BIO® SMRT® Analysis Barcoding Overview (v8.0)

Introduction

This document applies to Sequel[®] and Sequel II Systems using SMRT[®] Link v8.0.

Note: For information on SMRT Link v6.0, see the document **SMRT Analysis Barcoding Overview v6.0**, available here.

SMRT Link v8.0 includes a streamlined end-to-end barcoding workflow for automated analysis of multiplexed samples. It enables users to define barcodes and their associated sample names for an experiment at Run Design.

This workflow provides extensive QC metrics for evaluation of barcoding performance.

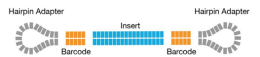
The SMRT Link graphical user interface supports up to 10,000 barcodes per sample. Support for more than 10,000 samples is available using the command-line. See **SMRT Tools Reference Guide (v8.0)**, available here.

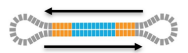
This overview covers:

- Barcoding designs, strategies and modes for preparing barcoded samples and data analysis for all the applications available with SMRT Link v8.0.
- Identifying barcodes using SMRT Link.
- Finding demultiplexed subread.bam files.

This document does **not** include information about barcoding designs and analysis for the Iso-Seq[®] and No-Amp application. For details about barcoding samples for Iso-Seq analysis, see **SMRT Link User Guide** (v8.0), available here.

PacBio Barcoding Background





Multiple passes around a SMRTbell® molecule allows barcode detection.

Barcoding Strategies

There are several options for barcoding samples for SMRT sequencing:

1. Barcoded Primers

 Target-specific primers are tailed with 16-bp barcodes. Use this method if your primers are already fully validated.

2. Barcoded Universal Primers

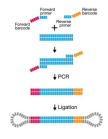
- This method requires two rounds of PCR. The first round uses target-specific primers tailed with universal forward and reverse sequences.
- For the second PCR round, round one PCR products are barcoded using F/R universal primers tailed with 16-bp barcodes.
- For symmetric barcoding, PacBio recommends using our commercially-available 96 Barcoded Universal Primers kit. For asymmetric barcoding, we recommend using the barcoded M13 Primer approach.

3. Barcoded Overhang Adapters

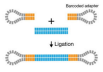
- PacBio barcodes are added to the sample through ligation of barcoded overhang SMRTbell[®] adapters.
- Recommended for validated PCR systems and smaller projects.
- Besides amplicons, barcoded overhang adapters are also used for microbial multiplexing.

4. Linear Barcoded Adapters for Probe-Based Captures

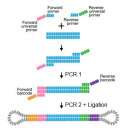
- Recommended for probe-based captures of genomic regions of interest.
- 1. Barcoded Target-Specific Primers



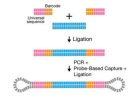
3. Barcoded Overhang Adapters



2. Barcoded Universal Primer



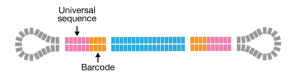
4. Barcoded Linear Adapter (Target Capture)



Description of Linear Barcoded Adapters for Multiplexing Target Capture Experiments

For targeted captures only, use Linear Barcoded Adapters:

Linear Barcoded Adapters with Universal Sequences



Linear barcoded adapters containing universal sequences are ligated to double-stranded DNA fragments for amplification and capture. This workflow supports up to 24 barcodes, which can be ordered from any oligo synthesis provider.

Demultiplexing of such experiments is supported through SMRT Link command-line options.

Barcode Scoring Modes

The demultiplexing algorithm included in SMRT Link v8.0 only requires information about the barcoding sequences – whether they are the **same** or **different** on either side of the read. Barcode orientation does **not** need to be specified. The demultiplexing algorithm scores **all** barcode sequences in both orientations - forward and reverse complement.

This section describes barcode scoring modes and provides information about the different experiment options depending on barcoding strategy.

Barcode Library Designs Input Sequence Leading Barcode **Output Sequence Trailing Barcode** Terminology Clip Trailing Clip Leading **Demultiplex Barcodes** library tailed same AAGG AAGG same AAGG symmetric CCTT asymmetric different AAGG TGGC TGGC AAGG Barcode 1: Barcode 2: GCCA CCTT Barcode 1 RC: < Barcode 2 RC:

The **Demultiplex Barcodes** application in SMRT Link supports demultiplexing of subreads and CCS data. The following terminology is based on the per (sub-) read view.

Symmetric Mode

For **symmetric** and **tailed** library designs, the **same** barcode is attached to both sides of the insert sequence of interest. The only difference is the orientation of the trailing barcode. For barcode identification, one read with a single barcode region is sufficient. Symmetric barcoding is used for samples constructed using Barcoded Overhang Adapters, Barcoded Universal Primer and target enrichment (linear). This is also the default scoring mode in SMRT Link v8.0 and later.

Asymmetric Mode

Barcode sequences are **different** on the ends of the SMRTbell template. Asymmetric mode is used with the M13 barcoding procedure. (See **Procedure & Checklist - Preparing SMRTbell Libraries using PacBio Barcoded M13 Primers for Multiplex SMRT Sequencing** for details.) Pacific Biosciences recommends using this mode **only** for small inserts (up to 5 kb) where both ends of the insert are expected to be sequenced. **Both** barcodes must be detected.

Note: For **both** Symmetric and Asymmetric modes, the limit for unique individual barcode sequences is 768, and the limit for the number of different barcode pairs is 384.

When running the **Demultiplex Barcodes** applications in SMRT Link, set the **Same Barcodes on Both Ends of the Sequence** option to **Off**.

Mixed Mode

Libraries with combined symmetric and asymmetric barcoding are **not** supported.

Which Barcoding Strategies to Use For application recommendations, see Overview Sequel Systems Application Options and Sequencing Recommendations.

Note: For multiplexing microbial genomes experiments, click here to see details about the workflows, analysis and barcoding kits available.

The following table includes the names of FASTA files used for analysis:

FASTA File	Sample Preparation		
	Barcoded Adapters	Barcoded Universal Primers	Custom Barcoded Primers
Sequel_16_barcodes_v1	✓		✓
Sequel_96_barcodes_v1		✓	✓
Sequel_384_barcodes_v1		√ ¹	✓

¹ Using oligos from PacBio recommended lists.

Custom Symmetric Barcode FASTA

Independent of the barcoding strategy and mode, if you are using custom barcodes, you **must** generate a FASTA file of barcode sequences. This file should:

- Include one entry per barcode sequence.
- · Include no duplicate sequences.
- Be orientation-agnostic (forward or reverse-complement, but not reversed.)

>bc1000 CTCTACTTACTTACTG >bc1001 GTCGTATCATCATGTA >bc1002 AATATACCTATCATTA

Name your barcodes with an alphabetic character prefix to avoid later confusion of barcode name and index. Duplicate names are **not** allowed.

Working with Barcoded Data in SMRT Link v8.0

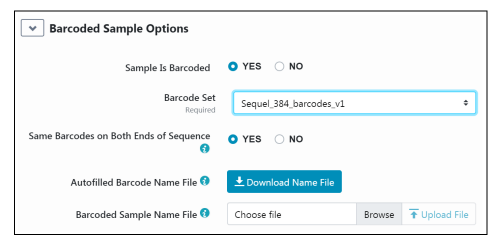
This section describes how to use SMRT Link to work with barcoded data. Demultiplex Barcodes analysis is powered by the Lima SMRT[®] Analysis tool.

The canned data provided with SMRT Link v8.0 includes 7 barcode sets:

- Sequel_16_barcodes_v1
- Sequel 16 barcodes v3
- Sequel 96 barcodes v1
- Sequel 384 barcodes v1
- RSII 96 barcodes
- RSII 384 barcodes
- IsoSeqPrimers v2 (Includes the content of IsoSeqPrimers as well as support for NEB and Clontech primers.)

Step 1: Specify the Barcode Setup and Sample Names in a Run Design

 In SMRT Link, create a new run design as described in "Creating a New Run Design" on page 10 of the SMRT Link User Guide (v8.0). Before you finish the new Run Design, perform the following steps.



- 2. Click **Barcoded Sample Options** and then click **Yes** for **Sample is Barcoded**. Additional fields related to barcoding display.
- 3. Specify a **Barcode Set** using the dropdown list. **Note**: You can specify up to 384 samples.
- 4. Specify if the **same** barcodes are used on both ends of the sequences.
 - Selecting Yes specifies symmetric and tailed designs where all the reads have the same barcodes on both ends of the insert sequence.
 Barcode analysis of such experiments retains only data with the same barcode identified on both ends.
 - Selecting No specifies asymmetric designs where the barcodes are different on each end of the insert. Barcode analysis of such experiments retains any barcode pair combination identified in the Data Set.
- 5. SMRT Link automatically creates a CSV-format **Autofilled Barcode**Name File. The barcode name is populated based on your choice of barcode set, and if the barcodes are the same at both ends of the sequence. The file includes a column of automatically-generated Bio Sample Names 1 through N, corresponding to barcodes 1 through N, for the biological sample names.
 - To specify the biological sample names corresponding to each barcode, click **Download Name File**, enter the biological sample names associated with the barcodes in the second column, and save the file. (Use alphanumeric characters, spaces, hyphens, underscores, colons, or periods only, with a maximum of 40 characters.) If you did **not** use all barcodes in the Autofilled Barcode Name file in the sequencing run, **delete** unused rows.
 - Note: Open the CSV file in a text editor and check that the columns are separated by commas, not semicolons or tabs.
- 6. Select the Barcoded Sample Name File you edited in Step 5.
- 7. Click Save.

Step 2: Perform the Sequencing Run

Load the samples and perform the sequencing run, using the Run Design you created in Step 1. The demultiplexing analysis is performed automatically on the SMRT Link Server once the data is transferred from the Sequel/Sequel II System. This creates an analysis of type

Demultiplex Barcodes (Auto) in the SMRT Analysis module. You can click to select this analysis and review the reports and data created. If everything looks fine, you can continue to **Step 4** and use the demultiplexed Data Set(s) created by the run as input to further analysis.

Note: By default, <code>Demultiplex Barcodes</code> (Auto) creates one Data Set per autodetected barcode within the selected barcode set. It also applies a Data Set filter of a minimum barcode score greater than 26 for optimal results in secondary analyses. If used, the analysis parameter Filters to add to the Data Set overrides other barcode filtering, even if the barcode score set with it is lower than 26.

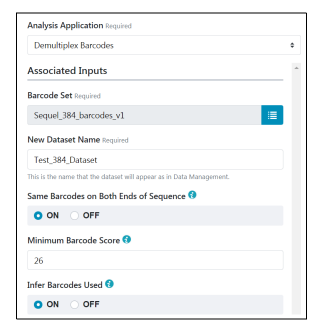
Step 3: (Optional) Run the Demultiplex Barcodes Application

If you did **not** specify the barcode setup in the Run Design, or if you need to change any of the parameters used in the **Demultiplex Barcodes** analysis automatically launched from Run Design, run the **Demultiplex Barcodes** application. This application separates reads by barcode and creates a new demultiplexed Data Set that you can then use as input to other secondary analysis applications.

- 1. Click + Create New Analysis.
- 2. Enter a **name** for the analysis.
- 3. Select the type of data to use for the analysis:
 - **Sequel Data**: Subreads from Sequel or Sequel II Systems.
 - CCS Data: Single Molecule consensus reads generated from the CCS analysis.

The Data Sets box displays the appropriate Data Sets available for the analysis.

- 4. In the Data Sets box, select one or more Data Sets to be analyzed together.
- 5. Click Next.
- 6. Select **Demultiplex Barcodes** from the Applications list.



- 7. Specify a Barcode Set (barcode sequence file.)

 Note: You can specify up to 384 samples.
- 8. Specify the name for the new demultiplexed Data Set that will display in SMRT Link.
- 9. Specify if the **same** barcodes are used on both ends of the sequences.
 - Selecting On specifies symmetric and tailed designs where all the reads have the same barcodes on both ends of the insert sequence.
 Barcode analysis of such experiments retains only data with the same barcode identified on both ends.
 - Selecting Off specifies asymmetric designs where the barcodes are different on each end of the insert. Barcode analysis of such data retains any barcode pair combination identified in the Data Set.
- 10. Specify the **Minimum Barcode Score**: Reads with barcode scores below the value are **not** included in downstream analysis. We recommend that you set this value to 26 for **all** applications.
- 11. Specify if you want to infer which barcodes were used:
 - On infers which subset of barcodes from the selected barcode set were used, and outputs one data set for each of those inferred barcodes.
 - Off outputs one data set with all barcodes in the selected barcode set.
- 12. Click **Start**. After the analysis is finished, a new demultiplexed Data Set is available.

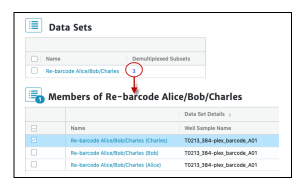
Note: For information about the reports generated by the Demultiplex Barcodes application, see "Demultiplex Barcodes Reports and Data Files" on page 11.

Step 4: Run Applications Using the Demultiplexed Data as Input

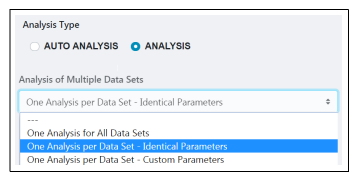
All secondary analysis applications except **Demultiplex Barcodes** can use demultiplexed Data Sets as input.

Note: For **Iso-Seq** analysis with barcoded samples, use the Iso-Seq application **instead** of the Demultiplex Barcodes application, as the Iso-Seq application already includes the demultiplexing step as part of the pipeline. When performing multiplexed Iso-Seq analysis, ensure that the Run Design **Sample Is Barcoded** option is set to **No** (the default setting). Then, in SMRT Analysis, go straight to the Iso-Seq application and, in the parameters section, select a Primer Set containing multiple primers, such **as** IsoSeq Primers_12_Barcodes_v1.

- 1. Select the secondary analysis application to use.
- 2. Click the number in the **Demultiplexed Subsets** column, then select the demultiplexed Data Set to use as input:



- You can select the **entire** Data Set as input, or one or more specific outputs from selected barcodes, to a maximum of 16 sub-Data Sets, 12 for Iso-Seq Analysis.
- 3. Additional **Analysis Type** options become available. You can select from the following options:



 One Analysis for All Data Sets: Runs one analysis using all the selected barcode Data Sets as input, for a maximum of 30 Data Sets.

- One Analysis per Data Set Identical Parameters: Runs one separate analysis for each of the selected barcode Data Sets, using the same parameters, for a maximum of 10,000 Data Sets.
 Optionally click Advanced Parameters and modify parameters.
- One Analysis per Data Set Custom Parameters: Runs one separate analysis for each of the selected barcode Data Sets, using different parameters for each Data Set, for a maximum of 16 Data Sets. Click Advanced Parameters and modify parameters. Then click Start and Create Next. You can then specify parameters for each of the included barcode Data Sets.
- Note: The number of Data Sets listed is based on testing using PacBio's suggested compute configuration, listed in SMRT Link Software Installation (v8.0).
- 4. Click **Start** to submit the analysis.

Demultiplex Barcodes Input and Output

Given an input set of barcodes and a BAM Data Set, the Demultiplex Barcodes application produces:

- A set of BAM files whose reads are annotated with the barcodes;
- A subreadset file that contains the file paths of that collection of barcode-tagged BAM files and their related files.

Barcode Set (Required):

Specify a barcode sequence file to separate the reads.

New Data Set Name (Required):

 Specify the name for the new demultiplexed Data Set that will display in SMRT Link.

Same Barcodes on Both Ends of Sequence (Default = ON)

- Specify On to retain all the reads with the same barcodes on both ends of the insert sequence, such as symmetric and tailed designs. (See "Barcode Scoring Modes" on page 3 for information on barcode designs.)
- Specify Off to specify asymmetric designs where the barcodes are different on each end of the insert.

Minimum Barcode Score (Default = 0)

 A barcode score measures the alignment between a barcode attached to a read and an ideal barcode sequence, and is an indicator of how well the chosen barcode pair matches. It ranges between 0 (no match) and 100 (a perfect match). Specifies that reads with barcode scores below this minimum value are not included in analysis.

Infer Barcodes Used (Default = ON)

The barcoding algorithm can detect the set of barcodes used. It infers
the barcodes used by looking at the first 35,000 ZMWs, then selecting
barcodes with ≥10 counts and mean scores ≥45. Specify ON to use
this mode.

Demultiplex Barcodes Reports and Data Files

Reports and Data Files

The Demultiplex Barcodes application generates the following reports:

Barcodes > Summary Metrics

- Unique Barcodes: The number of unique barcodes in the sequence data.
- Barcoded Reads: The number of barcoded reads in the sequence data.
- Mean Reads: The mean number of reads per barcode.
- Max. Reads: The maximum number of reads per barcode.
- Min. Reads: The minimum number of reads per barcode.
- Mean Read Length: The mean read length of reads per barcode.
- **Mean Longest Subread Length:** The mean length of the longest subread in each barcoded sample.
- Unbarcoded Reads: The number of reads without barcodes in the sequence data.

Barcodes > Barcode Data

- **Bio Sample Name:** The name of the biological sample associated with the barcode.
- Barcode Index: The index number associated with the barcode.
- Barcode Name: A string containing the pair of barcode indices for which the following metrics apply.
- Polymerase Reads: The number of polymerase reads associated with the barcode.
- **Subreads**: The number of subreads associated with the barcode.
- Bases: The number of bases associated with the barcode.
- Mean Read Length: The mean read length of reads associated with the barcode.
- Longest Subread Length: The longest subread length associated with the barcode.
- Mean Barcode Quality: The mean barcode quality associated with the barcode.
- Rank Order (Num. Reads): The rank order of this barcode in terms of the number of reads.

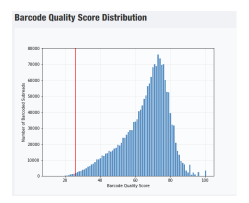
Barcodes > Barcoded Read Statistics

- Number of Reads per Barcode: Line graph displays the number of sorted reads per barcode.
 - Good performance: The Number of Reads per Barcode line (blue) should be mostly linear. Note that this depends on the choice of Y-axis scale. The mean Number of Reads per Barcode line (red) should be near the middle of the graph and should not be skewed by samples with too many or too few barcodes.
 - Questionable performance: A sharp discontinuity in the blue line, followed by no yield, with the red line way far from the center. Check the output file Inferred Barcodes, note the correct barcodes used, and consider reanalyzing the multiplexed samples with the correct Bio Sample names for the barcodes actually used. If you reanalyze the data, ensure that the Barcode Name File includes only the correct barcodes used.
- Barcode Frequency Distribution: Histogram distribution of read counts per barcode.
 - Good performance: A uniform distribution, which is most often a fairly tight symmetric normal distribution, with few barcodes in the tails.

- Questionable performance: A large peak at zero. This can indicate use of incorrect barcodes. Check the output file Inferred Barcodes, note the correct barcodes used, and consider reanalyzing the multiplexed samples with the correct Bio Sample names for the barcodes actually used. If you reanalyze the data, ensure that the Barcode Name File includes only the correct barcodes used.
- **Mean Read Length Distribution**: Histogram distribution of the mean polymerase read length for all samples.
 - Good performance: The distribution should be normal with a relatively tight range.
 - Questionable performance: A spread out distribution, with a mode towards the low end.

Barcodes > Barcode Quality Scores

- Barcode Quality Score Distribution: Histogram distribution of barcode Quality scores. The scores range from 0-100, with 100 being a perfect match. Any significant modes or accumulation of scores <40 suggests issues with some of the barcode analyses. The red line is set at 26 – the minimum default barcode score.
 - Good performance: Distributions with a mode >65 and the low-end tail tapering off below 40.



 Questionable performance: A bimodal distribution with a large second peak usually indicates that some barcodes that were sequenced were not included in the barcode scoring set.

Barcodes > Barcoded Read Binned Histograms

- Read Length Distribution By Barcode: Histogram distribution of the Polymerase read length by barcode. Each column of rectangles is similar to a read length histogram rotated vertically, seen from the top. Each sample should have similar Polymerase read length distribution. Non-smooth changes in the pattern looking from left to right might indicate suboptimal performance.
- Barcode Quality Distribution By Barcode: Histogram distribution of the perbarcode version of the Read Length Distribution by Barcode histogram. The histogram should contain a single cluster of hot spots in each column. All barcodes should also have similar profiles; significant differences in the pattern moving from left to right might indicate suboptimal performance.
 - Good performance: All columns show a single cluster of hot spots.
 - Questionable performance: A bimodal distribution would indicate missing barcodes in the scoring set.

Data > File Downloads

The following files are available on the Analysis Results page. Additional files are available on the SMRT Link server, in the analysis output directory.

- Analysis Log: Log information for the analysis workflow.
- SMRT Link Log: Server-level analysis log information. (This file is displayed when you choose Data > SMRT Link Log.)
- Barcode Files: Barcoded subread Data Sets; one file per barcode.
- Barcode Summary CSV: Data displayed in the reports, in CSV format.
- Barcodes Unassigned: Barcodes not assigned to a specific isoform.

Note: You can get the demultiplexed BAM files using the Data Management module's "Export Data Sets" feature. In the demultiplexed BAM output file a tag bc is added for each read, indicating the assigned barcode. The bc tag is the zero-based index of the barcodes in the FASTA file. For example, when using the barcodes RSII_96_barcodes, a subread with barcode lbc1 identified on both sides will have the tag bc:B:S,0,0 in the BAM output file.

A second bg tag corresponds to the barcode quality (0-100).

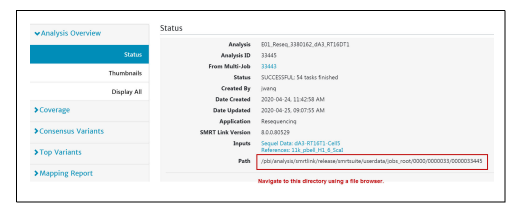
For details on running the Demultiplex Barcodes application using the command-line, see **SMRT Tools Reference Guide (v8.0)**, available here.

Working with Demultiplexing Results

Finding Demultiplexed subreads.bam Files

Subreads.bam files **cannot** be downloaded from the SMRT Analysis Report page. You can find these files using the command line by navigating to the SMRT Link server. To do so:

 Copy the job path (Analysis Overview > Status > Path) on the SMRT Link server shown on the Report page.



2. Open the command line, login into the SMRT Link Server and change directory to the path you copied:

\$ cd /pbi/analysis/smrtlink/release/smrtsuite/userdata/jobs_root/0000/000029/
0000029399

3. Display the folders and files in the directory of the Barcode Demultiplexing Run:

```
$ 1s -1
lrwxrwxrwx 1 fas Domain Users 140 Oct 10 09:36 cromwell-job -> /pbi/analysis/smrtlink/
release/smrtsuite/userdata/jobs_root.local/cromwell-executions/pb_demux_subreads/
c9cf0alc-5fef-46d9-97e4-3ad44d80b597
drwxr-xr-x 2 fas Domain Users 4096 Oct 10 09:36 entry-points
drwxr-xr-x 3 fas Domain Users 4096 Oct 10 10:04 logs
drwxr-xr-x 2 fas Domain Users 20480 Oct 10 10:04 outputs
-rw-r--r-- 1 fas Domain Users 0 Oct 10 09:36 pbscala-job.stderr
-rw-r--r-- 2 fas Domain Users 22778 Oct 10 10:05 pbscala-job.stdout
drwxr-xr-x 2 fas Domain Users 4096 Oct 10 10:04 workflow
```

Notice that the cromwell job is a hyperlink to another directory, designed to save space on your SMRT Link server.

4. Navigate to the subdirectory where the .bam files are stored:

\$cd cromwell-job/call-demultiplex_barcodes/demultiplex_barcodes/083b56b3-dee2-44aeaa97-e6656bb4763f/call-lima/execution

\$1s -1

-rw-r--r-- 1 fas Domain Users

-rw-r--r-- 1 fas Domain Users -rw-r--r-- 1 fas Domain Users

```
-rw-r--r-- 1 fas Domain Users 215561027 Oct 10 09:58 lima.bc1001--bc1001.bam
-rw-r--- 1 fas Domain Users 388923 Oct 10 09:58 lima.bc1001--bc1001.bam.pbi
-rw-r--r-- 1 fas Domain Users 12857 Oct 10 09:58 lima.bc1001--bc1001.subreadset.xml
-rw-r--r-- 1 fas Domain Users 227808959 Oct 10 09:58 lima.bc1002--bc1002.bam
-rw-r--r- 1 fas Domain Users 408837 Oct 10 09:58 lima.bc1002--bc1002.bam.pbi
-rw-r--r-- 1 fas Domain Users
                                  12857 Oct 10 09:58 lima.bc1002--bc1002.subreadset.xml
-rw-r--r-- 1 fas Domain Users 217156999 Oct 10 09:58 lima.bc1003--bc1003.bam
-rw-r--r-- 1 fas Domain Users 390196 Oct 10 09:58 lima.bc1003--bc1003.bam.pbi
                                  12857 Oct 10 09:58 lima.bc1003--bc1003.subreadset.xml
-rw-r--r-- 1 fas Domain Users
-rw-r--r-- 1 fas Domain Users 238209137 Oct 10 09:58 lima.bc1004--bc1004.bam
-rw-r--r-- 1 fas Domain Users 428762 Oct 10 09:58 lima.bc1004--bc1004.bam.pbi
-rw-r--r-- 1 fas Domain Users 43595 Oct 10 09:58 lima.json
-rw-r--r-- 1 fas Domain Users 2757 Oct 10 09:58 lima.lima.counts
-rw-r--r-- 1 fas Domain Users 2977 Oct 10 09:41 lima.lima.guess
-rw-r--r-- 1 fas Domain Users 433394971 Oct 10 09:58 lima.lima.report
-rw-r--r-- 1 fas Domain Users 791 Oct 10 09:58 lima.lima.summary
-rw-r--r-- 1 fas Domain Users 992123522 Oct 10 09:58 lima.removed.bam
-rw-r--r-- 1 fas Domain Users 2413108 Oct 10 09:58 lima.removed.bam.pbi
-rw-r--r- 1 fas Domain Users 569 Oct 10 09:58 lima.removed.json rw-r--r- 1 fas Domain Users 12542 Oct 10 09:58 lima.removed.subreadset.xml
                                  2 Oct 10 09:58 rc
4459 Oct 10 09:37 script
1162 Oct 10 09:37 script.submit
-rw-r--r-- 1 fas Domain Users
                                     0 Oct 10 09:37 stderr
```

For each demultiplexed barcode, the Barcode Demultiplexing workflow generates three files: subread.bam, subread.bam.pbi and subreadset.xml.

0 Oct 10 09:37 stderr.submit

63 Oct 10 09:37 stdout.submit

0 Oct 10 09:37 stdout

Other files are generated and display in the advanced report:

```
lima.lima.counts, lima.lima.guess, lima.lima.report,
lima.lima.summary,lima.removed.bam
```

The json files (lima.json and lima.remove.json) are useful for running a user-defined workflow.

Note: For information on working with barcoded data in pbcromwell, see page 60 of **SMRT Tools Reference Guide (v8.0)**.

R plotting scripts are available on github: https://github.com/pacificbiosciences/barcoding/#quality-control.

For information on working with barcoded data in lima, see: https://github.com/PacificBiosciences/barcoding.

Additional Information:

- Demultiplex Barcodes Algorithm: https://github.com/PacificBiosciences/barcoding
- PacBio BAM Format Specifications: http://pacbiofileformats.readthedocs.io/en/9.0/BAM.html

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