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How to use this guide

Text conventions
This guide uses the following conventions:

- **Bold** text indicates user action. For example:
  Type 0, then press **Enter** for each of the remaining fields.

- **Italic** text indicates new or important words and is also used for emphasis. For example:
  Before analyzing, *always* prepare fresh matrix.

- A right arrow symbol (▶) separates successive commands you select from a drop-down or shortcut menu. For example:
  Select **File ▶ Open ▶ Spot Set**.
  Right-click the sample row, then select **View Filter ▶ View All Runs**.

User attention words
Two user attention words appear in Life Technologies user documentation. Each word implies a particular level of observation or action as described below:

<table>
<thead>
<tr>
<th>Note:</th>
<th>– Provides information that may be of interest or help but is not critical to the use of the product.</th>
</tr>
</thead>
<tbody>
<tr>
<td>IMPORTANT!</td>
<td>– Provides information that is necessary for proper instrument operation, accurate chemistry kit use, or safe use of a chemical.</td>
</tr>
</tbody>
</table>
How to obtain support

For the latest services and support information for all locations, go to:

www.appliedbiosystems.com

At the Life Technologies web site, you can:

- Access worldwide telephone and fax numbers to contact Life Technologies Technical Support and Sales facilities
- Search through frequently asked questions (FAQs)
- Submit a question directly to Technical Support
- Order Life Technologies user documents, MSDSs, certificates of analysis, and other related documents
- Download PDF documents
- Obtain information about customer training
- Download software updates and patches
Chapter 1

SETS Software Overview

This chapter covers:
- SOLiD™ 4 System overview ........................................... 10
- SETS Software overview ........................................ 12
- SETS homepage ................................................ 13
SOLiD™ 4 System overview

IMPORTANT! The features presented in this document are related to the SOLiD™ 4 System SETS v4.0.1.

SOLiD™ system technology

The Applied Biosystems SOLiD™ 4 System provides parallel sequencing of clonally amplified DNA fragments linked to magnetic beads. The sequencing methodology is based on sequential ligation with dye-labeled oligonucleotides. All fluorescently labeled oligonucleotide probes are present simultaneously, competing for incorporation. After each ligation, fluorescence is measured before another round of ligation takes place.

This ligation-based chemistry eliminates dephasing. The use of a two-base encoding mechanism, which interrogates each base twice for errors during sequencing, discriminates between true polymorphisms and system noise. The SOLiD™ 4 System software allows customers to monitor runs in real-time, and provides basic data analysis tools.

The SOLiD™ 4 System consists of:

- SOLiD™ 4 Analyzer
- All ancillary equipment
- ICS software, which is described in the Applied Biosystems SOLiD™ 4 System ICS Software Help
- SOLiD™ Experiment Tracking System (SETS) software, which is described in this document
The following shows the naming convention for the sequencing runs.

- Fragment
- Paired-end
- Mate-pair
- Multiplex fragment
- Multiplex paired-end
- WFA (not shown below)

Selecting a Fragment run and then selecting Multiplexing will result in creating a barcoded fragment library. The same concept applies to the Paired-End run.
Overview

The SOLiD™ Experiment Tracking System (SETS) software is a web-based application for viewing real-time data and completed run analysis reports from the SOLiD™ 4 System. The SOLiD™ 4 Analyzer uses ligation-based sequencing, then filters data to remove missing calls, nonligations, and/or redundancies.

Data are analyzed at two levels:

• Primary analysis, which involves:
  – Image alignment
  – Color assignment
  – Color calls
  – Quality value assignments

• Secondary analysis, which compares color space calls to a reference sequence.

Note: Applied Biosystems recommends performing the secondary analysis off-instrument with BioScope™ Software. Refer to BioScope™ Software for Scientists Guide: Data Analysis Methods and Interpretation (PN 4448431).

IMPORTANT! Do not install BioScope™ Software on the SOLiD™ 4 Analyzer.

Opening SETS

Access SETS by typing in the SETS software URL supplied by the Applied Biosystems Field Service Engineer:
Chapter 1 SETS Software Overview

SETS homepage

This address opens the SETS software to the homepage and you are automatically logged-in as a “guest” user. As a guest, you can only view analyzed data and reports. Log in to the software to perform other functions. See “Log in to SETS” on page 21.

The SETS software opens the homepage where you can view real-time and completed-run data generated by the SOLiD™ 4 Analyzer.

The homepage is divided into three sections:

- **Top section**: Main menu and error messages (when present)
- **Middle section**: In-progress or the most recent runs (per flowcell)
- **Base section**: Recent analysis
Error messages

When there are error messages, the Error Messages section displays them at the top of the homepage after you click the View Messages arrow.

You can scroll through the messages, refresh the view for all messages received, or remove one message, or all messages from the field, as shown.

For more information on log messages see “System Logs” on page 61.

In progress or most recent run

This section displays the status of a current run or results for the most recently completed run. Flowcell 1 displays on the left-hand side of the window; Flowcell 2 displays on the right side.

To understand more about your run, click More Information within each flowcell section.
Run metrics is quality metrics in the form of traffic lights to help you assess the progress and measurable quality of the run. “More Information” provides a view of analyzed data for a particular run. For details on monitoring a run, see Chapter 3, Monitor the Run in SETS.

Note: At any time you can return to the SETS homepage by clicking the Home button on the main menu at the top of the screen.

Recent analyses
This section displays the progress of data analysis for individual samples.

For every analysis, you can see:
- Run name
- Sample name
- Date created
- Last closed stage
Chapter 2

Prepare Run Settings

This chapter covers:
- ICS, SETS, and BioScope™ Software applications ....... 18
- SOLiD™ 4 System Software workflow ................. 19
- Log in to SETS ........................................ 21
- Set analysis parameters .................................. 22
- Load a Reference Sequence ......................... 37
ICS, SETS, and BioScope™ Software applications

To understand the relationship between the three software components in the SOLiD™ 4 System and how a user would interact between them in a typical workflow, see the “SOLiD™ 4 System Software workflow” on page 19.

Analysis settings

Before you start a run on the SOLiD™ 4 Analyzer using the Instrument Control Software (ICS), open the SETS software and set your primary analysis and the secondary analysis settings. See “Set analysis parameters” on page 22.

IMPORTANT! For more complex genomes (such as the human genome), barcoded samples, and larger reference (and reads) the secondary analysis should be performed off-instrument with the BioScope™ Software. Refer to BioScope™ Software for Scientists Guide: Data Analysis Methods and Interpretation (PN 4448431). If a sample contains more than the maximum allowable number of reads, the SETS software selects only up to the maximum allowed number of reads and the rest will not be included in the results of the secondary analysis.

IMPORTANT! Do not install BioScope™ Software on the SOLiD™ 4 Analyzer.

The Secondary analysis (sequence data alignments to a reference sequence) will not occur without a preloaded reference sequence. Perform secondary and tertiary analysis with the BioScope™ Software. See “Load a Reference Sequence” on page 37.
SOLiD™ 4 System Software workflow

The following figure shows the relationship workflow between ICS, SETS, and BioScope™ Software.
Log in to SETS

Logging in  The default login status for the SOLiD™ SETS Software is a “guest” user. In the guest role, you can only view real-time run and completed-run data. To perform administrative functions (edit, reanalyze, or delete), log in as “admin” or “user.”

Note: The current instrument and user name always displays in the upper right corner of the SETS software homepage. For example:

To log in to SETS:

1. Click on the main menu to open the Login screen.

   ![Login Screen](image)

2. Enter the user name and assigned password in the Log In dialog.

   Note: A Field Service Engineer (FSE) will assign the Administrator name and password at the time of installation. The “admin” can then assign new users. For more information, see “Manage Users” on page 130.

3. Click Submit.
Login permissions

The permissions for each login status are:

- Default login as “guest” allows you to view real-time and completed-run data
- Log in as “user” allows you to:
  - View real-time and completed-run data
  - Edit analysis settings
  - Reanalyze data (primary and secondary reanalysis)
  - Export runs
  - Delete runs
- Log in as “admin” allows you to perform all user actions and:
  - Load reference sequences
  - Manage users

Set analysis parameters

You can set parameters by selecting primary analysis settings and the application type (or secondary analysis settings and the application type), before starting a run in ICS.

You can perform filtering during the primary analysis.

IMPORTANT! For more complex genomes (such as the human genome), barcoded samples, and larger reference (and reads) the secondary analysis should be performed off-instrument with the BioScope™ Software. Refer to BioScope™ Software for Scientists Guide: Data Analysis Methods and Interpretation (PN 4448431).

If a sample contains more than the maximum allowable number of reads, the SETS software selects only up to the maximum allowed number of reads and the rest will not be included in the results of the secondary analysis.

IMPORTANT! Do not install BioScope™ Software on the SOLiD™ 4 Analyzer.
Chapter 2  Prepare Run Settings

Prepare Run Settings

Set analysis parameters

The Secondary analysis (sequence data alignments to a reference sequence) will not occur without a preloaded reference sequence. Perform secondary and tertiary analysis with the BioScope™ Software. See “Load a Reference Sequence” on page 37.

<table>
<thead>
<tr>
<th>Sample Name</th>
<th>Library</th>
<th>Primary Analysis</th>
<th>Application</th>
<th>Secondary Analysis</th>
<th>Description</th>
<th>Remove</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sample1</td>
<td>default library</td>
<td>default primary</td>
<td></td>
<td>SINGLE_TAG_DIII...</td>
<td></td>
<td>✗</td>
</tr>
<tr>
<td>Sample2</td>
<td>default library</td>
<td>default primary</td>
<td></td>
<td>SINGLE_TAG_DIII...</td>
<td></td>
<td>✗</td>
</tr>
<tr>
<td>Sample3</td>
<td>default library</td>
<td>default primary</td>
<td></td>
<td>SINGLE_TAG_DIII...</td>
<td></td>
<td>✗</td>
</tr>
<tr>
<td>Sample4</td>
<td>default library</td>
<td>default primary</td>
<td></td>
<td>SINGLE_TAG_DIII...</td>
<td></td>
<td>✗</td>
</tr>
</tbody>
</table>

Define primary analysis settings

In SETS, you can define the traffic light thresholds as a part of your setup for primary analysis.

To modify the threshold settings:

1. Log into SETS.

2. Go to Main Menu
   > Analysis
   > Primary Analysis Settings.

3. Click [New...] to begin modifying the threshold settings.

   ![Primary Analysis Settings](New.png)

   **Note:** Click ✗ to delete an existing setting or ✔ to edit one.

4. Enter a new value for the Low or High percentage setting for any of the following parameters:
   - Stringency
   - Panel alignment
   - HQ bead
   - Bead retention
Chapter 2 Prepare Run Settings
Set analysis parameters

- Color balance
- Quality value
- Blur value
- Blur population
- Exposure value
- Exposure population
- Exposure time

**Note:** Refer to “Primary analysis parameters” on page 25 for details on these parameters.

<table>
<thead>
<tr>
<th>Primary Analysis Setting</th>
<th>Owner</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>None</td>
<td></td>
</tr>
</tbody>
</table>

**Minimum Calls**
- 25

**Stringency**
- 20

**Barcode Error Correction Level**
- 1

<table>
<thead>
<tr>
<th>Low and high thresholds for traffic light parameters:</th>
<th>Low</th>
<th>High</th>
</tr>
</thead>
<tbody>
<tr>
<td>Panel Alignment</td>
<td>70%</td>
<td>80%</td>
</tr>
<tr>
<td>Read Retention</td>
<td>80%</td>
<td>95%</td>
</tr>
<tr>
<td>HQ bead</td>
<td>15%</td>
<td>85%</td>
</tr>
<tr>
<td>Color Balance</td>
<td>5</td>
<td>8</td>
</tr>
<tr>
<td>Quality Value</td>
<td>15.0</td>
<td>20.0</td>
</tr>
<tr>
<td>Blur Value</td>
<td>2.2</td>
<td>2.8</td>
</tr>
<tr>
<td>Blur Population</td>
<td>75%</td>
<td>90%</td>
</tr>
<tr>
<td>Exposure Value</td>
<td>0.15</td>
<td>0.15</td>
</tr>
<tr>
<td>Exposure Population</td>
<td>9</td>
<td>90</td>
</tr>
<tr>
<td>Exposure Time</td>
<td>200.0</td>
<td>500.0</td>
</tr>
</tbody>
</table>

**Advanced parameters**
- Save
- Restore Defaults
- Cancel

* = required fields
5. Click **Save** to keep the new settings.

   **Note:** Click **Restore Defaults** at any time to reestablish the original (and recommended) default settings for primary analysis.

**Primary analysis parameters**

Primary analysis parameters control the traffic light “signals,” or thresholds that you will use in determining the quality of a run cycle, even to stop a run midstream if the traffic lights signal a poor quality run cycle.

The value for each setting is accessible in SETS and modifications can be made to increase or decrease each value for a custom run setting.

**Target values for parameters**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Target Value for Primary Analysis</th>
</tr>
</thead>
<tbody>
<tr>
<td>Stringency</td>
<td>20</td>
</tr>
<tr>
<td>Panel Alignment</td>
<td>≥90%</td>
</tr>
<tr>
<td>HQ Beads</td>
<td>≥35%</td>
</tr>
<tr>
<td>Bead Retention</td>
<td>≥95%</td>
</tr>
<tr>
<td>Color Balance</td>
<td>≤5%</td>
</tr>
<tr>
<td>QV Median</td>
<td>≥20</td>
</tr>
<tr>
<td>Blur Value</td>
<td>≤2.2</td>
</tr>
<tr>
<td>Blur Population</td>
<td>≤75%</td>
</tr>
<tr>
<td>Exposure Value</td>
<td>≤0.05</td>
</tr>
<tr>
<td>Exposure Population</td>
<td>0</td>
</tr>
<tr>
<td>Exposure Time</td>
<td>≤200</td>
</tr>
</tbody>
</table>
## Parameters descriptions

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Stringency</strong></td>
<td>Stringency is one criterion used to filter out junk reads in ReadBuilder stage. For filtered reads, csfasta and QV.qual are stored in the reject folder instead. The valid range of stringency is from 0 to 80. Any stringency less than 0 is treated as 0, where all reads will pass the stringency threshold. If stringency is set larger than 80, maximal possible stringency of 80 is used to avoid over filtering good beads.</td>
</tr>
<tr>
<td><strong>Panel Alignment</strong></td>
<td>Panel alignment refers to the ability of the primary analysis process to be able to take a channel ligation image and successfully match it to the corresponding focal map image. The stage does not reset to the exact same position in either the X, Y, or a combination of the two directions. If there is a Z plane difference, the focus can also change the characteristics of the image. A lack of signal from either a failure in chemistry or an error in reagents can also lead to insufficient bead information required to complete the analysis. There are alignment scores for each pairwise combination of ligation and focal map. A panel that fails to align has gone past a critical threshold. The scores, available in the underlying cycle *.PA.tab files, contain information about the panel specific information. The panel alignment metric is a measure of the panels that failed to align, resulting in the loss of a cycle’s worth of information for a given panel count of beads. When seeing a decrease in panel alignment, explore other image quality metrics reported such as: blur value and population, exposure value, and times.</td>
</tr>
</tbody>
</table>
HQ Beads

When visualizing the beads on a scatter plot, those that are on the axis and far from the origin are the beads that have a bright signal in only one channel. High quality beads are a summary of the “good” and “best” beads that are reported in the Cycle Scans Report (available in both ICS and SETS).

There are two concept definitions necessary: intensity and angle.

The concept of intensity is defined as the magnitude of the scaled intensity vector:

The concept of angle is defined as 1 - \( \cos(\theta) \), where \( \theta \) is the angle (in four-dimensional scaled intensity space) between the bead intensity vector and the axis of the called color.

The intensity value can increase due to the total intensity across all channels independent of the intensity of the primary channel alone.

Considering the angle and the intensity, a “good” bead is a usable bead that satisfies the following conditions:

- intensity >0.05
- 0.0125 < angle \( \leq 0.05 \)

A “best” bead is a usable bead that satisfies these conditions:

- intensity >0.05
- angle \( \leq 0.0125 \)

A “best” bead is not a “good” bead because these are exclusive classes.

The set of high quality beads, for this metric, is the merging of the two classes.

These values are also highly correlated to the quality values (QV) for calls. A drop in the percentage of beads falling into the HQ category indicates that an accurate color calling is becoming more difficult.

Bead Retention

The number of usable beads in the analysis of a panel for any particular ligation cycle is nearly always smaller than the total number of beads found in the corresponding focal map image.

Bead retention is simply the fraction of the total number of beads that are usable.

Because the stage can drift slightly from one slide scan to the next, beads near the edges of a panel might not be imaged in all four colors.

The software will call a color only for bead locations that are imaged in all four color-scans of a ligation cycle (i.e. any usable bead).

Since beads are identified, or found, only during the focal map analysis stage, a call is made for such a bead location, even if the bead has physically moved from that position. In such cases, since the bead location is most likely now vacant, subsequent color calls will be based on data dominated by noise or, if present, a close neighbor bead. As such beads are still classified as usable, bead retention is not sensitive to such dislocations.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>HQ Beads</td>
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</tr>
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<td></td>
<td>- intensity &gt;0.05</td>
</tr>
<tr>
<td></td>
<td>- 0.0125 &lt; angle ( \leq 0.05 )</td>
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</tr>
<tr>
<td></td>
<td>- angle ( \leq 0.0125 )</td>
</tr>
<tr>
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<tr>
<td></td>
<td>The software will call a color only for bead locations that are imaged in all four color-scans of a ligation cycle (i.e. any usable bead).</td>
</tr>
<tr>
<td></td>
<td>Since beads are identified, or found, only during the focal map analysis stage, a call is made for such a bead location, even if the bead has physically moved from that position. In such cases, since the bead location is most likely now vacant, subsequent color calls will be based on data dominated by noise or, if present, a close neighbor bead. As such beads are still classified as usable, bead retention is not sensitive to such dislocations.</td>
</tr>
</tbody>
</table>
Color Balance

Because of the 2-base encoding utilized in the SOLiD™ 4 System, changes in underlying base balance should not significantly skew the color balance. In fact, the complete absence of a base would still leave all four colors present in approximately equal distribution.

Color balance is the relative proportion of beads in a given cycle that are called as each of the four colors.

Changes in the color balance can indicate that there are problems in the sequencing run.

Signal drops, or absence in one channel, can be large enough that normalization does not compensate, resulting in a significant drop in beads called for a channel.

It is important to note that library preparation methods can create either positive or negative selections that do change the color distribution at significant cycles (cycles corresponding to primer attachments or at the ends of a tag).

If a shift in color balance is out of expectation, looking at imaging metrics—exposure in particular—for the channel that is either too high or low can explain the problem.

If the images, alignment, exposure, quality values, and HQ beads all look within norms, then the color balance shift may be the result of the sample.

For the color balance calculation, the colors are assumed to be in equal proportion.

The real expected color balance would more accurately be determined by the dinucleotide frequency of the underlying DNA source. A target value of 6% would mean that all channels remain within +/- 6% of the 25% “ideal.”

Quality Value

Every bead has a color call and a quality value assigned to it after every cycle.

The quality value is 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 and the higher the QV. This is similar to the HQ beads discussed previously, but where the HQ classification tracks the number of beads that fall into that threshold, the QV metric tracks the quality distribution of all the beads for a given cycle.

A quality value is associated with beads that are in panels that do not align, or with beads that have no color call.

The median QV of all calls is tracked and reported.

In general, a higher QV is assigned when a bead is brighter, because the brighter the bead, the greater the differences between the primary and secondary color calls.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Color Balance</td>
<td>Because of the 2-base encoding utilized in the SOLiD™ 4 System, changes in underlying base balance should not significantly skew the color balance. In fact, the complete absence of a base would still leave all four colors present in approximately equal distribution. Color balance is the relative proportion of beads in a given cycle that are called as each of the four colors. Changes in the color balance can indicate that there are problems in the sequencing run. Signal drops, or absence in one channel, can be large enough that normalization does not compensate, resulting in a significant drop in beads called for a channel. It is important to note that library preparation methods can create either positive or negative selections that do change the color distribution at significant cycles (cycles corresponding to primer attachments or at the ends of a tag). If a shift in color balance is out of expectation, looking at imaging metrics—exposure in particular—for the channel that is either too high or low can explain the problem. If the images, alignment, exposure, quality values, and HQ beads all look within norms, then the color balance shift may be the result of the sample. For the color balance calculation, the colors are assumed to be in equal proportion. The real expected color balance would more accurately be determined by the dinucleotide frequency of the underlying DNA source. A target value of 6% would mean that all channels remain within +/- 6% of the 25% “ideal.”</td>
</tr>
<tr>
<td>Quality Value</td>
<td>Every bead has a color call and a quality value assigned to it after every cycle. The quality value is 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 and the higher the QV. This is similar to the HQ beads discussed previously, but where the HQ classification tracks the number of beads that fall into that threshold, the QV metric tracks the quality distribution of all the beads for a given cycle. A quality value is associated with beads that are in panels that do not align, or with beads that have no color call. The median QV of all calls is tracked and reported. In general, a higher QV is assigned when a bead is brighter, because the brighter the bead, the greater the differences between the primary and secondary color calls.</td>
</tr>
</tbody>
</table>
Blur Value

Given a 1 micron bead and the optics and detector characteristics of the signal acquisition system, a bead should have an apparent radius of approximately 2.2 microns.

Deviations from this value are almost exclusively in the positive direction and can come from a number of sources.

The most obvious is that of focus where the focal plane and the bead plane are not aligned, causing the bead to blur and grow larger.

Another source of change can be in vibration of the sequencer or settling of the stage while the image is being acquired. These latter reasons are a motion blur while the first is a focal blur.

For the purposes of determining the blur value for a given panel, a composite model of all beads in a region of a ligation image is created. The radius is just the center-to-edge measurement of that model. The deviation of this value from its nominal 2.2 is the metric being tracked. This is important because when the bead size becomes too large, the panel will not align, but when it aligns with a larger value, the region of the bead integrated to determine its intensity is too small.

If only one channel of the three has a blur problem, the relative integrated signal for that channel is less accurate than the others, degrading the call quality.

The blur value reported is the largest mean blur value of the four channels over all panels in one sample.

Blur Population

Using the blur metric discussed in Blur Value, rather than providing the average blur, this parameter reports the population of panels that have a blur value above a desired threshold.

A larger than average size can be due to a small number of panels with large outlier values or a large number of panels with a larger size.

The two values taken together can estimate whether there is an endemic problem over the whole sample (larger blur value but no outliers), a regional problem (an outlier population that spans channels), or a transient problem (outliers in a single channel).
### Exposure Value

Typically the ideal exposure level of an image is one that maximizes the dynamic range of the content with some signal occupying the lowest (darkest) range and some information extending just to the saturation (highest) point. An overexposed image is one that compresses too much of the content at the top end of the range, creating a non-linearity in response and making it impossible to differentiate between the highest signals. An underexposed image is one that does not extend the content of the image across the possible range, thereby creating a low-end compression and failing to use the available range of the system.

The calculated exposure value is a measure of how close the system is to capturing an image that maximizes dynamic range. Looking at the tail behavior of the signal distribution makes the determination. If the tail does not extend to the saturation point, the image is underexposed. In the case of overexposure, the tail has a peak at the far right side that is a measurement of the saturation. To get an estimate of the overexposure, the slope of the curve prior to the peak is extrapolated to a distance equal to the area of the saturated peak. The computed exposure quality metric is two-sided, extending from an ideal of 0 in the negative direction for underexposure and in the positive direction for overexposure and uses a log (2) scale. To fit this value into the traffic light model, the distance from ideal performance (not necessarily a value of 0) is used. Applied Biosystems has found that for purposes of our color calling algorithms, slightly overexposing the images yields better results.

As with the blur value, the reported value is the largest exposure value of the four channels over all panels in the sample.

### Exposure Population

As with the blur population, rather than providing the average exposure quality value, the exposure population value reports the number of panels that have an exposure quality beyond a desired threshold.

A larger than average size can be due to a small number of panels with large outlier values or a large number of panels with a larger size.

The two values, taken together, can estimate whether there is an endemic problem over the whole sample (larger deviation with no outliers), a regional problem (outlier population that spans channels), or a transient problem (outliers in a single channel).

A difference from the blur metric, though, is that blur metric is invariant to the number of beads present in a panel where the exposure is more sensitive to the number of beads.

A baseline exposure value per channel is determined based on a sampling strategy, but panels with either significantly more or fewer beads can be affected differently.
Chapter 2 Prepare Run Settings

Set analysis parameters

There are two types of applications available for the secondary analysis settings:

- SingleTag
- IQ

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Exposure Time</td>
<td>In the SOLiD™ 4 System, every panel could technically have a different exposure value within a channel. This metric tracks the mean exposure time over all panels and reports the highest of the four channels. Increasing exposure times can indicate problems in the sequence. If the exposure time jumps too high too quickly, the signal on the beads is decreasing too quickly cycle over cycle, and there will likely be problems achieving longer read lengths. Vibration, or other issues, can also up the exposure time as the beads become larger due to blur and the exposures increase in order to get sufficient signal spread over a larger area. An increase in exposure time by itself is not a problem, but it is a warning sign that is valuable when taken into consideration with the other available run metrics to diagnose potential problems. Exposure times are ‘effective’ exposure times; they are a combination of both the exposure time and the gain.</td>
</tr>
</tbody>
</table>

Application Descriptions

There are two types of applications available for the secondary analysis settings:

- SingleTag
- IQ
These are the available SingleTag pipelines and their descriptions.

<table>
<thead>
<tr>
<th>Pipeline</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mapping</td>
<td>Used to find all hits between a set of reads and the reference sequence(s), based on the settings of anchor length, number of mismatches allowed, and anchor start.</td>
</tr>
<tr>
<td>MaToBAM</td>
<td>Writes a BAM file from a single-fragment match (.ma) file v4.0, and its associated quality value file, for SOLiD™ SETS Software.</td>
</tr>
<tr>
<td>Position Errors</td>
<td>Produces a file containing statistics on color mismatches vs read position. The output file has the format {primer.set}_positionErrors.txt.</td>
</tr>
</tbody>
</table>

These are the available IQ pipelines and their descriptions.

<table>
<thead>
<tr>
<th>Pipeline</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>IQ</td>
<td>Instrument qualification and troubleshooting. An independent pipeline, used by manufacturing and service, that is tied to a specific set of beads.</td>
</tr>
</tbody>
</table>
Define secondary analysis settings

In SETS, you can define the secondary analysis pipelines and parameters as a part of your setup for secondary analysis.

To modify any secondary analysis pipeline or parameter:

1. Log into SETS.

2. Go to Main Menu ➤ Analysis ➤ Secondary Analysis Settings.

3. Select the Application type (SingleTag or IQ, as shown) to create a secondary settings.

4. Click [New...] to begin modifying the threshold settings for your application.

**Note:** Click ✗ to delete an existing setting or ✏️ to edit one.
5. Provide a name for your new settings configuration, then select the reference genome to associate with this setting. See “Load a Reference Sequence” on page 37.

6. Click Next to view or select individual settings for each tag in a paired run.
   Different settings can be applied to each tag for paired applications, as shown below.

7. Mouse over a pipeline’s name to see its description.
8. Click **View** to see the rollover summary of current settings.

   ![Rollover summary screenshot]

   **Note:** Click ☑️ to activate or deactivate a pipeline.

9. Click **Edit** to the right of a name to edit any active pipeline.
Chapter 2 Prepare Run Settings

Set analysis parameters

10. Enter a new value or values, then click **Finish**. To edit more pipelines, click **Save and Edit Another Pipeline**.

**Note:** Click **☐** to select show first position as bases.

**IMPORTANT!** Some pipelines are dependent upon others. When you select or deselect one, you may be affecting the activation state of the others.
Load a Reference Sequence

Specify a New Reference

If a specific reference sequence is not in the Reference Genome drop-down list (Secondary Analysis Settings view), you can load a new reference sequence before you start a run. Open the SETS software and perform steps 1 to 4 below to add a new reference sequence name to the list of choices you see in SETS.

Secondary analysis (sequence data alignments to a reference sequence) would not occur without a preloaded reference sequence.

IMPORTANT! For more complex genomes (such as the human genome), barcoded samples, and larger reference (and reads) the secondary analysis should be performed off-instrument with the BioScope™ Software. Refer to BioScope™ Software for Scientists Guide: Data Analysis Methods and Interpretation (PN 4448431).

If a sample contains more than the maximum allowable number of reads, the SETS software selects only up to the maximum allowed number of reads and the rest will not be included in the results of the secondary analysis.

IMPORTANT! Do not install BioScope™ Software on the SOLiD™ 4 Analyzer.

The Secondary analysis (sequence data alignments to a reference sequence) will not occur without a preloaded reference sequence. Perform secondary and tertiary analysis with the BioScope™ Software. See “Load a Reference Sequence” on page 37.
To upload a new reference sequence:

1. In the Home page log in as “Administrator.”

   **Note:** The “Load reference” menu option under Analysis is not available unless you are logged in as an “Administrator.”

2. Click **Analysis ➤ Load Reference**.

   The Load Reference Genome page opens.

   **Load Reference Genome**

   The maximum file upload size is 2GB.

   Reference genome: 
   
   Name: 

   Submit Cancel

   * = required fields

   **Note:** For instructions on how to upload reference genomes larger than 2GB see “To upload reference genome files larger than 2GB” on page 39.
3. Click **Browse**, then select the appropriate fasta file for your reference sequence.

4. Enter a name for the selected file and click **Submit**.

   The reference genome name that you enter in step 4 appears in the Secondary Analysis Settings window in the drop-down list of reference genome.

   **IMPORTANT!** After uploading the genome, the SETS Software validates the reference. The validation may take several minutes, during which the genome is not available. If validation fails, the new genome is not available in the drop-down list in the Secondary Analysis Settings and an error log is displayed in the homepage (under Error Messages).

---

**To upload reference genome files larger than 2GB**

Reference genome files larger than 2GB cannot be uploaded through the SETS Software due to the limitations set by browsers. To upload reference genomes larger than 2GB:

1. Use an FTP tool to first place the file inside the home directory, on the cluster.

2. Run the reference validation perl script to produce a validated reference.

3. Name the validated output and save it with the ".valid" extension.

   **Example**

   ```
   /share/apps/corona/bin/reference_validation.pl -r
   MY_REFERENCE.fasta -o MY_REFERENCE.fasta.valid
   ```

4. To ensure that the reference genome appears in the Reference genome box of the Secondary Analysis Settings list, move the reference genome file to the genomes folder, inside the reference folder.

   **Example**

   ```
   mv MY_REFERENCE.fasta.valid
   /share/reference/genomes/MY_REFERENCE.fasta
   ```
5. From the Home page, navigate to the Secondary Analysis Settings and click **New**. Ensure that the reference genome file (MYREFERENCE.fasta) appears in the Reference genome drop-down list, as shown in the example below.
This chapter covers:

- Overview ............................................. 42
- View Run Metrics ................................. 42
- View Cycle Scans ................................ 46
- View sample data ................................. 49
- Troubleshoot failed jobs ...................... 51
- View History ....................................... 57
- Cancel analysis midstream ................ 58
- Cluster Status ..................................... 60
- System Logs ....................................... 61
- RSS feed ........................................... 62
Overview

While executing a sequencing run, the instrument performs a Pre-scan. In a Pre-scan, all of the beads are labeled and their positions on the slide are recorded to derive a focal map followed by a single ligation cycle. The run can be remotely monitored in SETS by checking the Run Metrics, Cycle Scans, Heat Maps, and System Logs.

Note: To set up e-mail notification regarding instrument run and system information, see Chapter 7, “Event notifications and email services” on page 125.

View Run Metrics

You can view a current run on the SOLiD™ 4 System and determine the quality of each cycle by looking at the traffic lights. The Run Metrics page in SETS displays all metrics per sample for the selected ligation cycle.

To view the quality of a run:

1. Open SETS and from the Run Details page, click Run metrics.
Note: For in-progress runs, you can also access the Run metrics link from the homepage.
2. Select the primer set, primer, cycle, and version you want to monitor, then click **Display** to update the traffic lights.

3. Determine the quality of your run by looking at the traffic light indicators for in-progress runs.
   - **Green** - when a traffic light for a metric displays green, that parameter has a passing quality value and is hitting its target threshold.
   - **Yellow** - when a traffic light for a metric displays yellow, that parameter has a value that falls between the low and high threshold and is questionable in quality.
   - **Red** - when a traffic light for a metric displays red, that parameter has a value that is above the high threshold limit and the quality could be considered poor enough to fail; if a sample displays all red traffic lights during a run, you may want to abort that run immediately.

4. Click **Get summary** when you want a full summary of all the metrics for an entire run. **Get latest status** displays by default.
Chapter 3 Monitor the Run in SETS

View Run Metrics

Note: Refer to the Imaging Metrics Report and the Exposure Time Report as additional resources when monitoring a run.
View Cycle Scans  

To view cycle scans of runs in progress or completed:

1. Go to Run Details ➤ Overall Reports.
2. Click Cycle Scans to see all the scans for the selected run.
A cycle can have multiple scan versions as indicated in the second column (Ver.), however, only one version is used in the latest analysis and is checked in the third column (Used).

To view cycle data:

1. Click on the version number hyperlink to display the cycle scan data for that particular version in a table at the bottom of the page.
2. Click \( \mathcal{H} \) to open a Heat Map for any particular scan. Click **Close** when you have finished viewing the Heat Map.

3. Click *Back to Run page* after you view cycle scans.
**Run Details Page**

The Run Details page displays all relevant Run Information that can be accessed by expanding individual samples in the sample tree in the left-hand panel of the page. Within each sample folder are analysis cycles that correspond to individual samples on a slide.

**Sample information**

1. Click a sample in the sample tree to expand the folder to show general information about the sample, its analyses, and libraries. When you select an analysis or library, an Analysis/Library data panel displays in the lower half of the right-hand panel.

2. Click the sample folder drop-down arrow or the sample link to view individual analysis, or reanalysis cycles for each sample.
3. Click one of the Overall Reports, or one of the Sample Reports (Satay or Color Balance Report) to view report data. For more information on reports, see “View Reports” on page 63.

The sample tree shows the active pipelines as defined in your secondary settings.

Click **Collapse All** to collapse all samples back to their original folder.

Click **Expand All Samples** to open all folders and view all samples at once.

**Note:** If a pipeline has been inactivated from within the settings, it will not display.
Troubleshoot failed jobs

Sample analysis cycles A blue folder indicates a completed analysis while a red folder indicates a failed analysis. To better understand why an analysis failed:

1. Click the arrow next to the analysis folder to view the analysis cycles for each sample (Run Details page).

2. Click the individual analysis job link to see details for that particular stage of analysis.

3. Select one of the job details for more information on each sample.
Chapter 3 Monitor the Run in SETS
Troubleshoot failed jobs

Job details Beadfinding

The beadfinding stage executes immediately after the Primer A cycle 1 scans have completed on-instrument.

Details for job: F3_PA_05_V3  (jobid: 36661)

Settings: default primary (v.1)
Stage: post ScanSlidePrimary
Status: finished
Created: 03/28/2010 03:11 PM
Started: 03/28/2010 03:02 PM
Completed: 03/28/2010 03:22 PM

job  code 1
focal.map.dir /yn_1545/yw_1545_FOCALMAP_V1
focal.map.ofg /yn_1545/yw_1545_FOCALMAP_V1/yw_1545_FOCALMAP_V1.0.7.0
focal.map.version 1
image.dir  /yw_1545/yw_1545_FOCALMAP_V1
instrument.name FLA480DA
num.bases 9
p1.channel CY3
p2.channel CY2
p3.channel PE
primerset F0
probe.set 12
run.name yw_1545
run.type run
sample.name Sample1
spots 2
version 1
Colorcall Jobs

The job details (colorcall jobs) of the Post Scan Slide Primary stage are activated after each cycle is scanned. Incremental image analysis is called during this stage.

Filter Fasta

The Filter Fasta stage, Post Primer Set Primary, is performed after all cycles are imaged and the colorcalled data are ready for aggregation into final results.
### Details for job: Filter_FastaF3 (Id=36665)

<table>
<thead>
<tr>
<th>Setting</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Settings:</strong></td>
<td>default primary(v.1)</td>
</tr>
<tr>
<td><strong>Stage:</strong></td>
<td>postPrimerSetPrimary</td>
</tr>
<tr>
<td><strong>Status:</strong></td>
<td>finished</td>
</tr>
<tr>
<td><strong>Created:</strong></td>
<td>03/28/2009 03:13 PM</td>
</tr>
<tr>
<td><strong>Started:</strong></td>
<td>03/28/2009 03:04 PM</td>
</tr>
<tr>
<td><strong>Completed:</strong></td>
<td>03/28/2009 03:04 PM</td>
</tr>
</tbody>
</table>

- **focal.map**: `yw_1565/yw_1565_FOCAL_MAP_V1
- **focal.map.version**: 1
- **images**: `yw_1565/yw_1565_F3_F2_01_V1
- **nrun**: 5
- **panels**: 2
- **prime.set**: F3
- **read.length**: 5
- **read.prefix**: T
- **run.name**: `yw_1565
- **sample.name**: Sample1
- **spots**: 2

---

Chapter 3  Monitor the Run in SETS

Troubleshoot failed jobs
Filter Fasta fails

The Filter_Fasta job details will show a red icon indicating that the job has failed.

This problem can occur either at the Filter Fasta stage or at the Barcording stage due to a bad hard-drive, a lack of disk space, or other system-related issues.

The software displays an error message and provides the Bead ID or Sequence number where the record is corrupted.

Example

Based on the original system-related issue, you will see an error message indicating that the “Fasta is empty” or the “Format is not right expecting sequence. Current line - >1_10_117_F3.”

Click OK to continue.

The error message will contain the problematic file’s location and record information.

To resolve, you need to fix any system-related issues first. Then, any of the following step may be applicable:

- Check the corrupted Fasta File manually; if possible fix it from the command line.
- Reanalyze primary analysis stage through SETS or manually by changing the database record/flag.
- If the SPCH files are also corrupted, report the error message to the Applied Biosystems Field Support Engineer.

Note: Report any critical error messages to the Applied Biosystems FSE.
Chapter 3 Monitor the Run in SETS

Troubleshoot failed jobs

Barcoding

**IMPORTANT!** Multiplexing reports can be created as soon as barcoding completes for a run.

Active only in multiplexing run; Barcoding job is performed after the FilterFasta job associated with each barcode tag. The barcoding job identifies and assigns each bead to a library. The output separates the other tag-based primary directories into library subdirectories within the Sample directory hierarchy.

---

**Details for job: BarcodingF3 (id=36667)**

**Settings:** default primary(v.1)

**Stage:** postBarcodePrimary

**Status:** standby

**Created:** 01/28/2009 03:13 PM

**Started:**

**Completed:**

---

**Analysis Report**

---

**Details for job: AnalysisReportF3 (id=596)**

---

Applied Biosystems SOLiD™ 4 System SETS Software User Guide
View History in SETS

1. Click **View History** on the Main Menu to view the history of runs performed on the SOLiD™ 4 System.

A list of all runs displays by name and indicates the start and end date of each run.

2. Click **Export View** to see the Run Export History view.

3. Click **Recalculate Disk Space** when needed.

   **Note:** Performing “Recalculate Disk Space” takes time to complete. Once started, disk space consumption (results data and image data) for all runs in Run History is calculated.
View Run Export History

The Run Export History view displays the runs that have been exported, as well as those runs currently being exported. This view shows the progress of the export and the specific location where each run was exported.

Cancel analysis midstream

Cancel in-progress analyses

You can cancel an analysis from the Run Details view in SETS. Only analyses in-progress are available for cancellation. The analysis will display as an active link if the analysis is in-progress and the run state is complete (no chemistry is running).

1. Click Cancel analysis for the selected run.
   A complete list of all cancellable analyses displays.

Note: If Cancel analysis is disabled, it is possible that the run is no longer in-progress. Mouse over to see the explanation.
2. Select the analysis to cancel, then click **Submit**.

3. Cancel more than one analysis at a time from the list by clicking **Check all**, then **Submit**.
   - All the jobs for selected analyses will display gray (cancelled).
   - Any job still processing in the queue will be stopped and removed from the queue.

![Cancel Analysis - r1ahead01_20060006_4](image)
Cluster Status

Use the Cluster Status view to monitor the available disk space and progress of a current analysis job, as well as jobs that are scheduled to run. Specifically, the Cluster Status view shows:

- **Average Load (5 min)** – Average computing work load for each 5 minutes. If this is always 95%±, then the cluster may be overloaded or a node may be down.

- **Free Disk Space** – Available disk space for images (/data/images) and available disk space for data (/data/results). For example, 1.27 TB / 8.86 TB (14.38%) (41,773 cycle-panels) indicates there are 1.27 TB of disk space out of 8.86 TB total space available to store images, and the disk is able to store up to 41,773 more cycle-panels.

- **Current Analysis Job Progress** – Displays which jobs are currently in progress and their completion percentage.

- **Job Queue Statistics** – Information about the job queues on the cluster. Search for blocked jobs.

### Cluster Status

Cluster name: soli3336m
Compute nodes up (ganglia): 3
Avg load (5min): 14.22%
Free disk space (images): 1.27 TB / 8.86 TB (14.38%) (41,773 cycle-panels)
Free disk space (results): 1.63 TB / 9.93 TB (14.43%)

Current analysis jobs

<table>
<thead>
<tr>
<th>Job Name</th>
<th>Job Id</th>
<th>Date Started</th>
<th>Progress</th>
<th>Analysis Name</th>
<th>Run Name</th>
<th>Sample Name</th>
</tr>
</thead>
<tbody>
<tr>
<td>F3_P5_01_VL</td>
<td>743</td>
<td>01/26/2010 01:01 PM</td>
<td>green</td>
<td>auto-analysis 2</td>
<td>20100108_F20100109_D106_b5x5_F12</td>
<td>somset</td>
</tr>
<tr>
<td>F3_P5_05_VL</td>
<td>742</td>
<td>01/26/2010 12:09 PM</td>
<td>green</td>
<td>auto-analysis 2</td>
<td>20100108_F20100109_D118_b5x5_F12</td>
<td>somset</td>
</tr>
</tbody>
</table>

Queue statistics

- Total Jobs: 4
- Active Jobs: 2
- Idle Jobs: 0
- Blocked Jobs: 0

<table>
<thead>
<tr>
<th>Queue</th>
<th># of running jobs</th>
<th># of queued jobs</th>
</tr>
</thead>
<tbody>
<tr>
<td>primary</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>forwarding</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>feedback</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>latch</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>interleave</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>interactive</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>secondary</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>
System Logs

The System Log View (Admin ▶ System View) allows you to view error logs on specific software components and to check the severity and the date the problem occurred.

Make your selections, then click Search or Display Last 24 Hours.
RSS feed

You can observe real-time data from a current, in-progress run and/or from instrument logs by using any available RSS feed reader (Internet Explorer 7.0, Firefox, Sage, etc.). The feed reader links are located on the right side of the page above the “In-progress or most recent runs” section on the homepage.

To set up the RSS feed reader:

1. Click the appropriate Feed button:

2. Copy the web address for each feed, then paste it into the appropriate location in your browser.
Chapter 4

View Reports

This chapter covers:
- View overall run reports ........................................ 64
- View sample reports ............................................. 77
- View Analysis Reports ........................................... 85
- Master Report Tool ............................................... 93

Applied Biosystems SOLiD™ 4 System SETS Software User Guide
You can view information about a completed run by opening one of the overall run reports:

- Imaging Metrics
- Heat Map Report
- Cycle Heat Map Report
- Cycle Scans (View)
- Exposure Time
- Flowcell Mask
- WFA Report (will display only when applicable)

**Note:** Most reports can be viewed in both real-time and completed run data.
The Imaging Metrics report displays the cycle scan heat map view for all four channels (FAM™, CY3®, Texas Red®, and CY5® dyes) for the selected metric type, (for example, exposure).

Definitions of all the metric types can be found in “Primary analysis parameters” on page 25.

Click Imaging Metrics Report (Run Details ➔ Overall Reports).

### Imaging Metrics Report

<table>
<thead>
<tr>
<th>Run: solid3008_20090109_OD11B_21-50_F01</th>
<th>Fixed Scale:</th>
<th>Primer Set:</th>
<th>Primer:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mineral Mask</td>
<td>FAM</td>
<td>CY3</td>
<td>Texas Red</td>
</tr>
</tbody>
</table>

**R3_PA_01_Y1**

**R5_PA_02_Y1**
Heat Map Report  

Heat maps are generated from the results of the analysis of the focal map images. Heat maps give a colorized view of the selected metric across all panels for a run. Use this report to view experimental data mapped to the physical layout of the flowcell slide.

Click **Heat Map Report (Run Details ➤ Overall Reports)**.

---

Look for uniform deposition of beads on the slide. The actual average bead deposition density/panel value should be similar in value to the targeted average bead deposition density. A large number of missing panels could indicate a deposition problem.

**Note:** Heat Map Reports are available after beadfinding occurs.
Select a heat map type using the drop-down menu:

- **Bead Count** - number of beads per panel (Focal Map)
- **Bead Signal** - average signal intensity of the beads in a panel
- **Image Signal** - overall image signal intensity for a panel

**Cycle Heat Map Report**

This report displays the Cycle Scan Heat Map view for selected beads. See “Bead definitions” on page 68.

Click **Cycle Heat Map Report (Overall Reports)**.
### Bead definitions

<table>
<thead>
<tr>
<th>Bead Type</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Usable Beads</td>
<td>Beads that, in a particular ligation cycle, are mapped in each of the four images. Typically, most beads in a panel are usable.</td>
</tr>
<tr>
<td></td>
<td>The ones that are not lie near the edge of the panel, which for current purposes is defined as the area imaged by the corresponding reference or focal map image.</td>
</tr>
<tr>
<td></td>
<td>The ligation fluorescence images must be shifted a few to tens of pixels to register them with the reference image. If, for example, a ligation image must be shifted 10 pixels to the left to register with the reference image, beads within 10 pixels of the right edge of the panel will not be mapped in that ligation image.</td>
</tr>
<tr>
<td></td>
<td>Another common case is a panel-cycle where one of the ligation images fails to register. In this case, none of the beads are usable.</td>
</tr>
<tr>
<td>Good Beads</td>
<td>Usable beads that satisfy all of the following conditions:</td>
</tr>
<tr>
<td></td>
<td>intensity &gt;0.05</td>
</tr>
<tr>
<td></td>
<td>0.0125 &lt; angle ≤ 0.05</td>
</tr>
<tr>
<td>Best Beads</td>
<td>Usable beads that satisfy all of the following conditions:</td>
</tr>
<tr>
<td></td>
<td>intensity &gt;0.05</td>
</tr>
<tr>
<td></td>
<td>angle ≤ 0.0125</td>
</tr>
</tbody>
</table>
Chapter 4 View Reports

View overall run reports

Cycle Scans view

Use the Cycle Scans window to distinguish normal runs from problematic runs.

Click Cycle Scans (Run Details ➤ Overall Reports).

The top section of the Cycle Scans window lists all the scans per ligation cycle for the slide, and the bottom section shows scan information sorted by sample for the scan selected in the top section. Use the parameters shown in the table on the next page to assess the progress of the sequencing run.

See “View Cycle Scans” on page 46 for more details.
### Parameter

<table>
<thead>
<tr>
<th></th>
<th>Normal Run</th>
<th>Problematic Run</th>
</tr>
</thead>
<tbody>
<tr>
<td>Failed panels</td>
<td>In general, the number of failed panels should be relatively small and</td>
<td>Run begins with extremely high number of failed panels or dramatic increase in</td>
</tr>
<tr>
<td></td>
<td>consistent.</td>
<td>any subsequent ligation cycle for each sequencing primer.</td>
</tr>
<tr>
<td>Fraction of Best + Good</td>
<td>The fraction can vary depending on the quality of the library, the</td>
<td>A significant drop in the fraction of good beads in the initial ligation cycles is</td>
</tr>
<tr>
<td>Good beads/Usable</td>
<td>efficiency of the PCR, and the enrichment process. As a guideline, the</td>
<td>a reason to pause the run and troubleshoot the performance.</td>
</tr>
<tr>
<td></td>
<td>fraction is around 0.5 to 0.6 in the first ligation cycle of each primer</td>
<td></td>
</tr>
<tr>
<td></td>
<td>and drops to 0.2 to 0.3 in the last cycle.</td>
<td></td>
</tr>
<tr>
<td>Effective exposure times</td>
<td>Gradual increase from ligation cycles 1 to 5 or higher for each sequence</td>
<td>250 ms or greater in ligation cycle 1 or when instrument times out when the</td>
</tr>
<tr>
<td></td>
<td>primer. Performance varies from slide to slide and as a function of the</td>
<td>effective exposure time exceeds 500 ms. Long exposure times may indicate</td>
</tr>
<tr>
<td></td>
<td>age of the SOLiD™ Light Source. As a guideline, the effective exposure time</td>
<td>replacement of the SOLiD™ Light Source.</td>
</tr>
<tr>
<td></td>
<td>is typically 20 to 40 ms in the first ligation cycle and increases to 100 to</td>
<td></td>
</tr>
<tr>
<td></td>
<td>300 ms in the fifth cycle.</td>
<td></td>
</tr>
<tr>
<td>Satay plots</td>
<td>The first cycle of any primer should show a relatively “clean” Satay plot,</td>
<td>An abnormal “fuzzy” Satay plot in the first cycle is a reason to pause the run</td>
</tr>
<tr>
<td></td>
<td>with most points clustered on the four color axes and with minimal fraction of</td>
<td>and troubleshoot the performance.</td>
</tr>
<tr>
<td></td>
<td>the points clustered around the origin. The quality of the Satay plot</td>
<td></td>
</tr>
<tr>
<td></td>
<td>typically degrades gradually with each ligation cycle for a single primer</td>
<td></td>
</tr>
<tr>
<td></td>
<td>cycle, becoming more “fuzzy” in the last cycle.</td>
<td></td>
</tr>
</tbody>
</table>
Exposure Time Report

The Exposure Time report displays the average exposure time of run cycle scans for each of the four channels (6-FAM™, Cy3®, Texas Red®, and Cy5® dyes).

In this report, you can:
- Mouse over any data point to see the called value.
- Export the Average Exposure Time table in a CSV format.

Click Exposure Time Report (Run Details → Overall Reports).
**Flowcell Mask view**  
Use the Flowcell Mask report to view a representation of the sample slide. Each numbered area on the flowcell mask is a spot that corresponds to a specific sample location on the slide.

<table>
<thead>
<tr>
<th>Flowcell Mask</th>
<th>1: 180K_1202</th>
<th>2: 100K_1202</th>
<th>3: 220K_1202</th>
<th>4: 190K_1202</th>
</tr>
</thead>
</table>

**Note:** The numbered areas on the Flowcell Mask also correspond to the numbered sample names in the sample tree, which appears at the bottom left of the Run Details page.
WFA report

A WFA run on the SOLiD™ 4 System is uniquely labelled with a green WFA stamp and titration sample folders. When there is a valid WFA run, then the WFA report will be accessible in the Run Details page. Click **WFA Report (Overall Reports)**.

This example shows one WFA report for a single 4-well slide.
Click on individual titration links for more detail.

The Results Details for each titration opens to view.
Determine the optimal titration point

Determine which titration has the highest titration metric. This titration is the optimal titration point.

Determine the bead deposition density for a sequencing run

1. Calculate the concentration of P2-positive beads using the formula below, where $X$ is the volume of templated beads used for the WFA sample (equivalent to 15 million beads) and P2# is given by the WFA report:

$$X \, \mu\text{L} = \frac{15 \times 10^6 \text{ beads}}{\# \text{ beads/}\mu\text{L (according to NanoDrop®)}}$$

$$Y \text{ P2-positive beads/}\mu\text{L} = \frac{\text{P2# beads/panel} \times 426 \text{ panels}}{X \, \mu\text{L}}$$

**Example:**

For a sample with a concentration of 500,000 beads/\mu\text{L} measured by NanoDrop® ND-1000, where the WFA report indicates a P2# value of 20,000 beads/panel.

$$X \, \mu\text{L} = \frac{15 \times 10^6 \text{ beads}}{500,000 \text{ beads/}\mu\text{L}} = 30 \, \mu\text{L}$$

$$Y \text{ P2-positive beads/}\mu\text{L} = \frac{20,000 \text{ beads/panel} \times 426 \text{ panels}}{X \, \mu\text{L}}$$

$$= 284,000 \text{ P2-positive beads/}\mu\text{L}$$

2. Use the calculated concentration in place of the value determined by the NanoDrop® ND-1000 Spectrophotometer for more accurate deposition densities when preparing slides for sequencing. It is recommended to use this resulting calculated bead concentration to determine the volume of beads for deposition and multiply that volume by 120%:
Chapter 4 View Reports

View overall run reports

\[ Z \mu L \text{ bead solution} = \frac{\text{Desired } \# \text{ P2-positive beads to be deposited}}{Y \text{ P2-positive beads}/\mu L} \]

**Example:**

For a sample with a concentration of 284,000 P2-positive beads/\mu L to be deposited in one well of an 8-well SOLiD™ Deposition Chamber at a density of 300,000 beads/panel.

\[ Z \mu L \text{ bead solution} = \frac{56 \text{ million P2-positive beads}}{284,000 \text{ P2-positive beads}/\mu L} \times 120\% \]

\[ = 237 \mu L \text{ bead solution} \]
View sample reports

**Satay Reports**

Use this report to view analysis cycle signals, which are converted into X,Y plots. For each cycle of sequencing, these plots show the relative intensities of the signals for each bead and how well separated the signal is across the four dye channels.

Each data point is an individual color call (i.e., an individual bead). Because total data points can exceed 20 million, only a subset of randomly selected points is displayed for each primer set (F3 and R3) and for various types of data (raw, scaled, angled, intensity, and N2S).

**IMPORTANT!** A satay report that collapses towards the origin or drifts away from the axes often indicates low signals or multiple DNA templates. If you observe such a satay report in the *early* cycles, consider aborting the run.

1. Click **Satay Report** *(Run Details ➤ Overall Reports).*
2. Select the Primer Set and Data type of interest:
   - Raw – “Raw data” on page 80
   - Scaled – “Scaled data” on page 81
   - N2S – “N2S data” on page 82
   - Angle – “Angle data” on page 83
   - Intensity – “Intensity data” on page 83
3. Select the Dye Channel of interest:
   - 6-FAM™
   - Cy3®
   - Texas Red®
   - Cy5®

4. Click **Display**. The window opens to display a thumbnail plot for each cycle.

5. Click each thumbnail to see a larger, more detailed view.

6. In each data window, scroll down to view graphs for all cycles.
Resizing images

When you view the Satay images in SETS, your browser may automatically resize the large image to fit the screen. You may need to make an extra mouse click to view the full size image. The following instructions allow you to turn the browser’s automatic image resizing function ON or OFF.

To resize image window in Internet Explorer:

1. On the Tools menu, click Internet Options.
2. Click the Advanced tab, then scroll down to the multimedia section.
3. Select or clear the Enable Automatic Image Resizing check box.

To resize image window in Firefox:

1. In the Firefox address bar, type about:config.
2. In the filter bar, type browser.enable.
3. Double-click browser.enable_automatic_image_resizing. The value changes to False.
**Raw data**

Raw data images display the Satay plot for beads using raw signal intensities from the image analysis algorithm.

![Satay Reports](image)

**Note:** The plot is calculated by subtracting the CY3 dye intensity from the CY5 dye intensity, then subtracting the FAM dye intensity.

![Satay Report](image)

**Note:** The CY3 dye is offset from the axis in this typical raw data view compared to the graph showing processed data.
Scaled data

Scaled data images display the Satay plot for beads after signal intensities in each channel are normalized.
N2S data

N2S data plots show the distribution of high- and low-quality beads using a measure of the noise-to-signal (N2S) ratio.
Angle data

Angle data histograms show the distribution of the angle between the 4-color signal vector and the canonical signal for each dye channel (6-FAM™, Cy3®, Texas Red®, and Cy5® dyes).

Note: Acceptable or good beads have angles close to zero.

Intensity data

Intensity data histograms show the distribution of the intensities for each bead in each of the four dye channels (6-FAM™, Cy3®, Texas Red®, and Cy5® dyes).

Note: Good beads have high intensities.
Use this report to view the number of beads that have a particular color call for a specific ligation cycle. For most samples, the distribution of colors is relatively uniform across the four channels. However, a strong deviation from uniformity or an imbalance trend could indicate problems in the sequencing chemistry or imaging process, or a bias in the library preparation.

In this report, you can:

- Mouse over any data point to see the called value.
- Export the Scan Bead Summary table to a CSV file.
View Analysis Reports

View sample files

Auto-analysis creates fasta files and various report graphs from the analyzed sample data. Primer sets F3 and R3 are represented by links to csfasta files for F3 and R3.

Note: When the analysis is successful for F3 and R3 primer sets, a csfasta file is generated. The “.csfasta” format is a text-based format that displays the base-pair sequences in color space (that is, bases display as 0, 1, 2, 3).

Click each auto-analysis folder’s expansion arrow to see its contents.
Select a library hyperlink to view the details.
View analysis Data Files

In the sample folder, you can view and download the following data files for each sample.

- Mapping Summary F3
- Mapping Summary R3
- Mapping Summary F5-P2
- Mapping Summary F5-BC
- csfasta file F3
- csfasta file R3
- csfasta file F5-P2
- csfasta file F5-BC
- QV file F3
- QV file R3
- QV file F5-P2
- QV file F5-BC

View Analysis Reports

Sample analysis creates various reports for individual sample files. You can view and download the following reports:

- Multiplexing Assignment Report – page 92
- Quality Values Report – page 87
- Auto-Correlation Report – page 89
- Error Profiles Report – page 91

Quality Values Report

The Quality Values Report contains the quality values for each base sequenced.

Click Quality Values Report (Run Details ➤ Analysis Reports).
The bar chart shows the proportion of color calls with a particular quality value.
With quality values, it is possible to predict the expected number of mismatches to the reference genome caused by measurement errors as depicted in the bar chart above. The blue line shows the accumulative read proportion with predicted mismatches. For this run example, about 28% of the color space calls are predicted to have 0 to 0.5 mismatches. About 80% of the bases are predicted to have approximately 0 to 3 mismatches.

The final graph included with the Quality Values Report shows the proportions of six groups of quality values versus the primers and cycles in the run. The proportion of the highest quality bases tend to decrease in later cycles and increase after the primer resets. (PAC1 represents Primer A Cycle 1, PBC5 represents Primer B Cycle 5).

**Auto-Correlation Report**

Use this report to view the correlation between pairs of color calls in the raw reads.

Click **Auto Correlation Report (Run Details ▶ Analysis Reports)**.
Auto-Correlation Report

Run: r1ahead01_20100113_FRAG Analysis primary_reanalysis
Sample: Sample1
Library: 102

Raw Data: F3_autoCorrelation.txt (2.02 KB)

Correlation Scale

The Auto-correlation of 'base' position "i" with position "j" for Tag F3 displayed as a heat map, with high correlation as dark red/bright green, and low correlation as dark blue. The correlation is calculated as an "r" coefficient between each possible pair of positions.
The Same data as the one on the previous page but sorted in the order of the chemistry cycles on the sequencing instrument.

Light blue shading or dark red shading shows stronger correlations between the two cycles. Such correlations indicate potential problems with swapped reagents, incomplete cleavage, and other sequencing chemistry problems.

**Error Profiles Report**

The Error Profiles Report gives the frequency of errors as a function of base position in the read and type of error produced for a particular tag.

Click the **Error Profiles Report** link (Run Details Analysis Reports) to view the graphs that show the number of mismatches to the reference sequence, shown by position on read (top) and by cycle (bottom). Mouse over to view the calculated values for each bar.
Multiplex Assignment Report

To see an example of multiplex assignment report, go to “Multiplexing Assignment Report” on page 108.
Master Report Tool

Generate a Master Report

SETS provides a powerful report building tool to assist you in generating a summary of several reports bundled into a comprehensive one, customized with only the data that you want to see.

1. From the main menu, click **Report**.
2. Select a report setting from the drop-down menu.
3. Select at least one run from the full list of runs, then click **Generate Report**.
4. To specify which analyses are included in the Master Report, click **Select Samples** and check only the samples that you want included in the report.

To select multiple runs, use the Control key and the scroll bar.
Create a custom Report Setting

To create a custom report setting:

1. Click New in the Report Settings view (next to the Name field).

   **IMPORTANT!** You must be logged in before the software associates the new report setting to your user profile.

2. Select the reports that you want in your custom setting by selecting one at a time, then click Add Report.

3. Click Save.

Edit and save new Report Settings

To edit report parameters, click Edit. To delete a report from your custom report setting, click Delete.
Click **Save** when you are finished making edits to the report setting.

**IMPORTANT!** You must be logged in to save a new report setting or edit an existing setting. If you are not logged in, any changes that you make are temporary and will apply only to the current session.

**Save reports**  
After creating a Master Report with the comprehensive report-building tool (see “Master Report Tool” on page 93), use your browser menu to save the report.

Go File ▶ Save Page As ▶ Save as type ▶ Web Page.

Click **Save** to save the entire page as one html file and create a folder by the same name containing the associated report graphics.
Print Reports  Use the Print function on your browser to print reports.

Note: In some cases, you may need to adjust your browser to print the data reports.
This chapter covers:

- Overview .................................................. 98
- Primary reanalysis ................................. 98
- Secondary reanalysis ......................... 100
Overview

You can perform two types of reanalysis on the SOLiD™ 4 Analyzer.

- Primary
- Secondary

When you perform primary reanalysis of a sample, SETS keeps the original focal map and color-calling data and recollects the filter FASTA and run analysis data. When you perform secondary reanalysis of a sample, only the run analysis data are collected again.

**Note:** You must be logged in with “user” or “admin” privileges in order to perform reanalysis.

Primary reanalysis

**Overview**  
Primary reanalysis starts at the Filter Fasta stage after color calling. After you select scans and reanalyze, a new analysis workflow is created in the database resulting in a new set of FASTA files to be generated from the pipeline. Reports that are based on FASTA file inputs reflect changes created by reanalysis.

**Note:** Make sure to use the correct version of the cycle scans that you want to reanalyze. Reanalysis is applied to all samples in a run.

1. Click **Reanalyze Primary** to reanalyze the primary analysis data.  
   Click the icon for more information.
Selecting the **Mask** column allows the analysis to mask, or ignore, this cycle. Masked cycles will be represented by a '.' in the FASTA output file.

If the cycle has more than one scan version, click the radio button next to the version that you want to select for reanalysis.
Secondary reanalysis

Overview

Secondary reanalysis starts at the Single Tag Analysis stage. This workflow stage generates application-specific results such as base space sequences, alignment, tag counting from the colorspace sequence reads, and ".csfasta" file. The external analysis pipelines will also be executed if they are plugged-in and turned-on in the selected analysis setting.

The secondary reanalysis allows you to selectively apply secondary analysis settings to individual samples. You are not forced to reanalyze all samples as you are in primary reanalysis. The downstream effect is that a new analysis is created. All Single Tag analyses are rerun.

To perform a secondary reanalysis:

1. Click Reanalyze Secondary to reanalyze the secondary analysis data. Click the icon for more information.
2. Check the sample, or samples to reanalyze, specify a library keyword (optional), then click Add Libraries.

3. Select the library or libraries from the left panel to apply the Secondary Analysis Settings to, then click Add.

4. Verify that libraries, samples, and settings are set up correctly, then click Reanalyze.

   **Note:** Upon clicking Reanalyze, you will be warned that reanalysis might affect the performance of the instrument system. At this point, you can cancel reanalysis.

**Effect on the current run**

After you initialize SETS to reanalyze your samples, you are returned to the Run Portal page where a notification of reanalysis is displayed.

**Results**

If you continue with reanalysis, you can monitor progress by expanding samples in the sample tree. In the Run Details page, you will see a new analysis under each sample.
Chapter 5  Perform Reanalysis

Secondary reanalysis
This chapter covers:

- Fragment and Paired-End Multiplexing ................. 104
- View Multiplexing Series in SETS ....................... 105
- Set up Multiplexing run in ICS ......................... 107
- Multiplexing Assignment Report ....................... 108
- Reassign Run ............................................ 111
Fragment and Paired-End Multiplexing

Overview of Multiplexing and Barcodes

Multiplexing in general, and barcodes in particular, enable the SOLiD™ 4 System to target many applications, from deep sequencing of a single source to extreme parallel sequencing of a large number of samples over limited regions - sometimes on the same slide.

The additional stretches of DNA sequence, known as barcodes, can be used to segregate pooled library samples during sequencing back into their original (logical) single library groupings. Barcodes—in their simplest form—allow the pooling of samples on a slide without the loss of surface area creating geographic isolations. Multiplexing occurs at both the whole slide level as well as at the geographically isolated (mask) region level.

A barcode, by definition, is a tag with a known sequence that is attached to a sequencing primer on a bead during library preparation. When sequenced, it is the barcode that identifies and classifies the bead to a named sample. This barcoding, or indexing, approach allows multiplexing at the sequencing level.

Following sequencing of the target DNA, additional rounds of ligation-based sequencing are performed using primer sets complimentary to the barcode. The resulting reads can then be sorted by the barcode and aligned in groups to the reference sequence.

Barcodes are assigned to a library both conceptually and physically (as specific tags) as part of the library preparation. If more than one barcode is assigned to a single library during the library preparation step, the conceptual grouping is referred to as a barcode group. All barcodes in the pool will assign the underlying beads to the same original library for analysis.

The SOLiD™ 4 System supports multiplexing for both fragment sequencing and paired-end sequencing. See “Naming convention for sequencing reads” on page 11.

Note: Multiplexing is not available for mate-paired libraries.
View Multiplexing Series in SETS

The Multiplexing Series view in SETS shows all the barcode kits in the system to date. When a multiplexing series run is created in ICS, only the multiplexing series names are displayed.

To view multiplexing series details, click View ▶ Multiplexing Series.

Select the multiplexing series run that you want to view.

Every multiplexing series file includes:

- Name of the multiplexing kit
- Colorspace code
- Basespace sequence
### Chapter 6 Support Multiplexing

**View Multiplexing Series in SETS**

<table>
<thead>
<tr>
<th>Barcode</th>
<th>Name</th>
<th>Colorspace</th>
<th>Basespace</th>
</tr>
</thead>
<tbody>
<tr>
<td>0000</td>
<td>0000</td>
<td>0000000000</td>
<td>0000000000</td>
</tr>
<tr>
<td>0010</td>
<td>0010</td>
<td>0010000000</td>
<td>0010000000</td>
</tr>
<tr>
<td>0020</td>
<td>0020</td>
<td>0020000000</td>
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<tr>
<td>0090</td>
<td>0090</td>
<td>0090000000</td>
<td>0090000000</td>
</tr>
</tbody>
</table>

**Note:** Refer to the advanced topic appendices in this guide to learn more on colorspace and basespace.
Set up Multiplexing run in ICS

Create a Fragment and Paired-End Multiplexing run in ICS

There are two ways to set up a multiplexing run in the ICS:

- Creating a multiplexing run using the Run Wizard
- Importing a multiplexing run

Note: Refer to the Applied Biosystems SOLiD™ 4 System ICS Software Help or Applied Biosystems SOLiD™ 4 System Instrument Operation Guide (PN 4448379) for detailed instructions on these procedures.

Assign Libraries in ICS

Continuing in the Run Wizard, you manually configure each barcode library to generate a new run definition. You need to save the run definition before you assign the run to one of the two flowcells.

Using the Run Wizard, assign libraries to the appropriate barcode from the appropriate multiplexing series by creating a run definition file.

Note: Refer to the Applied Biosystems SOLiD™ 4 System ICS Software Help or Applied Biosystems SOLiD™ 4 System Library Preparation Guide (4445673) for detailed instructions on these procedures.
To view multiplexing run analysis results, select the sample folder ▶ analysis folder ▶ library folder.

**Note:** Enter a library name in the Search field to automatically expand the sample tree to that library.

There is a wide range of reports that you can generate from the results data, one of which is the Multiplexing Assignment Report.
The Multiplexing Assignment report is a report specific to barcode runs. There is one report generated per analysis.

When you have selected your target analysis folder, click

### Multiplexing Assignment Report

**Run**  
ND_20361122_v2.3_Bcux

**Sample** bcSample

**Barcode Statistics**

<table>
<thead>
<tr>
<th>Barcode</th>
<th>Library</th>
<th># Matches</th>
<th>1 Match</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>L1</td>
<td>SOLB C4</td>
<td>3976</td>
<td>1364</td>
<td>3970</td>
</tr>
<tr>
<td>L1</td>
<td>SOLB C5</td>
<td>3968</td>
<td>1742</td>
<td>3980</td>
</tr>
<tr>
<td>L1</td>
<td>SOLB C6</td>
<td>3968</td>
<td>1473</td>
<td>3964</td>
</tr>
<tr>
<td>L1</td>
<td>SOLB E0</td>
<td>3187</td>
<td>1640</td>
<td>4157</td>
</tr>
<tr>
<td>L1</td>
<td>SOLB E1</td>
<td>3186</td>
<td>738</td>
<td>4134</td>
</tr>
<tr>
<td>L1</td>
<td>SOLB E2</td>
<td>3456</td>
<td>4462</td>
<td>3770</td>
</tr>
<tr>
<td>L1</td>
<td>SOLB E3</td>
<td>3271</td>
<td>1142</td>
<td>4413</td>
</tr>
<tr>
<td>L1</td>
<td>SOLB E4</td>
<td>3271</td>
<td>369</td>
<td>3760</td>
</tr>
<tr>
<td>L2</td>
<td>SOLB C6</td>
<td>3445</td>
<td>1013</td>
<td>4456</td>
</tr>
<tr>
<td>L2</td>
<td>SOLB C7</td>
<td>3445</td>
<td>1279</td>
<td>4624</td>
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<tr>
<td>L2</td>
<td>SOLB E0</td>
<td>3206</td>
<td>1407</td>
<td>4613</td>
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<td>L2</td>
<td>SOLB E1</td>
<td>13245</td>
<td>5968</td>
<td>12273</td>
</tr>
<tr>
<td>unassigned</td>
<td>unlocked</td>
<td>Na</td>
<td>Na</td>
<td>2765</td>
</tr>
<tr>
<td>unassigned</td>
<td>NotIdentified</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>All Read1</td>
<td>Total</td>
<td>13018</td>
<td>2675</td>
<td>13283</td>
</tr>
</tbody>
</table>

**Expanded Library**

- Barcoding3
- AnalysisReport3
- Filter_Fastq3
- F3_PE_01_V1
- F3_PD_01_V1
- F3_PC_01_V1
- F3_PB_01_V1
- F3_RA_DL_V1
- BarcodingBc
- AnalysisReportBc
- Filter_FastqBc
- BC_PE_01_V1
- BC_PD_01_V1
- BC_PC_01_V1
- BC_PB_01_V1
- BC_RA_DL_V1
- BeadfindingBc

- bcSample2
The Multiplexing Assignment report shows the number of barcode reads assigned to each library, along with the subtotals for all libraries. It also shows the number of reads that were not assigned to any specified library. These numbers include both passed and failed reads from the filter fasta stage.

The value in the “0 Mismatch” column is the number of barcode reads that match exactly to the barcode shown in the barcode column. The “1 Mismatch” column is the number that missed matching that barcode by one.

Reads that did not match any of the barcodes at the specified level are counted in the unassigned library with the name unresolved. It is possible that barcodes in the kit that were not assigned to a library are also matched. In that case, the unassigned library may show barcodes named with the color sequence of the matched barcode. Two additional libraries may appear:

- For beads that have barcode read but no F3 read (missing-f3).
- For reads that have an F3 read but no barcode read 9missing-bc).

The “Total % Assigned” is the sum of the library subtotals divided by the “All Beads Total” expressed as a percent.

**Import a Multiplexing Series**

Follow the steps that use the Command line to import a multiplexing series found in the Applied Biosystems SOLiD™ 4 System ICS Software Help.

**Multiplexing report**

Multiplexing report can be attained as soon as barcoding is complete for a run.
Reassign Run

1. Download the sample sheet of the run that you want to reassign.

2. Open the sample sheet in excel and edit the barcoding reassignment or spot reassignment.

**Note:** When reassigning the spots, ensure that the new, or edited, sample sheet has the same number of spots.

3. Log in as Administrator.

4. From Home page, click **Reassign Run**.

5. Click **Log on** enter your name **enter password**.

6. From the Home page, click on the analyses result of your choice, then click **Reassign Run**. Importing a new run definition for an existing run will force regeneration of all results.
Chapter 7

Manage Administrative Tasks

This chapter covers:

- Export Data with SOLiD™ 4 System Software .................. 114
- Auto export in SETS ........................................ 115
- Manual export in SETS ....................................... 124
- Event notifications and email services .......................... 125
- Alerts .................................................................. 128
- Delete runs ....................................................... 129
- Manage Users ..................................................... 130
Export Data with SOLiD™ 4 System Software

The SOLiD™ 4 System software supports:

- Auto export
- Manual export

**Auto export**

You can set up SETS to auto-export data (.spch files) on a cycle-by-cycle basis for every run, or per run. Enable auto-export in SETS prior to a run.

**IMPORTANT!** Ensure that the off-instrument has the auto-export software installed.

**Manual export**

When opting to perform a manual export, you can configure the export to be all data or specific data. See “Manual export in SETS” on page 124 for details.
Auto export in SETS

To auto-export run data to the offline cluster, log into SETS, then go to Admin ▶ Change Preferences.

1. On the Edit User Preferences and Profile page, enter the host name or IP address of the server where you will export your runs to.

**IMPORTANT!** Prior to performing auto export, ensure that the off-instrument cluster has enough disk space.

2. Check the **Auto-export after primary analysis** box.

3. Click Save.

**IMPORTANT!** When a run is created, the “Export to Host” and “Auto-export after primary analysis” preferences are saved to that run. Any more changes to the preferences in the Edit User Preferences page will not affect the values of previous runs.

**IMPORTANT!** RSA keys must be set up between machines in order for the host name to be accepted. See “Manually configure the RSA keys” on page 121.
**Note:** Once auto export is configured in the SETS software, ensure that the auto export option is selected in the ICS, before starting the run.

**IMPORTANT!** If run export fails displaying a similar message to the one above, it is usually due to a change in the host IP address or a loss in the RSA key files. Manually configuring the RSA keys should resolve the problem.

The following diagram shows the deployment diagram on remote machine (the offline cluster).
Auto export can work in two modes:

1. With (Java Messaging Service) JMS broker and Export Daemon.

2. Without JMS broker and Export Daemon.

Auto export with JMS broker and Export Daemon

In this mode, all the delta spch is automatically transferred to remote location. On the remote side, the auto export daemon automatically runs readbuilder and barcoding. See the diagram below.
In this mode, the user can manually run readbuilder and barcoding. When auto-export is performed on a run, all the jobs folders are also exported to the remote cluster.

Jobs folders are typically located at:

```
/data/results/<instrument name>/<run name>/<sample name>/jobs.
```

To run read builder:

1. Log in to the linux cluster as user which can submit jobs to pbs (pipeline).
2. Change directory (cd) to the postPrimersetPrimary under jobs folder.
3. Run the following command:

   jprimaryanalysis.sh -dir /data/results/<instrument name>/<run name>/<sample name>/jobs /postPrimerSetPrimary.<job id> --ini=parameters.ini

   This will run readbuilder and generate csfasta output.

To run barcoding:

1. Log in to the linux cluster as user which can submit jobs to pbs (pipeline)

2. Change directory (cd) to the postBarcodePrimary under jobs folder.

3. Run the following command:

   jprimaryanalysis.sh -dir /data/results/<instrument name>/<run name>/<sample name>/jobs /postBarcodePrimary.<job id> --ini=parameters.ini

   This will run barcoding and split the csfasta into libraries as defined in the original run configuration. See the diagram below.
Example of saving preferences

1. User A logs in, and enters **Host 1** as the host name and checks **Auto-export after primary analysis** (Edit User Preferences page).

2. Then User A logs into ICS and creates a run “Run1_FC1.” User A’s preferences are applied to this run.

3. Next, User A goes back to SETS and changes the preferences to Export to Host 2.

4. User A returns to ICS and creates another run called “Run2_FC2.” This run copies the most recent preferences, including the new host name, and exports data once the run is complete.

   With the current setup, User A has created Run1 on Flowcell 1 auto-exporting to one host (Host 1) and Run 2 on Flowcell 2 to a different host (Host 2).
Manually configure the RSA keys

When you export, your runs will be exported to the server called “hostName”. For example, assume you are exporting a run from server_src to server_dest. Enter server_dest in the hostName field and check the Auto-export box.

If you get an error message, follow the steps below:

1. From the command line, confirm that each machine can see the other by the hostname on the network:
   - Use the secure shell (ssh) to log into the server_src, then type ping server_dest.
   - Use the secure shell (ssh) to log into the server_dest, then type ping server_src.
   In both cases, you should see bytes returned.

2. If you have access to remoteKey.sh, execute the following commands: (If you do not have access, go to step 3).
   - Use the secure shell (ssh) to log into the server_src as corona, then type remoteKey.sh pipeline@server_dest.
   - At the prompt for corona's password of server_dest, type in the password of corona user.
   - Use the secure shell (ssh) to log into the server_src as pipeline, then type remoteKey.sh pipeline@server_dest.
   - At the prompt for pipeline's password of server_dest, type the password of pipeline user.
   - Go to step 4.

3. If you do NOT have access to remoteKey.sh, execute these commands:
   - Use the secure shell (ssh) to log into the server_dest as pipeline, then go to the ssh directory
     - ssh pipeline@server_dest.
     - cd ~/.ssh.
   - Execute the following commands which copy the appropriate public keys onto server_dest:
     - scp corona@server_src:~/.ssh/id_rsa.pub/remote.pub then enter corona’s password.
     - cat remote.pub >> authorized_keys.
Chapter 7  Manage Administrative Tasks
Auto export in SETS

– scp
  pipeline@server_src:~/.ssh/id_rsa.pub./remote.pub
  then enter pipeline’s password
– cat remote.pub >>authorized_keys.
– Exit.

4. Test

Execute the following commands:
• ssh corona@server_src
• ssh pipeline@server_dest
• Exit.
• Exit from server_src
• ssh pipeline@server_src
• ssh pipeline@server_dest
• Exit.
To modify the export preferences for the current run only and keep run data stored locally:

1. Go to **Run Details ▶ Action box ▶ Host Configuration.**

2. Uncheck **Auto-export after primary analysis.**
   Any subsequent secondary analysis for the current run will now occur on the local cluster and run data will not be exported.

3. Click **Save.**
Manual export in SETS

Set up manual export

1. Log into SETS, then go to Run Details ➤ Action box.
2. Click Set Up Export...

Note: If the Host has not yet been configured in the Preferences, this link will be disabled.

3. Select one of the two export options, then the host location:

4. Click OK to save your settings.
5. On the Run Details page, click Export (action box) to manually export the run.
Event notifications and email services

Configuration setup

There is a one-time setup requirement for notifications and emails to work properly. Work with your IT representative to configure the correct SMTP host and port information for your network.

1. Log in to the SETS software.
2. Click **Admin** ➤ **Email Service**.
3. Check the **Activate Service** checkbox.
4. Enter your SMTP host number.
5. Enter you SMTP port number, or leave the port number blank to allow the system to assign the port number 25 by default.

6. Click **Save**.
Event notification

Event notification includes:

- Email notification
- Default actions when an event occurs

To set up event notifications:

1. Log in to the SETS software, if you are not already logged in.
2. Click **Admin ➤ Change Preferences**.
3. Enter the email address, or addresses, where you want to receive the event notifications.

**Note:** If you enter multiple email addresses, separate each with a comma.

4. Select the run-specific events you want to be notified about and the default actions to take under each category shown (System Errors, Fatal Run Errors, Informational Updates, and Completion Updates).
### Event notifications and email services

**Note:** When you select “My Runs Only” you will receive email notifications for the runs created only by you. When you select “All Runs” you will receive email notifications for all runs on the instrument.

5. Click **Save** to begin receiving email notifications.

**Default actions**

Default actions are available for these events:

- Run metrics (stop export)
- Run metrics (pause run)
- Low signal in spots (pause run)

Run Metrics event notifications are triggered whenever one or more of the quality or other metrics falls into the red zone during a run.
Alerts

Job Manager  If the Job Manager daemon service is not running, you may see this alert in SETS:

![Alert Center](image1)

Results disk space alert  If you are working in SETS, and you see this alert, you need to clean up disk space by either exporting or deleting data.

![Alert Center](image2)

Note: To change the default setting of your minimum space threshold, change the property file (runDirectory.properties). After making changes, restart ICS and the SETS server.
Delete runs

When you need to clean up disk space and want to delete runs, go to a particular run, then click **Delete** (Run Details page).

Select a delete option when prompted.

- Selecting intermediate results files deletes:
  - RUN_FOLDERS/SAMPLE_NAME/"jobs" folder
  - RUN_FOLDERS/SAMPLE_NAME/"results.*"/"colorcalls"/
    All folders except "CACHE"
  - RUN_FOLDERS/SAMPLE_NAME/"results.*"/"intermediat e.*" folder
  - RUN_FOLDERS/SAMPLE_NAME/"results.*"/"primary.*"/
    "reject" folder
  - RUN_FOLDERS/SAMPLE_NAME/"results.*"/"libraries"/
    LIBRARY_NAME/"intermediate.*" folder
  - RUN_FOLDERS/SAMPLE_NAME/"results.*"/"libraries"/
    LIBRARY_NAME/"primary.*"/"reject" folder
Chapter 7 Manage Administrative Tasks

Manage Users

- Selecting images deletes all imaging data associated with that run.
- Selecting all run data except database records deletes all results and imaging data associated with that run.
- Selecting all run data deletes all results, imaging data, and database records associated with that run.
- Selecting database records deletes the database records for that run, but nothing on the file system.

Note: After deletion, the run will not be viewable in SETS.

Manage Users

An administrator can add new users, create passwords, change user preferences, and inactivate current users under Admin ▶ Manage Users.

See “Log in to SETS” on page 21 for information about “admin”, “user”, and “guest” (default) log in permissions.

Create a New user

1. In the SETS Home page, go to Admin ▶ Manage Users. The Current Users window opens.

2. Click New to open the Create a New User dialog box.
3. Enter a name in the User name field to add a new user.

4. Select the Role – “guest”, “user”, or “admin” – (See “Log in to SETS” on page 21 for information about each role.)

5. Enter a password for that user, then re-enter the same password.

6. Click Submit.

**Inactivate a user** In the Current Users table, under the Activation column, click the check mark next to a current user name to inactivate that user.

Click **Inactivate** at the “Are you sure” prompt.
Deleting a current user deletes all analysis settings associated with that user.
Web Services

Overview

Within SETS, you can view the status of runs and control the analysis. SETS also exposes some of its functionalities through web services. Web services provides an open channel that allows you to integrate directly with the system.

Client code

Web-based API is a method used to send information from one computer system to another. Write web service client code, then point the client (computer/analyzer system) to the SETS software WSDL.
WSDL defines an API that can be used to programmatically retrieve information from the SOLiD™ 4 System. The WSDL definition (for SETS web services) is available through the SETS About menu.

The URL is:

[SETS service URL]/webservice/solid?wsdl

For example, if SETS is installed on a host named "foo," the [SETS service URL] might be:

http://foo:8080/sets.

The corresponding WSDL definition is available at:

http://foo:8080/sets/webservice/solid?wsdl

API documentation

On the iExternalService, go to the Method Summary table to see the services available.
### Method Summary

<table>
<thead>
<tr>
<th>Method</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>getStatus()</code></td>
<td>Get information about the status of the cluster.</td>
</tr>
<tr>
<td><code>getLastStartedRun()</code></td>
<td>Get the run last started on server1 and server2.</td>
</tr>
<tr>
<td><code>getWebServicesLog()</code></td>
<td>Get a list of log messages.</td>
</tr>
<tr>
<td><code>getRunByInstrumentName()</code></td>
<td>Get a run by instrument name.</td>
</tr>
</tbody>
</table>

### Binding

A Java binding to the web services is available on request. Bindings for other languages can be readily generated based on the SETS web services WSDL definition.
Advanced Topic: Data Analysis

Overview

The topics provided in this appendix are intended for advanced users of the SOLiD™ 4 System and do not apply to the typical user.

Fundamentals of color-space analysis

The Applied Biosystems SOLiD™ 4 System sequencing technology is based on sequential ligation of dye-labeled oligonucleotides. This technology makes possible massive parallel sequencing of clonally amplified DNA fragments. Features of this system, such as mate-paired analysis and 2-base encoding, enable studies of complex genomes by providing a greater degree of accuracy. This section describes the principles of 2-base encoding and the benefits of performing analysis in the di-base alphabet, known as color-space analysis.

The 2-base color coding scheme

Until recently, most DNA sequencing was performed using the chain termination method developed by Frederick Sanger. (Refer to the paper by Sanger F., Coulson A. R., 1975, A rapid method for determining sequences in DNA by primed synthesis with DNA polymerase. *J Mol Biol.* 94(3): 441-448.) This type of sequencing is often referred to as Sanger sequencing. Sanger sequencing data is also encoded in color-space by the four fluorescent dyes used in the sequencing chemistry and displayed as peaks in an electropherogram. In Sanger sequencing, each color, representing only a single nucleotide, is automatically translated to A, C, G, or T. With the SOLiD™ 4 System, each color represents four potential 2-base combinations (see Figure 1). The conversion into nucleotide base space is usually done after the sequence is aligned to a reference genome transcribed in color-space. As an alternative, translation can occur following the generation of a consensus sequence.
Appendix B Advanced Topic: Data Analysis Overview

Fundamentals of color-space analysis

<table>
<thead>
<tr>
<th>[code]</th>
<th>0</th>
<th>1</th>
<th>2</th>
<th>3</th>
</tr>
</thead>
<tbody>
<tr>
<td>[dye]</td>
<td>FAM</td>
<td>Cy3</td>
<td>TXR</td>
<td>Cy5</td>
</tr>
<tr>
<td>(XY)_1</td>
<td>AA</td>
<td>AC</td>
<td>AG</td>
<td>AT</td>
</tr>
<tr>
<td>(XY)_2</td>
<td>CC</td>
<td>CA</td>
<td>GA</td>
<td>TA</td>
</tr>
<tr>
<td>(XY)_3</td>
<td>GG</td>
<td>GT</td>
<td>CT</td>
<td>CG</td>
</tr>
<tr>
<td>(XY)_4</td>
<td>TT</td>
<td>TG</td>
<td>TC</td>
<td>GC</td>
</tr>
</tbody>
</table>

Figure 1 SOLiD™ 4 System’s 2-base color-coding scheme

Color-coding process

The column under code \( i \) (0, 1, 2, or 3) lists the corresponding dye and the di-base probes (adjacent nucleotides) encoded by color \( i \). For example, GT is labeled with Cy3® dye and coded as “1”.

The instrument takes the following steps to encode a DNA sequence. Consider the example ATCAAGCCTC:

1. Start at the 5’ end.
2. Replace the di-base AT at this position with its corresponding code 3 from the table.
3. Advance by one base, which shows the TC di-base (code 2).
4. Continue to advance by one base, as shown below.

Base Sequence: A T C A A G C C T C
Color String: 3 2 1 0 2 3 0 2 2

This process encodes a \( k \)-mer of bases as a \((k-1)\)-mer of colors. Although this color string codes for four different \( k \)-mers, knowledge of the type and position of any of its \( k \) bases helps to encode the sequence. For SOLiD™ 4 System sequencing applications, the
instrument prepends the leading base A in the example above to result in a $k$-mer A321023022 to reconstruct the base sequence. The SOLiD™ 4 System generates its reads in this encoded form, as shown in Figure 2.
Figure 2  Sequence by ligation using di-base labeling probes on the SOLiD™ 4 System
Required properties for a 2-base color code scheme

The 2-base color-coding scheme satisfies the requirements in the following list. This conclusion can be observed by treating the properties as requirements and constructing the color code from them. This section addresses only bases, not other International Union of Biochemistry (IUB) codes. Let \( B = \{A, C, G, T\} \).

The color code should satisfy the following requirements. For all bases \( b, d, e \) in \( B \):

1. The available colors are 0, 1, 2, and 3.  
   \[ \text{color} \ (bd) \in \{0, 1, 2, 3\}. \]

2. Two different di-base probes that have the same first base result in different colors.  
   \[ \text{color} \ (bd) \neq \text{color} \ (be) \text{ if } d \neq e. \]  
   For example, \( \text{color} \ (AC) \neq \text{color} \ (AG) \).

3. A di-base probe and its opposite result in the same color.  
   \[ \text{color} \ (bd) = \text{color} \ (db). \]  
   For example, \( \text{color} \ (AC) = \text{color} \ (CA) \).

4. Mono- and di-base probes result in the same color.  
   \[ \text{color} \ (bb) = \text{color} \ (dd). \]  
   In other words, \( \text{color} \ (AA) = \text{color} \ (CC) = \text{color} \ (GG) = \text{color} \ (TT) \).

   The following are interesting properties that follow from the above four requirements. Property 5 follows from requirements 2 and 3 and makes constructing the color code easier.

5. Two different di-base probes that have the same second base result in different color codes: \( \text{color} \ (bd) \neq \text{color} \ (cd), \text{ if } b \neq c. \)  
   For example, \( \text{color} \ (AC) \neq \text{color} \ (TC) \).

   Property 6 also follows from requirements 1-4, but it is most easily verified against the completed code (see Figure 3, Panel E).
6. A di-base probe and its complement result in the same color.
   \[ \text{color (bcdc)} = \text{color (dbdc)}. \]
   For example, \( \text{color (AC)} = \text{color (TG)} \).

How the system satisfies 2-base coding system requirements

Figure 3 lists the colors for each di-base probe. For example, the value in row C and column T is the color 2 for di-base CT.

Requirements 1 and 2 require that all colors are present in the first row. Because the system can use any one-to-one mapping between the actual dyes and the labels 0, 1, 2, and 3 (provided that requirements 1 and 2 are satisfied), the first row can be labeled as shown in Figure 3, Panel B.

Requirement 3, that \( \text{color (bd)} = \text{color (db)} \), gives a unique labeling for column A (see Figure 3, Panel C).

Requirement 4, that \( \text{color (bb)} = \text{color (AA)} \), gives a unique labeling for the diagonal (see Figure 3, Panel D).

Finally, requirements 1, 2, and 5 state that every color must appear in every row and every column exactly once (see Figure 3, Panel E).

The table (see Figure 3, Panel E) is easy to memorize and work with because, by virtue of Property 3, di-base probes can be thought of as two-element sets for assigning colors. The di-base probes that start with A result in colors 0, 1, 2, and 3 respectively.

Therefore:

- AA, CC, GG, TT all are assigned color 0.
- AC and CA are assigned color 1, and so must GT and TG.
- AG and GA are assigned color 2, and so must CT and TC.
- AT and TA are assigned color 3, and so must CG and GC.
Principles of ligation-based chemistry and 2-base encoding

The SOLiD™ 4 System’s sequencing technology is based on sequential ligation of dye-labeled oligonucleotide probes. Each probe assays two base positions at a time (Figure 2). The system uses four fluorescent dyes to encode for the sixteen possible 2-base combinations. Multiple ligation cycles of probe hybridization, ligation imaging, and analysis are performed to extend the strand from a primer hybridized to a ligated adaptor by the immobilized bead (P1 adaptor). The resulting product is then removed and the process repeated for 5 more rounds with primers hybridized to positions n-1, n-2, and so on, in the P1 adaptor.

There are several fundamental properties unique to ligation-based sequencing. These properties contribute to the high accuracy inherent in the SOLiD™ 4 System. The advantages of these properties and their contribution to data quality are:

- Two bases are interrogated in each ligation reaction, increasing specificity.
• The primer is periodically reset for five or more independent rounds of extension reactions, improving the signal-to-noise ratio of the system.
• Each base is interrogated twice in two independent primer rounds, increasing confidence in each call.
• Four dyes are used to encode for sixteen possible 2-base combinations. The design of the encoding matrix enables built-in error-checking capability.
Color-space and base space as applied in the SOLiD™ 4 System

The final files generated by the SOLiD™ 4 System are in base space. To maximize the built-in error correction of 2-base encoding, all file analysis is conducted in color-space prior to final results.

Color-space data

Rather than reading one base per cycle, the software measures information on two bases simultaneously. In each cycle, the software calls one of four colors (color-space call). Because each ligation measurement event measures two bases, all bases (except for the final base of a read) are interrogated twice, providing for an additional level of error correction.

Relationship between cycle and base position

Figure 4 shows the relationship between color-space, base position, and the sequencing chemistry. Base number refers to base position. Base 0 is the last base of the adapter and is not part of the target sequence. The five primer lines show the order in which the data was generated (0-indexed).

Note: In all files, only processed data refers to a color-space position.
**Figure 4** Relationship between color-space, base position, and sequencing chemistry

**Color-space formats**

Color-space data are presented in three slightly different formats. In two of the formats, a base (A, C, G, or T) is appended to the color-space calls.

**Note:** Color-space data are self-complementary: In some situations, when you might expect to see complemented data (for example, reverse), the data appear the same. For example, AC = 1, TG = 1.

The different types of color-space data are:

- **Processed color-space data:** Consists of a numeric string prefixed (suffixed if reversed) by a single base. The base that precedes the numeric (color code) data is the first base of the actual sequence (in base space, not color-space).

- **Unprocessed color-space data:** Consists of a numeric string prefixed by a single base. This base is the final base of the sequencing adapter and is not part of the target sequence. It is included to disambiguate the first color call.
Complementing color-space data

Color-space data are self-complementary as shown in Figure 5.

<table>
<thead>
<tr>
<th>1st Nucleotide</th>
<th>A</th>
<th>C</th>
<th>G</th>
<th>T</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>C</td>
<td>1</td>
<td>0</td>
<td>3</td>
<td>2</td>
</tr>
<tr>
<td>G</td>
<td>2</td>
<td>3</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>T</td>
<td>3</td>
<td>2</td>
<td>1</td>
<td>0</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>2nd Nucleotide</th>
<th>A</th>
<th>C</th>
<th>G</th>
<th>T</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>C</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>G</td>
<td>2</td>
<td>3</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>T</td>
<td>3</td>
<td>2</td>
<td>1</td>
<td>0</td>
</tr>
</tbody>
</table>

**Sequence:**

<table>
<thead>
<tr>
<th>Base</th>
<th>A</th>
<th>G</th>
<th>C</th>
<th>T</th>
<th>C</th>
<th>G</th>
<th>T</th>
<th>G</th>
<th>C</th>
<th>A</th>
<th>G</th>
</tr>
</thead>
<tbody>
<tr>
<td>Color-space</td>
<td>2</td>
<td>3</td>
<td>2</td>
<td>2</td>
<td>3</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>1</td>
<td>1</td>
<td>3</td>
</tr>
</tbody>
</table>

**Complemented:**

<table>
<thead>
<tr>
<th>Base</th>
<th>T</th>
<th>C</th>
<th>G</th>
<th>A</th>
<th>G</th>
<th>C</th>
<th>A</th>
<th>G</th>
<th>C</th>
<th>A</th>
<th>G</th>
<th>T</th>
<th>C</th>
</tr>
</thead>
<tbody>
<tr>
<td>Color-space</td>
<td>2</td>
<td>3</td>
<td>2</td>
<td>2</td>
<td>3</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>1</td>
<td>1</td>
<td>3</td>
<td>1</td>
<td>2</td>
</tr>
</tbody>
</table>

**Figure 5  Color-space data**
2-base encoding and error recognition

The error-checking abilities of the 2-base encoding schemes have not been used to correct any of the data provided in these files.

Example:

Reference = 2 3 2 2 \textbf{3} 1 2 3 1 3 1 2

Observed = 2 3 2 2 \textbf{0} 1 2 3 1 3 1 2

\textbf{Note:} Notice the single color-space error.

In this example, a single color-space error occurs. The most likely explanation for the observed 0 is that it is a measurement error. Because a single color-space change is not allowed, a change to one of the adjacent bases is needed for a real SNP. This correction requires multiple measurement errors, leaving the most likely explanation that the 0 is a 3. The fact that the two surrounding bases are the same as the reference is further evidence that correcting the 0 to a 3 is acceptable. Any single color error can likely be corrected, especially when using multiple aligned reads.

Processing

Typically, a panel is imaged four times per cycle (once per channel). The color-space calls are carried out on a per-cycle basis. Each data point represents an individual bead with four intensity values. Beads are assigned a color-space call using a clustering algorithm. As part of the clustering process, data are scaled and baselined, quality values are assigned, and color call is assigned.

Results of primary analysis for each panel are stored in files with the “.spch” extension (Solid Panel Cache HDF5). The “.spch” file is used as a cache; that is, primary analysis both writes results to the file and reads results from the file as analysis proceeds on a cycle-by-cycle basis. Data within the spch file are organized by cycle. When all cycles have been analyzed, a final processing step is run to convert the data from the cycle-based format to the “read-based” ASCII formats (“.csfasta” and “.qual”).
The spch file is in the HDF5 format. Details on the format are available at:

http://www.hdfgroup.org/HDF5/.

Tools to view the contents and extract results are available at: http://www.hdfgroup.org/hdf-java-html/hdfview/.

Prior to secondary analysis, results are filtered by removing all tags with missing data. Filtering removes all incomplete beads and reads with missing calls. The filtered data are then processed to generate the xxxx.csfsta file. As part of this process, a known base (from the adapter sequence) is prepended to the color-space data, allowing disambiguation of the first color call. Each read is treated separately, even if it is part of a mate-paired tag. The file contains the color-space read for each tag, and the color-space data are used as the basis of all subsequent results. The data are then aligned in color-space to the color-space reference sequence.

During analysis and alignment:

- Color-space reference sequences are derived by converting the base-space reference sequence to color-space (this process occurs internally; you supply the reference sequence in base space).
- All alignment is done in color-space.
- All the alignments are with individual tags; the mate-paired information is not used in the initial alignments.
- After the single reads are analyzed, a further round of analysis is performed in which the mate-pair information is used.
- The paired tags are analyzed in a two-step process:
  a. The tags where both the forward and reverse tags passed (both tags were matched to a reference sequence) are analyzed.
b. Those mate-pairs where only one of the tags matched to the reference sequence (in color-space) are re-analyzed, and the non-matching tag is compared to the reference sequence.

Because analysis can be constrained by the knowledge of the allowed distances, this process allows alignment with more disagreements and the use of a more CPU-intensive alignment algorithm.

**Note:** Where possible, all data are presented in a fasta-compatible format to facilitate use in a variety of applications. Final output files are in base space and relative to the submitted reference (for example, consensus file, variation file).
Data analysis considerations

Understand Ns and color-space

In the SOLiD™ 4 System, unknown color-space calls are represented as ‘.’ or N. By default, the system filters out bases with a single ‘.’ in the sequence and leaves only reads with color-space calls present at all positions. You can change this behavior by masking a specific position of a read. For example, if position 21 of a 35 bp run is missing color-space calls, you can mask this position.

Find Single Nucleotide Polymorphisms (SNPs)

SNPs are single base-pair mutations that usually consist of two alleles. Color-space rules allow the software to detect only valid SNP sites. According to the color-space rules, one isolated color change is always an error when you consider single nucleotide changes. Two adjacent color changes can be either an error or a valid SNP. To detect two adjacent SNPs, look for three adjacent color changes. There are very specific rules to determine which color changes are valid and which are not.

Color-space rules for SNP detection

In the simplest case, where you have a single base change, the following rules apply (Figure 6):

- For any given reference, there are only three valid double-color changes.
- For the other 12 possibilities, the 6 single-color and 6 two-color changes are invalid, since they would require multiple changes within that genomic region. For these complicated genomic changes, the software uses a more sophisticated algorithm.
Appendix B  Advanced Topic: Data Analysis Overview

Data analysis considerations

Figure 6  Color-space rules for single nucleotide polymorphism (SNP) sites

Color-space rules are automatically applied during data analysis using the SOLiD™ 4 System’s software analysis tools. You do not need to apply these filters manually.
SNPs and errors

Because a SNP generates two adjacent and valid color-space mismatches to the reference sequence, the software allows a minimum of two color-space mismatches during alignment to the reference sequence. Depending on the complexity of the alignment, the optimum number of mismatches in alignment is at least three. If only two errors are allowed, then whenever there is a SNP (two mismatches) and a measurement error, the third error causes omission of that read. You can override the omission by using an alignment program setting that counts two adjacent mismatches as a single mismatch. Therefore, two adjacent mismatches and a measurement error are classified as two errors, whereas three non-adjacent errors are classified as three errors.

High accuracy for SNP detection

2-base encoding provides high system accuracy and built-in error checking because it enables discrimination between measurement errors and true sequence polymorphisms. The SOLiD™ software interrogates each base twice in two independent reactions. Information about each base is included in two adjacent pieces of color-space data. Because the system uses four fluorescent dyes, there are 16 possible two-color combinations (Figure 7).

<table>
<thead>
<tr>
<th></th>
<th>Blue</th>
<th>Green</th>
<th>Yellow</th>
<th>Red</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blue</td>
<td>BB</td>
<td>BG</td>
<td>BG</td>
<td>BR</td>
</tr>
<tr>
<td>Green</td>
<td>GB</td>
<td>GG</td>
<td>GY</td>
<td>GR</td>
</tr>
<tr>
<td>Yellow</td>
<td>YB</td>
<td>YG</td>
<td>YY</td>
<td>YR</td>
</tr>
<tr>
<td>Red</td>
<td>RB</td>
<td>RG</td>
<td>RY</td>
<td>RR</td>
</tr>
</tbody>
</table>

Figure 7  Color-space transitions
A SNP must have two adjacent color changes. Single-color mismatches are not evidence of a SNP. This feature allows SOLiD™ software to distinguish errors in measurement from true SNPs.

**SNP error rates**

When you use 2-base encoding, any SNP in the original sequence is represented as two adjacent mismatches in color-space. Only three of nine possible adjacent mismatches can correspond to a real SNP. Suppose the raw sequencing error rate for a species is 1% per base, and for the same species that is sequenced, the SNP rate is about 0.1%. For a sequencing project of \( M \) total bases, there are about 0.001\( M \) real SNP occurrences in the data set. Among these SNPs, 0.001\( M \times 0.02 \) total cases appear as single mismatches or two invalid mismatches because a sequencing error happens at one of the alleles of the SNP. Only 2% of the real SNPs fail to appear as two adjacent, valid mismatches. If only two adjacent, valid mismatches are treated as candidates for SNP detection, then there is a 2% false negative rate. For any data set, two adjacent and valid mismatches can be caused by sequencing errors. However, for a total of \( M \) bps sequenced, there are 0.00003\( M \) total occurrences of two adjacent mismatches. Note that there are about 0.001\( M \) adjacent valid mismatches from real SNPs, among all two adjacent, valid mismatches observed in a particular data set. Of these, 97% are from real SNPs, and only 3% may be from sequencing errors. This is a 97% true discovery rate for that particular data set. Because there are about 0.01\( M \) total sequencing errors in the data set caused by 2-base encoding and the software’s ability to remove all single base mismatches, the error is reduced from 0.01\( M \) to 0.00003\( M \). This is a reduction of 300 times, making the effective error rate 0.003%. This calculation illustrates the power of 2-base encoding in resequencing and finding SNPs.
Sampling

If coverage at a genome position is low, the second allele might often not be sampled, even if it is present. Heterozygotes are expected to be underrepresented at low coverage. (They can be called as a homozygote for one of the two alleles.) It is important to have a low false positive rate for SNP detection, because SNPs are expected to exist at only 1 in 1,000 positions for humans and some other species. The false positive rate in an individual species should be at least an order of magnitude lower than this, to avoid a high false discovery rate. An error model that incorporates quality values (QV) can increase the overall accuracy of SNP detection. The SNP detection algorithms of the SOLiD™ 4 System use explicit error models generated from each run and biologically meaningful prior probabilities to evaluate evidence of a SNP. 2-base encoding provides a built-in way to distinguish errors from true alleles to manage the false positive rate.

Allele ratio

If the sample preparation method or some other cause produces results in allele ratios that are considerably different from the expected 50:50 ratio, it is more difficult to detect heterozygosity. Detecting heterozygosity then requires more sequence coverage.

Find polymorphisms in color-space

The SOLiD™ 4 System can detect complicated genomic variations such as adjacent SNPs, insertions, deletions, and structural rearrangements.

For other analysis tools, visit the AB SOLiD™ Software Community website:

http://www3.appliedbiosystems.com/AB_Home/applicationstechnologies/SOLiDSystemSequencing/SoftwareCommunityDataAnalysisResourcesforScientists Developers/index.htm

An example of various polymorphisms in color-space is shown in the figure below.
Figure 8  Examples of polymorphisms in color-space
Advanced Topic: Data Management

The topics provided in this appendix are intended for advanced users of the SOLiD™ 4 System and do not apply to the typical user.

Saving data

This appendix describes data storage for the following:

- All data
- Raw image data
- Primary analysis results data

Approximate storage space for all data

The following table shows the size of all data for archival and storage consideration. The following is estimated file sizes.

<table>
<thead>
<tr>
<th>50 nt tag/300k/panel, 2357 panels</th>
<th>Image data size‡</th>
<th>Primary analysis results size in “.spch” format</th>
<th>Primary Analysis Data size (flatfile: “.csfasta”, “.QV.qual”, “.stats”)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 slide - 1 tag</td>
<td>1.84 TB</td>
<td>646 GB</td>
<td>170 GB</td>
</tr>
<tr>
<td>1 slide - 2 tags</td>
<td>3.6 TB</td>
<td>1.29 TB</td>
<td>340 GB</td>
</tr>
<tr>
<td>2 slides - 1 tag</td>
<td>3.6 TB</td>
<td>1.29TB</td>
<td>340 GB</td>
</tr>
<tr>
<td>2 slides - 2 tags</td>
<td>7.2 TB</td>
<td>2.58 TB</td>
<td>680 GB</td>
</tr>
</tbody>
</table>

‡ Image data is not needed after analysis is complete.
Raw image data

The data should be kept until the full completion of primary analysis is verified and primary analysis results are archived. After confirming that the image files are no longer needed, you can delete these image files to make room for additional sequencing runs.

Primary analysis results data

The image analysis results are stored in “.spch” files, which are located in:

```
$sampleResultFolder/colorcalls/CACHE/.
```

Table 2  Image analysis results files

<table>
<thead>
<tr>
<th>File type</th>
<th>File size</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>.spch</td>
<td>646 GB</td>
<td>Should be archived for the lowest-level non-image backup. An estimate for the disk space needed for one “.spch” file (per panel) is ( n \times (18m + 17) ) bytes, where ( n ) is the number of beads, and ( m ) is the number of cycles (F3 and R3). For 50 cycles (1x50) and 300K beads, this yields 275 MB (per panel). A full slide has 2357 panels, which means that 646 GB are required for a full slide (50 bp single tag run).</td>
</tr>
</tbody>
</table>

The color call results, including color-space raw reads and QV data, are located in:

```
$sampleResultFolder/primary.[17-digit timestamp]/reads/.
```

Table 3  Color call results files

<table>
<thead>
<tr>
<th>File type</th>
<th>File size</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>.csfasta</td>
<td>56 - 80 GB</td>
<td>Should be archived.</td>
</tr>
<tr>
<td>_QV.qual</td>
<td>112 - 140 GB</td>
<td>Should be archived.</td>
</tr>
<tr>
<td>.stats</td>
<td>&lt; 10 KB</td>
<td>Should be archived.</td>
</tr>
<tr>
<td>postPrimerSetPrimary.[JobID]</td>
<td>364 - 420 GB</td>
<td>Temporary files. They are removed after completion. Located in $sampleJobFolder/.</td>
</tr>
</tbody>
</table>
There are also color-space raw reads that were filtered in the sequence generation process. These raw reads include duplicate reads and reads with less than the expected read length. They can be found in:

$sampleResultFolder/primary.[17-digit timestamp]/reject/

<table>
<thead>
<tr>
<th>File type</th>
<th>File size</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>.csfasta.reject</td>
<td>&lt; 1 GB</td>
<td>Should be archived.</td>
</tr>
<tr>
<td>.QV.qual.reject</td>
<td>&lt; 1 GB</td>
<td>Should be archived.</td>
</tr>
</tbody>
</table>

The following files can be ignored. They can be archived if needed.

<table>
<thead>
<tr>
<th>File type</th>
<th>File size</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intensity files</td>
<td>$4 \times 336 - 420 = 1344-1680$ GB</td>
<td>Optional. Not generated by default. $sampleResultFolder/primary.[17-digit timestamp]/reads/ or $sampleResultFolder/primary.[17-digit timestamp]/reject/</td>
</tr>
</tbody>
</table>
Data transfer

Data transfer from the SOLiD™ 4 System is supported and tested on a 1-GB LAN. Applied Biosystems strongly recommends that you set up a dedicated 1-GB network between the SOLiD™ 4 System and the rest of your network. To ensure the appropriate transfer speed, you need to configure your network according to one of the suggested solutions:

<table>
<thead>
<tr>
<th>Current network</th>
<th>Solution</th>
</tr>
</thead>
<tbody>
<tr>
<td>1-GB LAN (shared)</td>
<td>On a shared network, the effective network speed is much lower than 1GB. Use or install a dedicated 1-GB network between the instrument and the rest of your network to ensure top transfer speeds.</td>
</tr>
<tr>
<td>No Gigabit LAN</td>
<td>Install a gigabit switch with the shortest path spanning tree. Connect all the SOLiD™ 4 Analyzer outgoing connections to the switch so that data transfer between the SOLiD™ 4 Analyzer and the rest of the network only occurs through the extra switch at 1 Gbps.</td>
</tr>
</tbody>
</table>

Exporting data

You export data from the SOLiD™ 4 Analyzer using SOLiD™ 4 Experiment Tracking System (SETS) software. You can set up SETS to auto-export data (csfasta and “.spch” files) for every run, or per run. You can also export files manually to a specified destination.

For detailed procedures, see Chapter 7, “Manage Administrative Tasks” on page 113.
Advanced Topic: Primary Analysis

The topics provided in this appendix are intended for advanced users of the SOLiD™ 4 System and do not apply to the typical user.

What is primary analysis?

Primary analysis is the process of extracting color calls or assignments (.csfasta file) from images captured from a sequencing run, calculating a quality value for each color call, and writing the color-call value to a QV file (.qual).

Primary analysis also provides real-time feedback about a sequencing run.
Primary analysis overview

Table 4 lists the stages in primary analysis, their inputs, and their outputs.

<table>
<thead>
<tr>
<th>Analysis stage</th>
<th>Inputs</th>
<th>Outputs</th>
<th>Steps</th>
</tr>
</thead>
<tbody>
<tr>
<td>postBeadFinding</td>
<td>Focal map and first ligation cycle &quot;tiff&quot; images</td>
<td>• Bead location (in <em>.spch</em> files)</td>
<td>For each panel, analyze the focal map and first ligation cycle images to identify beads.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Image statistics and metrics (in *.FOCALMAP_summary.tab files)</td>
<td></td>
</tr>
<tr>
<td>postScanSlide</td>
<td>.tiff images from each ligation cycle</td>
<td>• Intensities, Color calls, Quality metrics, QV, in <em>.spch</em> files</td>
<td>1. Align images.</td>
</tr>
<tr>
<td></td>
<td>Focal map &quot;tiff&quot; images</td>
<td>• Satay &quot;.png&quot; files</td>
<td>2. Extract intensities.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Summary data (tab)</td>
<td>3. Classify colors (color calls).</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Run status info (traffic lights) (tab).</td>
<td>4. Calculate quality value (QV).</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>5. Generate satay images.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>6. Accumulate run status information.</td>
</tr>
<tr>
<td>postPrimerSet</td>
<td>All <em>.spch</em> files</td>
<td>• Color-space raw reads (&quot;.csfasta&quot;)</td>
<td>1. Assemble color-space reads per panel.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• QV (&quot;.qual&quot;)</td>
<td>2. Concatenate per-panel reads into one single &quot;.csfasta&quot; file (optional).</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Reads statistics file (&quot;.stats&quot; file).</td>
<td>3. Filter reads shorter than specified read length or duplicate reads (optional).</td>
</tr>
</tbody>
</table>

Table 4 Primary analysis stages, inputs, and outputs
**Primary analysis workflow**

In the primary analysis workflow, each spot on the sample slide is broken down into a number of panels. Each panel is defined by the area that is imaged by the camera in one exposure. Primary analysis handles each panel independently of the others. For each panel, primary analysis comprises a sequence of steps. When the Instrument Control Software (ICS) has finished collecting the images for each step, it creates a job workflow in the SOLiD™ 4 System database. The Job Manager spots the job workflows and prepares and submits the needed jobs to the PBS resource manager. When a job completes, the Job Manager updates the SOLiD™ 4 System database with the appropriate analysis status.

Each primary analysis step falls into one of the following three classes:

- **Bead finding**: Focal map and first ligation cycle images are analyzed to identify and locate beads. Additionally, various image statistics and metrics are computed and recorded.
- **Color calling**: After a ligation cycle, the four color images are registered to the focal map image. The fluorescent intensity for each color is estimated for each bead. These color intensities are analyzed to determine a color call and quality value for each bead for that cycle.
- **Read building**: After a complete run, the cycle-wise results are assembled into color reads. After some optional filtering, these reads are then ready for subsequent secondary analysis (e.g., mapping to a reference sequence).

See Figure 9 for the primary analysis workflow. Multiple primary analysis jobs can be run on the Linux computer cluster. The analysis results are written to various files on this cluster. Upon completion, the primary analysis output includes a summary statistics file that shows a summary of the overall color calls. This file can be used to evaluate whether primary analysis completed successfully.
Appendix D  Advanced Topic: Primary Analysis

Primary analysis workflow

Figure 9  Primary analysis workflow
Primary analysis inputs and outputs

The input images are in the following files:

**Focal Map**
/data/images/{run.name}/{run.name}_FOCALMAP_V1

**Ligation Cycle**
/data/images/{run.name}/{run.name}_F3_P1_01_V1
/data/images/{run.name}/{run.name}_F3_P1_02_V1
...
... 
/data/images/{run.name}/{run.name}_R3_P1_01_V1
...
...

$sampleJobFolder =
/data/results/{instrument.name}/{run.name}/{sample.name}/jobs

$sampleResultFolder =
/data/results/{instrument.name}/{run.name}/{sample.name}/results

**Job analysis parameter and temporary files for the Focal Map**
$sampleJobFolder/postBeadFindingPrimary.[JobID]

**Job analysis parameters and temporary files for ligation cycle color calls**
$sampleJobFolder/postScanSlidePrimary.[JobID]
$sampleJobFolder/postScanSlidePrimary.[JobID]
...
...

**Job analysis parameters and temporary files for postPrimerSetPrimary**
$sampleJobFolder/postPrimerSetPrimary.[JobID]
Image analysis results ".spch" files (SOLiD Panel Cache HDF5)

$sampleResultFolder/colorcalls/CACHE/{run.name}_{panelID}.spch (one full slide has 2357 panels)

Satay ".png" files

$sampleResultFolder/cycleplots

Summary data

$sampleResultFolder/colorcall_summary/{run.name}_FOCALMA_P_V1.tab
$sampleResultFolder/colorcall_summary/{run.name}_F3_P1_01_V1.tab
.. $sampleResultFolder/colorcall_summary/{run.name}_R3_P1_01_V1.tab
.. $sampleResultFolder/colorcall_summary/{run.name}_BC_P1_01_V1.tab

Run status information (traffic light)

$sampleResultFolder/traffic_lights/

Color-space F3 raw reads (.csfasta)

$sampleResultFolder/primary.[17-digittimestamp]/reads/{run.name}_{sample.name}_F3.csfasta

Color-space BC raw reads (.csfasta)

$sampleResultFolder/primary.[17-digittimestamp]/reads/{run.name}_{sample.name}_BC.csfasta
**Color-space F3 raw reads for library (.csfasta)**

`$sampleResultFolder/libraries/{library.name}/primary.[17 - digittimestamp1]/reads/{run.name}_{sample.name}_F3.csfasta`

**Color-space BC raw reads for library (.csfasta)**

`$sampleResultFolder/libraries/{library.name}/primary.[17 - digittimestamp]/reads/{run.name}_{sample.name}_BC.csfasta`

**F3 QV (.qual) file for library**

`$sampleResultFolder//libraries/{library.name}/primary.[17-digittimestamp]/reads/{run.name}_{sample.name}_F3_QV.qual`

**BC QV (.qual) file for library**

`$sampleResultFolder//libraries/{library.name}/primary.[17-digittimestamp]/reads/{run.name}_{sample.name}_BC_QV.qual`

**F3 Primary analysis statistics file (.stats)**

`$sampleResultFolder/primary.[17-digittimestamp]/reads/{run.name}_{sample.name}_F3.stats`

**BC Primary analysis statistics file (.stats)**

`$sampleResultFolder/primary.[17-digittimestamp]/reads/{run.name}_{sample.name}_BC.stats`

**F3 Primary analysis statistics file (.stats) for library**

`$sampleResultFolder/libraries/{library.name}/primary.[17 - digittimestamp]/reads/{run.name}_{sample.name}_F3.stats`
BC Primary analysis statistics file (.stats) for library

$sampleResultFolder/libraries/{library.name}/primary.[17-digit timestamp]/reads/{run.name}_{sample.name}_BC.stats

Reads that have less than expected read length or duplicated

$sampleResultFolder/primary.[17-digit timestamp]/reject

Reads that have less than expected read length or duplicated for library

$sampleResultFolder/libraries/{library.name}/primary.[17-digit timestamp]/reject

Reads without barcode reads

$sampleResultFolder/libraries/{library.name}/missing-bc
$sampleResultFolder/libraries/{library.name}/missing-f3

Reads that could not be assigned to a library

$sampleResultFolder/libraries/{library.name}/unassigned

Barcode assignment statistics file/report

$sampleResultFolder/libraries/BarcodeStatistics.[17-digit timestamp].txt
Storage requirements for primary analysis

The storage requirements values given in the table below assume a 50-base tag run in one flowcell. If you use a tag size that is larger or smaller than 50 bases, the storage requirements differ from those given in Table 5.

Table 5  Primary analysis storage requirements

<table>
<thead>
<tr>
<th>Files</th>
<th>Storage required</th>
<th>Notes about archiving</th>
</tr>
</thead>
<tbody>
<tr>
<td>.csfasta</td>
<td>56 - 80 GB</td>
<td>Should be archived.</td>
</tr>
<tr>
<td>_QV.qual</td>
<td>112 - 140 GB</td>
<td>Should be archived.</td>
</tr>
<tr>
<td>.spch</td>
<td>646 GB</td>
<td>Should be archived for the lowest-level non-image backup. An estimate for the disk space needed for one &quot;spch&quot; file (per panel) is $n(18m+17)$ bytes, where $n$ is the number of beads, and $m$ is the number of cycles (F3 and R3). For 50 cycles (1x50) and 300K beads, this yields 275 MB (per panel). A full slide has 2357 panels, which means that 646 GB are required for a full slide (50 bp single tag run).</td>
</tr>
<tr>
<td>postPrimerSetPrimary.[JobID]</td>
<td>364 - 420 GB</td>
<td>Temporary files. Most of these are removed upon completion.</td>
</tr>
<tr>
<td>Intensity files</td>
<td>$4 \times 364 - 420 = 1344 - 1680$ GB</td>
<td>Optional. Not generated by default. It is strongly recommended that intensity files be generated with offline ReadBuilder, which is part of the BioScope™ Software.</td>
</tr>
</tbody>
</table>
Perform primary re-analysis

To perform primary analysis manually, use the following command:

```
jprimaryanalysis.sh --ini=PrimaryParameters.ini
```

**Note:** Only buildReads tasks are executed for re-analysis. Neither beadFinding nor colorCalling are enabled for re-analysis in the pipeline.

The PrimaryParameters.ini file can be found inside the JobID folder with the file named `parameters.ini`.

$sampleJobFolder/postPrimerSetPrimary

To prepare for the re-analysis:

1. Create a new analysis directory, such as:
   
   `postPrimerSetPrimary.m1`

2. Copy the following file onto the new directory.
   
   `parameters.ini`

3. Edit the following file to ensure that all the paths are correct.
   
   `PrimaryParameters.ini`

4. Remove parameters marked for daemon property.
   For example:
   
   `analysis.job.id`
   
   (Default behavior does not accept non-integer.)
   
   `update.solid.script`
   
   (Used by the daemon for updating the SOLiD™ 4 System database.)
   
   `logging.socket`
   
   (Used by the daemon for logging analysis progress.)

5. Enter

   ```
   jprimaryanalysis.sh --ini=PrimaryParameters.ini
   ```
The following is an example of the PrimaryParameters.ini file:

```
# Primary Analysis Settings
instrument.name=Solid
run.name=Solid_20080310_1
sample.name=DH10B
panels=1-2357
primer.set=F3
read.length=35
read.prefix=T

# Analysis Job Parameters
focal.map.dir=Solid_20080310_1/Solid_20080310_1_FOCALMAP_V1
focal.map.stg=Solid_20080310_1/Solid_20080310_1_FOCALMAP_V1/Solid_20080310_1_FOCALMAP_V1.STG
focal.map.version=1
images.dir.1=Solid_20080310_1/Solid_20080310_1_F3_P5_01_V1
images.dir.2=Solid_20080310_1/Solid_20080310_1_F3_P4_01_V1
images.dir.3=Solid_20080310_1/Solid_20080310_1_F3_P3_01_V2
```
images.dir.4=Solid_20080310_1/Solid_20080310_1_F3_P2_01_V1
images.dir.5=Solid_20080310_1/Solid_20080310_1_F3_P1_01_V1
images.dir.6=Solid_20080310_1/Solid_20080310_1_F3_P5_02_V1
images.dir.7=Solid_20080310_1/Solid_20080310_1_F3_P4_02_V1
images.dir.8=Solid_20080310_1/Solid_20080310_1_F3_P3_02_V1
images.dir.9=Solid_20080310_1/Solid_20080310_1_F3_P2_02_V1
images.dir.10=Solid_20080310_1/Solid_20080310_1_F3_P1_02_V1
images.dir.11=Solid_20080310_1/Solid_20080310_1_F3_P5_03_V1
images.dir.12=Solid_20080310_1/Solid_20080310_1_F3_P4_03_V1
images.dir.13=Solid_20080310_1/Solid_20080310_1_F3_P3_03_V1
images.dir.14=Solid_20080310_1/Solid_20080310_1_F3_P2_03_V1
images.dir.15=Solid_20080310_1/Solid_20080310_1_F3_P1_03_V1
images.dir.16=Solid_20080310_1/Solid_20080310_1_F3_P5_04_V1
images.dir.17=Solid_20080310_1/Solid_20080310_1_F3_P4_04_V1
images.dir.18=Solid_20080310_1/Solid_20080310_1_F3_P3_04_V1
images.dir.19=Solid_20080310_1/Solid_20080310_1_F3_P2_04_V1
images.dir.20=Solid_20080310_1/Solid_20080310_1_F3_P1_04_V1
images.dir.21=Solid_20080310_1/Solid_20080310_1_F3_P5_05_V1
images.dir.22=Solid_20080310_1/Solid_20080310_1_F3_P4_05_V1
images.dir.23=Solid_20080310_1/Solid_20080310_1_F3_P3_05_V1
images.dir.24=Solid_20080310_1/Solid_20080310_1_F3_P2_05_V1
images.dir.25=Solid_20080310_1/Solid_20080310_1_F3_P1_05_V1
images.dir.26=Solid_20080310_1/Solid_20080310_1_F3_P5_06_V1
images.dir.27=Solid_20080310_1/Solid_20080310_1_F3_P4_06_V1
images.dir.28=Solid_20080310_1/Solid_20080310_1_F3_P3_06_V1
images.dir.29=Solid_20080310_1/Solid_20080310_1_F3_P2_06_V1
images.dir.30=Solid_20080310_1/Solid_20080310_1_F3_P1_06_V1
images.dir.31=Solid_20080310_1/Solid_20080310_1_F3_P5_07_V1
images.dir.32=Solid_20080310_1/Solid_20080310_1_F3_P4_07_V1
images.dir.33=Solid_20080310_1/Solid_20080310_1_F3_P3_07_V1
images.dir.34=Solid_20080310_1/Solid_20080310_1_F3_P2_07_V1
images.dir.35=Solid_20080310_1/Solid_20080310_1_F3_P1_07_V1

#################################################################
###- Daemon Properties
###- These are optional
###-
analysis.ini=/data/results/Solid/Solid_20080310_1/DH10B/jobs/postPrimerSetPrimary.m1/PrimaryParameters.ini
analysis.name=Filter_Fasta
Appendix D Advanced Topic: Primary Analysis

Primary analysis workflow

```
analysis.pipeline=Class:com.apldbio.aga.analysis.primary.core.PrimerSetPrimaryStage
analysis.stage.name=postPrimerSetPrimary

#############################
##- Run Directories
##- The analysis output directories. Make sure that these are correct.
##-
analysis.run.dir=/data/results/Solid/Solid_20080310_1/DH10B
analysis.results.dir=/data/results/Solid/Solid_20080310_1/DH10B/results.01
summary.dir=/data/results/Solid/Solid_20080310_1/DH10B/results.01/basecall_summary
colorcallfolder=/data/results/Solid/Solid_20080310_1/DH10B/results.01/colorcalls
cycleplots.dir=/data/results/Solid/Solid_20080310_1/DH10B/results.01/cycleplots
analysis.sample.dir=/data/results/Solid/Solid_20080310_1/DH10B
analysis.work.dir=/data/results/Solid/Solid_20080310_1/DH10B/jobs/postPrimerSetPrimary.m1
reads.result.dir.1=/data/results/Solid/Solid_20080310_1/DH10B/results.01/primary.m1/reads
read.dir=/data/results/Solid/Solid_20080310_1/DH10B/results.01/primary.m1/reads
```
Redo the read-filtering process

Redo the read-filtering process by trimming the last several bases:

1. Go to the following location inside the JobID folder:
   
   `$sampleJobFolder/postPrimerSetPrimary`

2. Enter the following modification in the parameters.ini file:
   
   `read.length={your desired read.length}`

3. Enter the following command:
   
   `jprimaryanalysis.sh --ini=parameters.ini`

Check the completion of primary analysis

The summary of the primary analysis results is written in a “.stats” file. This file is called:

 `{run.name}_{sample.name}_F3.stats`

for F3 tag analysis

or

 `{run.name}_{sample.name}_R3.stats`

for R3 tag analysis.

The file is located in:

 `/data/results/{instrument.name}/{run.name}/{sample.name}/results/primary.[17-digit timestamp]/reads/`

Example of a “.stats” file:

```
/share/apps/corona/bin/filter_fasta.pl --noduplicates --output=/data/results/Solid/Solid_20080310_1/DH10B/results.01/primary.20080328155539890 --name=Solid_20080310_1_DH10B_ --tag=F3 --minlength=35 --mask=11111111111111111111111111111111111 --prefix=T /data/results/Solid/Solid_20080310_1/DH10B/jobs/postPrimarySetPrimary.847/rawseq
# Cwd: /home/pipeline
# Title: Solid_20080310_1_DH10B_

Totals (2357 p): 81935898 / 90577909 (90.5%):
(Duplicates: 8085 [0.0%] (TooManyErrors: 8633926 [9.5%])
```
Appendix D Advanced Topic: Primary Analysis
Primary analysis workflow

Average: 34762.8 / 38429.3 (90.6%): (Duplicates: 3.4 [0.0%]) (TooManyErrors: 3663.1 [9.4%]) Usable (2251 p) (95.5%): 81935898 / 86395269 (94.8%): (Duplicates: 8085.0 [0.0%]) (TooManyErrors: 4451286.0 [5.2%]) Usable Average: 36399.8 / 38380.8 (94.9%): (Duplicates: 3.6 [0.0%]) (TooManyErrors: 1977.5 [5.1%])

The last several lines show the summary of the primary analysis. If this file is empty or the last four lines are missing, the primary analysis has not finished, or it encountered a problem and terminated.
Image metrics

Image metrics are measures of the quality of the images produced by the SOLiD™ 4 Analyzer. The principal image metrics are the blur metric and the exposure metric.

**Blur metric** is a real number greater than zero that provides a measure of the average apparent radius of the beads in an image. The blur metric is computed for every image that is analyzed by primary analysis. The blur metric supports diagnosing instrument focusing and vibration issues. Figure 10 gives two examples of the blur metric:

![Blur metric images](image)

**Figure 10 Two examples of the blur metric**

The *blur population* reports the population of panels that have a blur value above a given threshold. A larger average size can be caused by a small number of panels with large outlier values or a large number of panels with a large size. The two values taken together can indicate whether there is a localized problem over the whole sample (larger blur value but no outliers), a regional problem (an outlier population that spans channels), or a transient problem (outliers in a single channel).
Exposure metric is a real number that measures whether and by how much an image is over- or under-exposed, relative to a particular definition of ideal exposure. A value of zero indicates ideal exposure. A value greater than zero indicates over-exposure, and one less than zero indicates under-exposure. The scale is logarithmic with base two; a value of 1 indicates that the image is exposed twice as long as ideal.

The typical ideal exposure level of an image is one that maximizes the dynamic range of the content, with some signal occupying the lowest (darkest) range and some information extending to the saturation (highest) point.

- An overexposed image compresses too much of the content at the top end of the range. Overexposure creates a non-linearity in response and makes it impossible to differentiate between the highest signals.
- An underexposed image does not extend the content of the image across the possible range. Underexposure creates a low-end compression and fails to use the available range of the system.

**Note:** For purposes of color-calling algorithms, slightly overexposing the images yields better results.
Figure 11 gives two examples of the exposure metric:

![Figure 11 Two examples of the exposure metric](image)

The exposure value calculated is a measure of how close the system is to capturing an image that maximizes the use of the dynamic range. Looking at the tail behavior of the pixel distribution makes the determination.

- If the tail does not extend to the saturation point, the image is underexposed.
- If the tail has a peak at the far right side that is a measurement of the saturation, the image is overexposed.

The *exposure population* reports the population of panels that have an exposure quality beyond a given threshold. A large average size can be caused by a small number of panels with large outlier values or a large number of panels with a large size. The two values taken
together can estimate whether there is a localized problem over the whole sample (larger deviation with no outliers), a regional problem (outlier population that spans channels), or a transient problem (outliers in a single channel).

A difference with the blur metric, however, is that whereas the blur metric is constant over the number of beads present in a panel, the exposure is more sensitive to the number of beads. A baseline exposure value per channel is determined based on a sampling strategy, but panels with either significantly more or fewer beads can be affected differently. As with the blur value, the reported value is the largest mean exposure value of the four channels over all panels in the sample.

**Understanding quality values in the SOLiD™ System**

The SOLiD™ primary analysis software analyzes the raw image data collected by the system and generates a color-space read for each tag. The software generates a quality value (QV) for each color call, which is an estimate of confidence for that color call.

The SOLiD™ 4 System QVs are similar to those generated by Phred and the KB™ Basecaller software for capillary electrophoresis sequencing. The quality value \( q \) for a particular call is mathematically related to its probability of error \( p \).

\[
q = -10 \log_{10} p
\]

The algorithm used to predict QVs for the SOLiD™ 4 System is similar to Phred and the KB™ Basecaller software. (Refer to the paper, Ewing B., Green P., 1998, Base-calling of automated sequencer traces using Phred. II. Error probabilities. Genome Research 8:186-194.)

The algorithm relies on training or calibration to a large set of control data and color calls for which the correct call is known. In the SOLiD™ 4 System, the correct call is determined by mapping the read to a known reference sequence.
Usage

Given a read with QVs, the expected number of errors in the read can be estimated. If the QV of the i-th call is $q_i$, then the expected number of errors in the read is:

$$m = \sum_{i=1}^{n} p_i$$

where

$$p_i = 1.00 \cdot 10^{-q_i/10}$$

$p_i = 1.00 \cdot 10^{-q_i/10}$ is the predicted probability of error for the i-th call. This probability includes the fact that QVs are conventionally rounded to the nearest integer. The average error rate for this read is $m/n$, which can be converted into an overall quality score for the whole read.

$$Q = -10 \log_{10} \frac{m}{n}$$

Validation and observed quality

The conventional method of validating QVs involves computing a so-called observed quality for all the calls with a particular predicted quality in a set of control data. The following figure shows such a validation.
Figure 12  Statistical characteristics of quality values (QV) for the SOLiD™ 4 System

The top panel shows distribution of QVs. The bottom panel shows deviation of observed from predicted QVs.
Key files generated by primary analysis

Table 6  Key files generated by primary analysis

<table>
<thead>
<tr>
<th>File name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>xxxx_sequence.csfasta</td>
<td>The raw data from all beads</td>
</tr>
<tr>
<td>xxxx_sequence.QV.qual</td>
<td>The quality values file</td>
</tr>
<tr>
<td>.spch files</td>
<td>Cache files for primary analysis</td>
</tr>
</tbody>
</table>

xxx_sequence.csfasta

The “.csfasta” file is a color-space fasta file that contains the color calls generated for each tag, with the last base of the primer prepended.

The format is:

>TAG_ID
Color_space

For example:

>1_88_1830_R3
G32113123201300232320
>1_89_1562_R3
G23133131233333101320

This file contains all the data that passed filtering, for complete reads (for example, 35 base read has 35 bases). This file is the input to the alignment tool.
Appendix D Advanced Topic: Primary Analysis

Key files generated by primary analysis

**xxx_sequence.QV.qual**

The QV.qual files are fasta-like files that list the quality values in sequence order (not cycle order).

The format is:

```
TAG_ID
Quality Values:
```

For example:

```
>97_2040_1850_P3
38 36 26 33 41 26 24 33 28 31 27 23 5 35 32 31 11 10 24
38 22 24 7 12 15 21 12 18 34 31 27 11 15 26 13 14 17 17
13 12 8 5 17 5 12
>97_2040_1898_P3
41 41 41 38 32 29 39 24 23 36 32 38 25 30 28 21 27 33 34
33 24 27 9 35 34 14 30 18 33 8 13 32 10 31 24 7 22 5 27
30 21 5 0 27 9
```

All primary data (with exception of images) is stored in the file `xxxx.spch`. This file is in the HDF5 format. Tools to view and extract this data are available at:

[http://www.hdfgroup.org/HDF5/](http://www.hdfgroup.org/HDF5/)

The data within the "spch" file is organized by cycle. You can extract data in any format for your own analysis pipelines.
.spch files

Results of primary analysis for each panel are stored in files with the ".spch" extension (Solid Panel Cache HDF5). The ".spch" file is used as a cache; that is, primary analysis both writes results to the file and reads results from the file as analysis proceeds on a cycle-by-cycle basis. Data within the ".spch" file is organized by cycle. When all cycles have been analyzed, a final processing step converts the data from the cycle-based format to the “read-based” ASCII formats (“.csfasta” and “.qual”).

".spch" files use the hierarchical binary format HDF5. Details on this format are available at:

http://www.hdfgroup.org/HDF5/.

Tools to view the contents and extract results are available at:

http://www.hdfgroup.org/hdf-java-html/hdfview/
Procedures to verify primary analysis status

To verify primary analysis status:

1. From SETS, click the **History** menu to access Run History view, then click the run name of interest.
   The numbered items in the data tree on the left side of the page relate to the individual samples on the slide.

2. Click one of the data tree sample numbers on the left of the page to see the Library file name, Spot number, Sample name, and Reports information for that particular sample.

3. Click the drop-down arrow to the left of the sample folder to view individual analysis cycles for each sample.

4. Check that all the analysis completed successfully. A **blue dot** before the analysis name means that the analysis completed successfully.
   Click the name of the analysis library to see additional information about the analysis.
Appendix D Advanced Topic: Primary Analysis

Procedures to verify primary analysis status

Troubleshooting analysis failure

If there is a red dot before the analysis job name, the analysis job failed. To see details:

1. Click the name of the failed analysis job. The following example shows 2 failed analysis jobs: F3_PC_01_V1 and F3_PB_10_V1. The right portion of the figure shows the details of analysis F3_PC_01_V1.

2. Check Filter_FastaF3 and Filter_FastaR3 (if paired-end library) to see if the analysis job was successful. Check the details for the analysis job and each cycle to ensure that they are correct. See this example:
Appendix D  Advanced Topic: Primary Analysis

Procedures to verify primary analysis status

The above example shows detailed information for one Filter_FastaF3 run. This run uses 5 primers (PA-PE) and for each primer, 5 cycles. This analysis used all 5 primers (PA-P5); for each primer, 5 cycles were included in the analysis job, indicating no missing images in the analysis.

Sometimes, an analysis job might be reported as successful (no red x or warning dot next to the analysis name), but some of the primer/cycle combinations were not included in the analysis. Even if the analysis job is successful, if it is incomplete, it should be treated as a failed analysis job.
3. Log in to the instrument and look in the following directory:

```
{SampleResultsFolder}/primary.{17-digit timestamp}/reads
```

There should be a file named `{run.name}_F3.stats` for F3 tag, or `{run.name}_R3.stats` for R3 tag. This file contains information about the color-space reads generation for each panel. It contains the number of reads and an indication of whether they have too many errors, are too short, or are duplicates.

The Totals (### p) line summarizes the number of panels found and the percentage of error, short, and duplicate reads.

After this line is a line beginning with Usable (### p) (###%). This line summarizes how many panels were usable and the percentage of erroneous, short, or duplicate reads among the usable panels.

If these two lines are missing, the primary analysis did not complete.
Appendix D Advanced Topic: Primary Analysis
Procedures to verify primary analysis status

Procedure for re-analyzing the image (primary analysis)

If there was failure in the automatically-generated primary analysis pipeline, there are two ways to do primary re-analysis of the image. Before doing so, look at the primary log files to find out reason of failure. The log files are in the folder:

/data/result/{instrument.name}/{run.name}/{sample.name}/
jobs/{stage.name}.[JobId]/analysis/

Also, ensure that all images are present at /data/images/{run.name}.

After finding out the reason, change primary analysis parameters.

To re-analyze through SETS:

1. Log in to SETS using an administrative account.
2. Modify the primary analysis settings. Either modify added parameters or add your new values in Additional parameter field.
3. Open the Run Details page for that run and do primary re-analysis.

To perform primary analysis manually:

1. Access the job folder at:

   /data/result/{instrument.name}/{run.name}/{sample.name}/jobs/{stage.name}.[JobId]/

2. Modify the parameters.ini file with the desired value.
3. Enter the following command:

   jprimaryanalysis.sh --dir=<parameter file directory>
   --ini=<parameter file name>

   For example:

   jprimaryanalysis.sh
   --
   dir=/data/results/DAEMONAC2/LAST_MP_1245_041008/Sample1/jobs/  postPrimerSetPrimary.3793 --ini=parameters.ini
APPLIED BIOSYSTEMS END USER SOFTWARE LICENSE AGREEMENT

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Applied Biosystems SOLiD™ 4 System SETS Software

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## Glossary

**Alignment Matrix**
The alignment matrix is displayed as:
- AA is encoded as 0
- CG is encoded as 3
- AACG is encoded as 0 1 3

**Color Codes**
Four dye colors, encoded as follows:
- 6-FAM™ = 0
- CY3® = 1
- Texas Red® = 2
- CY5® = 3

**Cycle Order**
The order in which the data are generated.

**Location**
For processed files, describes the location of the prepended base on the base-space reference sequence.
Mate-Pair

Fragments from the genome paired during sequencing and separated by the same number of bases. For a known reference sequence of size 5 Mbp and a read length of 25 bp per tag, the system allows a total of six mismatches when placing the F3 and R3 tags. These mismatches can be distributed in any allocation across the F3 and R3 tags as long as the total number of mismatches does not exceed six. Any beads with F3 and R3 tags, which map under this mismatch allowance and which satisfy the mate-pair constraints, are considered to be a successfully placed mate-paired read. The mate-pair constraints are as follows:

The distance between the F3 and R3 tags must be within a prescribed range of distances determined by the insert size and library preparation.

The tags must map such that R3 comes before F3 reading 5’ > 3’.

The tags must map such that F3 and R3 map to the same strand on the reference sequence.

Multiplex sequencing

Sequencing of multiple barcoded fragment libraries on a single spot on a slide.

Panel

The region of the array that the camera can image at one time. Successive cycles result in four images for each panel (one per color). Beads are identified by position within a panel, that is, x,y coordinates are not unique values across panels. The combination of panel and x,y coordinates uniquely identifies a bead.
Paired-End
Paired-end libraries are standard fragment libraries. Sequence reads are generated from both ends of the fragment (5’>3’ and 3’>5’). These reads are paired during sequencing and separated by the same number of bases (typically 50-200 bases).

The paired-end constraints are as follows:
- The distance between the F3 and F5 tags must be within a prescribed range of distances determined by the insert size and library preparation.
- The tags must map such that F3 comes before F5 reading 5’ > 3’.
- The tags must map such that F3 and F5 map to opposite strands on the reference sequence.

Quality Value
Accuracy estimation of color calling using metrics similar to Phred or KB™ Basecaller.

Sequencing Run
During a SOLiD™ 4 System sequencing run, two probe sets are used to maximize the fraction of mappable beads, read length, and sequencing throughput. (Mappable beads are beads, amplified with template, that map to the reference genome.) This protocol must be used for sequencing 50-bp reads of both mate-paired and fragment libraries. Unlike terminator-based sequencing, the SOLiD™ 4 System does not collect base-sequencing information. Instead, five rounds of primers (Primers A, B, C, D, and E) are used to sequence template by ligation of di-base labeled probes. For sequencing of fragment libraries, the set of primers used is specific to the P1 adaptor.

SOLiD™ 4 Analyzer
The sequencing instrument and its supporting computer cluster.

SOLiD™ 4 System
The SOLiD™ 4 Analyzer, its instrument control software (ICS), and ancillary equipment.

Spot
A series of panels, each panel with unique numbers for ordering across spots. When multiple samples are run on a single array, each array has its own spot. This is shown in the file name format S#. For example, S1 are files for Spot 1.
Stringency  
Stringency is one criterion used to filter out junk reads in ReadBuilder stage. For filtered reads, csfasta and QV.qual are stored in the reject folder instead. The valid range of stringency is from 0 to 80. Any stringency less than 0 is treated as 0, where all reads will pass the stringency threshold. If stringency is set larger than 80, maximal possible stringency of 80 is used to avoid over filtering good beads.

TAG-ID  
A unique identifier for every tag, which consists of four components: `panel_xpixel_ypixel_tagtype`. Each tag has a TAG-ID. For example, `1_567_321_F3` describes a bead in panel 1 at coordinates 56, 321 (X,Y) with the R3 tag (second tag in a mate-pair).

**Note:** A Tag ID describes the bead and its data within files.

Values  
A dot (.) is used in color space to show that there is no call for this position. Although low values may have been measured, during clustering it was determined that no call could be made.

A minus one (−1) in a set of calculated values (for example, N2S) indicates that there is no data for this point; that is, a dot (.) is in the color space for this position.

A zero (0) in a set of calculated values (for example, N2S) means zero (0). The calculated result after rounding is zero (0).
WFA

A Workflow Analysis (WFA) run analyzes a quadrant of a slide that undergoes a single ligation cycle. The quadrant contains beads deposited at a lower density than the density of beads deposited for a sequencing run. You can optimize sequencing results by performing workflow analysis (WFA) runs.

A WFA run determines the:

- **Optimal library concentration**: The library concentration for optimal preparation of templated beads using the library. You use this library concentration for any preparation of templated beads for that library when the scale of templated bead preparation is the same.

- **Bead enrichment efficiency**: The proportion of beads that have been successfully amplified using emulsion PCR (ePCR) as a fraction of the total number of beads prepared. You use this value to more accurately deposit successfully amplified beads for a sequencing run.

WFA runs require the same materials as those materials needed for sequencing runs. If you perform multiple WFA runs routinely, order additional SOLiD™ Instrument Buffer Kits.

Zero and One Indexing

For preprocessed data, the first color space call is position 1, which refers to the transition between the last base of the adapter and the first base of the read. For processed data, the positions are 0-based so that the first position (the prepended base) is 0 on both strands. The $n^{th}$ position of the tag is numbered $(n - 1)$ in the forward direction and $-(n - 1)$ in the reverse direction.
### Related documentation

The following related documents are shipped with the system:

<table>
<thead>
<tr>
<th>Document</th>
<th>Part number</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>BioScope™ Software for Scientists Guide: Data Analysis Methods and Interpretation</td>
<td>4448431</td>
<td>Provides a bioinformatics analysis framework for flexible application analysis (data-generated mapping, SNPs, and count reads) from sequencing runs.</td>
</tr>
<tr>
<td>Applied Biosystems SOLiD™ 4 System ICS Software Help</td>
<td>Not applicable</td>
<td>Describes the software and provides procedures for common tasks (see the Instrument Control Software).</td>
</tr>
<tr>
<td>Applied Biosystems SOLiD™ 4 System Instrument Operation Guide</td>
<td>4448379</td>
<td>Describes how to load and run the SOLiD™ 4 Analyzer for sequencing.</td>
</tr>
</tbody>
</table>

A portable document format (PDF) version of this guide is also available on the Applied Biosystems SOLiD™ 4 System SETS v4.0.1 Software Help menu.

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