arbitrary name-value pair attributes. For the SOLiD<sup>TM</sup> GFF, attributes are used to detail color reads and quality values, base space translations, mate-pair information, and other aspects of the aligned read. Increased throughput of second-generation sequencing technologies resulted in the definition of the SAM file format and a compressed, indexed binary format (BAM) that expands the basic alignment information of a GFF file to include paired-read information and a more structured attribute set. A number of tools exist for basic manipulation of BAM files, and an array of viewers and analysis tools are available.

LifeScope<sup>™</sup> Software secondary analysis (mapping and pairing) now produces a BAM file as the main alignment format. Mate-pair and paired-end analysis directly produces a BAM file, while a single file conversion is needed for fragment libraries. Depending on the output filter selected, unmapped and secondary alignments can be included.

LifeScope<sup>™</sup> Software supports the XSQ sequence data file format introduced with the 5500 Series SOLiD<sup>™</sup> Sequencer.

## XSQ file format

overview

XSQ (eXtensible SeQuence) is an extensible file format for storing sequence data. The XSQ format supports multiple independent reads at the same position in a fragment. XSQ is a binary format based on the open HDF format.

Sequencing run data are automatically exported from the 5500 Series SOLiD<sup>™</sup> Sequencer in XSQ binary file format. If an ECC primer round has been performed, the XSQ output also includes the sequence information in base space, in addition to color space.

#### XSQ file content 5500 Series SOLiD<sup>™</sup> Sequencer data

XSQ files generated by the 5500 Series SOLiD<sup>™</sup> Sequencer contain:

- **Base-space data** Reference-free base-space data from primary analysis (if the optional ECC round is performed).
- Color-space data Referred to as 2+4 color, as follows:
  - The 2BE (Base Encoding) data is the data for first 5 primer rounds of regular 2BE probes. This data is in the same format as a data created by a SOLiD<sup>™</sup> 4 System non-ECC run.
  - The 4BE data is the data for the 6th primer round, the optional ECC round.

Data for optional ECC primer rounds is supported in an XSQ file.

Paired-end libraries do not support ECC data. Fragment libraries support ECC data, and mate-pair libraries support ECC data on both tags.

Trimming is allowed with base-space reads, and can occur at the beginning as well as the end of a read. The trimming is typically an output of an ECC run. The mapping of trimmed reads requires that the untrimmed portion be at least as long as the mapping seed/index length.

#### SOLiD<sup>™</sup> 4 System data

SOLiD<sup>™</sup> 4 System CSFASTA and QUAL files can be used with LifeScope<sup>™</sup> Software, but first must be converted to XSQ format. Converted XSQ files contain only:

- Color-space 2BE data
- Color for both tags (one tag for fragment and both tags for mate-pair and pairedend data).

See Appendix C, "XSQ Tools" on page 479 for information on the XSQ converter.



#### This section describes how physical aspects of the 5500 Series SOLiD<sup>™</sup> Sequencer Relation to instrument affect the output XSQ files. sequencing instrument Each lane of the 5500 instrument results in a single XSQ output file. Different models of the 5500 support from 6 to 12 lanes, so typically multiple XSQ files are generated per sequencing run. You can direct your LifeScope<sup>™</sup> Software analyses to data from specific barcodes run in different lanes, or group the output of multiple lanes or multiple sequencing runs to be processed as a single specimen. The most granularity possible for a LifeScope<sup>™</sup> Software analysis is the data from one barcode in one lane. For information on specifying barcodes and specimen groups, see the following material: The add xsq command in Table 5 "LifeScope™ Software shell commands" • on page 77. • "Run a grouped analysis" on page 88 in Chapter 6, "Run a Command Shell Analysis" on page 71

# XSQ file format<br/>descriptionThis section describes several tables within an XSQ file. The complete XSQ file format<br/>specification is available on the SOLiD<sup>™</sup> Software Community website. At the time of<br/>this writing the specification is this site:

#### http://solidsoftwaretools.com/gf/project/xsq

The LibraryDetails table describes the properties of each library used in the XSQ file. String fields are limited in length to 255 characters.

Column name	Description
LibraryName	Describes the name of one library preparation, which may be used across several indexes in an indexing run.
Application	Describes how this sequence is intended to be analyzed. Allowed values are:
	Whole genome resequencing
	Targeted resequencing
	Whole transcriptome (Fragment)
	Whole transcriptome (PairedEnd)
	• Small RNA
	• ChIP-Seq
	Methylation
ProjectName	Describes the project within which this sample is being run.
Sample0wner	Names the owner of this sample.
SampleIdentifier	A unique name for this sample.
Description	Describes the sample.
Comments	Optional additional information about this sample or its preparation.
ERCC	Whether internal controls have been used. Reserved for future use.

 Table 124
 XSQ library details table

Column name	Description
Species	The intended species for mapping. Example values include:
	Homo sapiens
	Mus musculus
	• other
Assembly	The intended assembly for mapping (may be blank if unknown). Example values include:
	• hg18
	• hg19
	• mm9
	Other valid assembly names are accepted. Spaces are not allowed.
LibraryInsertSize Minimum	The expected minimum insert size in nucleotides. Used with paired-end and mate-pair libraries. Allowed values: Integers >= 0. Set to 0 for fragment libraries, or when the size is unknown.

Table 124	XSQ librar	y details	table	(continued)
-----------	------------	-----------	-------	-------------

Each read type (tag) has a tag-level properties in the XSQ file. Table 125 lists the attributes describing the base-space and color-space data for a single type of read. Table 126 on page 459 lists tag names for current library types.

#### Table 125 XSQ tag details table

Column name	Default	Description
TagSequence	_	The full sequence of the tag used for primer annealing. It is used in conjunction with the read offset to determine the nucleotide or nucleotides for phasing the color-space reads. When the full sequence is not available, the last base(s) of the tag are sufficient as long as they cover the largest offset in the DatasetColorEncoding table. Conditional: Required when color space is present.
IsColorPresent	0	Whether color-space call data is present. Values:
		• <b>0</b> : No calls are reported in color space (causing the ColorCallQV dataset to be missing).
		• Non-zero: Color calls are reported.
IsBasePresent	1	Whether base call data is present. Values:
		• <b>0</b> : No calls are reported in base space (causing the BaseCallQV dataset to be missing).
		• <b>Non-zero</b> : Base calls are reported.
NumBaseCalls	_	The maximum untrimmed read length in base space across all reads for this tag. This length is the width of the BaseCallQV dataset. This field is not used when the BaseCallQV dataset is not present, but is required when BaseCallQV is present.
MinTrimmedReadLength	_	The minimum trimmed read length across all unfiltered reads for this tag. Allowed values: Integers >= 0.
DoesRepresentedStrandMatchSource	1	Whether the reads are represented on the same strand as the source. Values:
		• <b>0</b> : The synthesis strand is not the same as the source strand.
		• <b>Non-zero</b> : The synthesis strand is the same as the source strand.



Table 125	XSQ tag	details	table	(continued)
-----------	---------	---------	-------	-------------

Column name	Default	Description
RelativeOrder	_	Describes the relative order of a read on the sense strand of the source nucleic acid, in relation to all of the other reads from the same fragment. 1 based. Required only when there is more than one tag per fragment.
SynthesisFromThreePrime	0	( <i>Optional</i> ) Whether the synthesis direction of the synthesized strand is from the 3' end. Values:
		<ul> <li>0 or missing: synthesis from the 5' end to the 3' end.</li> <li>Non-zero: The synthesis direction is from the 3' end to the 5' end.</li> </ul>

#### Table 126 lists tag names for current library types

Table 126 Tag information for current library types

Instrument	Library type	Indexing	Tag name	Base	Does- Represented- StrandMatch- Source	Relative- Order	Synthesis- FromThree- Prime
S0LiD™	Fragment	Yes   no	F3	Т	1	1	1
S0LiD <sup>™</sup> and	Mate-pair	No	F3	Т	1	2	1
5500			R3	G	1	1	1
S0LiD™	Paired-end	No	F3	Т	1	1	1
			F5-P2	Т	0	2	0
S0LiD™	Paired-end	Yes	F3	Т	1	1	1
			F5-BC	G	0	2	0
5500	Paired-end	Yes   no	F3	Т	1	1	1
	(DNA)		F5-DNA	Т	0	2	0
5500	Paired-end	Yes   no	F3	Т	1	1	1
	(RNA)		F5-RNA	G	0	2	0

The DatasetColorEncoding table in the XSQ file is required when color space is present. The table describes how reads are encoded as colors by the instrument. Indexing reads are not included in this table. The DatasetColorEncoding table is described in Table 127.

Table 127 The DatasetColorEncoding tabl	able 127	The DatasetColorEncoding	g table
---	----------	--------------------------	---------

	Column name	Description
1	DataSetName	Must match the name of a read dataset under a tag group. This table describes the encoding scheme used to interpret that read dataset.
2	Offset	Describes the distance and direction of the primer end as compared to the tag end. The offset is used to relate individual calls to the bases they encode.

	Column name	Description
3	Encoding	Contains a set of integers that direct the conversion from nucleotides to colors as described in the full XSQ file format document on the SOLiD <sup>™</sup> Software Tools website.
		(2BE is '11', 4BE is '1303').
4	Stride	Used to pack discontinuous reads into a continuous array. Stride is used to determine the relative positioning of the start nucleotides between color calls.
5	NumColorCalls	The maximum untrimmed read length in base space across all reads for this tag. This value is the width of the associated CallQV dataset.

#### Table 127 The DatasetColorEncoding table

For more information on the XSQ file format refer to this site:

http://solidsoftwaretools.com/gf/project/xsq

## BAM header usage

The BAM file generates all of the header information required by the SAM format specification, including @HD, @SQ, and @RG lines.

To view the content of the BAM file header, use the following command:

samtools view -H <BAMfilename>

Sequence Sequence header lines include the reference file URL, for example, file://share/ reference/genomes/hg18.fa in the optional UR field of the reference file. This value might become invalid if you relocate files.

**Read group (@RG)** Read groups receive an arbitrary ID and sample name. The library field (LB) contains information that is important to downstream algorithms that use pairing information. A library name, which is specified by the tool parameter library.name, and the library type, are separated by a dash in the LB field. The library type is a structured value that details the nominal length of the two tags and the protocol used as shown in the following syntax example:

11(x12)[F|MP|RR|RRBC]

In the syntax example, *l*<sup>1</sup> is the nominal length of the first read and *l*<sup>2</sup> is the nominal length of the second read. There will only be one number for fragment libraries. The library types correspond to fragment (*F*), mate-pair (*MP*), reverse read (*RR*), and reverse read-bar coded (*RRBC*).

Detecting structural variations, particularly large insertions and deletions, depends on the range of pairing insert (PI) sizes. The pairing tool generates the information about PI sizes. The information has been used to define the three-letter pairing category, specifically the third letter. (See Table 20 on page 156 and Table 21 on page 157 for descriptions of the three-letter categories.) The PI field in the read group captures the range of pairing insert sizes with a range of the form shown in the following example:

PI:low-high

In the PI example, low is the lower bound of the pairing range in nucleotides, and high is the upper bound. The LifeScope<sup>™</sup> Software SNPs module requires that BAM headers of its input files Header (@HD) sort contain an @HD line with a SO sort-order field, even if the BAM file is sorted properly. The required header line is: анр VN:1.0 GO:none SO:coordinate This header contains the information that the BAM file is sorted by genomic coordinates. BFAST mapping, for example, by default sorts the BAM file by coordinates, but does not update the header to reflect the sort order. This section describes how to update the @HD SO field. If your BAM file is sorted properly but does not contain the SO sort-order field, follow these instructions before using a BAM file generated outside of LifeScope<sup>™</sup> Software with the SNPs module: 1. Write the BAM file header to a file samtools view -H input.bam > input\_header.sam

2. Edit the BAM header file (input\_header.sam created in the previous step). Insert the following as the first line of the header:

анр VN:1.0 GO:none SO:coordinate

Make sure that the fields are tab-separated.

3. Update your BAM file with this new header. This command requires samtools-0.1.8 or later.

samtools reheader input\_header.sam input.bam

These instructions are not required with BAM files generated by LifeScope<sup>™</sup> Software mapping modules.

In order to be used by LifeScope<sup>TM</sup> Software tertiary analysis tools, the BAM file must **BAM** file validation have the @RG field populated in the BAM header. If you are using BFAST mapping analysis, you must run it with the -R option in the post-processing step to create the @RG field in the BAM file header. BAM files created outside of LifeScope<sup>™</sup> Software must pass Picard validation in order to be used with LifeScope<sup>™</sup> Software tertiary analysis tools.

> Note: Picard validation and samtools are supported by the open source project SourceForge.net. Applied Biosystems is not responsible for the availability or functionality of the Picard utility.

At the time of this writing the syntax for Picard validation is available at the following URL:

http://picard.sourceforge.net/command-line-overview.shtml#ValidateSamFile

The Picard utility, picard-tools-version.zip, (where version is the current version number) can be downloaded at:

http://sourceforge.net/projects/picard/files

The suggested Picard validation command is:

java -Xmx8g -jar ./ValidateSamFile.jar I=./input.bam \ IGNORE=MISSING\_TAG\_NM

order

where *input*. *bam* is the name of the BAM file to be validated.

#### XSQ metadata in BAM headers LifeScope<sup>™</sup> Software secondary analysis modules write metadata contained in XSQ reads files to their mapping output files, as metadata in BAM header comment (@CO) lines. The new metadata is not in the BAM specification, but BAM files with this metadata conform to the BAM specification. Table 128 lists the XSQ file fields with both the new @CO field names and the existing BAM header fields that contain the XSQ metadata.

#### Tie to existing BAM header lines

Each @CO metadata line with metadata information references one of the following types of BAM header lines:

• **Program group** – @PG lines capture information about the program that generated the data originally. The sequencing instrument and primary analysis software are considered to be the initial program. An example @PG line is:

@PG ID:Mordor\_201103161921\_0\_3 {...}

- **Read grou**p @RG lines contain information on library type and read length. (See "Read group (@RG)" on page 460 for more information.) An example @RG line is:
  - @RG ID:1 SM:Stooges LB:Larry DS:"75x35PE" PI:225 {...}
- General header @HD lines are general header lines.

All the attributes on the @CO metadata line then apply to referenced line.

The following example shows how @CO lines containing BAM metadata information refer to read group and program group lines. Except for @CO HD lines, the second field of the @CO metadata lines tie its attributes to either @RG or @PG lines. In the example, the RG:1 fields tie the metadata in those @CO lines to the @RG read group with an ID of 1. The PG:Mordor\_201012161921\_0\_3 field ties the metadata in that @CO line to the @PG program group with an ID of Mordor\_201012161921\_0\_3.

#### Metadata table

Table 128 lists the existing BAM header fields that contain the XSQ metadata, the new @CO metadata field names, and the XSQ file fields. See also "Input read count fields" on page 464.

Table 128 XSQ-related BAM metadata field	Table 128	XSQ-related	BAM	metadata	field
--	-----------	-------------	-----	----------	-------

Field name	Public BAM 1.3 spec field	New field in @CO line	XSQ field name	Optional or mandatory
Reference	(asq sp	_	Species	Mandatory
			{ 'Homo sapiens'   'Mus musculus'   'other'  }	



Table 128 X	(SQ-related	BAM	metadata	fields
-------------	-------------	-----	----------	--------

Field name	Public BAM 1.3 spec field	New field in @CO line	XSQ field name	Optional or mandatory
Assembly	ା ପରୁ AS	_	Assembly { 'hg18'   'hg19'   'mm9'  }	Mandatory
_	-	ଉCO HD UF	FileUUID	Optional
Program	@PG ID	_	InstrumentName_Date_FlowcellAssignment_ LaneNumber	Mandatory
Analysis Software	-	CO PG:x AS	AnalysisSoftware	Optional
InstrumentSerial	-	CO PG:x PS	InstrumentSerial	Optional
InstrumentName	-	CO PG:x PN	InstrumentName	Optional
InstrumentVendor	-	CO PG:x PV	InstrumentVendor	Optional
InstrumentModel	-	@C0 PG:x PM	InstrumentModel	Optional
ReadGroupID	(arg ID	_	LibraryName or LibraryName_IndexName (if present)	Mandatory
_	-	CO RG:x IX	IndexName (if present)	Optional
_	_	CO RG:x II	IndexID (if present)	Optional
Sequencing Center	(arg cn	_	SequencingCenter	Optional
Description	(arg ds	_	l1(xl2)[F MP RR RRBC]	Mandatory
Library Description	-	@CO RG:x LD	LibraryDetails.Description	Optional
Library Type	_	ାପିCO RG:x LT	LibraryType { 'MatePair'   'PairedEnd'   'Fragment' }	Mandatory
Application Type	_	ଉCO RG:x AT	ApplicationType { 'Whole Genome Resequencing'   'Targeted Resequencing'   'Whole Transcriptome (Fragment)'   'Whole Transcriptome (PairedEnd)'   'Small RNA'   'ChIP-Seq'   'Methylation' }	Mandatory
_	(arg dt	-	RunStartDate	Optional
_	-	@CO RG:x DE	RunEndTime	Optional
Library	(arg lb	_	LibraryName	Mandatory
Predicted Insert Size	(arg pi	_	(LibraryInsertSizeMin + LibraryInsertSizeMax) / 2	Conditional
Calculated Average Insert Size	_	@CO RG:x IA	<calculated></calculated>	Mandatory
Calculated Insert Size Std Dev	-	ଉCO RG:x IS	<calculated></calculated>	Mandatory
Min Insert Size from User Input	-	ଉCO RG:x IN	LibraryInsertSizeMinimum	Conditional
Max Insert Size from User Input	-	ାର୍ପCO RG:x IM	LibraryInsertSizeMaximum	Conditional
Input Read Count Passing All Filtering Steps	_	ଉCO RG:x CU	The sum across all ImageUnits of ImageUnitID.Fragments.NumFragmentsPassed (passes all filtering steps)	Optional



#### Table 128 XSQ-related BAM metadata fields

Field name	Public BAM 1.3 spec field	New field in @CO line	XSQ field name	Optional or mandatory
Input Read Count Total	-	@CO RG:x CT	The sum across all ImageUnits of ImageUnitID. FragmentCount (filtered and unfiltered)	Mandatory
			Not used in data converted from pre-5500 systems. Mandatory for 5500 data.	
Specimen		@CO RG:x SP	SampleIdentifier	Optional
Platform name	(arg pl	_	"SOLID"	Mandatory
Lane Id	(arg pu	_	FlowcellAssignment_LaneNumber	Optional
Sample (Pool)	(arg sm	_	SequencingSampleName	Optional
	_	@CO RG:x SD	SequencingSampleDescription	Optional
Tag1 Was Base Present In XSQ	-	@CO RG:x BX	Tag1.IsBasePresent	Mandatory
Tag1 Min Read Len	_	ଉCO RG:x TN	Tag1.MinTrimmedReadLength	Mandatory
Tag1 Max Read Len	-	@CO RG:x TX	Tag1.NumColorCalls or Tag1.NumBaseCalls	Mandatory
Tag2 Was Base Present In XSQ	_	@CO RG:x BY	Tag2.IsBasePresent	Mandatory for MP, PE.
Tag2 Min Read Len	-	@CO RG:x UN	Tag2.MinTrimmedReadLength	Optional for
Tag2 Max Read Len	-	CO RG:x UX	Tag2.NumColorCalls or Tag2.NumBaseCalls	
ECC run	-	@CO RG:x EC	<whether (inferred)="" an="" ecc="" run="" this="" was=""></whether>	Mandatory
ERCC	-	@CO RG:x ER	ERCC	Optional
_	-	@CO RG:x CO	Operator	Optional
_	_	@CO RG:x UU	LibraryIndexUUID	Optional
_	_	CORG:x PN	Application	Optional
_	-	(aco RG:x PJ	ProjectName	Optional
_	-	ଉCO RG:x SO	SampleOwner	Optional

#### Input read count fields

This section describes aspects of the Input Read Count fields from the viewpoint of the sequencing instrument and the viewpoint of secondary analysis.

The Input Read Counts fields are:

- CU Input Read Count Passing All Filtering Steps, @CO RG:x CU
- CT Input Read Count Total, @CO RG:x CT

The number of *total beads*, before any filtering on the instrument, is given by these fields:

- NumUnfilteredBeads In mapping statistics output
- FragmentCount In an XSQ file
- **CT** In BAM header metadata

The number of *beads that pass filtering* on the instrument is given by these fields:

- NumFilteredBeads In mapping statistics output
- NumFragmentsPassed In an XSQ file
- CU In BAM header metadata

The number of *beads filtered out* on the instrument is given by:

NumUnfilteredBeads - NumFilteredBeads

#### Example of @CO metadata in a BAM header

The following is an example of a BAM header with an @CO comment line containing metadata information. The @CO line applies to the @RG ID:12345678 line below it. (The @SQ lines for chromosomes 2–22 are deleted.)

```
@HD VN:1.0 GO:noneSO:coordinate
```

@SQ SN:chr1 LN:247249719 UR:file:/ref/hg18.fasta	
SP:Homosapiens AS:hg18	
<pre>@SQ SN:chrX LN:154913754 UR:file:/ref/hg18.fasta</pre>	
SP:Homosapiens AS:hg18	
<pre>@SQ SN:chrY LN:57772954 UR:file:/ref/hg18.fasta</pre>	
SP:Homosapiens AS:hg18	
<pre>@SQ SN:chrM LN:16571 UR:file:/ref/hg18.fasta</pre>	
SP:Homosapiens AS:hg18	
@PG ID:SOLiD_12321_1	
<pre>@CO RG:12345678 LT:MatePair AT:WholeGenomeResequencing</pre>	
LN:<50X50MP> PI:1500 IA:1500 IS:300 IN:1200 IM:1800	
PU:1234 SM:sample BX:0 TN:50 TX:50 BY:1 UN:50 UX:60 EC	:0
@RG ID:12345678 DS:50x50MP LB:MP PI:1500 PU:1234 SM:sampl	е

#### **@CO** syntax and rules

This section describes the syntax rules and conventions for @CO metadata lines. The @CO metadata rules are:

- The second field of each metadata @CO line either notes that the line is a general header (HD), or refers to a single @RG ID or @PG ID.
- Each metadata @CO line contains one or more tab-separated attributes.
- For a metadata @CO line that refers to either a @RG or @PG line, all of the @CO attributes must apply to the same @RG or @PG.
- The beginning of metadata @CO lines must match the regular expression "^@CO\t[PR]G:" and contain no other non-metadata text. LifeScope<sup>™</sup> Software ignores and does not change @CO lines that do not match the regular expression.
- The @CO PG:*id* line describing the instrument is required and must refer to a valid @PG ID:*id* program group line. This @CO line contains information about the primary analysis done on the instrument.

According to the BAM specification, the @HD line is optional. If present, the @HD line is required to be the first line of the header. The order of all other header lines is not defined. The following order of @CO metadata lines is preferred when possible (but not required):

- Place the @CO HD:UF comment immediately after the @HD line.
- Place @CO RG: and @CO PG: comment lines before other @CO lines for ease of visibility.

• Place @CO RG: and @CO PG: comment lines before the @RG and @PG blocks that the comment lines refer to.

## **Color-space** attributes

The SAM format specification includes the attribute tags CS, CQ and CM. All BAM files support color-space reads (see Table 129).

Attribute tag	Description
CS	Color-space (CS) read. The CS field contains the original color-space read, which includes the primer base, in the orientation of the CSFASTA file. CS entries are not manipulated to be top-strand relative.
CQ	Color qualities. Color qualities are encoded according to the ascii-33 scheme used for the QUAL field. The orientation is the same as the orientation used for the CSFASTA file.
СМ	The number of color-space mismatches.

 Table 129
 Color attribute tag description

## Pairing information in a BAM file

The BAM file that is produced by the pairing tool supports both mate-pair and pairedend protocols using the standard SAM format fields, in particular the ISIZE and FLAG fields.

#### Calculation of tag names

The paired libraries use tag names to refer to members of the pair. The mate-pair libraries use F3 and R3 as the tag names. The paired-end libraries use F3 and F5 as the tag names. Use the FLAG field and information from the LB field of the read group to recapitulate tag names (see Table 130).

 Table 130
 Calculating tag names

FLAG bit	Library type	Tag name
0x0040 (first read in a pair)	LMP	F3
0x0080 (second read in a pair)	LMP	R3
0x0040	PE	F3
0x0080	PE	F5
0x040	F	F3

**Proper pairs** Legacy file formats, such as \*.mates, and GFF, described pairs using a three-letter category. Pairs in the AAA category correspond to the "proper pair" concept in the SAM format. The pairs reflect pairings that are not altered by a structural variation such as an inversion or deletion (see Figure 46). The BAM file field values for proper pairs are different for mate-pair and paired-end libraries:

#### Mate-pair libraries

- Strand flag is equal for both mates (both 0 or both 1).
- ISIZE is between the lower and upper limit of the insert range.
- For forward strand hits R3 POS < F3 POS.
- For reverse strand hits F3 POS < R3 POS.

#### Paired-end

- Strand flag is opposite for the mates.
- ISIZE is between the lower and upper limit of the insert range. In the case of paired-end libraries, the ISIZE might be smaller than the sum of the alignment lengths.
- F3 POS < F5 POS if F3 is on the forward strand.
- F5 POS < F3 POS if F5 is on the forward strand.



Figure 46 Example of proper pairs for mate-pair and paired-end protocols

## Single read mapping quality

As described in the SAM format specification, the MAPQ field for paired results contains a pairing quality value. Under some circumstances. it is valuable to include the original single-read alignment quality value. The original single-read alignment value is maintained in the SM:i attribute in BAM files.

## Hard clipping of incomplete extensions

LifeScope<sup>™</sup> Software mapping uses a seed-extend algorithm. The algorithm increases mapping throughput by matching a seed, usually 25 bp, and extending the alignment until mismatches drive down the alignment score. Many alignments do not completely cover the color-space read. Because the base-space sequence of color reads cannot be precisely known in the absence of alignment, incomplete extensions are represented as a hard-clip (H) operation in the BAM CIGAR string (see Figure 47).

T33232030301212311201322311232302131021221120112222

#### AAGGCCTCTGAACCCACTCAGGTACTTAGCTGTAGATGGACATCAGTTAATTCGATGAC 22221102112212013120323211322310211321210303023233T \* \* \*

8H42M

Figure 47 Example of hard clipping from a color alignment

Figure 34 shows the read in normal orientation (see the top section) and aligned in reverse orientation to the reference top strand (see the middle section). The lines below the alignment show the extent of the seed (top horizontal line) and extension (bottom horizontal line) phases of mapping. The extension only results in 42 bases of alignment. The remaining portion of the color alignment has a number of mismatches that prevent extension. These are coded as hard-clipped regions. The CIGAR field in the BAM file is top-strand relative, so even though the hard clipping is on the end of the reversed color read, it is on the beginning of the CIGAR string.

## Visualize BAM output

	You can use third-party software visualization tools to view BAM files in a browser.				
Integrative Genomics View (IGV)	The Integrative Genomics Viewer (IGV) available from the Broad Institute is a visualization tool for interactive exploration of large, integrated datasets. The IGV reads BAM files directly, which allows for easy viewing and inspection of alignments against the genome (see Figure 47).				
	For more information, go to the following site:				
	www.broadinstitute.org/igv/				
	If you use IGV to visualize the BAM files, verify that the BAI file is present. The BAI file is the index that is built for BAM files and is a standard part of the public SAM specification. If the pairing and MaToBam tools do not automatically create the BAI file:				
	<b>1.</b> Log in to the LifeScope <sup><math>TM</math></sup> Software cluster.				
	<b>2.</b> At a command prompt, enter:				
	samtools index <bam file="" name="">.bam</bam>				
	Indexing only works if the file is sorted in coordinate order. If the file is not sorted in coordinate order, at a command prompt, run the following command to sort the file in coordinate order:				
	<pre>samtools sort <unsorted bam="" name="">.bam <sorted bam="" name=""></sorted></unsorted></pre>				
UC Santa Cruz (UCSC) genome browser	The UCSC Genome Browser serves as an interactive web-based microscope that allows researchers to view all 23 chromosomes of the human genome at any scale, from a full chromosome down to an individual nucleotide.				
	For more information, go to the Genome Browser website:				
	www.cbse.ucsc.edu/research/browser				


Figure 48 An example of a BAM file visualized in the IGV viewer

### Indel alignments

The alignments are included in the secondary analysis BAM file. Indel alignments are included in quality calculations and the selections of primary alignments and pairs.

Nearly all of the fields in the PAS file are represented in SAM format. However, there is no allowance for ambiguous locations in the CIGAR string or elsewhere. Ambiguity occurs when a repeat element is inserted or deleted with respect to the reference sequence. Indel alignments include a user-defined attribute (XW:Z) to specify the range of possible locations within the read for an indel in a repeat region (see Figure 49).

GTCTCACCAAGTTTTTTACATTTTCT' 12221101021.00000211300022

Figure 49 Example indel in a repeat region with ambiguous placement

Referring to Figure 49, in the alignment shown of a reference sequence stretch with a color read, the deletion of a single T, which is represented as a dot in the color sequence, cannot be placed precisely because of the repeated Ts. The XW attribute would span the homopolymer region (30\_35).

Table 131 PAS file c	column descri	ptions
----------------------	---------------	--------

Field Name	Example	Description
Genome position	4294973387	The genome position given by this formula:
		C * 232 + P – 1
		where $C$ is the chromosome number and $P$ is the position on that chromosome.



#### Table 131 PAS file column descriptions

Field Name	Example	Description
Indel size	-7	Number of bases in the indel. A negative value means a deletion, a positive value is an insertion, for example, $-8$ is a deletion of size 8, and 3 is an insertion of size 3.
Number of errors	2	Number of errors in the tag where the indel was found.
Alignment	See below.	Details of the alignment in the form of a number of concatenated fields.

The PAS file is a legacy format containing four tab-separated columns as described in Table 131. The fourth column contains the full alignment information. One example of an alignment is illustrated as follows.

The following is an example of PAS file content:

>600\_16\_579\_14\_Lib1\_1\_50\_1\_runName\_sampleName,1\_6074.51.2(17:17
\_17)[G20]| 1\_7297.1:(44.8.0)[T02] !8B52/
:B;\*%=7+'539&)4>455++60+0<9.(2</pre>

The alignment is in the form of a number of concatenated fields. The text represents:

<data source>,<R3 tag details>|<F3 tag details> R3\_FASTQ F3\_FASTQ

where the indel may be either F3 or R3, and the other tag would be the anchor, ungapped alignment. The R3\_FASTQ and F3\_FASTQ is the color quality strings for those tags in FASTQ base quality format. For fragment, there is only one tag:

<data source>,<tag details> FASTQ

Table 132 explains the fields in the example PAS file content.

Table 132 PAS file field descriptions

Field	Meaning	Notes		
Data source fields	Data source fields			
600_16_579	The bead ID.	_		
14	the unmatched length for that bead if an ungapped extended alignment was found for that bead.	The value 999999 is used if there was no ungapped alignment found.		
Lib1_1	A library indicator.	Deprecated.		
50	The full read-length.	—		
1	The run ID.	—		
runName_sampleName	The concatenation of the run name and the sample name.	_		
Indel tag fields				
1_6074.26.2	A sequence 1 hit at position 6074.	_		
	26 is the match length with 2 mismatches.			



Field	Meaning	Notes
(17:17_17)	The read position found was position 17 (zero-based).	insertion size = read_length – match_length – 1
	17_17 is the range of other possible positions.	Negative values are deletions, and positive values are insertions. In this example, the read
	[G20] is the read sequence with the	length was 50, so the indel is a deletion of 2.
	gap alignment.	If an indel is not detected in a tag, the Tag indel field contains the mapping hit information:
		<seq index="">_<alignment start="">.<num mismatches&gt;</num </alignment></seq>
Anchor (non-indel) tag fields		
1_7297.1:(44.8.0)[T02]	The alignment and the sequence of the anchor non-indel tag.	_
	1: The chromosome index (not necessarily the chromosome number).	
	7297: The chromosome position of the alignment.	
	1: The number of mismatches in the anchor part of the alignment.	
	44: The alignment extended length.	
	8: The overall number of mismatches.	
	0: The start of the alignment.	
	[T02]: The color-space sequence.	

 Table 132
 PAS file field descriptions (continued)

**Note:** The positions in the PAS files are 0-based. This contrasts with the positions reported in the TXT and GFF files, which have been adjusted to be 1-based.

## **BED** file format

The Browser Extensible Display (BED) format was developed to extend the UCSC Genome Browser with user- defined tracks. BED is used to visualize the splice and fusion junctions in the UCSC Genome browser and in the IGV browser (see Figure 50 on page 472). For general documentation about the BED format, including information about all of the BED fields, go to:

#### genome.ucsc.edu/FAQ/FAQformat.html

For information about visualization software, see "Pairing information in a BAM file" on page 466.

Each line in the track defines a junction where chromStart is the smaller of the coordinates and chromEnd is the greater.

There are two blocks because a junction typically contains two exons. BlockSizes are the lengths of the exons. The block starts the beginning of the exons. When fusions on different strands or chromosomes, two lines are added to the output, with each line representing one chromosome. Different colors are used to color-code different types: Figure 50 on page 472 shows the Upstream Hypersensitive Region (UHR) gene region displayed with the Integrative Genomics Viewer (IGV) for positions 3,530,193 to 3,548,355 of Human Chr-1. The following sections describe the tracks in Figure 50.

#### WIG (x2) tracks

The top two tracks show the genomic coverage using the negative strand and positive strand generated by the Bam2Wig tool (Max: 100 coverage).

#### **BAM track**

The middle track shows the alignments from the BAM file. For display purposes, reads are filtered with MAPQ threshold of 45 (a stringent filter). Bases with quality value 5 to 20 are shaded.

#### **BED track**

The fifth track shows the junctions detected by the Junction Finder (BED file). As shown in the figure, all junctions detected are "known" and so are shaded in green.



Figure 50 BED-IGV extended track example

# **BEDGRAPH** file format

Table 133 describes the BEDGRAPH format fields. These are tab-separated text fields.

 Table 133
 Description of BEDGRAPH fields

Field name	Example content	Description
Track annotation headers	track type=bedGraph name="Coverage"	Annotations used for UCSC browser display purposes.

В

Field name	Example content	Description
Contig name	chr1	Contig name of the feature. This value comes from the sequence name (SN) in the BAM header. However, in order to be viewed in the UCSC browser, these values must be of the form chr1, chr2, etc.
Feature start	148	Start location of the feature. Zero-based, inclusive.
Feature end	150	End location of the feature. Zero-based, exclusive (half-open ranges).
Coverage	1	Depth of coverage over the given (start-end) range.

Table 133 Description of BEDGRAPH fields

# Reference file data overview

	Nearly all LifeScope <sup>™</sup> Software analysis modules use reference sequence data and, in some cases, the modules also use reference annotations and metadata.	
	Reference data takes a number of forms, which are described in the following sections.	
Contig multi-fasta file	A single, multiple-entry FASTA file where each entry corresponds to a genomic contig or chromosome.	
Single contig FASTA file	A single-entry FASTA file containing one contig or chromosome.	
CMAP file	A tab-delimited text file used in previous versions of the software. Each line specifies the path to a single contig FASTA file. Columns in the CMAP file provide annotation and paths to other chromosome specific support files.	
GTF file	A genome annotation file. This file describes gene models used by the whole transcriptome and small RNA modules. It is similar in form to the GTF files downloaded from the UCSC website, but includes some changes for WT or small RNA processing.	
	Genomic reference files downloaded from UCSC; for example, you can download hg18 from:	
	http://hgdownload.cse.ucsc.edu/goldenPath/hg18/bigZips/chromFa.zip	
	The file can be used as a basis for the reference sequence data in LifeScope <sup>™</sup> Software tools. A few small transformations are required to prepare the GTF files for use by LifeScope <sup>™</sup> Software applications. See "Add new reference files" on page 517 and "Convert a GTF file" on page 520.	

A GTF file is an ASCII text file composed of lines of text separated by line feed characters (UNIX-style new lines). Each line of text represents a comment or a tab delimited content about a genomic feature. A comment line always starts with a "#" character. GTF does not specify any format of structured content in comments. Comments containing parsable meta-data should precede any lines containing genomic features. For more detail about the format, refer to the following site:

http://genome.ucsc.edu/FAQ/FAQformat.html#format4

VCF file	The Variant Call Format, in a text format. The file contains meta-information lines, a header line, and data lines that each contain information about a position in the genome. This file format was developed as part of the 1000 Genomes project, and is described here:	
	http://vcftools.sourceforge.net/specs.html	
	Recent releases of dbSNP are available in as VCF files. An example is:	
	ftp://ftp.ncbi.nih.gov/snp/organisms/human_9606/VCF/v4.0/00-All.vcf.gz	
Reference sequence data validation	The reference_validation.pl script provided with LifeScope <sup>™</sup> Software checks and corrects common issues with FASTA sequence files that can cause errors in LifeScope <sup>™</sup> Software runs. Examples of errors generated by an uncorrected GTF file can include extra spaces and mixed case.	
Concatenation	The contig multi-FASTA reference file can be generated by concatenating the individual chromosome files together.	
Select a reference file	Be sure to use the correct reference file type for your requirements. The identifiers in the FASTA definition line are carried through many tools, including the BAM file generated by mapping and pairing, and the GFF files typically written by tertiary analysis tools.	
	For some downstream viewing and integration tools, for example, the UCSC genome browser, the identifiers can be used to connect sequence references with other sources of annotation.	

# Read-set file format

An RRS (Read Range Specifier) file, also called a read-set file, is an internal file that defines which reads data is processed in a mapping run. LifeScope<sup>™</sup> Software LifeScope<sup>™</sup> Software automatically creates the RRS file for your mapping analysis. This format is considered an internal file format and specification for this file format can change in future versions of LifeScope<sup>™</sup> Software.

An RRS file allows a mapping analysis to process, in a single run:

- Data from a single XSQ reads file.
- Data from multiple XSQ reads files.
- Data for a range of barcodes (a subset of XSQ file data).
- Data for a range of panels within a barcode (also a subset of XSQ file data).



The RRS file is a tab-separated file with three columns which describe the input XSQ file, the sample name, and optionally a barcode range and panel range (see Table 134 on page 475). The following rules apply to RRS files:

- One read-set (barcode) cannot be part of two samples.
- For a single analysis, mapping can be in color or base. If the mapping is in color, all XSQ files in the RRS file must have color available. If the mapping is in base, all XSQ files in the RRS file must have base available. Mapping fails if these restrictions are not met.
- For barcoded XSQ files, the RRS file can specify a range of barcodes. If some of specified barcodes are missing, mapping will log a warning for the missing barcode IDs.
- One read-set is specified per content line.
- At least one of the read-sets needs to be valid for mapping module to complete.

The requirements for RRS fields are described in Table 134. The read-set file also accepts comments, which contain a '#' character at the beginning of the line.

 Table 134
 Read-set file description

Column name	Description
Sample name	Multiple read-sets with the same sample name are treated as a single sample (and are processed together).
	Allowed values: A string of alpha-numeric characters up to 255 chars longs. Spaces are not allowed.
	Examples: Human_101, Bob, Sample19
	(This sample name does not need to match any information provided on the instrument.)
File id	A positive integer that distinguishes between XSQ files according to these rules:
	Two different XSQ files must have different file IDs.
	<ul> <li>Two identical XSQ files must have same file ID.</li> </ul>
	(For read-set files created with the UI, LifeScope <sup>™</sup> Software assigns this number.)
read-set range URL	A string specifying one or more XSQ data files and optionally also barcode and panel specifiers. The format is a URI scheme: <i>fileName</i> ?start= <i>barcodeID.panelstart&amp;end=barcodeID.panelend</i>
	<i>fileName</i> : either a read in the reads repository or the absolute path to an XSQ file on the Linux file system (and that file must be a valid XSQ file for which the user has read permission).
	<i>barcodeID</i> : integers from 1–96.
	<i>panelstart, panelend</i> : integers from 1–4000. These fields correspond to panels on the sequencing instrument.
	The start= section must appear before the end= section. Start <i>barcodeID</i> value must be <= the end <i>barcodeID</i> value. The <i>panelstart</i> value must be <= the <i>panelend</i> value.

#### Example RRS file

The following is a simple example of an RRS file which defines an entire XSQ file as the read-set. The XSQ file in this example has been imported into the reads repository, so an absolute path to the local file system is not required.

#SampleName FileID ReadSets
Huref 1 reads/solid0054\_20110102\_RD\_HuRef100\_F3.xsq

# Legacy format translation

Table 135 and Table 136 map the GFF file field name to the corresponding BAM file field name for header fields and alignment data.

Table 135 GFF file header fields

GFF field	Corresponding BAM field	Comments
##reference-name	@SQ UR field	This header field contains the reference file name in the UR field.
##history	-	-
##color-code	-	Only a single color encoding is used.
##primer-base	_	This field is not needed. The primer base is stored with the color read attribute.
##max-num-mismatches	-	-
##max-read-length	@RG LB field	The read lengths in the file are determined from the library type information in the LB field of the read group.
##line-order	@HD SO field	The sort order field in the BAM header indicates the line order. Valid values are:
		None
		Coordinate
		• Qname

#### Table 136 Alignment data

GFF field	Corresponding BAM field	Comments
seqid	RNAME	GFF files use an ordinal number to indicate the reference contig. The ordinal number could be translated to names by a header table. The BAM file uses the QSQ lines to perform a similar lookup.
source	@RG PL	The read group platform (@RG PL field) is used in a manner similar to the SOLiD™ GFF source.
type	_	All records are alignments.
start	POS	The 1-based, left-most, top-strand position of the alignment.
end	Calculated from CIGAR	The BAM file does not support an explicit end point. An explicit end point can be calculated from the CIGAR string.
score	-	A read quality is not stored in the BAM alignment record. Mapping and pairing quality is stored in the BAM alignment record.

Table 136	Alignment	data	(continued)
-----------	-----------	------	-------------

GFF field	Corresponding BAM field	Comments
strand	5th FLAG bit (query strand)	The fifth bit of the FLAG field indicates the strand of the alignment.
GFF attributes		
aID ( <i>bead id</i> )	QNAME	The bead id.
at ( <i>adaptor type</i> )	Calculated value	The adaptor type or primer set field can be calculated as described in "Calculation of tag names" on page 466.
b ( <i>bases</i> )	SEQ	Base space representation of the aligned read. Bases are all uppercase. Equal signs "=" are not used.
c (category)	2nd FLAG bit (proper pair) and calculation	The most commonly-used category value, AAA, corresponds to the proper pair bit in the FLAG field. Use POS, MPOS, and strand bits in the FLAG field to calculate other categories. You can find C** pairs (different contigs) by interrogating the mate reference (MRNM). The D** is indicated by the fourth FLAG bit.
g (color-space read)	CS attribute	The color read is stored in the CS attribute. However, unlike the 'g' GFF attribute, the color read is stored in an unaltered CSFASTA form that includes the last primer base.
mq (mq [mapping quality])	MAPQ (or SM for pairs)	The mapping quality is stored in the MAPQ field for fragment libraries and in the SM attribute for paired results.
o (offset)	CIGAR (hard-clipping)	The offset for an alignment can be computed by checking the number of hard clip (H) operations on the CIGAR string.
p (mappability ambiguity)	_	Use mapping quality for uniqueness determinations.
pq (pairing quality)	MAPQ (for pairs)	The pairing quality is stored in the MAPQ field for paired results.
q (color qualities)	CQ	The CQ attribute stores color quality. The values are not represented as integers, but are represented by ascii-33 encoding.
r (reference color at mismatches)	Calculate reference color at mismatches using the samtools fillmd command.	CIGAR strings do not distinguish between matches and mismatches. Calculate this information using the samtools fillmd command, though in a form that is different from the rb GFF attribute.
s (color annotations)		-
u (accumulated mismatch count)	Per-alignment mismatches supported with CM attribute	The CM attribute specifies the number of color mismatches for the current alignment. You can use a combination of the number of color mismatches across alignments in the BAM file to reconstruct values like the 'u' attribute.

# Legacy CMAP file format description

CMAP is a legacy file format normally used to describe the chromosomes of an organism. A CMAP file provides a mapping between:

- Individual FASTA reference files
- An integer index
- A short name

Additional annotation is added in subsequent columns. File paths can be relative or absolute. FASTA reference files must be single entry. See Table 133 for a description of CMAP file fields.

CMAP files generated by the CNV tool contain additional information, pointing to the mappability files for each chromosome arm. See Table 134 on page 445 for a description of the Human CNV CMAP file fields.

The CMAP file may begin with one or more comment lines beginning with a pound sign (#). The comments can include file variables. Variable names use dot separated syntax and must appear immediately after the pound sign. Variable values are separated from the name by an equal sign (=) which may not be surrounded by white-space. The following fields are currently supported:

- data.dir You can use this optional field to define a root directory. If defined, relative file paths used in the data columns are relative to the root directory.
- Header This optional field may be used to define column headers for the columns. Column headers should be tab-separated.

Column	Field name	Туре	Example	Description
1	Seq id	Non-negative integer	2	The seq id is an ordinal index of the sequence.
2	Seq name	String	2	The name of the sequence. The name is typically a chromosome name and might include X, Y,M, etc. The *.chr suffix is added to the sequence name in a number of tools to support the UCSC browser function.
3	Fasta file path	String (path)	_	The path to a FASTA record containing a single sequence entry. The path can be relative or absolute. If the path is relative, you must define the data.dir field in the comments.
4	Double- encoded file path	String (path)	_	If there is a double-encoded version of the FASTA reference file, the path is specified here. Like the FASTA file path, if a relative path is provided, data.dir must be specified.
5, 6, 7, etc.	Additional data	Any	_	Any number of additional fields can be specified after the first four columns. These fields can be integers, strings, file paths, etc. If additional fields are file paths, the rules for relative file paths apply.

Table 137 CMAP file parameters

# XSQ Tools

# С

#### This appendix covers:

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# **Overview**

As sequencing technologies evolve and their output capacity increases, large amounts of new sequence data are straining current data storage and analysis systems in the genomics community. One approach to mitigating this pain is to store the instrument data in a more compact format, replacing the older text based formats — FASTQ, CSFASTA, and QUAL — with more efficient binary encoding. The Extensible Sequence (XSQ) file format has been developed to store each call and quality value in a single byte, which results in file sizes that are up to 75% smaller. There are many other benefits of this format, including embedded metadata, hierarchical structure of the file allowing simplified access to its contents, and the ability to store parallel datasets for easier integration of results.

Each new data format, however, comes with costs. Learning the intricacies of a new format can be easier with full descriptions and simplified examples. Integrating the new format with legacy data or legacy workflows can be more challenging, but this can be mitigated through the use of appropriate data format converters. The goal of the XSQ Tools package is to lower the activation energy associated with adopting the XSQ format by providing documentation, examples, and converters.

For command-line users, the XSQ Tools package is downloaded and run separately from LifeScope<sup>™</sup> Software. The LifeScope<sup>™</sup> Software UI handles the file conversion automatically.

Utilities in the XSQ Tools package perform these functions:

- For SOLiD<sup>™</sup> 4 System users, convert CSFASTA format files into the XSQ format required by LifeScope<sup>™</sup> Software.
- For 5500 Series SOLiD<sup>™</sup> Sequencer users running BioScope<sup>™</sup> Software or thirdparty tools, convert XSQ files into the CSFASTA format required by BioScope<sup>™</sup> Software.
- For LifeScope<sup>™</sup> Software users, optionally split an XSQ file into multiple XSQ files, each containing the data for one library.

More information about XSQ files and the XSQ file format is available on the SOLiD<sup>™</sup> Software Tools website:

#### http://solidsoftwaretools.com/gf/project/xsq

This website provides:

- The XSQ file format specification document
- Slides describing XSQ and its relation to the 5500 Series SOLiD<sup>™</sup> Sequencer
- A link to the XSQ Tools package

Table 138 lists the platforms supported by the XSQ Tools package.

Platform	Conversion type			
rtationin	To XSQ	From XSQ		
Linux <sup>®</sup> CentOS 4.7	Yes	No		
Linux <sup>®</sup> CentOS 5.5	Yes	Yes		

 Table 138
 Supported platforms for the XSQ conversion tools

# The XSQ Tools package

Package components

The following components are included in the XSQ Tools package:

- The convertToXSQ.sh wrapper script, which sets paths and executes tools for converting from CSFASTA and QUAL to XSQ.
- The convertFromXSQ.sh wrapper script, which sets paths and executes tools for the following:
  - Convert from XSQ to CSFASTA and QUAL.
  - Convert from XSQ to FASTQ.
  - Split an indexed (barcoded) XSQ file by library into new XSQ files.
  - Filter an XSQ file.
- Operating system-specific software libraries to support standalone executables.
- Example data sets, which show the data format using simulated reads. (These do not represent data generated by an instrument). We recommend you use the example data to verify your setup and to practice a conversion. This data is based on chromosome 6.
- Example usage scripts.
- End User License Agreement (EULA).
- Third-party licenses.
- Documentation.

DownloadTo download the XSQ Tools package, visit the following site:instructionshttp://solidsoftwaretools.com/gf/project/xsq

You must accept the EULA and follow the download directions on the site. After downloading the file, follow these steps to install the XSQ Tools package:

- 1. Move the file to an appropriate location.
- 2. Change directory to the tarball location.
- **3.** Execute the command tar -xvfz XSQ\_Tools.tgz. This creates a directory named XSQ\_Tools with the package contents.
- **4.** Add the XSQ\_Tools directory to your path, with one of the following commands:
  - bash: export PATH=\$PATH:\$tarlocation/XSQ\_Tools
  - csh/tcsh: set PATH = ( \$PATH: \$tarlocation/XSQ\_Tools ) where *tarlocation* is the directory containing the package tarball.

# Conversion to the XSQ format

This section describes converting CSFASTA and QUAL files from earlier SOLiD<sup>TM</sup> Systems to the XSQ format. This conversion is also supported in the LifeScope<sup>TM</sup> Software UI.

The convertToXSQ.sh script converts a pair of CSFASTA and QUAL files to an XSQ file. The command options vary by library type, as shown below:

• To convert SOLiD<sup>™</sup> fragment data:

```
sh convertToXSQ.sh \
    --mode=Fragment \
    --c1=<csfastafile> --q1=<qualfile> \
    --xsqfile=<output XSQ path and filename> \
    --libraryName=<LibraryName> \
    --runStartTime="yyyy-mm-dd hh:mm:ss"
• To convert SOLiD<sup>™</sup> mate-pair or paired-end data:
sh convertToXSQ.sh \
    --mode=LMP OR --mode=PE \
    --c1=<csfastafile> --q1=<qualfile> \
    --c2=<csfastafile> --q2=<qualfile> \
    --libraryInsertSizeMinimum=<Minimum insert size> \
    --libraryInsertSizeMaximum=<Maximum insert size> \
```

- --xsqfile=<output XSQ path and filename> \
- --libraryName=<LibraryName> \
- --runStartTime="yyyy-mm-dd hh:mm:ss"
- To convert SOLiD<sup>™</sup> barcoded fragment data (a single barcoded fragment data set):

```
sh convertToXSQ.sh \
```

```
--mode=BC \
```

--cl=<csfastafile> --ql=<qualfile> \

- --bc1=<index csfastafile> --bq1=<index qualfile> \
- --xsqfile=<output XSQ path and filename> \
- --libraryName=<LibraryName> \
- --runStartTime="yyyy-mm-dd hh:mm:ss"
- To convert SOLiD<sup>™</sup> barcoded paired-end data (a single barcoded paired-end data set):
  - sh convertToXSQ.sh  $\$ 
    - --mode=BCPE  $\$
    - --c1=<csfastafile> --q1=<qualfile> \
    - --c2=<csfastafile> --q2=<qualfile> \
    - --bc1=<index csfastafile> --bq1=<index qualfile> \
    - --libraryInsertSizeMinimum=<Minimum insert size> \
    - --libraryInsertSizeMaximum=<Maximum insert size> \
    - --xsqfile=<output XSQ path and filename> \
    - --libraryName=<LibraryName> \
    - --runStartTime="yyyy-mm-dd hh:mm:ss"

Table 139 lists the options required by the convertToXSQ.sh script. Table 140 on page 483 lists the remaining options.

Table 139 Required arguments for the convertToXSQ.sh script (to create an XSQ file)

Option	Required with	Description
mode	All	Allowed values are Fragment, LMP, PE, BC, BCPE:
		<ul> <li>Fragment: Convert fragment CSFASTA and QUAL files to XSQ format.</li> </ul>
		• LMP: Convert mate-pair CSFASTA and QUAL files to XSQ format.
		• <b>PE</b> : Convert paired-end CSFASTA and QUAL files to XSQ format.
		• <b>BC</b> : Convert single barcoded fragment CSFASTA/QUAL with index information to XSQ format.
		• <b>BCPE</b> : Convert single barcoded paired-end CSFASTA/QUAL with index information to XSQ format.
libraryName	All	Name of the library preparation.
runStartTime < <i>arg</i> >	All	Time when the instrument run was executed. This information is entered into the output XSQ file. Use one of these formats:
		<ul> <li>'yyyy-mm-dd hh:mm:ss' (including the quotes, either double or single).</li> </ul>
		<ul> <li>yyyy-mm-dd\ hh:mm:ss (escape the space with a backslash).</li> </ul>
xsqfile <arg> or -x <arg></arg></arg>	All	Requested name of output XSQ file. The name must include the extension $.xsq$ .
c1 < <i>arg</i> >	All SOLiD <sup>™</sup> System data	The CSFASTA input file for the first tag.
q1 <i><arg></arg></i>	All SOLiD <sup>™</sup> System data	The QUAL input file for the first tag.
c2 <arg></arg>	LMP, PE, or BCPE S0LiD <sup>™</sup> System data	The CSFASTA input file for the second tag.

Option	Required with	Description
q2 <i><arg></arg></i>	LMP, PE, or BCPE S0LiD <sup>™</sup> System data	The QUAL input file for the second tag.
bc1 <i><arg></arg></i>	BC or BCPE S0LiD <sup>™</sup> System data	The index (barcoded) CSFASTA input file.
bq1 <i><arg></arg></i>	BC or BCPE S0LiD™ System data	The index (barcoded) QUAL input file.
libraryInsertSizeMinimum	LMP or PE S0LiD <sup>™</sup> System data	The expected minimum insert size, in nucleotides.
libraryInsertSizeMaximum	LMP or PE S0LiD <sup>™</sup> System data	The expected maximum insert size, in nucleotides.

 Table 139
 Required arguments for the convertToXSQ.sh script (to create an XSQ file) (continued)

Table 140 lists options for the convertToXSQ.sh script. See Table 139 on page 482 for the required options.

Fable 140 Optional argume	ts for the	convertToXSQ.	sh script (t	to create an	XSQ file)
---------------------------	------------	---------------	--------------	--------------	-----------

Option		Description
-a <arg></arg>	application <arg></arg>	The type of analysis intended for this input data. Allowed values are listed below. Quotes are required around values that contain spaces.
		Whole genome resequencing
		Targeted resequencing
		Whole transcriptome (Fragment)
		Whole transcriptome (PairedEnd)
		Small RNA
		• "ChIP-Seq"
		"Methylation"
	assembly < <i>arg</i> >	The intended assembly for mapping. May be blank if unknown. Examples:
		• hg18
		• hg19
		• mm9
		Other assembly names are allowed. Spaces are not allowed in these names.
-c <arg></arg>		Use the configuration file <i><arg></arg></i> as a list of conversion parameters. The parameters are specified as nested key value pairs. Example file content:
		q1=/data/results/reads/uhr.150.F3.qual
		c1=/data/results/reads/uhr.150.F3.csfasta
		x=./out.xsq mode=Fragment
		runStartTime=2011-04-01\ 12:01:02
		libraryName=150_UHR
		Option hyphens ('-', '') are not used in the configuration file.
	comments	Additional information about this sample or its preparation.
-d <i><arg></arg></i>	description <arg></arg>	A description of the sample.
-е	ercc	Whether internal controls have been used. Reserved for future use.

С

Table 140	Optional arguments	for the convertToXSQ.	sh script (to create a	n XSQ file) <i>(continued)</i>
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Option		Description		
	flowCellAssignment <arg></arg>	The position of the instrument flowcell in which this flowchip was placed.		
-h	help	Display command usage and options.		
	instrumentName < <i>arg</i> >	The name of the instrument.		
	instrumentSerial < <b>arg</b> >	The unique serial number identifying the instrument.		
-k <arg></arg>	indexKitName <i><arg></arg></i>	The name of the kit used to add the indexing nucleotides.		
	laneNumber < <i>arg</i> >	Lane number in which this sequencing sample was loaded within the flowchip layout.		
	libraryName <i><arg></arg></i>	The name of library preparation.		
-n	noqualfile	Use this attribute to specify that the QUAL file is not available.		
-o <arg></arg>	operator < <i>arg</i> >	The name or ID of person who set up the instrument run.		
-p < <i>arg</i> >	projectName < <i>arg</i> >	Project within which this sample is being run. This value is entered into the projectName field in the libraryDetails table, and may be different from the LifeScope <sup>™</sup> Software project name.		
	runEndTime <arg></arg>	Time when the instrument run completed. This information is entered into the output XSQ file. Use one of these formats:		
		• 'yyyy-mm-dd hh:mm:ss' (including the quotes, either double or single).		
		• yyyy-mm-dd\ hh:mm:ss (escape the space with a backslash).		
	runName < <i>arg</i> >	The name of a flowchip run.		
	sampleIdentifier < <i>arg</i> >	The unique name for this sample.		
	sampleOwner < <i>arg</i> >	The owner of this sample.		
	-sequencingSampleDescription <arg></arg>	Description of the sample loaded in a lane on a flowchip.		
	sequencingSampleName <arg></arg>	The name of the sample loaded in a lane on a flowchip.		
	species <arg></arg>	The intended species for mapping. Example values include:		
		Homo sapiens		
		Mus musculus		
		• other		
-V	version	Print version information.		



# Conversion from the XSQ format

	-C	Converts the input XSQ file into CSFASTA, QUAL, and FASTQ files.
	Option	Description
	Table 141 Optio	ons for the convertFromXSQ.sh script
Options table	Table 141 lists options includ	the options supported by the convertFromXSQ.sh script. These le functionality for XSQ file-splitting and for version information.
	By default con- converted out during conver written in the	nvertFromXSQ. sh ignores the filtering flags and writes all reads to the put files. With the -f option, the converter applies the filtering flags sion, which suppresses the filtered reads and quality values from being output files.
	In the XSQ file for filtering. C not written to	e, each read contains a field of filtering flags, which may mark the read only when the -f option is used, reads that are marked for filtering are the CSFASTA file.
	convert	FromXSQ.sh -c -f <i>xsqfile</i>
Filter option	Use the – f opt of the comma	tion to filter (remove) marked reads during conversion, with this version
	FASTQ files a	re created only if the input XSQ file contains base-space data.
	• .fastq	
	• .0V.qua	1
		a
	where out he	the following values:
		LibraryName> <tagname> ext</tagname>
	The cutout fil	res/~reads/~reads/
	Ine converted	ines are created in the following directory:
	where <i>xsqfile</i> g	rives an existing XSQ file on the file system.
	convert	FromxSQ.sn -c xsqiile
Conversion syntax	The convert QUAL files. T	FromXSQ.sh script converts an XSQ file into a pair of CSFASTA and he usage for this script is:
	If the input XS space data inte	Q file includes base-space data, the conversion also exports the base- o a FASTQ file.
	The conversio This conversio Software com	n program is available as a standalone script to be run on the Linux shell. on is not available within the LifeScope <sup>TM</sup> Software UI or the LifeScope <sup>TM</sup> nand shell.
	This section de Sequencer into BioScope <sup>™</sup> So are analyzing	escribes converting XSQ format files from the 5500 Series SOLiD <sup>TM</sup> o CSFASTA and QUAL files, in the format required by earlier version of ftware. This section does not apply to LifeScope <sup>TM</sup> Software users who XSQ data files generated by the 5500 Series SOLiD <sup>TM</sup> Sequencer.

Option	Description
-f	(Optional) During conversion, removes reads that are marked for filtering.
-0	(Optional) Specifies an output directory partial path.
-S	Splits the input XSQ file into multiple output files, one for each library in the parent XSQ file.
-V	Displays the package version.

Table 141 Options for the convertFromXSQ.sh script

# XSQ file splitting

The convertFromXSQ.sh script with the -s option splits an input XSQ file into multiple XSQ files, one file per library. For example, if a paired-end library is created with indices for each of lib1, lib2, and lib3 and run in the same lane, a single XSQ file will be created. Using the -s option creates three separate XSQ files - one for each of lib1, lib2, and lib3. Each XSQ file contains data for both the F3 and F5 tags.

To split an XSQ file, use the following command:

convertFromXSQ.sh -s xsqfile -o outputpath

The following output files are produced:

outputpath/ lib1.xsq lib2.xsq

Only one input XSQ file is allowed per run.

# HDF5 tools

The XSQ format conforms to the HDF5 specification, and HDF5 tools work on XSQ files. Figure 51 is an example of how HDFView displays an XSQ file, showing Groups, Tables, and Datasets.







### Internal conversion parameters

This section describes the parameters used internally by the LifeScope<sup>™</sup> Software UI when converting a file to the XSQ format from the older CSFASTA and QUAL formats used in earlier versions of the software. These parameters are for internal use.

**Parameters table** Table 142 lists and describes the conversion parameters used internally by the LifeScope<sup>™</sup> Software UI.

 Table 142
 Internal XSQ conversion parameters

Parameter	Default	Description
xsqconverter.run	_	For internal use.



#### Table 142 Internal XSQ conversion parameters (continued)

Parameter	Default	Description
xsq.application xsq.assembly		Describes how this sequence is intended to be analyzed. Allowed values are: • Whole genome resequencing • Targeted resequencing • Whole transcriptome (Fragment) • Whole transcriptome (PairedEnd) • Small RNA • ChIP-Seq • Methylation The intended assembly for mapping. May be blank if unknown.
		Example values: • hg18 • hg19 • mm9
xsq.comments	_	Additional information about this sample or its preparation.
xsq.description	_	Sample description.
xsq.ercc	0	<ul> <li>Whether or not internal controls have been used. Allowed values:</li> <li>0: No</li> <li>1: Yes</li> </ul>
xsq.flowCellAssignment	255	The position of the instrument flowcell in which this flowchip was placed.
xsq.indexKitName	-	Name of the kit used to add the indexing nucleotides.
xsq.input.barcodedCsfasta1	-	Barcoded CSFASTA file (used only in the case of indexed runs).
xsq.input.barcodedQual1	_	Barcoded QUAL file (used only in the case of indexed runs).
xsq.input.csfasta1	_	Input CSFASTA file.
xsq.input.csfasta2	_	Second CSFASTA file (use only in the case of LMP or PE runs).
xsq.input.qual1	_	Input QUAL file.
xsq.input.qual2	_	Second QUAL file (use only in the case of LMP or PE).
xsq.instrumentName	_	Name of the sequencing instrument.
xsq.instrumentSerial	_	Unique serial number to identify a particular sequencing instrument.
xsq.laneNumber	255	Lane number within the flowchip layout in which this sequencing sample was loaded.
xsq.libraryInsertSizeMaximum	_	Expected maximum insert size in nucleotides. (Required for LMP and PE conversion).
xsq.libraryInsertSizeMinimum	_	Expected minimum insert size in nucleotides. (Required for LMP and PE conversion).
xsq.libraryName	_	The name of library preparation.

$\cap$

	Table 142	Internal XSQ	conversion	parameters	(continued)
--	-----------	--------------	------------	------------	-------------

Parameter	Default	Description
xsq.mode	_	XSQ mode of operation. Allowed values are:
		• Fragment: Convert fragment CSFASTA and QUAL formats to XSQ.
		LMP: Convert mate-pair CSFASTA and QUAL formats to XSQ.
		• <b>PE</b> : Convert paired-end CSFASTA and QUAL formats to XSQ.
		• <b>BC</b> : Converting single barcoded fragment CSFASTA and QUAL formats with index information to XSQ.
		• <b>BCPE:</b> Convert single barcoded paired-end CSFASTA and QUAL formats with index information to XSQ.
xsq.noqualfile	false	Specify that the input QUAL file is not available. Allowed values:
		• false: The QUAL file is available.
		• <b>true</b> : The QUAL file is missing.
xsq.operator	_	The name of person or ID who set up the instrument run.
xsq.output.xsqfile	_	The requested name for output XSQ file. Must use the ".xsq" extension.
xsq.projectName	—	Project within which this sample is being run.
xsq.runEndTime	_	Time when the instrument run completed. Required format: 'yyyy-mm-dd hh:mm:ss' (including quotes).
xsq.runName	_	Name of a flowchip run.
xsq.runStartTime	_	Time when the instrument run started. Required format: 'yyyy-mm-dd hh:mm:ss' (including quotes).
xsq.sampleIdentifier	_	Unique name for this sample.
xsq.sampleOwner	_	Owner of this sample.
xsq.sequencingSampleDescription	_	Description of the sample loaded in a lane on a flowchip.
xsq.sequencingSampleName	_	Name of sequencing sample loaded in a lane on a flowchip. This value is the bead emulsion loaded on the instrument. The emulsion may be a collection of barcoded libraries from multiple samples.
xsq.species	—	The Intended species for mapping. Example values include:
		Homo sapiens
		Mus musculus
		• Other

**Internal INI file** The following is an example of an INI file used internally by the LifeScope<sup>™</sup> Software UI during a conversion to the XSQ format from the older CSFASTA and QUAL formats used in earlier versions of the software. Refer to the *LifeScope<sup>™</sup> Genomic Analysis* Software User Guide (Part no. 4465696) for information about the UI. This INI file is for internal use.

##Run parameter xsqconverter.run=1 ##How this sequences is intended to be analyzed xsq.application=Unknown ##Intended assembly for mapping xsq.assembly=Unknown ##Aditional information about this sample or its preparation xsq.comments=Unknown

##Sample description
xsq.description=Unknown

##Whether internal controls have been used (0=No, 1=Yes)
xsq.ercc=0

##The position of the instrument flowcell in which this flowchip
was placed
xsq.flowCellAssignment=255

##Name of the kit used to add the indexing nucleotides
xsq.indexKitName=Unknown

##Barcoded csfasta file (should be used only in case of indexed runs) xsq.input.barcodedCsfasta1=

##Barcoded qual file (should be used only in case of indexed runs) xsq.input.barcodedQual1=

##Csfasta file
xsq.input.csfasta1=

##Second csfasta file (should be used only in case of LMP or PE)
xsq.input.csfastac2=

##Qual file
xsq.input.qual1=

##Second qual file (should be used only in case of LMP or PE)
xsq.input.qual2=

##Name of instrument
xsq.instrumentName=Unknown

##Unique serial number to identify a particular instrument
xsq.instrumentSerial=Unknown

##Lane number within the flowchip layout in which this sequencing sample was loaded xsq.laneNumber=255

##Expected maximum insert size in nucleotides. (Required for LMP and PE conversion) xsq.libraryInsertSizeMaximum=

##Expected minimum insert size in nucleotides. (Required for LMP and PE conversion) xsq.libraryInsertSizeMinimum=



# The name of library preparation
xsq.libraryName=DefaultLibrary

# Allowed values are: Fragment, LMP, PE, BC, BCPEUse 'Fragement' for converting fragement csfasta/qual to XSQ. Use 'LMP' for converting mate pair csfasta/qual's to XSQ. Use 'PE' for converting paired end csfasta/qual's to XSQ. Use 'BC' for converting single barcoded fragment csfasta/qual with index information to XSQ. Use 'PE' for converting single barcoded paired end csfasta/qual with index information to XSQ xsq.mode=Fragment

##Specify that qual file is not available (Use 'true' - to specify qual file missing) xsq.noqualfile=false

##The name of person/ID who set up the instrument run xsq.operator=Unknown

##Desired name for output XSQ file
xsq.output.xsqfile=pluginfrag.xsq

##Project whithin which this sample is being run
xsq.projectName=Unknown

##Time when run completed. The run end time should be in 'yyyymm-dd hh:mm:ss' format xsq.runEndTime=

##Name of a flowchip run
xsq.runName=Unknown

##Time when run was executed. The run start time should be in 'yyyy-mm-dd hh:mm:ss' format xsg.runStartTime=2011-01-11 20:10:20

##Unique name for this sample
xsq.sampleIdentifier=Unknown

##Owner of this sample
xsq.sampleOwner=Unknown

##Description of the sample loaded in a lane on a flowchip
xsq.sequencingSampleDescription=Unknown

##Name of sample loaded in a lane on a flowchip
xsq.sequencingSampleName=Unknown

##Intended species for mapping
xsq.species=Unknown



### Links to resources

This section lists resources related to XSQ file format and converters.

For the XSQ Tools package and documentation, XSQ file format specification, XSQ webinar slides, and example XSQ files, visit this site:

http://solidsoftwaretools.com/gf/project/xsq/

HDFView is a Java<sup>®</sup> tool for browsing HDF-based files, including XSQ files. For information about HDFView, visit this site:

http://www.hdfgroup.org/hdf-java-html/hdfview/

HDF5 is a technology suite for managing of large, complex data collections. HDF5 APIs are available at this site:

http://www.hdfgroup.org/HDF5/release/obtain5.html

Exact Call Chemistry (ECC) is a unique ligation-based sequencing methodology, introduced with the 5500 Series  $SOLiD^{TM}$  Sequencer. The Exact Call Chemistry white paper is available through this link and the instructions that follow:

http://www.appliedbiosystems.com/absite/us/en/home/applicationstechnologies/solid-next-generation-sequencing/publications-literature.html

- 1. Under the Publication & Literature section, click the Product Literature tab.
- 2. Scroll down to the White Papers section.
- **3.** Click on the "SOLiD<sup>™</sup> System Accuracy with the Exact Call Chemistry Module" link.

# FAQ - XSQ Tools

#### What software accepts the XSQ file format?

Initially, only LifeScope<sup>™</sup> Software supports the new format. Life Technologies Corporation is working with third-party developers to adapt their workflows to support the new chemistry and data format.

2

1

#### I have pipelines that require CSFASTA and QUAL files as input. What should I do?

Life Technologies Corporation provides tools on the SOLiD<sup>™</sup> Software Tools website to convert XSQ files into CSFASTA and QUAL files. See "Links to resources" on page 492, and download the XSQ Tools package.



#### I have pipelines that require FASTQ files as input. What should I do?

When the ECC module is used during sequencing, base-space data is available in the XSQ file and can be exported into a FASTQ file. See "Conversion from the XSQ format" on page 485.

Can I use data from both a SOLiD<sup>™</sup> 4 System and a 5500 Series SOLiD<sup>™</sup> Sequencer for data analysis?

Yes. There are two options and both involve converting the data into a common format, either XSQ or CSFASTA+QUAL. One way is to convert the SOLiD<sup>™</sup> 4 System CSFASTA and QUAL output to the XSQ format (see "Conversion to the XSQ format" on page 481). Then supply both the converted XSQ file and the 5500 XSQ file as input to your LifeScope<sup>™</sup> Software analysis.

Another method is to convert the XSQ data into CSFASTA+QUAL for use with older analysis tools.

5

6

3

4

Is there a converter to change CSFASTA and QUAL files to the XSQ format?

Yes, standalone converters are provided in the XSQ Tools package at the following site: http://solidsoftwaretools.com/gf/project/xsq/

How does the XSQ format handle multiplexing?

Multiple libraries may be included in a single XSQ file, such as are generated for each lane of a SOLiD<sup>™</sup> 5500 Series Sequencer run. When multiple different users are providing samples to be run in the same lane, the resulting file may be split by library for distribution to individual users, without sharing all of the data with each user. The convertFromXSQ.sh script provides the -s splitting option to separate libraries into separate XSQ files when necessary. See "XSQ file splitting" on page 486.

**Note:** The –s splitting option only works for indexing files. Non-indexing files already contain only a single library (and are not split with the –s option).

The convertToXSQ. sh script is run once for each barcode index, and each run results in a single XSQ file. Multiple barcode indices cannot be joined in a single XSQ file with the current converter.

### 7

Are APIs available for accessing XSQ data?

#### Yes. HDF5 APIs work with the XSQ format, and are available at this site: http://www.hdfgroup.org/HDF5/release/obtain5.html

#### Is an XSQ file viewer available?

Yes. XSQ data is can be viewed with the HDFView browser, which can be obtained at this site:

#### http://www.hdfgroup.org/hdf-java-html/hdfview/

#### 9

8

#### Can XSQ files be converted in the LifeScope<sup>™</sup> Software UI and command shell?

Yes. The LifeScope<sup>TM</sup> Software UI supports the conversion of CSFASTA and QUAL files to the XSQ format. If you are running the LifeScope<sup>TM</sup> Software command shell, use the standalone XSQ conversion tool that is part of the XSQ Tools package.

### 10

#### Is library index correction supported?

Not in the current version. An XSQ library index reassignment tool is under consideration.

# 11

#### Why does the converter not work on my variant of Linux.

The table in the "Overview" on page 479 section describes that conversion *to* XSQ requires CentOS 4.7 or later, and conversion *from* XSQ requires CentOS 5.5 or later. Other versions of Linux are not supported.

### 12

#### What are the reserved quality values (QVs) in the XSQ file?

From the XSQ File Format Specification:

For the QV, the range of the 6 bits is between 0 and 63. Valid quality values will be from 3 to 62, represented as PHRED scores [QV = -10\*log10P(error)]. 0 and 1 are reserved for future usage; 2 indicates a missing quality value (not trimming); 63 refers to an uninformative or missing call.



# 13

What values should I use for libraryInsertSizeMinimum and libraryInsertSizeMaximum when they are not known during the conversion of mate-pair and paired-end files?

Any values will work as long as libraryInsertSizeMinimum is less than or equal to libraryInsertSizeMaximum and both values are nonnegative.

# 14

#### How can I collect statistics on an XSQ file?

Not supported currently. An XSQ Statistics utility to extract the number of reads by file, barcode, tag, etc., in addition to library type and read length, is under consideration.

# 15

#### How can I tell if my XSQ file is valid or corrupt?

Not supported currently. An XSQ validator utility is under consideration.

# 16

# I have a CSFASTA file to convert to XSQ format, but I do not have the QUAL file. Can I create the XSQ file?

When converting CSFASTA to XSQ, if the QUAL file is missing or not found, all reads are marked for filtering in the output XSQ file. As a result, the reads are ignored (not processed) when used as input into LifeScope<sup>TM</sup> Software analyses.

The workaround is to create a dummy QUAL file to use as input with the convertToXSQ.sh script. Create a QUAL file with the same bead ids (and in the same order) as the CSFASTA file. Use QV values at least high enough to pass default thresholds.



Appendix C XSQ Tools FAQ - XSQ Tools



# **Command Shell Control Files**

This appendix covers:

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Set up your own analysis run	502
LifeScope <sup>TM</sup> Software parameters	503
(Optional) The Linux mail command	504

# **Overview**

This chapter describes the configuration files that LifeScope<sup>™</sup> Genomic Analysis Software uses to control its execution. These files control which LifeScope<sup>™</sup> Software modules are executed and their order of execution, and also define the run-time parameters that customize each module's behavior.

When you use the LifeScope<sup>™</sup> Software UI or the LifeScope<sup>™</sup> Software command shell, the configuration files are automatically generated for you.

The following types of control files are used with your LifeScope<sup>™</sup> Software analyses:

 Read-only module XML files – Contain the parameters accepted for each module and default settings for those parameters. The XML files, one per module, are in the <installdir>/etc/plugins/pipelines directory.

Example: <installdir>/etc/plugins/pipelines/dibayes.xml

- Module properties files Define settings used in all runs for a particular module. These files are located in <*installdir*>/etc/plugins/properties.
   Example: <*installdir*>/etc/plugins/properties/ \dibayes.properties
- A global.ini file Defines settings which by convention are imported into other INI files in the current directory only. For example, in a workflow tertiary directory, the other INI files in that tertiary directory import global.ini and all its parameters. In the case of conflicting settings, the setting in the global.ini files takes precedence over settings in the *<installdir>* module properties file

Example: <installdir>/etc/workflows/small.rna/tertiary/ \
global.ini

• LifeScope<sup>™</sup> Software modules' INI files – Define settings for a single module run. In the case of conflicting settings, the module INI file setting takes precedence over settings in both your global.ini file and the *<installdir>* module properties file.

Example: <installdir>/etc/workflows/ \
targeted.resequencing.frag/tertiary/dibayes.ini



• PLN files – Define a sequence of analyses to be executed. A PLN file lists one or more INI or PLN files. Each INI file listed corresponds to a LifeScope<sup>™</sup> Software module to be executed.

```
Example: <installdir>/etc/workflows/ \
targeted.resequencing.frag/tertiary/tertiary.pln
```

# **Configuration files**

#### PLN files

A PLN file defines your analysis through a list of tasks, which is a list of one or more INI or PLN files. For each listed INI file, LifeScope<sup>™</sup> Software executes an analysis run of the corresponding module.

The following is an example of a simple PLN file, which causes a SNPs module analysis to be run:

diBayes.ini

Tasks (INI or PLN files) are listed one per line. When there are multiple tasks listed, every task without a dependency is executed in parallel (assuming there are sufficient compute nodes).

#### Dependencies

A PLN file specifies a directed acyclic graph of module dependencies. The less-than sign ('<') signifies a dependency. A dependency means that task on the left of the less-than sign is executed only if the task on the right of the sign completes successfully.

For example, the following is a typical workflow PLN file:

```
secondary.pln
tertiary.pln < secondary.pln</pre>
```

This PLN file specifies that tertiary.pln is dependent on secondary.pln. First the modules specified in secondary.pln are executed. Only if those modules completed successfully, then the modules specified in tertiary.pln are executed.

A task with no dependencies takes this form in a PLN file:

x.ini

Multiple dependencies are added after the less-than sign, separated by commas:

```
x.ini < a.ini, b.ini, c.ini
```

Forward references are not allowed in the dependency list, and a task cannot list itself as a dependency.

#### Other syntax

Blank lines and comment lines are ignored in PLN files. Comment lines must begin with a "#" as the first character.

#### On error

When one module in a sequence fails, the remaining modules specified in the PLN file continue. However, any module that depends in the failed module is not attempted.



#### INI files Module INI files

Module INI files are the main source of configuration parameters requiring your customization for LifeScope<sup>™</sup> Software runs. A module INI file defines run-time parameters which control the behavior of the module's algorithm.

Many examples of module INI files are included in the LifeScope<sup>™</sup> Software standard workflows, under the following directories:

- <installdir>/etc/workflows/\*/secondary
- <installdir>/etc/workflows/\*/tertiary

The following is example content of a module INI file, in this case the CNV INI file:

import global.ini

##Run parameter
cnv.run=1

#cnv.mappability.dir=\${analysis.mappability.dir}

##Size of the window block to be considered as a region for writing coverage output. Mean coverage of all bases in each of these windows will be written #cnv.coverage.wsize=1000

##The maximum p-value for the regions to be shown as copy number deletions #cnv.deletions.max.pval=1.0

##The maximum ratio between the coverage of the region and the expected coverage, for the region to be called as cnv deletion #cnv.deletions.max.ratio=0.5

##The minimum mappability percentage for the regions to be shown as copy number deletions #cnv.deletions.min.mappability=50

##The minimum number of windows for the regions to be shown as copy number deletions #cnv.deletions.min.windows=2

##To set the ploidy of all chromosomes for Humans
#cnv.gender=Male

##The maximum p-value for the regions to be shown as copy number insertions #cnv.insertions.max.pval=1.0

##The minimum mappability percentage for the regions to be shown as copy number insertions #cnv.insertions.min.mappability=10

##The minimum ratio between the coverage of the region and the expected coverage, for the region to be called as cnv insertion #cnv.insertions.min.ratio=1.25



##The minimum number of windows for the regions to be shown as copy number insertions #cnv.insertions.min.windows=2

##To indicate whether genome wide normalization or local normalization should be perfomed. false for genome wide and true for chromosome-arm local normalization #cnv.local.normalization=0

##The minimum quality value of the alignments
#cnv.min.quality=2

##The general ploidy of the genome
#cnv.ploidy=2

##List of {contig id: ploidy of the contig} of all the contigs
whose ploidy is different to the general ploidy of the genome
#cnv.ploidy.exception=None

##To set the stringency setting for calling CNVs
#cnv.stringency.setting=Medium

##The distance in kilo bases to be trimmed from the extreme ends
of the chromosome arms
#cnv.trim.distance=1000

##The size of the window block to be considered as a region
#cnv.window.size=5000

##To indicate whether coverage files should be output or not. Enter 1 to write coverage output in \*.wig format. Keep the default value if you do not want to output coverage #cnv.write.coverage=0

**Note:** The parameters for CNV are described in Table 31 on page 189 in Chapter 10, Run a Human CNVs Analysis.

#### Import command

The module INI file typically imports the global.ini file. The effect of the import statement is to include the content of the global.ini file, with all its parameters and defined values, into the module INI file.

**Note:** An imported file must be in the same directory as the INI file. No pathnames are allowed with INI file import statement.

#### .run parameters

Within a INI file, the \* . run parameters control which modules are run. For example, these .run parameters appear in INI files:

saet.run
bamstats.run



fragment.mapping.ini
dibayes.run

The run parameters are set automatically by the shell during standard workflows. Accept the default for most use.

The allowed values for .run parameters are 0 and 1:

- **0**: Do not run the module.
- 1: Run the module analysis (default).

Because mapping is a prerequisite for all other runs, mapping.run may not be set to 0 unless it has already been successfully run to completion. You may set the .run parameters for other modules to 0, under these conditions:

- You do not want to run that module analysis.
- That module analysis has already been completed (for instance, if you are restarting the workflow run and repeating a particular module is not necessary).

When you run in the project repository, the LifeScope<sup>TM</sup> Software command shell resume command does not repeat modules that have successfully completed.

#### The global.ini file

The global.ini file defines parameters for one or more INI files.

If a parameter in the global.ini also appears in an analysis module's INI file, then the value set in the global.ini file is ignored and the analysis module's INI's definition is used. The import global.ini statement must be placed at the top of the an analysis module's INI file.

Paths are not allowed in the import global.ini statement. The global.ini file must be in the same directory as the analysis module INI file that imports it.

**Properties files** A properties file sets the default parameters used by every run of its module type, for all users of the LifeScope<sup>™</sup> Software installation. Properties files are located in the directory <installdir>/etc/plugins/properties.

For example, the file fragment.mapping.properties contains the settings for genomic mapping runs. Example contents for the fragment.mapping.properties file are shown below.

```
wall.time=120
fragmap.number.of.nodes=4
mapping.np.per.node=8
fragmap.minreads.per.node=4000000
fragmap.maxreads.per.node=150000000
fragmap.max.number.of.nodes=100
mapping.memory=15gb
java.heap.space=1500
```

Parameter values defined in a .properties file apply to all runs of that type of module, except that parameter definitions in a INI file override the value set in the properties file. The relationship between the two types of files is explained in "Property files vs. INI files".

XML files

#### Property files vs. Parameter values set in a properties file affect all runs of that module, but may be reset in the module INI file. **INI** files Modifying a setting in the properties file affects all future runs of that module, by all users. Modifying the setting in the fragment.mapping.ini file affects only the current run and any subsequent runs using the INI file. Settings present in the fragment.mapping.ini file take precedence over settings in the properties file. Table 143 compares the influence of parameters in a .properties file and an INI file. Mapping module files are used as a example to demonstrate the general principle relating .properties files and INI files.

File and location	Coverage	Behavior on parameter conflicts
<installdir>/etc/plugins/ properties/ mapping.properties</installdir>	Affects all mapping runs.	Values may be overridden by settings in fragment.mapping.ini.
your global.ini file	Affects all runs (of all modules) if an INI file imports this global.ini file.	Overrides settings in mapping.properties. Values may be overridden by settings in fragment.mapping.ini.
your fragmant.mapping.ini file	Affects all fragment mapping runs that use this INI file.	The parameter value defined in fragment.mapping.ini is used, overriding settings from the mapping.properties and global.ini files.

Table 143 Comparing module properties and INI files (using mapping as an example)

Analysis module In the directory <installdir>/etc/plugins/pipelines are module XML files, one per module, which list all parameters accepted by each module, along with the parameters' default values. Use the XML files as a reference for allowed parameters and their default values.

> Do not edit these XML files in order to change a parameter's value. Instead, modify one of these two files, depending on your purpose:

- **The module's .properties file** To change a parameter for all subsequent module runs for all users, edit the parameter in the module's properties file under <installdir>. See "Properties files" on page 501.
- **Your module INI file** To change a parameter for only your own runs of a module, edit the parameter in your module's INI file. This change affects only analyses that use that INI file. See "PLN files" on page 498.

In either on the two cases above, you may add a parameter that did not previously exist in the file, only if that parameter appears in the module's XML file.

# Set up your own analysis run

The most direct way to customize your own LifeScope<sup>™</sup> Software analysis is to copy an existing workflow. You can then change the workflow to execute your preferred module or sequence of modules. See "Create a new workflow" on page 106 and "Run a standard workflow" on page 103 for more information.

The optional examples download provides single-module analyses. See Appendix E, "Demo Analyses" on page 507 for information on these analyses.



The shell get workflow command copies a workflow's INI files, PLN files, and their directory structure to the local file system. These files are copied to the directory you are in when you open the shell with the lscope.sh command. You can modify the INI and PLN files to customize your own analysis. See Table 5 on page 77 and "Create a new workflow" on page 106.

# LifeScope<sup>™</sup> Software parameters

This section describes parameters the software uses to control your analysis runs. When you run an analysis in the command shell, these parameters can be set automatically by the software. If you create your own INI files, you often need to specify some of these parameters (depending on the type of analysis you are running).

# Table 144 lists parameters that LifeScope<sup>™</sup> Software sets when you run an analysis in the command shell.

Table 144 General parameters

Parameter	Description
analysis.input.readset.file	An RRS file built by the system specifying the XSQ readsets being analyzed (only applies to secondary analysis). RRS is an internal file format.
analysis.genome.reference	Path to the genome reference file.
analysis.sample.name	User-defined sample name.
analysis.input.bams	Comma-separated paths to input BAM files (for tertiary analysis).

# Reference parameters

General

parameters

Table 145 lists reference-related parameters that LifeScope<sup>™</sup> Software sets for you when you use the set reference command with a recognized assembly name. Table 145 Reference-related parameters

Parameter	Description
annotation.dbsnp.file	The file used to annotate SNPs and small Indels. The reference repository includes dbSNP_b130.tab (hg18) and dbSNP_b132_00-All.vcf (hg19).
annotation.gtf.file	The file containing gene and exon annotations corresponding to the genome assembly used in the analysis.
analysis.filter.reference	The file containing a collection of sequences that are used to filter out commonly-occurring motifs and contaminating sequences prior to mapping.
analysis.mirbase.mature.file	The file containing mature form microRNA database annotations.
analysis.mirbase.precursor.file	The file containing precursor microRNA database annotations.
analysis.assembly.name	The name of the genome assembly used in current analysis. Examples: hg18, hg19
analysis.species.name	The name of the species used in current analysis.
	Example: human



#### Table 145 Reference-related parameters (continued)

Parameter	Description
analysis.mappability.dir	The directory containing binary mappability files used by the Human CNV module.

# Execution control parameters

 Table 146 lists parameters used for job management.

 Table 146
 Job management parameters

Parameter	Description
task.temp.dir.delete	Toggles cleanup of intermediate files created in the temp directory.
task.scratch.dir.delete	Toggles cleanup of intermediate files created in the scratch directory
scratch.dir	Location of the scratch directory.
*.run	Parameters that toggle the execution of analysis modules. Examples:
	• cnv.run
	• dibayes.run

# Read-only parameters

Table 147 lists *read-only* parameters that you can use in your INI files to set other parameters. Do *not* set any of these parameters.

Table 147 Read-only parameters

Parameter	Description
analysis.dir	The base analysis directory.
analysis.name	The name of the base analysis.
analysis.output.dir	\${analysis.dir}/outputs.
analysis.temp.dir	\${analysis.dir}/temp.
task.name	The name of the current task. Same as base name of the task's INI file.
task.working.dir	\${analysis.dir}/\${task.name}, the task working directory,.
task.temp.dir	<pre>\${analysis.temp.dir}/\${task.name}, the task temp directory where task writes temporary files. Deleted at the end of a successful run, unless task.temp.dir.delete is set to false.</pre>
task.scratch.dir	The task-specific scratch directory. For cluster installations, this directory is local to the compute node.
task.output.dir	\${analysis.output.dir}/\${task.name}/\${analysis.sample.name}, the task's sample-specific output directory.

# (Optional) The Linux mail command

The LifeScope<sup>TM</sup> Software command shell protects your currently running analyses as well as the configuration and definition of both your projects and your analyses. The running jobs and the configuration are not affected if you close your LifeScope<sup>TM</sup> Software shell and exit your Linux session. You do not need to use Linux commands such as screens or nohup in order to keep your analysis or session window open.
D

The UNIX mail command can be used to send an email notification of a script's completion. Your UNIX mail system must be configured properly. Add the following to your lscope.shrun or lscope.sh resume command:

&& mail -s "subject" <email\_address>

where *subject* is the subject of the email message to be sent and *email\_address* is the email account to be sent the completion message. You must follow UNIX command conventions. Keep the *subject* string short and avoid special characters.

An example of the complete command is:

lscope.sh run my.pln -u user && mail -s "done" me@xyz.com



Appendix D Command Shell Control Files (Optional) The Linux mail command

# Demo Analyses

#### This appendix covers:

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How a demo is setup	508
Run all demos	509
Run a single demo analysis	509
Output files and logs	510

### **Overview**

This appendix describes the demo modules included in an optional download. The demos show how to run an individual LifeScope<sup>TM</sup> Software analysis module, such as mapping or SNPs, using the LifeScope<sup>TM</sup> Software command shell.

The demo analyses are mostly self-contained. Input data and references files are provided. The exception is that some demos use hg18 references files in the LifeScope<sup>™</sup> Software reference repository.

The standard workflows, which are sequences of analyses combined into one run of LifeScope<sup>™</sup> Software, are recommended for general use instead of the demos. See Chapter 7, "Run a Standard Workflow Analysis" on page 97 for information on standard workflows. The demo analyses are based on execution methods used in earlier versions of the software, and have been adapted for the LifeScope<sup>™</sup> Software command shell.

### List of demo modules

The following analyses are included in the demos download:

- Mapping fragmentMapping
- Paired mapping pairMapping
- SNPs diBayes
- Human CNV cnv
- **Inversion** inversion
- Large indel largeIndel
- Small indel smallIndel
- SAET saet
- Enrichment enrichment

- WT fragment wholeTranscriptomeFrag
- WT paired-end wholeTranscriptomePE

Except for whole transcriptome, the demos are configured to run an individual analysis module. The whole transcriptome demos are similar to the whole transcriptome standard workflows, but the demos include the input files and reference files.

### **Demos location**

The LifeScope<sup>™</sup> Software demos are a separate download not included in the main installation. Ask your LifeScope<sup>™</sup> Software administrator for the location of the demos.

### How a demo is set up

This section describes the mechanics of how each demo and its files and directories are organized.

Each demo analysis directory contains a run.sh script, which invokes lscope.sh in scripted mode. The scripted lscope.sh shell commands are contained in a file named run.txt. Here are example run.txt contents, for SNPs:

```
cd projects
mk examples
cd examples
rm diBayes
mk diBayes
cd diBayes
set workflow ./analysis.pln
add bam ./bam/F3-R3-Paired.twoSNPs.bam
add bam ./bam/F3-R3-Paired.twoSNPsmerged.bam
set reference hg18
run
wait
ls
exit
```

Table 148 explains the purpose of the steps shown above.

Table 148	Description of	sample shell	commands	to run a workflow
-----------	----------------	--------------	----------	-------------------

Command	Purpose
cd /projects mk examples cd examples rm diBayes mk diBayes	In the projects repository, create a project named examples, and in that project create an analysis named dibayes. Open the dibayes analysis. <b>Note:</b> The open analysis (cd dibayes, in this case) command is important. The data, reference, and other configuration commands which follow apply only to the current analysis.
set workflow ./analysis.pln	Define the workflow used with this analysis.



 Table 148
 Description of sample shell commands to run a workflow (continued)

Command	Purpose
add bam ./bam/F3-R3-Paired.twoSNPs.bam add bam ./bam/F3-R3-Paired.twoSNPsmerged.bam	Define the BAM input files containing data for this analysis.
set reference hg18	Define the reference for this analysis. The string hg18 matches an assembly name in the reference repository.
run	Start the analysis run.
wait	Wait until the current analysis run completes.
ls	Display status information about the analysis run.

Workflow	The workflow defined in this demo is a PLN file, analysis.pln. A PLN file defines your analysis through a list of tasks, which is a list of one or more INI or PLN files. For each listed INI file, LifeScope <sup>™</sup> Software executes an analysis run of the corresponding module.
	In the SNPs demo, analysis.pln calls tertiary.pln, which contains the entry dibayes.ini to invoke the SNPs analysis module
Input files	The demo uses BAM input, because SNPs is a tertiary analysis. Demos which do mapping analysis instead take XSQ files as input.

### Run all demos

This section describes how to run all the demo analyses in one run. The combined run should complete in under 30 minutes.

The demos require the default reference repository, which includes hg18 and hg19 assemblies.

Follow these steps to execute all the demos in one run:

- 1. Cd to the demo directory.
- **2.** Execute this command in the Linux shell:
  - runall.sh

This script runs each demo one at a time.

The runall.sh script by default executes the run under the lifescope user account. To use a different user name, use the -u and -w options:

runall.sh -u username -w password

### Run a single demo analysis

Each demo is executed with the same command. To execute any demo, follow these steps:

- 1. Cd to the general demo directory, and then cd to the specific analysis directory.
- 2. Execute this command in the Linux shell:

run.sh

The run.sh invokes lscope.sh in scripted mode. The scripted lscope.sh shell commands are contained in a file named run.txt, and are similar to the commands in Table 148.

The major differences in run.txt commands are the following:

- The analysis name
- The input commands, add xsq or add bam
- The set reference command

The runall.sh script by default executes the run under the lifescope user account. To use a different user name, use the -u and -w options:

runall.sh -u username -w password

### **Output files and logs**

The results files are generated in the projects repository, which is set during installation. Within the projects repository directory, the demos results are generated in the following directories:

user\_name/examples/analysis\_name/outputs/sample\_name

This list describes the folder names

- *user\_name* The LifeScope<sup>™</sup> Software user name of the person running the demo.
- examples The project name used by all demos.
- *analysis\_name* The name of the analysis module used by a particular demos. For example: cnv, dibayes, enrichment.
- outputs A hard-coded subfolder name.
- *sample\_name* Determined by the sample name in the input data.

In its results directory, each of the demos analyses creates an output file named summary.log. Check for a success message in the last statement of the log.

The results files are overwritten on subsequent runs executed by the same user. You can view the results of your own demo runs in the LifeScope<sup>TM</sup> Software UI.

# **Administration**

#### This appendix covers:

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### **Overview**

This appendix describes administrative functions that can be performed on either in the Linux shell or the LifeScope<sup>TM</sup> Software command shell.

Most administrative functions performed in the LifeScope<sup>™</sup> Software UI. Refer to the *LifeScope<sup>™</sup> Genomic Analysis Software User Guide* (Part no. 4465696) for more information.

### **Reference file conflicts**

In order to avoid conflicts with other reference files in the reference repository, always add new reference files to the correct assembly subfolder under the internal directory. See "Add new reference files" on page 517 in Appendix G, "The Reference Repository" on page 515.

# Troubleshooting

Message: Read- only db connection	If power was lost, causing the LifeScope <sup>™</sup> Software server to shut down, and the server generates the message "read-only db connection," type the following commands:	
	1. lscope-server.sh stop	
	2. rm /share/apps/lifescope/server/UserDB/db*.lck	
	3. lscope-server.sh start	
	If the problem persists, or if the error message is different (indicating a corrupt db), type the following commands to delete the UserDB, which will remove all the current users in the system. Then recreate the users.	
	1. lscope-server.sh stop	
	2. rm -rf /share/apps/lifescope/server/UserDB	
	3. lscope-server.sh start	
Sequence name not found in	An analysis fails with a reference error, but the reference file is in the reference repository. The error message is similar to the following:	
reference	12 Apr 2011 16:17:41,750 ERROR [0x2a96825860] reference_sam_check: 49 - sequence name chr6 was not found in reference file;	
	If the error is seen with an analysis that previously ran without error, a conflict in the reference repository is possibly the cause. LifeScope <sup>™</sup> Software first looks in the reference repository by assembly name, and then in the assembly folder searches for a reference file that includes the assembly name in its file name. If a file with the assembly name is not found, the software then uses the first file in that folder (alphabetically). The reference selected may or may not contain the required sequence.	
	The avoid this type of error, when adding a new reference file to the repository, always add it under the subfolder for the correct assembly name. Create a new assembly name subfolder under referenceData/internal when adding a new genome. See Appendix G, "The Reference Repository" on page 515 for more information.	
	To recover from this error, you could use the set reference command with an absolute path to the reference file on the Linux file system.	
Read-set repository path: notify users	If you change the path to the read-set repository, notify users to restart LifeScope <sup>™</sup> Software so that the repository can be updated to the new path.	

## Shell admin commands

The table below lists the LifeScope<sup>™</sup> Software shell admin commands. These commands cover user administration and read repository index. These commands require the admin role privilege.

Only admin users have access to the /users category. The admin user commands must be executed at the /users level.

Command	Description		
Administrative user co	Administrative user commands:		
ls	Lists all users.		
mk <i>name</i>	Creates a new user named <i>name</i> , with the password <i>name</i> and the role user.		
rm <i>name</i>	Deletes the LifeScope <sup><math>M</math></sup> Software shell user named <i>name</i> .		
loggedin	Shows logged in users.		
set username <i>role</i> [password]	Sets a user's role and optionally set the user's password. Valid roles are: admin, user		
makepasswdfile	Creates an encrypted password file for non-interactive login. The file is named <pre>lscope.passwd, and is created in the directory where your <pre>lscope.sh shell</pre> command.is originally issued.</pre>		
Other administrative c	ommands:		
rebuild	In /reads, rebuilds the read repository index, which is used to display the list of read-sets in the repository. Recent changes (additions or deletions) to the read repository are not reflected in the 1s output until the index is rebuilt.		

### Reset the admin password

The resetpwd.sh script is provided as an emergency way to reset the administrator's password to its default value. To reset the admin password, follow these steps (at a Linux prompt):

**1.** Stop the LifeScope<sup>TM</sup> Software server:

lscope-server.sh stop

- Reset the admin password: resetpwd.shlifescope
- **3.** Restart the LifeScope<sup>™</sup> Software server.
- 4. In the LifeScope<sup>™</sup> Software UI, change the admin password to a secure password, according to your local policy.

The use of the resetpwd.sh script is not recommended for any user account except the LifeScope<sup>TM</sup> Software admin. Passwords reset for other users may become out of sync the authenticating realm.



# The Reference Repository

This appendix covers:

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Reference file conflicts	516
Add new reference files	517
Prepare reference files	520
Initial repository contents	521

### **Overview**

This appendix describes the LifeScope<sup>™</sup> Genomic Analysis Software reference repository. The repository is created during the pre-installation and installation process, using files shipped on a separate data drive.

You can add new reference files to the repository at any time after installation, but you must follow conventions described in this appendix.

# **Repository structure**

The files on disk that make up the reference repository have the following top-level organization:

- **lifetech** Contains files that have been prepared or formatted to support customizations within LifeScope<sup>™</sup> Software
- **external** Contains files that have been downloaded and placed here verbatim, such as annotations from public repositories.
- **internal** Intended to hold locally derived files to support new genomes and annotations. The internal structure mirrors the lifetech directory.

The next directory level are assembly names such as hg18 and hg19.

For more detail see "Initial repository contents" on page 521.

## **Repository location**

The location (path) of the reference repository is set during the pre-installation and installation process. The location is determined by where the data drive contents are copied, and must match the location answered to the installer question "Enter the location for reference files."

The reference repository location after installation is given by the parameter lifescope.references in the file <*installdir*>/server/server.properties. For example:

```
# Define reference directory
lifescope.references=/data/results/referenceData
```

The location /data/results is an example only. The reference repository location is set during "Copy data drive content" on page 32 in Chapter 2, LifeScope<sup>TM</sup> Genomic Analysis Software Installation, and is the location <LSDF>/referenceData, where <LSDF> is the LifeScope<sup>TM</sup> Software Data Folder that the LifeScope<sup>TM</sup> Software administrator creates during the "Copy data drive content" step.

## **Reference file conflicts**

In order to avoid conflicts with reference files, follow the instructions in "Add new reference files" to create a new folder when adding references for a new genome to the repository.

The behavior for a set reference *assembly\_name* command, such as set reference hg19, is the following:

- LifeScope<sup>™</sup> Software first looks in the reference repository for the *assembly\_name* folder.
- The software searches in the *assembly\_name/reference* folder for a genomic reference files that includes the assembly name in its file name
- In the *assembly\_name* folder, the software searches for reference files that include the assembly name in its file name.
- If a file with the assembly name is not found, or if multiple files are found that both have the assembly name and have the expected file extension, the software then uses the first file (alphabetically) in that folder.

The reference selected may or may not be the one intended.

Assembly names and reference parameters Users are recommended to use the assembly name option with the set reference command, when possible. Using a recognized assembly name automates the specification of reference-related parameters. See Table 5 on page 77 for more information on the set reference command.

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# Add new reference files

	You use the Linux file system, not Lifes reference files. Depending on the local admin permissions to add new reference	Scope <sup>™</sup> Software shell commands, to add new Linux file system configuration, you may need ce files to the repository.
Add by assembly name	The reference repository is organized b choice of folder name is important beca name and is used with the set reference-related parameters.	y subfolders for each assembly name. The ause the folder name matches the assembly ence <i>assembly</i> command to automate setting
	The internal and external folders Follow this convention:	are available for your new reference files.
	<ul> <li>external – Use for files that have b placed here verbatim.</li> <li>internal – Use for locally derived</li> </ul>	een downloaded from public repositories and
	• Internal – Use for locally-derived	files to support new genomes and annotations.
	Example references at internal/uc0 repository location, provide a structure	1 and external/uc01, in the reference e to be followed for new references.
	Follow these steps to add a new referer	nce file to the reference repository:
	<b>1.</b> Find the location of the reference <b>1</b>	repository on the local Linux file system.
	<ol> <li>If necessary, create a new subfolde external) for the new reference the name for the new subfolder.</li> </ol>	er under referenceData/internal (or being added. Use a unique assembly name as
	Mimic the structure of the exampl uc01, or referenceData/exte location, for your new references.	e references at referenceData/internal/ rnal/uc01, in the reference repository
	For example, to add references for following directory: referenceD external/uc02.	an example assembly named uc02, create the ata/internal/uc02 or referenceData/
	<b>3.</b> Follow the instructions in "Prepar new reference files.	e reference files" to validate and convert your
Example steps to add a new assembly	This section lists example steps to add reference repository. This example deso repository. In most installations, file per administrator perform these steps.	a new assembly or a new species to the cribes how to add the rat assembly rn4 to the rmissions require that the LifeScope <sup>™</sup> Software
	Repository location and directories	
	<ol> <li>Identify the reference repository lo <lsdf>/referenceData.</lsdf></li> </ol>	ocation, which in the installation instructions is
	2. Under <lsdf>/referenceData</lsdf>	, create these directories for the new assembly:
	external/rn4	
	external/rn4/ucsc	
	external/rn4/reference	
	external/rn4/dbSNP	(only required for annotations)

external/rn4/mirbase (only required for small RNA analyses) external/rn4/targetEnrichment (only required for targeted analyses) The tables in "Initial repository contents" on page 521 explain the repository directory structure.

#### FASTA file

- 1. Find a FASTA file describing the reference sequence for rat, for instance from an NCBI, UCSC, or Ensembl site.
- 2. Copy the rat FASTA file to a local directory.
- 3. Validate the FASTA file with the reference\_validation.pl script (see "Validate new reference files" on page 520).
- Copy or move the validated FASTA file to external/rn4/reference, for instance, as the file:

external/rn4/reference/rat\_rn4.fa

#### RefGene file

1. Find a RefGene file for the rat assembly. RefGene files are downloaded directly from UCSC, from this website:

http://hgdownload.cse.ucsc.edu

The files have this filename pattern: refGene.\${ASSEMBLY}.\${DATE}.gtf. Convert the RefGene file to GTF, using the refgene2gff.sh described in "Convert a GTF file" on page 520. The output is in the same filename pattern:

refGene.rn4.20110517.gtf

**2.** This step is only required for whole transcriptome analyses. In conjunction with the reference FASTA file, the WT exon extractor and junction extractor modules generate new FASTA files for mapping transcripts to exons and junctions:

external/rn4/reference/rat\_rn4.fa

Copy the new FASTA files to:

external/rn4/refGene/refGene.rn4.20110517.exon.fa
external/rn4/refGene/refGene.rn4.20110517.junctions.fa

#### Regions of interest file

This section is only required for targeted resequencing analyses.

- 1. Find or develop a regions of interest file, in BED format (see "BED file format" on page 471).
- Copy the regions of interest file to the following location and file name: external/rn4/targetEnrichment/targetRegionsRn4.bed

#### miRBase files

This section is only required for small RNA analyses.

1. Find the species-specific microRNA database annotations (precursor file) on the miRBase website:

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#### http://www.mirbase.org/ftp

- Download the mature forms together as an Excel document: http://mirbase.org/pub/mirbase/CURRENT/miRNA.xls
- Save the Excel document as a text file in this location and file name: external/rn4/mirbase/miRNA.txt

#### Filter reference file

- **1.** Find or develop a filter reference file.
- **2.** Copy the filter reference file to a local directory.
- **3.** Validate the filter reference file with the reference\_validation.pl script (see "Validate new reference files" on page 520).
- 4. Copy or move the validated FASTA file to external/rn4, for instance, as the file:

external/rn4/rat\_filter\_reference.fa

### Genomic annotations

Annotation source files such as dbSNP are only required to run the LifeScope<sup>™</sup> Software genomic annotations module.

- **1.** See "Annotation sources" on page 272 in Chapter 14, "Add Genomic Annotations to Analysis Results" on page 267 for information on obtaining these files.
- Copy the dbSNP and other annotation source files to the following directory: external/rn4/dbSNP

### Shell command

When you configure your LifeScope<sup>™</sup> Software command shell analysis, use the following shell command to specify your new reference files:

set reference rn4

This command sets the following parameters with default values for rn4:

- annotation.dbsnp.file
- annotation.gtf.file
- analysis.mirbase.precursor.file
- analysis.mirbase.mature.file
- analysis.filter.reference
- analysis.assembly.name
- analysis.species.name

**Note:** If an assembly of the same name exists in the reference repository lifetech directory, the software finds the lifetech assembly first. In this case you must set the reference parameters individually.

## Prepare reference files

You must perform these procedures on new reference files that you add to the reference repository, before you can use them in a LifeScope<sup>™</sup> Software analysis. Each of these steps is described below. 1. Validate the reference file to a format that complies with  $LifeScope^{TM}$  Software. Convert new GTF files and refGene files that are required for your analyses. **3.** Concatenate the reference files into a single file, if you have more than one, such as when the sequence is separated into single chromosome files. Validate new Reference validation is not performed automatically. You must validate the reference files that you add to the reference repository so that reference files are presented in the reference files correct format to each LifeScope<sup>™</sup> Software module that uses a reference file. To validate a reference file, follow these steps: 1. Log in to the LifeScope<sup>™</sup> Software cluster. Be sure that you log in to Linux with a user name that has execute ('x') privileges on the directory (<installdir>/ bin) that contains the reference\_validation.pl script. The script is under the bin folder where the LifeScope<sup>™</sup> Software is installed. 2. Navigate to the directory that contains the reference\_validation.pl file. **3.** At a command prompt, enter: <installdir>/bin/reference\_validation.pl -f <InputFasta.ext> If the input FASTA file contains IUB codes, the default is to replace the IUBs with Ns. Preserving IUBs is not recommended for use with some tertiary modules. To keep IUBs unchanged, use the -i (keep IUBs) option: reference\_validation.pl -i TRUE -f < InputFasta.ext> The output file is named < InputFasta>\_validated.fa. Convert a GTF file Not all species have exactly the same format of GTF file. Before using a GTF file from a public source with a LifeScope<sup>™</sup> Software module, you must convert the GTF file into a format that is compatible with LifeScope<sup>TM</sup> Software. Convert an Ensembl GTF file The Ensembl website has GTF-formatted genome annotations available for many popular assemblies. Ensembl GTF files are properly normalized by gene and transcript IDs. The site is at: www.ensembl.org Ensembl GTF files use gene accession numbers instead of HUGO-style gene names. Ensembl GTF files also use unprefixed sequence identifiers, such as 1,2,3....X,Y,MT. The Ensembl GTF files are incompatible with genome reference FASTA files that have

UCSC-style sequence IDs with the prefix "chr", for example, chr1, chr2, chr3....chrX,

chrY, chrM.

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### Reformat a Ensembl GTF file

- 1. Log in to the cluster where LifeScope<sup>™</sup> Software is installed.
- **2**. Run the following command:

reformat\_ensembl\_gtf.pl Homo\_sapiens.GR

### Convert a refGene.txt.gz file

The UCSC Genome Browser has genome annotations available for many assemblies at

#### hgdownload.cse.ucsc.edu/goldenPath/

The GTF-formatted annotations available for download are not properly normalized by gene ID. The required content is present for each assembly in the file export of the refGene database table database/refGene.txt.gz.

For example, annotation for human genome build 18 is available at:

hgdownload.cse.ucsc.edu/goldenPath/hg18/database/refGene.txt.gz

**Note:** The GTF-formatted annotation is not in the GTF format required by LifeScope<sup>TM</sup> Genomic Analysis Software. You must convert the annotation before using it in WTA.

Run the bin/refgene2gff.sh script to convert the refGene.txt.gz file:

```
gunzip refGene.txt.gz
refgene2gff.sh -i refGene.txt -o refGene.gff
```

Genome annotations that are downloaded from the UCSC Genome Browser and converted by the annotation conversion script are optimal because they contain Human Genome Organization (HUGO)-style gene names. HUGO-style gene names allow interpretation when using a genome browser or reading reports.

The annotation conversion script works with the latest format of refGene.txt files. Assemblies, such as the rat genome, use an alternative format for the refGene.txt file. The refgene2gff.sh script does not convert alternative formats.

## Initial repository contents

The initial contents of the repository are created from the data drive during the installation process ("Copy data drive content" on page 32).

The following tables describe the files and directories that make up the initial reference repository:

- Table 149 lifetech/hg19
- Table 150 lifetech/hg18
- Table 151 external/hg18 and hg19
- Table 152 external/uc01
- Table 153 internal/uc01

See "Add new reference files" on page 517 to add new reference files to the repository.



### Table 150 describes the reference files in the lifetech/hg19 directory.

Table 149 Contents and purpose of lifetech/hg19 reference files and directories

lifetech/hg19 file or directory	Description
hg19/human_filter_reference.fasta	The file human_filter_reference.fasta is a collection of sequences that are used to filter out commonly occurring motifs and contaminating sequences prior to mapping.
hg19/reference	This directory contains a validated FASTA file describing the reference sequence for human hg19: human_hg19.fa
	An index file with the following name has also been created to describe each entry: human_hg19.fa.fai
	This file contains five columns, with one row for each entry:
	4. Chromosome or contig name (no spaces)
	5. Chromosome or contig length
	6. Cumulative chromosome or contig length
	7. Number of characters per line
	8. Line width (including carriage return)
	The other files are all derivatives of the original file and contain the hash tables used for mapping:
	human_hg19.fa.1001110100111010011101
	human_hg19.fa.101100111110111101
	human_hg19.fa.10110111011101
	human_hg19.fa.11010011101001110100111
	human_hg19.fa.1110100111010011101001
	human_hg19.fa.11101001110100111010011
	human_hg19.fa.11101110110111011
	human_hg19.fa.111110101110101101
hg19/refGene	RefGene files are downloaded directly from UCSC, from this website:
	http://hgdownload.cse.ucsc.edu/goldenness/hg19/database/refGene.txt.gz
	These files have this filename pattern: refGene.\${ORGAMISM}.\${DATE}.gtf.
	RefGene files are converted to GTF (preserving the filename pattern):
	refGene.hg19.20101221.gtf
	A refGene file is then used, in conjunction with the reference FASTA file,
	referenceData/lifetech/hg19/reference/human_hg19.fa
	to generate new FASTA files for mapping transcripts to exons and junctions:
	referenceData/lifetech/hg19/refGene/refGene.hg19.20101221.exon.fa
	referenceData/lifetech/hg19/refGene/refGene.hg19.20101221.junctions.fa



lifetech/hg19 file or directory	Description
hg19/dbSNP/dbSNP_b132_00-All.vcf	This file is from the National Center for Biotechnology Information (NCBI) dbSNP database, which contains information on SNPs and indels already found by other studies. Recent releases of dbSNP are available in VCF (Variant Call Format), such as this file for Human:
	ftp://ftp.ncbi.nih.gov/snp/organisms/human_9606/VCF/v4.0/00-All.vcf.gz
	The VCF file format was developed as part of the 1000 Genomes project. The format is described at this website:
	http://vcftools.sourceforge.net/specs.html
hg19/mirbase/hsa.gff	The microRNA database annotations are derived directly from the miRBase website: http://www.mirbase.org/ftp
	The species-specific precursor GFF file is downloaded from this page:
	http://www.mirbase.org/ftp.shtml
	which resolves into a file such as this (for human):
	ftp://mirbase.org/pub/mirbase/CURRENT/genomes/hsa.gff
hg19/CNV/mappability/ fragment_50.h5	The CNV hg19 mappability file.

Table 149 Contents and purpose of lifetech/hg19 reference files and directories (continued)

### Table 150 describes the reference files in the lifetech/hg18 directory.

Table 150 Contents and purpose of lifetech/hg18 reference files and directories

lifetech/hg18 file or directory	Description
hg18/human_filter_reference.fasta	The file human_filter_reference.fasta is a collection of sequences that are used to
	filter out commonly occuring motifs and contaminating sequences prior to mapping.



lifetech/hg18 file or directory	Description
hg18/reference	This directory contains a validated FASTA file describing the reference sequence for human hg18: human_hg18.fa
	An index file with the following name has also been created to describe each entry: human_hg18.fa.fai
	This file contains five columns, with one row for each entry:
	1. Chromosome or contig name (no spaces)
	2. Chromosome or contig length
	3. Cumulative chromosome or contig length
	4. Number of characters per line
	5. Line width (including carriage return)
	The other files are all derivatives of the original file and contain the hash tables used for mapping:
	human_hg18.fa.111011101101111011
	human_hg18.fa.10110111011101
	human_hg18.fa.11010011101001110100111
	human_hg18.fa.101100111110111101
	human_hg18.fa.1110100111010011101001
	human_hg18.fa.11101001110100111010011
	human_hg18.fa.1001110100111010011101
	human_hg18.fa.111110101110101101
hg18/refGene	RefGene files are downloaded directly from UCSC, from this website:
	http://hgdownload.cse.ucsc.edu/goldenPath/hg18/database/refGene.txt.gz
	These files have this filename pattern: refGene.\${ORGAMISM}.\${DATE}.gtf.
	RefGene files are converted to GTF (preserving the filename pattern):
	refGene.hg18.20090513.gtf
	A refGene file is then used, in conjunction with the reference FASTA file, referenceData/lifetech/hg18/reference/human_hg18.fa
	to generate new FASTA files for mapping transcripts to exons and junctions: referenceData/lifetech/hg18/refGene/refGene.hg18.20090513.exon.fa referenceData/lifetech/hg18/refGene/refGene.hg18.20090513.junctions.fa

### Table 150 Contents and purpose of lifetech/hg18 reference files and directories (continued)

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lifetech/hg18 file or directory	Description
hg18/dbSNP/dbSNP_b130.tab	This file is from the National Center for Biotechnology Information (NCBI) dbSNP database, which contains information on SNPs and indels already found by other studies. This file contains the data from dbSNP version 130, and has the following tab-separated fields:
	1. rsID, the refSNP identifier
	2. Chromosome number
	3. Chromosome start coordinate
	4. Chromosome end
	5. Is SNP?
	6. Function code (consequence to transcript if any)
	The function code is a comma separated list of pairs: locus_id:function code). The FunctionCode attribute is valid only for SNPs and small indels, and are used to annotate SNPs. Codes are based on functional codes from dbSNP, or refSNPs in gene features (from the dbSNP handbook).
hg18/mirbase/hsa.gff	The microRNA database annotations are derived directly from the miRBase website: http://www.mirbase.org/ftp
	The species-specific precursor GFF file is downloaded from this page:
	http://www.mirbase.org/ftp.shtml
	which resolves into a file such as this (for human):
	ftp://mirbase.org/pub/mirbase/CURRENT/genomes/hsa.gff
hg18/CNV/mappability/	The CNV hg18 mappability files.
fragment_25.h5	
fragment_50.h5	
matepair_50x50.h5	

Table 150 Contents and purpose of lifetech/hg18 reference files and directories (continued)

Table 152 describes the hg18 and hg19 reference files in the external directory.

 Table 151 Contents and purpose of external hg18 and hg19
 reference files and directories

external file or directory	Description
hg18/ucsc/refGene.txt.gz,	RefGene files are downloaded directly from UCSC, from these websites:
hg19/ucsc/refGene.txt.gz	http://hgdownload.cse.ucsc.edu/goldenPath/hg18/database/refGene.txt.gz http://hgdownload.cse.ucsc.edu/goldenPath/hg19/database/refGene.txt.gz

external file or directory	Description	
hg18/targetedEnrichment/ targetRegionsHg18.bed	Targeted Regions files are specific to each target capture procedure used during the sample preparation process before the library generation step. As there are many way	
hg19/targetedEnrichment/ targetRegionsHg19.bed	to enrich for targeted regions, including custom preparations, these files are to be provided by the user. This directory is the default location for such files, and BED format is typically used here.	
	Here are example locations and filenames:	
	external/hg18/targetedEnrichment/targetRegionsHg18.bed	
	external/hg19/targetedEnrichment/targetRegionsHg19.bed	
	The BED file format was developed to support the UCSC Genome Browser, and the format is described here:	
	http://genome.ucsc.edu/FAQ/FAQformat.html#format1	
	Note: The target regions BED files on the data drive are 0-length placeholders.	
hg18/mirbase/miRNA.txt,	The microRNA database annotations are derived directly from the miRBase website:	
hg19/mirbase/miRNA.txt	http://www.mirbase.org/ftp	
	The species-specific mature forms are downloaded together as an Excel document:	
	http://mirbase.org/pub/mirbase/CURRENT/miRNA.xls	
	and written as text files on the data drive:	
	referenceData/external/hg18/mirbase/miRNA.txt	
	referenceData/external/hg19/mirbase/miRNA.txt	

 Table 151 Contents and purpose of external hg18 and hg19
 reference files and directories (continued)

### Table 152 describes the uc01 example reference files in the external directory.

### Table 152 Contents and purpose of external uc01 reference files and directories

external uc01 file or directory	Description
dbSNP/b131_SNPChrPosOnRef_37_1.bcp	This set of three files is available from the NCBI dbSNP database, which
dbSNP/b131_SNPContigLoc_37_1.bcp	contains information on SNPs and indels already found by other studies. These files contain the data from dbSNP version 131, and are available from
dbSNP/b131_SNPContigLocusId_37_1.bcp	this website:
	http://www.ncbi.nlm.nih.gov/snp
	Three files are used together and all three are needed.
uc01/targetedEnrichment/ targetRegionsUc01.bed	Targeted Regions files are specific to each target capture procedure used during the sample preparation process before the library generation step. As there are many ways to enrich for targeted regions, including custom preparations, these files are to be provided by the user. This directory is the default location for such files, and BED format is typically used here.
	Here is an example location and filename: external/hg18/targetedEnrichment/targetRegionsHg18.bed
	The BED file format was developed to support the UCSC Genome Browser, and the format is described here:
	http://genome.ucsc.edu/FAQ/FAQformat.html#format1
	<b>Note:</b> The target regions BED files on the data drive are 0-length placeholders.

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Table 152 Cont	ents and purpose	of external <sup>•</sup>	uc01 reference	files and directo	ories (continued)
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external uc01 file or directory	Description
uc01/mirbase/	The microRNA database annotations are derived directly from the miRBase website:
	http://www.mirbase.org/ftp
	which resolves into a file such as this (for human):
	ftp://mirbase.org/pub/mirbase/CURRENT/genomes/hsa.gff
	The mature forms are downloaded together as an Excel document: ftp://mirbase.org/pub/mirbase/CURRENT/miRNA.xls

Table 153 describes the uc01 example reference files in the internal directory. The uc01 is an example assembly. Copy this directory structure to add your own locally-derived assembly to the reference repository, and replace the string uc01 with the name of your new assembly.

Table 153 Contents and purpose of internal uc01 reference files and directories

internal uc01 file or directory	Description	
reference/unicorn_uc01.fa	This directory is for a validated FASTA file describing the genomic reference sequence	
reference/unicorn_uc01.fa.fai	tor ucU1: ucU1.ta	
	( <i>Optional</i> ) An index file with the following name can be created to describe each entry: uc01.fa.fai	
	This file contains five columns, with one row for each entry:	
	1. Chromosome or contig name (no spaces)	
	2. Chromosome or contig length	
	3. Cumulative chromosome or contig length	
	4. Number of characters per line	
	5. Line width (including carriage return)	



**Appendix G** The Reference Repository *Initial repository contents* 



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# Glossary

alignment	The process of mapping sequencing reads to a reference genome or sequence.
alignment browser	An interactive software in which to view alignments of sequencing reads with the reference genome or sequence.
alignment score	Matching score; an optimal alignment is an alignment giving the highest score of matches.
allele	One of two or more alternative nucleotide sequences at the same location on homologous chromosomes.
analysis	A data analysis run in LifeScope <sup><math>TM</math></sup> Software; may include secondary analysis on data from multiple sequencer runs.
analysis type	An <i>analysis</i> based on common genetic-analysis experiment types. LifeScope <sup>™</sup> Software provides three: Genomic Resequencing Analysis, Targeted Resequencing Analysis, and Whole Transcriptome Analysis.
annotated gene	Within one of several reference databases, a gene sequence that has biological attributes attached that describe structure or function, such as coding regions or biochemical function.
annotation	Biological attributes or metadata that are attached to sequence data or files. Examples include: genes and protein-coding features, and verified variants.
BAM file	The compressed binary version of the Sequence Alignment/Map (SAM) format, a compact and index-able representation of nucleotide sequence alignments. The BAM file is generated from the mapping step and has special attributes to support SOLiD <sup>TM</sup> data.
barcode	A short, unique sequence that is incorporated into a library that enables identification of the library during multiplex sequencing.
barcode group, barcode pool	The conceptual grouping used during data analysis if more than one barcode is used during libration preparation from a single biological specimen.
barcoded library	A library that has a unique barcode sequence incorporated that enables identification of the library during multiplex sequencing.
base-space analysis	Analysis of SOLiD sequencing data that has been converted from color space to a nucleotide sequence.

Bayesian	An algorithm that is used to evaluate the probability of the existence of a heterozygote or a non-reference homozygote when the coverage at the position is not high.
bead	The substrate to which the individual strands of DNA molecules in a SOLiD <sup>™</sup> library are attached. The covalently attached strand serves as the template for SOLiD sequencing after the bead is immobilized on the flowchip surface.
BED file	BED format provides a flexible way to define the data lines that are displayed in an annotation track. The required BED fields are:
	chrom - name of the chromosome
	• chromStart - starting position of the feature in the chromosome (the first base in a chromosome is numbered 0).
	<ul> <li>chromEnd - ending position of the feature in the chromosome</li> </ul>
BEDGRAPH	In targeted resequencing, a text file in BEDGRAPH format that describes the depth of coverage within targets by on-target alignments.
bisulfite read	A sequence read of bisulfite-converted DNA, in which unmethylated cytosine residues have been converted to uracil. Used in methylation analysis of genomic DNA.
BLAST	In bioinformatics, Basic Local Alignment Search Tool, or BLAST, is an algorithm for comparing primary biological sequence information, such as the amino-acid sequences of different proteins or the nucleotides of DNA sequences.
call stringency	The criteria for calling a nucleic acid species or genetic variant present in the biological sample.
calling threshold	The criteria for calling a nucleic acid species or genetic variant present in the biological sample.
ChIP-Seq	Sequence analysis of genomic regions associated with DNA-binding proteins.
classic mapping	An algorithm that searches for matches at the full read length, allowing up to a user- specified number of mismatches.
clear zone	The threshold to decide whether a read is mapped uniquely in the reference.
clipping, hard clipping	An operation that codes a portion of a color alignment with a number of mismatches that prevent seed or anchor extension.
clone	A single bead with a clonal population of templates for sequencing, generated during templated bead preparation/emulsion PCR.
color balance	The relative proportion of beads in a given sequencing cycle that are called as each of the four colors.
color call	For each SOLiD chemistry cycle, the color that is determined for each bead.

color space, color- space analysis	Color space data is a sequence of colors obtained from 2-base encoded probes in SOLiD System sequencing, representing a series of overlapping dinucleotides. Color space data is converted to a nucleotide sequence by aligning to a reference.
complexity	The size of the genome that is composed of unique sequences. With respect to SOLiD <sup>TM</sup> libraries, the number of independent molecules derived from the original DNA input, not from amplification.
consensus calls	A file that lists the SNPs called with the SNP Finding algorithm, and provides general information about each position.
contig	A nucleotide sequence that has been assembled from shorter, overlapping sequences.
core	A single core is equivalent to a processor on a traditional computer. Multiple cores increase the performance and efficiency of a computer that is running multiple programs.
counting	Expression analysis that focuses on relative or absolute quantification of RNA molecules in a biological specimen.
copy number variation (CNV)	A variation in the number of copies of DNA segments among individuals in a population. The length of the segment can vary, and it is typically more than a few base pairs. Some use a broader definition that includes any insertion, deletion, or duplication longer than a few base pairs.
CountTags	A tool to count the number of reads that align within genomic features.
counting	Sequence analysis that generates read or tag counts for annotated regions of a reference sequence.
coverage	• In genomic analysis, the number of aligned (or mapped) sequencing reads that span a position in the reference genome.
	• In RNA analysis, this term is sometimes used to describe the fraction of the reference sequence that has sequencing reads aligned (or mapped), at a certain calling threshold.
csfasta file	The SOLiD System produces color-space sequence reads in a fasta-format labeled as CSFASTA. These reads can be retained and analyzed in color space.
dbSNP	A Single Nucleotide Polymorphism database repository for single nucleotide polymorphisms and short insertion and deletion polymorphisms, hosted by the NCBI.
<i>de novo</i> sequencing	The initial generation of the primary genetic sequence of a particular organism, without use of a reference genome or sequence.
deletion	A gap in a nucleotide sequence with respect to a reference genome or sequence.
SNP Finding	Analysis module to identify SNPs from mapped and processed SOLiD System color space reads.

discordant	In large indel analysis, mate-paired reads that have been mapped to a reference genome, and whose inter-read distance deviates significantly from the expected insert size of the mate-paired library as a whole.
	Discordant pairs have insert sizes that deviate significantly from the expected value. Discordant pairs containing a putative deletion appear larger when mapped to the reference.
EBI	European Bioinformatics Institute.
enriched data	A targeted resequencing tool, sequencing data that has been filtered for the target regions of interest.
enrichment statistics	A targeted resequencing tool, the parameters used to assess variations in coverage across all enrichment targets and on a per-target basis.
ENSEMBL	A genome browser. www.ensembl.org.
error correction	(SAET) This step takes each read in the input file and attempts to correct it if any of its k-mers are not present in the spectrum. Use of multi-threaded versioning may be recommended for speed of process.
Error expectation metric (EEM)	A whole transcriptome pipeline, a parameter that is used in the calculation of the Junction Confidence Value metric, to determine the statistical significance of detected variant fusion transcripts.
evidence graph	A visual representation of the data structure of a candidate fusion transcript.
exon	In eukaryotic organisms, a segment of a gene that encodes part or all of a protein. Exons may be separated by introns that are spliced out of the primary transcript to produce a mature mRNA for translation into protein.
exon mapping	A whole transcriptome pipeline, a step in which reads are mapped to the set of exon sequences defined by the genome annotation.
F3 tag	Sequencing data derived from the P1 end of the template in the SOLiD <sup>TM</sup> templated bead, using forward ligation chemistry. The F3 tag is generated using the SOLiD <sup>TM</sup> FWD1 Seq. Primers or the SOLiD <sup>TM</sup> Small RNA Seq. Primers. See the 5500 Series SOLiD <sup>TM</sup> Sequencers: Reagents and Consumables Ordering Guide (Part no. 4465650) for an illustration.
F5 tag	Sequencing data derived from the P2 end of the template in the SOLiD <sup>™</sup> templated bead, using reverse ligation chemistry. The F5 tag is generated using the SOLiD <sup>™</sup> REV1 (DNA) Seq. Primers or the SOLiD <sup>™</sup> REV1 (RNA) Seq. Primers. See the 5500 Series SOLiD <sup>™</sup> Sequencers: Reagents and Consumables Ordering Guide (Part no. 4465650) for an illustration.
false positive	Sequence variants that are present in the output data from a sequencing run but are not present in the biological source.

filter	A reference sequence of interest that is used to select reads that align with or map to the reference sequence for further analysis.
filter mapping	Alignment of sequencing reads to a reference sequence of interest, to select reads for further analysis.
Frequentist	An algorithm for finding SNPs (alternative to SNP Finding algorithm).
function code (dbSNPs)	A code in the dbSNP database that indicates the consequence, if any, of a SNP to the transcript in which it is located.
fusion transcript	An RNA molecule that results from transcription of a gene fusion. See also <i>gene fusion</i>
gapped alignment	A read alignment to the reference sequence that indicates an insertion or deletion. The pairing algorithm searches for gapped alignments (indels) when one of the tags (F3/R3/F5-P2) maps to the reference genome and the other tag does not map to the genome within the insert-size range. Small indel analysis calls indels from a consensus of gapped alignments, using the BAM file as input.
gene fusion	A section of the genome that maps to an exon from one gene followed by an exon from another gene. It can occur as the result of a translocation, deletion, or chromosomal inversion. A gene fusion junction excludes exon-to-exon boundaries that arise from alternative splicing of a transcript.
genomic classification table	For mate-paired or paired-end reads, a code that describes the strandedness, distance, and orientation of the two reads.
genomic mapping	See mapping.
genomics	Global analysis of the genome to discern elements involved in regulation of gene activity or expression, with an emphasis on genetic variation such as single nucleotide polymorphisms, small and large insertions and deletions, and other structural variants such as translocations and inversions. Some use the term genomics to as an umbrella term that includes transcriptomics, epigenomics, and analysis of the genome.
group	A collection of read-sets that are required to be analyzed together, in groups that you define. Reads that are grouped together are treated as one specimen, even if the reads are in different *.xsq input files.
hemizygous	A small indels tool, a parameter that indicates that the reference allele and the small indel are both present.
hg18, hg19	Reference sequence assemblies of the human genome using the nomenclature used by the UCSC Genome Browser.
highly expressed junctions	A whole transcriptome pipeline, high-count sequencing reads that span an exon-exon junction.
homozygous	In diploid organisms, having two identical alleles in the corresponding genes.

HUGO-style gene names	Gene names following the Human Genome Organization style.
HuRef	A diploid human genome sequence of one individual (J. Craig Venter).
indel	A difference in sequence due to either an insertion or a deletion event; especially used when the evolutionary direction of the change is unspecified.
insert size	The physical size of the genomic DNA segments or RNA molecules represented in a SOLiD <sup>™</sup> library.
	• Fragment libraries: the size of the sheared DNA fragments.
	<ul> <li>Mate-paired libraries: the length of the genomic DNA segment spanned by the corresponding mate-pair tags.</li> </ul>
	• Whole transcriptome libraries: the size of the RNA fragments.
insertion	An insertion of nucleotide sequence with respect to the reference genome or sequence.
intron	The genomic sequence between two exons that is spliced out of a primary transcript prior to translation. See <i>exon</i> .
inversion	A segment of DNA that is in its native location but is in the reverse orientation.
junction	A place where two regions that are not contiguous in the genomic sequence are joined in a single sequenced region under consideration.
Junction Confidence Value (JCV) metric	Whole transcriptome pipeline, a statistical confidence metric to detect false positives and assign a confidence level to a detected junction.
junction mapping	Whole transcriptome pipeline, a mapping step that incorporates exon, gene, and transcript definitions in the genome annotation for that region.
junk seed	An initial alignment in a seed-and-extend algorithm that is not aligned with the correct sequence
large indel	An insertion or deletion, with respect to a reference sequence, of more than 200 bp.
LB field	A field in a BAM file that contains library type information.
library	A set of DNA or cDNA molecules prepared from the same biological specimen and prepared for sequencing on the SOLiD System.
local mapping	The initial alignment step that starts with locating short matches between a read and the reference sequence.
locus-spanning	Pairs of reads derived from mate pairs or paired-end reads that span a relatively large distance in the reference sequence.

Map Fragment data tool	Tool that maps sequence data from fragment libraries.
mappability	In the CNVs analysis module, the fraction of mappable bases in the candidate CNV region.
mapping	The process of aligning sequencing reads to a reference genome or sequence.
mate alignment	The process of mapping paired-end or mate-paired reads to the reference sequence, taking into account the proximity of the reads on a template clone.
mate-paired library	Library consisting of two DNA segments that reside a known distance apart in the genome, linked by an internal adaptor, and with P1 and P2 Adaptors ligated to the 5' and 3' ends of the template strand, respectively.
mates	Sequencing reads that are linked through their origin in a mate-paired library or a paired-end sequencing run.
mer	The number of nucleotides in a sequencing read.
merging (WTA)	Algorithm parameter in paired-end WTA analysis that enables the merging of genomic and junction mappings.
methylation analysis	The study of how methylation of nucleic acids is involved in DNA structure and control of gene expression.
module	A virtual piece of LifeScope <sup>TM</sup> Software, such as mapping, SNP Finding, or CNV.
NCBI	National Center for Biotechnology Information.
node	A node is a single computer that is connected to the network. A multi-node cluster has several nodes (computers).
normalization	In general, the process of comparing an experimental measurement to a reference measurement.
	In the CNV analysis module, the process of comparing the relative coverage of a region of interest to the global coverage, based on the human reference genome.
	In targeted resequencing, the process of adjusting the amplicon amounts after PCR enrichment and before library preparation, so that amplicons are represented equally in the library.
	In the whole transcriptome pipeline, the parameter RPKM is a normalized measure of gene expression.
paired-end sequencing	Sequencing runs that acquire sequence from each end of the insert in a DNA fragment or whole transcriptome library, using both forward and reverse reads.
pairing	An algorithm that occurs after mapping in the LifeScope <sup>™</sup> Software workflow

pairing distances	Pairing distances (sometimes called insert sizes) for each pair are assigned during the mapping/pairing pipelines and subsequently used by the large indel tool to determine indel candidacy.
parallelization	A tertiary analysis tool by which you can split spectrum generation into multiple jobs, where each job generates a subspectrum from the subset of reads.
pipeline	Sets of program tools that are used in sequence for different data analysis goals, such as whole transcriptome analysis or methylation analysis.
ploidy	The number of chromosome sets in a cell.
polyploidy	The number of chromosome sets in a cell.
polymorphism	A genetic variant in a population of individuals that may or not may be associated with an observable (phenotypic) trait.
primer set	In the SOLiD System, the set of primers that are used sequentially to initiate ligation sequence chemistry.
project	A project is a container for your analysis runs. You create your projects. Within each project, you create one or more analyses. Each project is private to an individual LifeScope <sup>™</sup> Software user.
properties file	
p-value	The statistical significance of an alignment score is frequently assessed by its p-value, which is the probability that this score or a higher one can occur simply by chance, given the probabilistic models for the sequences.
quality value	An empirically defined value based on a phred-like score equating to the confidence that the color called for that cycle is the correct one. In general, the brighter the bead, the greater the difference in signal between the primary and secondary colors, the higher the quality value (QV).
R3 tag	The R3 tag applies only to mate-paired libraries; sequencing data derived from the mate-pair tag closer to the P2 end of the SOLiD <sup>™</sup> templated bead, using forward ligation chemistry. The R3 tag initiates in the IA sequence using the SOLiD <sup>™</sup> FWD2 Seq. Primers. See the <i>5500 Series SOLiD<sup>™</sup> Sequencers: Reagents and Consumables Ordering Guide</i> (Part no. 4465650) for an illustration.
raw reads	A format in which the nucleotide sequence appears without headers or comments.
read, sequencing read	Sequencing data from a single bead with a single primer set.
read-set	A group of reads belonging to one barcode from one *.xsq file.
read-set group	A collection of similar read-sets designated by a user.

read-set repository	A storage place in LifeScope <sup>™</sup> Software for instrument data intended to be input data for LifeScope <sup>™</sup> Software analyses.
reference, reference genome, reference sequence	A sequence against which sequencing reads are aligned before further bioinformatics analysis.
reference file	A file containing the reference sequence information, usually in the *.fasta format.
RefSeq	A multi-organism database archive of DNA, RNA, and protein sequences, hosted by the NCBI.
repository	A virtual container with your reference files, reads files, and projects. Repositories include:
	• Projects, which contains your projects.
	• Reads (read-sets), which contains your .*.xsq data files.
	• References, which contains the reference genomes used in your analyses.
rescue	An alignment method that is applied to read pairs that have at least one alignment, but no pair of alignments occurring within an expected range. Use the rescue method to find additional alignments.
rescue distance	The maximum distance x for a pair considered to be properly paired (SAM flag 0x2) is calculated by solving equation Phi((x-mu)/sigma)=x/L*p0, where mu is the mean, sigma is the standard error of the insert size distribution, L is the length of the genome, p0 is prior of anomalous pair and Phi() is the standard cumulative distribution function.
resequencing	The process of genomic sequencing in cases where a reference sequence already exists, and the new sequence is compared to the reference.
resource manager	Software for executing batch and interactive jobs on a cluster of networked computers.
RNA-Seq	Gene expression analysis using sequence-based approaches. RNA-Seq can include whole transcriptome analysis, small RNA analysis, and SOLiD SAGE <sup>TM</sup> analysis.
RPKM	The number of reads mapping to a transcript per kilobase of transcript length per million mappable reads. RPKM is used to set a threshold for calling a transcript or new RNA species or isoform "present." 1 RPKM is equivalent to 20 reads mapping to a 1 kb transcript per $20 \times 10^6$ mappable reads.
SAET	SOLiD Accuracy Enhancer Tool for correcting spectral-alignment errors in raw data; reduces color-calling error rate without alignment to reference genome. SAET takes as input a file with SOLiD reads in .csfasta format and outputs corrected reads and quality values into new .csfasta and .qual files, respectively.

SAGE <sup>™</sup> analysis	(Serial Analysis of Gene Expression) Nucleotide sequence analysis seeking to find specific gene expression information using short stretches of cDNA (also known as <i>tags</i> ) from the 3' ends of RNA molecules. In the SOLiD System, the SAGE tag is 25–27 bp in length.
sample, barcoded sample	In the 5500 Series SOLiD <sup>™</sup> ICS, the set of templated beads that will be sequenced in a single flowchip lane. A barcoded sample contains templated beads from up to 96 barcoded libraries.
SAMtools	SAMtools provide various utilities for manipulating alignments in the SAM format, including sorting, merging, indexing and generating alignments in a per-position format. SAMtools is hosted by SourceForge.net.
Satay plot	A cross-axis graph displaying the spectral purity and signal intensity of beads in a sequencing run that meet defined thresholds in each parameter ( <i>on-axis beads</i> ). <i>Best beads</i> meet the highest stringency thresholds; <i>good beads</i> meet less stringent thresholds.
secondary analysis	Sequence alignment and mapping to a reference sequence.
seed	A field: size of seed used in spectrum construction. Seeding: Using a heuristic method, finding homologous sequences, not by comparing either sequence in its entirety, but rather by locating short matches between two sequences.
seed-and-extend	Seed-and-extend algorithms map reads to a reference genome. Certain softwares can map reads with any number of differences or mismatches using the observation that for an r bp read to align to the reference with at most k differences, the alignment must have a region of length s=r/k+1 called a seed that exactly matches the reference.
sense strand	The strand of DNA with the same nucleotide sequence as that of the corresponding mRNA. Also called the <i>coding strand</i> .
sequence trimming	See <i>trimming</i> .
sequencing sample	A physical pooling of one or more library preparations before templated bead preparation. The sequencing sample can be used on one or more lanes on a flowchip.
single read	Forward-only sequence data that generates the F3 tag.
small indel	An insertion or deletion, with respect to a reference sequence, of less than 10 base pairs in length.
small RNA analysis	Global sequence analysis of the small RNA population of a cellular RNA sample; small RNA includes microRNAs (miRNAs), short interfering RNAs (siRNAs), piwi- interacting RNAs (piRNAs), and repeat-associated siRNAs (rasiRNAs).
small RNA library	A SOLiD <sup>™</sup> System-compatible library that is prepared from the small RNA fraction of a total RNA sample.
SNP	Single Nucleotide Polymorphisms (SNPs); single base pair variants in genomic DNA or the corresponding RNA transcript.

spectrum building	(SAET) When a temporary file with spectrum constructed from all reads generates in the output directory. This process is fast, resource less, and creates temporary swap files when memory usage is above 2GB.
splice junction	Exon-to-exon boundaries that arise from alternative splicing of a transcript. See also <i>gene fusion</i> .
splicing	The process whereby introns are removed from a primary mRNA, resulting in a mature mRNA that is ready for translation into protein.
strandedness	The polarity or orientation of a nucleic acid strand with respect to being sense or antisense. Libraries prepared using the SOLiD Total RNA-Seq Kit preserve the strandedness of the original RNA molecule such that F3 tag reads align to the sense strand and F5 tag reads align to the antisense strand.
stringency	Stringency is one criterion used to filter out junk reads in ReadBuilder stage. Specifically, in the SNP Finding tool, the stringency parameter (call.stringency) defines the criteria by which to report SNPs in analyzing genomic data. There are four empirical stringency settings (low, medium, high, highest).
tag	A tag is:
	<ul> <li>Sequencing data from a single bead with a single primer set; sometimes used interchangeably with read, sequencing read.</li> </ul>
	• A length of DNA or cDNA to be sequenced; especially, a relatively short stretch of DNA or cDNA that is used to infer information about the longer native molecule from which it is derived, such as in mate-paired library sequencing and SAGE <sup>TM</sup> analysis, respectively.
tags: BC, F3, F5, R3	Sequencing data derived from specific locations on the SOLiD <sup>™</sup> templated bead. See the 5500 Series SOLiD <sup>™</sup> Sequencers: Reagents and Consumables Ordering Guide (Part no. 4465650) for an illustration.
targeted resequencing	Comparative sequence analysis of selected candidate genes or regions to discover genetic variants and mutations.
template strand	The strand of each DNA molecule that is covalently attached to SOLiD <sup>™</sup> P1 beads during templated bead preparation and that serves as a template for SOLiD <sup>™</sup> System sequencing.
tertiary analysis	Data analysis that takes place after mapping and alignment.
threading	Threading is a technique that tries to match a target sequence on a library of known three-dimensional structures by "threading" the target sequence over the known coordinates. In this manner, threading tries to predict the three-dimensional structure starting from a given protein sequence. It is sometimes successful when comparisons based on sequences or sequence profiles alone fail due to a too low sequence similarity.
transcript rescue distance	See <i>rescue distance</i> .

transcriptome	The compilation of all transcribed sequences from a genome, both coding and non-coding.
transition	A single nucleotide (point) mutation that changes a purine nucleotide to another purine nucleotide (for example, A to G), or a pyrimidine nucleotide to another pyrimidine nucleotide (for example T to C).
translocation	A mutation in which a chromosomal segment changes position, usually moving from one chromosome to a different, nonhomologous chromosome.
transversion	A single nucleotide (point) mutation that changes a purine nucleotide to a pyrimidine nucleotide (for example, A to C), or a pyrimidine nucleotide to a purine nucleotide.
trimming	Filtering noise algorithmically which removes 5' overhangs and shortens 3' overhangs, typically to 4–5 bases. Trimming reads requires both modifying the .csfasta reads file to replace the color to be trimmed with a dot ("."), and modifying the mapping parameters to account for the alignment change.
trusted frequency (SAET)	(SAET) Use this developer option (-trustfreq freq) to overwrite estimated frequency cutoff of trusted seeds. All seeds with frequency < "freq" are filtered out of spectrum.
trusted seed	(SAET) If globally computed cutoff for frequency of trusted seeds does not meet your purpose, e.g., it is too low and too many junk seeds are considered correct or it is too high and many correct but low frequency seeds are filtered out, then use -trustfreq option to overwrite estimated frequency cutoff.
ungapped alignment	See gapped alignment.
uniquely placed reads	A read that is mapped only once in a genome with a given number of mismatches.
variant	A difference in the nucleotide sequence of interest, with respect to the reference sequence.
visualization	Viewing mapped reads on any number of publicly available genome browsers.
weighting	A weighting scheme can outperform a simple binary scheme traditionally applied in genomic analysis, independent of the organism, and used to improve the quality of the data.
whole transcriptome analysis	Global sequence analysis of coding and non-coding RNA transcripts along their entire length.
whole transcriptome library	A SOLiD <sup>™</sup> System-compatible library that is prepared from total or poly(A) RNA that enables sequence analysis of the transcripts along their entire length.
workflow	Configuration files that are used for common end-to-end analysis runs, including multiple modules. LifeScope <sup>™</sup> Software runs these common workflows: resequencing, targeted resequencing, whole transcriptome, Methyl Miner, and ChIP-Seq.

WTA	See whole transcriptome analysis.
*.xsq	eXtensible SeQuence, an extensible file format for storing sequence data. A binary sequence output file generated by the instrument. This file contains primary analysis results for a single lane in a flowchip.
z-normalize	Parameters called moving statistics, such as the number of locus-spanning pairs, and the sample average insert size, are calculated from a subset of pairs at each genomic position, yielding the absolute insert size deviation between the sample and the population in units of standard deviation. This normalization step allows multiple libraries with variable insert sizes to be combined into one analysis.
zygosity	The combination of <i>alleles</i> at a site in a nucleotide sequence; for example, homozygous reference allele, homozygous non-reference allele, or heterozygous.

# Documentation

# **Related documentation**

For the latest documentation on LifeScope<sup>™</sup> Genomic Analysis Software, go to:

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Document	Part number	Description
LifeScope™ Genomic Analysis Software User Guide	4465696	Describes the bioinformatics analysis framework for flexible application analysis (data-generated mapping, SNPs, count reads, etc.) from sequencing runs.

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