PacBi

SMRT® Tools reference guide (v12.0)



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SMRT® Tools reference guide (v12.0)

Introduction

This document describes the command-line tools included with SMRT[®] Link v12.0. These tools are for use by bioinformaticians working with secondary analysis results.

• The command-line tools are located in the \$SMRT_ROOT/smrtlink/smrtcmds/bin subdirectory.

Installation

The command-line tools are installed as an integral component of the SMRT Link software. For installation details, see **SMRT Link software installation guide (v12.0)**.

- Note: SMRT Link v12.0 is for use with Revio[™] systems, Sequel[®] II systems and Sequel IIe systems only.
- To install only the command-line tools, use the --smrttools-only option with the installation command, whether for a new installation or an upgrade. Examples:

```
smrtlink-*.run --rootdir smrtlink --smrttools-only
smrtlink-*.run --rootdir smrtlink --smrttools-only --upgrade
```

Supported chemistry

SMRT Link v12.0 supports all chemistry versions for Revio systems, Sequel II systems and Sequel IIe systems.

PacBio command-line tools

Following is information on the PacBio-supplied command-line tools included in the installation. For information on third-party tools installed, see "Appendix B - Third party command-line tools" on page 131.

Tool	Description
bam2fasta/ bam2fastq	Converts PacBio® BAM files into gzipped FASTA and FASTQ files. See "bam2fasta/bam2fastq" on page 4.
bamsieve	Generates a subset of a BAM or PacBio Data Set file based on either a list of hole numbers, or a percentage of reads to be randomly selected. See "bamsieve" on page 5.
ccs	Calculates consensus sequences from multiple "passes" around a circularized single DNA molecule (SMRTbell® template). See "ccs" on page 8.
dataset	Creates, opens, manipulates and writes Data Set XML files. See "dataset" on page 17.

Tool	Description
Demultiplex Barcodes	Identifies barcode sequences in PacBio single-molecule sequencing data. See "Demultiplex Barcodes" on page 24.
export-datasets	Takes one or more PacBio dataset XML files and packages all contents into a single ZIP archive. See "export-datasets" on page 36.
export-job	Takes one SMRT Link analysis job and packages all contents into a single ZIP archive. See "export-job" on page 37.
gcpp	Variant-calling tool which provides several variant-calling algorithms for PacBio sequencing data. See "gcpp" on page 39.
Genome Assembly	Generates <i>de novo</i> assemblies using HiFi reads. See "Genome Assembly" on page 42.
HiFiViral SARS-CoV-2 Analysis	Analyzes multiplexed viral surveillance samples for SARS-CoV-2. See "HiFiViral SARS-CoV-2 Analysis" on page 51.
ipdSummary	Detects DNA base-modifications from kinetic signatures. See "ipdSummary" on page 54.
isoseq3	Characterizes full-length transcripts and generates full-length transcript isoforms, eliminating the need for computational reconstruction. See "isoseq3" on page 58.
juliet	A general-purpose minor variant caller that identifies and phases minor single nucleotide substitution variants in complex populations. See "juliet" on page 67.
Microbial Genome Analysis	Generates <i>de novo</i> assemblies of small prokaryotic genomes between 1.9-10 Mb and companion plasmids between 2 – 220 kb and performs base modification detection, using HiFi reads. See "Microbial Genome Analysis" on page 75.
motifMaker	Identifies motifs associated with DNA modifications in prokaryotic genomes. See "motifMaker" on page 79.
pbcromwell	PacBio's wrapper for the cromwell scientific workflow engine used to power SMRT Link. For details on how to use pbcromwell to run workflows, see "pbcromwell" on page 81.
pbindex	Creates an index file that enables random access to PacBio-specific data in BAM files. See "pbindex" on page 86.
pbmarkdup	Marks or removes duplicates reads from CCS reads. See "pbmarkdup" on page 87.
pbmm2	Aligns PacBio reads to reference sequences; a SMRT wrapper for minimap2. See "pbmm2" on page 90.
pbservice	Performs a variety of useful tasks within SMRT Link. See "pbservice" on page 97.
pbsv	Structural variant caller for PacBio reads. See "pbsv" on page 102.
pbvalidate	Validates that files produced by PacBio software are compliant with PacBio's own internal specifications. See "pbvalidate" on page 106.
pigeon	Classifies and filters full-length transcript isoforms into categories against a reference annotation. See "pigeon" on page 109.
primrose	Performs motif-calling to detect 5mC CpG sites in HiFi BAM files. See "primrose" on page 115.
runqc-reports	Generates Run QC reports. See "runqc-reports" on page 116.
skera	Splits arrayed HiFi reads at adapter positions, generating read-segments. See "skera" on page 117.

Tool	Description
summarizeModifi cations	Generates a GFF summary file from the output of base modification analysis combined with the coverage summary GFF generated by resequencing pipelines. See "summarize Modifications" on page 121.

bam2fasta/ bam2fastq

The bam2fasta and bam2fastq tools convert PacBio BAM or Data Set files into gzipped FASTA and FASTQ files, including demultiplexing of barcoded data.

Usage

Both tools have an identical interface and take BAM and/or Data Set files as input.

```
bam2fasta [options] <input>
bam2fastq [options] <input>
```

Options	Description
-h,help	Displays help information and exits.
version	Displays program version number and exits.
-o,output	Specifies the prefix of the output file names. – implies streaming.
	Note : Streaming is not supported with compression or with the <code>split_barcodes</code> option.
-с	Specifies the Gzip compression level. Values are [1,2,3,4,5,6,7,8,9]. (Default = 1)
-u	Specifies that the output FASTA/FASTQ files are not compressedgz is not added to the output file names, and -c settings are ignored.
split-barcodes	Splits the output into multiple FASTA/FASTQ files, by barcode pairs. Note: The bam2fasta/bam2fastq tools inspect the bc tag in the BAM file to determine the 0-based barcode indices from their respective positions in the barcode FASTA file.
-p,seqid-prefix	Specifies the prefix for the sequence IDs used in the FAST/FASTQ file headers.

Examples

```
bam2fasta -o projectName m54008_160330_053509.subreads.bam
```

 $\label{lem:bam2fastq-omyEcoliRuns m54008_160330_053509.subreads.bam m54008_160331_235636.subreads.bam}$

bam2fasta -o myHumanGenomem54012 160401 000001.subreadset.xml

Input files

- One or more *.bam files
- *.subreadset.xml file (Data Set file)

Output files

- *.fasta.gz
- *.fastq.gz

bamsieve

The <code>bamsieve</code> tool creates a subset of a BAM or PacBio Data Set file based on either a list of hole numbers to include or exclude, or a percentage of reads to be randomly selected, while keeping all subreads within a read together. Although <code>bamsieve</code> is BAM-centric, it has some support for dataset XML and will propagate metadata, as well as scraps BAM files in the special case of SubreadSets. <code>bamsieve</code> is useful for generating minimal test Data Sets containing a handful of reads.

bamsieve operates in two modes: **list** mode where the ZMWs to keep or discard are explicitly specified, or **percentage/count** mode, where a fraction of the ZMWs is randomly selected.

ZMWs may be listed in one of several ways:

- As a comma-separated list on the command line.
- As a flat text file, one ZMW per line.
- · As another PacBio BAM or Data Set of any type.

Usage

Required	Description
input_bam	The name of the input BAM file or Data Set from which reads will be read.
output_bam	The name of the output BAM file or Data Set where filtered reads will be written to. (Default = $None$)

Options	Description
-h,help	Displays help information and exits.
version	Displays program version number and exits.
log-file LOG_FILE	Writes the log to file. (Default = None, writes to stdout.)
log-level	Specifies the log level; values are [DEBUG, INFO, WARNING, ERROR, CRITICAL]. (Default = WARNING)
debug	Alias for setting the log level to DEBUG. (Default = False)
quiet	Alias for setting the log level to CRITICAL to suppress output. (Default = False)
-v,verbose	Sets the verbosity level. (Default = NONE)
show-zmws	Prints a list of ZMWs and exits. (Default = False)
include INCLUDE LIST	Specifies the ZMWs to include in the output. This can be a comma-separated list of ZMWs, or a file containing a list of ZMWs (one hole number per line), or a BAM/ Data Set file. (Default = NONE)

Options	Description
exclude EXCLUDE LIST	Specifies the ZMWs to exclude from the output. This can be a comma-separated list of ZMWs, or a file containing a list of ZMWs (one hole number per line), or a BAM/Data Set file that specifies ZMWs. (Default = \mathtt{NONE})
percentage PERCENTAGE	Specifies a percentage of a SMRT $^{\otimes}$ Cell to recover (Range = 1-100) rather than a specific list of reads. (Default = NONE)
-n COUNT,count COUNT	Specifies a specific number of ZMWs picked at random to recover. (Default = NONE)
-s SEED,seed SEED	Specifies a random seed for selecting a percentage of reads. (Default = NONE)
ignore-metadata	Discard the input Data Set metadata. (Default = False)
barcodes	Specifies that the include/exclude list contains barcode indices instead of ZMW numbers. (Default = False)

Examples

Pulling out two ZMWs from a BAM file:

bamsieve --include 111111,222222 full.subreads.bam sample.subreads.bam

Pulling out two ZMWs from a Data Set file:

bamsieve --include 111111,222222 full.subreadset.xml sample.subreadset.xml

Using a text list:

bamsieve --include zmws.txt full.subreads.bam sample.subreads.bam

Using another BAM or Data Set as a list:

bamsieve --include mapped.alignmentset.xml full.subreads.bam mappable.subreads.bam

Generating a list of ZMWs from a Data Set:

bamsieve --show-zmws mapped.alignmentset.xml > mapped_zmws.txt

Anonymizing a Data Set:

bamsieve --include zmws.txt --ignore-metadata --anonymize full.subreadset.xml
anonymous sample.subreadset.xml

Removing a read:

bamsieve --exclude 111111 full.subreadset.xml filtered.subreadset.xml

Selecting 0.1% of reads:

bamsieve --percentage 0.1 full.subreads.bam random sample.subreads.bam

Selecting a different 0.1% of reads:

bamsieve --percentage 0.1 --seed 98765 full.subreads.bam random sample.subreads.bam

Selecting just two ZMWs/reads at random:

bamsieve --count 2 full.subreads.bam two reads.subreads.bam

Selecting by barcode:

bamsieve --barcodes --include 4,7 full.subreads.bam two barcodes.subreads.bam

Generating a tiny BAM file that contains only mappable reads:

bamsieve --include mapped.subreads.bam full.subreads.bam mappable.subreads.bam bamsieve --count 4 mappable.subreads.bam tiny.subreads.bam

Splitting a Data Set into two halves:

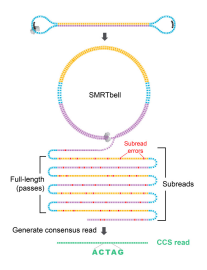
bamsieve --percentage 50 full.subreadset.xml split.1of2.subreadset.xml
bamsieve --exclude split.1of2.subreadset.xml full.subreadset.xml
split.2of2.subreadset.xml

Extracting Unmapped Reads:

bamsieve --exclude mapped.alignmentset.xml movie.subreadset.xml unmapped.subreadset.xml

ccs Circular Consensus Sequencing (CCS) Analysis computes consensus sequences from multiple "passes" around a circularized single DNA molecule (SMRTbell® template). CCS analysis uses the Arrow framework to achieve optimal consensus results given the number of passes available.

Note: CCS Analysis in SMRT Link v12.0 is **only** for **subread data** generated by Sequel II and Sequel IIe systems. Revio systems **always** run CCS oninstrument and **cannot** generate subreads directly.



CCS analysis workflow

- 1. Initial filtering
 - Filter ZMWs: Remove ZMWs with signal-to-noise ratio (SNR) below
 --min-snr.
 - Filter subreads: Remove subreads with lengths <50% or >200% of the median subread length. Stop if the number of full-length subreads is fewer than --min-passes.

2. Generate draft

- The polish stage iteratively improves upon a candidate template sequence. Because polishing is very compute-intensive, it is desirable to start with a template that is as close as possible to the true sequence of the molecule to reduce the number of iterations until convergence. The ccs software does not pick a full-length subread as the initial template to be polished, but instead generates an approximate draft consensus sequence using our improved implementation of the Sparc graph consensus algorithm. This algorithm depends on a subread-to-backbone alignment that is generated by the pancake mapper developed by PacBio, using edlib as the core aligner. Typically, subreads have accuracy of around 90% and the draft consensus has a higher accuracy, but is still below 99%.
- Stop if the draft length is shorter than --min-length and longer than --max-length.

3. Alignment

 Align subreads to the draft consensus using pancake with KSW2 for downstream windowing and filtering.

4. Windowing

 Divide the subread-to-draft alignment into overlapping windows with a target size of 22 bp with ±2 bp overlap. Avoid breaking windows at simple repeats (homopolymers to 4-mer repeats) to reduce edge cases at window borders. Windowing reduces the algorithm run time from quadratic to linear in the insert size.

5. Single-strand artifacts

Identify heteroduplexes, where one strand of the SMRTbell differs significantly from the reverse complement of the other strand.
 Subread orientation is inferred from the alignment. Small heteroduplexes, such as single base A paired with a matching G, are retained and the ambiguity is reflected in base quality. Molecules with a single difference longer than 20 bp between the strands are removed and recorded as heteroduplexes in the

<outputPrefix>.ccs report.txt file.

6. Trim large insertions

Insertions in the subreads relative to the draft that are longer than

 -max-insertion-size, default 30 bp, are trimmed since they
 typically represented spurious sequencing activity.

7. Filter candidates

 For each window, a heuristic is used to find those positions that likely need polishing. In addition, homopolymers are always polished. Skipping unambiguous positions makes the polishing at least twice as fast.

8. Polishing

- The core polishing uses the arrow algorithm, a left-right Hidden Markov-Model (HMM) that models the enzymatic and photophysical events in SMRT sequencing. Emission and transition parameters are estimated by a dinucleotide template context. Transition parameters form a homogeneous Markov chain. The transition parameters do **not** depend on the position within the template, only the pulse width of a base call, the dinucleotide context of the template, and the SNR of the ZMW. Arrow computes the loglikelihood that the subread originates from the template, marginalizing over all possible alignments of the subread to the template. For every position in the template that is a candidate for polishing, arrow tests if the log-likelihood is improved by substituting one of the other three nucleotides, inserting one of the four nucleotides after the position, or deleting the position itself. Once arrow does not find any further beneficial mutations to the template in an iteration, it stops.

9. OV calculation

 The log-likelihood ratio between the most likely template sequence and all of its mutated counterparts is used to calculate a quality for each base in the final consensus. The average of the per-base qualities is the read accuracy, rq.

10. Final steps

The per-window consensus template sequences and base qualities are concatenated and overhangs (overlaps between adjacent windows) are trimmed. If the predicted read accuracy is at least --min-rq, then the consensus read is written to the output.

Input files

One .subreads.bam file containing the subreads for each SMRTbell[®] template sequenced.

Output files

A BAM file with one entry for each consensus sequence derived from a productive ZMW. BAM is a general file format for storing sequence data, which is described fully by the SAM/BAM working group. The CCS analysis output format is a version of this general format, where the consensus sequence is represented by the "Query Sequence". Several tags were added to provide additional meta information. An example BAM entry for a consensus as seen by samtools is shown below.

```
m64009 201008 223950/1/ccs
                          4 * 0
                                        255
                                             * * 0
                                                          0
                                                                ATCGCCTACC
          RG:Z:a773c1f2
                          fi:B:C,26,60,21,41,33,26,63,45,73,33
~|~t~R~~r~
                                                               fn:i:6
fp:B:C,11,18,21,35,8,18,31,8,23,11 np:i:12
ri:B:C,17,37,24,4,70,21,12,44,21,32 rn:i:6
                                           rp:B:C,16,56,17,9,23,19,10,10,23,12
                sn:B:f,10.999,16.2603,3.964,7.17746 we:i:9816064
 rq:f:0.999651
zm:i:1
```

Following are some of the common fields contained in the output BAM file:

Field	Description
Query Name	Movie Name / ZMW # /ccs
FLAG	Required by the format but meaningless in this context. Always set to 4 to indicate the read is unmapped.
Reference Name	Required by the format but meaningless in this context. Always set to *.
Mapping Start	Required by the format but meaningless in this context. Always set to 0.
Mapping Quality	Required by the format but meaningless in this context. Always set to 255.
CIGAR	Required by the format but meaningless in this context. Always set to *.
RNEXT	Required by the format but meaningless in this context. Always set to *.
PNEXT	Required by the format but meaningless in this context. Always set to 0.
TLEN	Required by the format but meaningless in this context. Always set to 0.
Consensus Sequence	The consensus sequence generated.
Quality Values	The per-base parametric quality metric. For details see "Interpreting QUAL values" on page 13.
RG Tag	The read group identifier.
bc Tag	A 2-entry array of upstream-provided barcode calls for this ZMW.

Field	Description
bq Tag	The quality of the barcode call. (Optional: Depends on barcoded inputs.)
np Tag	The number of full passes that went into the subread. (Optional : Depends on barcoded inputs.)
rq Tag	The predicted read quality.
zm Tag	The ZMW hole number.
ma Tag	Bitmask storing whether a SMRTbell adapter is missing on either side of the molecule that produced a CCS read. 0 indicates that neither end has a confirmed missing adapter.
ac Tag	An array containing the counts of detected and missing SMRTbell adapters on either side of the molecule that produced a CCS read:
	Detected adapters on left/start
	Missing adapters on left/start
	Detected adapters on right/end
	Missing adapter on right/end

Usage

ccs [OPTIONS] INPUT OUTPUT

Example

ccs --all myData.subreads.bam myResult.bam

Required	Description
Input File Name	The name of a single subreads.bam or a subreadset.xml file to be processed. (Example = myData.subreads.bam)
Output File Name	The name of the output BAM file; comes after all other options listed. Valid output files are the BAM and the Dataset .xml formats. (Example = myResult.bam)

Options	Description
version	Prints the version number.
report-file	Contains a result tally of the outcomes for all ZMWs that were processed. If no file name is given, the report is output to the file ccs_report.txt. In addition to the count of successfully-produced consensus sequences, this file lists how many ZMWs failed various data quality filters (SNR too low, not enough full passes, and so on) and is useful for diagnosing unexpected drops in yield.
min-snr	Removes data that is likely to contain deletions. SNR is a measure of the strength of signal for all 4 channels (A, C, G, T) used to detect base pair incorporation. This value sets the threshold for minimum required SNR for any of the four channels. Data with SNR < 2.5 is typically considered lower quality. (Default = 2.5)
min-length	Specifies the minimum length requirement for the minimum length of the draft consensus to be used for further polishing. If the targeted template is known to be a particular size range, this can filter out alternative DNA templates. (Default = 10)
max-length	Specifies the maximum length requirement for the maximum length of the draft consensus to be used for further polishing. For robust results while avoiding unnecessary computation on unusual data, set to ~20% above the largest expected insert size. (Default = 50000)
min-passes	Specifies the minimum number of passes for a ZMW to be emitted. This is the number of full passes. Full passes must have an adapter hit before and after the insert sequence and so do not include any partial passes at the start and end of the sequencing reaction. It is computed as the number of passes mode across all windows. (Default = 3)

Options	Description
min-rq	Specifies the minimum predicted accuracy of a read. CCS analysis generates an accuracy prediction for each read, defined as the expected percentage of matches in an alignment of the consensus sequence to the true read. A value of 0.99 indicates that only reads expected to be 99% accurate are emitted. (Default = 0.99)
num-threads	Specifies how many threads to use while processing. By default, CCS analysis uses as many threads as there are available cores to minimize processing time, but fewer threads can be specified here.
log-file	The name of a log file to use. If none is given, the logging information is printed to STDERR. (Example: mylog.txt)
log-level	Specifies verbosity of log data to produce. By settinglogLevel=DEBUG, you can obtain detailed information on what ZMWs were dropped during processing, as well as any errors which may have appeared. (Default = INFO)
skip-polish	After constructing the draft consensus, do not proceed with the polishing steps. This is significantly faster, but generates less accurate data with no RQ or QUAL values associated with each base.
by-strand	Separately generates a consensus sequence from the forward and reverse strands. Useful for identifying heteroduplexes formed during sample preparation.
chunk	Operates on a single chunk. Format i/N , where i in [1,N]. Examples: $3/24$ or $9/9$.
max-chunks	Determines the maximum number of chunks, given an input file.
model-path	Specifies the path to a model file or directory containing model files.
model-spec	Specifies the name of the chemistry or model to use, overriding the default selection.
all	Generates one representative sequence per ZMW, irrespective of quality and passesmin-passes 0min-rq 0max-length 0 are set implicitly and cannot be changed;all also deactivates the maximum draft length filter. Filtering has to be performed downstream.
	The ccsall option changes the workflow as follows:
	 There is special behavior for low-pass ZMWs. If a ZMW has fewer than 2 full-length subreads, use the subread of median length as representative consensus, optionally with its kinetic information as forward orientation usingall-kinetics, and do not polish.
	2. Only polish ZMWs with at least two full-length subreads mapping back to the draft. Otherwise, set predicted accuracy rq tag to -1 to indicate that the predicted accuracy was not calculated, and populate per-base QVs with + (QV 10) the approximate raw accuracy. Kinetic information is not available for unpolished drafts.
	3. Instead of using an unpolished draft without kinetic information as a representative consensus sequence, ifsubread-fallback is used, fall back to a representative subread with kinetic information.
	How isall different from explicitly settingmin-passes 0min-rq 0?
	 Settingmin-passes 0min-rq 0 is a brute-force combination that polishes every ZMW, even those that only have one partial subread, with polishing making no difference. In contrast,all is a bit smarter and only polishes ZMWs with at least one full-length subread and one additional partial subread.

Options	Description
hifi-kinetics	Generates averaged kinetic information for polished reads, independently for both strands of the insert. Forward is defined with respect to the orientation represented in SEQ and is considered to be the native orientation. As with other PacBio-specific tags, aligners will not re-orient these fields.
	Base modifications can be inferred from per-base pulse width (PW) and inter-pulse duration (IPD) kinetics.
	Minor cases exist where a certain orientation may get filtered out entirely from a ZMW, preventing valid values from being passed for that record. In these cases, empty lists are passed for the respective record/orientation, and number of passes are set to zero.
	To facilitate the use of HiFi reads with base modifications workflows, we added an executable in pbbam called ccs-kinetics-bystrandify which creates a pseudoby-strand BAM with corresponding pw and ip tags that imitates a normal, unaligned subreads BAM.
all-kinetics	Adds kinetic information for all ZMWs, except for unpolished draft consensus.
subread-fallback	When combined withall, uses a subread instead of a draft as representative consensus.
suppress-reports	Suppresses the generation of default reports and metric files.

Interpreting QUAL values

The QUAL value of a read is a measure of the posterior likelihood of an error at a particular position. **Increasing** QUAL values are associated with a **decreasing** probability of error. For indels and homopolymers, there is ambiguity as to which QUAL value is associated with the error probability. Shown below are different types of alignment errors, with an * indicating which sequence BP should be associated with the alignment error.

Mismatch

* accma:

ccs: ACGTATA
ref: ACATATA

Deletion

ccs: AC-TATA

ref: ACATATA

Insertion

* *CCS: ACC

ccs: ACGTATA
ref: AC-TATA

Homopolymer insertion or deletion

Indels should always be left-aligned, and the error probability is only given for the first base in a homopolymer.

ccs: ACGGGGTATA ccs: AC-GGGTATA ref: AC-GGGTATA ref: ACGGGGTATA

CCS Analysis Yield report

The CCS Analysis Yield report specifies the number of ZMWs that successfully produced consensus sequences, as well as a count of how many ZMWs did **not** produce a consensus sequence for various reasons. The entries in this report, as well as parameters used to increase or decrease the number of ZMWs that pass various filters, are shown in the table below.

The first part is a summary of inputs and outputs:

ZMW results	Parameters affecting results	Description
ZMWs input	None	The number of input ZMWs.
ZMWs pass filters	All custom processing settings	The number of CCS reads successfully produced on the first attempt, using the fast windowed approach.
ZMWs fail filters	All custom processing settings	The number of ZMWs reads that failed to produce a CCS read.
ZMWs shortcut filters	-all	The number of ZMWs having fewer than 2 full-length subreads.
ZMWs with tandem repeats	min-tandem-repeat- length	The number of ZMWs with repeats larger than the specified threshold.

The second part explains in detail the exclusive ZMW count for those ZMWs that were filtered:

ZMW results	Parameters affecting results	Description
No usable subreads	None	The ZMW had no usable subreads. Either there were no subreads, or all subreads had lengths outside the range <50% or >200% of the median subread length.
Below SNR threshold	min-snr	The ZMW had at least one channel's SNR below the minimum threshold.
Lacking full passes	min-passes	There were not enough subreads that had an adapter at the start and end of the subread (a "full pass").
Heteroduplexes	None	The SMRTbell contains a heteroduplex. In this case, it is not clear what the consensus should be and so the ZMW is dropped.
Min coverage violation	None	The ZMW is damaged on one strand and cannot be polished reliably.
Draft generation error	None	Subreads do not match the generated draft sequence, even after multiple tries.
Draft abovemax-length	max-length	The draft sequence was above the maximum length threshold.
Draft belowmin-length	min-length	The draft sequence was below the minimum length threshold.
Lacking usable subreads	None	Too many subreads were dropped while polishing.
CCS analysis did not converge	None	The consensus sequence did not converge after the maximum number of allowed rounds of polishing.

ZMW results	Parameters affecting results	Description
CCS read below minimum predicted accuracy	min-rq	Each CCS read has a predicted level of accuracy associated with it. Reads that are below the minimum specified threshold are removed.
Unknown error during processing	None	These should not occur.

How do I read the ccs_report.txt file?

By default, each CCS analysis generates a <code>ccs_report.txt</code> file. This file summarizes how many ZMWs generated HiFi reads and how many ZMWs failed CCS reads generation because of the listed causes. For those failing, each ZMW contributes to exactly one reason of failure; percentages are with respect to number of failed ZMWs.

Does CCS analysis dislike low-complexity regions?

Low-complexity comes in many shapes and forms. A particular challenge for CCS analysis are highly-enriched tandem repeats, such as hundreds of copies of AGGGGT. Prior to $\cos v5.0$, inserts with many copies of a small repeat likely did not generate a consensus sequence. Since $\cos v5.0$, every ZMW is tested if it contains a tandem repeat of length

--min-tandem-repeat-length 1000. For this, we use symmetric DUST, specifically this sdust implementation, but slightly modified. If a ZMW is flagged as a tandem repeat, internally --disable-heuristics is activated for only this ZMW, and various filters that are known to exclude low-complexity sequences are disabled. This recovers most of the low-complexity consensus sequences, without impacting run time performance.

Can I produce one consensus sequence for each strand of a molecule?

Yes, use --by-strand. Make sure that you have sufficient coverage, as --min-passes are per-strand in this case. For each strand, CCS analysis generates one consensus read that has to pass all filters. Read name suffix indicates strand. **Example**:

```
m64011_190714_120746/14/ccs/rev
m64011_190714_120746/35/ccs/fwd
```

How does --by-strand work? For each ZMW:

- Determine orientation of reads with respect to the one closest to the median length.
- Sort reads into two buckets, forward and reverse strands.
- Treat each strand as an individual entity as we do with ZMWs:
 - Apply all filters per strand individually.
 - Create a draft for each strand.

- Polish each strand.
- Write out each polished strand consensus.

BAM tags generated

Tag	Туре	Description
ec	f	Effective coverage
fi	B,C	Forward IPD (Codec V1)
fn	i	Forward number of complete passes (zero or more)
fp	В,С	Forward PulseWidth (Codec V1)
np	i	Number of full-length subreads
ri	B,C	Reverse IPD (Codec V1)
rn	i	Reverse number of complete passes (zero or more)
rp	В,С	Reverse PulseWidth (Codec V1)
rq	f	Predicted average read accuracy
sn	B,F	Signal-to-noise ratios for each nucleotide
zm	i	ZMW hole number
RG	Z	Read group

dataset

The dataset tool creates, opens, manipulates and writes Data Set XML files. The commands allow you to perform operations on the various types of data held by a Data Set XML: Merge, split, write, and so on.

Usage

```
dataset [-h] [--version] [--log-file LOG_FILE]
        [--log-level {DEBUG,INFO,WARNING,ERROR,CRITICAL} | --debug | --quiet | -v]
        [--strict] [--skipCounts]
```

{create, filter, merge, split, validate, summarize, consolidate, loadstats, newuuid, loadmetada ta, copyto, absolutize, relativize}

Options	Description
-h,help	Displays help information and exits.
<command/> -h	Displays help for a specific command.
-v,version	Displays program version number and exits.
log-file LOG_FILE	Writes the log to file. (Default = None, writes to stdout.)
log-level	Specifies the log level; values are [DEBUG, INFO, WARNING, ERROR, CRITICAL]. (Default = INFO)
debug	Alias for setting the log level to DEBUG. (Default = False)
quiet	Alias for setting the log level to CRITICAL to suppress output. (Default = False)
-A	Sets the verbosity level. (Default = NONE)
strict	Turns on strict tests and display all errors. (Default = False)
skipCounts	Skips updating NumRecords and TotalLength counts. (Default = False)

create command: Create an XML file from a fofn (file-of-file names) or BAM file. Possible types: SubreadSet, AlignmentSet, ReferenceSet, HdfSubreadSet, BarcodeSet, ConsensusAlignmentSet, ConsensusReadSet, ContigSet.

Example

The following example shows how to use the dataset create command to create a barcode file:

dataset create --generateIndices --name my_barcodes --type BarcodeSet
my_barcodes.barcodeset.xml my_barcodes.fasta

Required	Description
outfile	The name of the XML file to create.
infile	The fofn (file-of-file-names) or BAM file(s) to convert into an XML file.

Options	Description
type DSTYPE	Specifies the type of XML file to create. (Default = NONE)
name DSNAME	The name of the new Data Set XML file.
generateIndices	Generates index files (.pbi and .bai for BAM, .fai for FASTA). Requires samtools/pysam and pbindex. (Default = FALSE)
metadata METADATA	A metadata.xml file (or Data Set XML) to supply metadata. (Default = NONE)
novalidate	Specifies not to validate the resulting XML. Leaves the paths as they are.
relative	Makes the included paths relative instead of absolute. This is not compatible withnovalidate.

filter command: Filter an XML file using filters and threshold values.

- Suggested filters: alignedlength, as, astart, bc, bcf, bcq, bcr, bq, cx, length, mapqv, movie, n_subreads, pos, qend, qid, qname, qstart, readstart, rname, rq, tend, tstart, zm
- More resource-intensive filter: [qs]

Note: Multiple filters with different names are ANDed together. Multiple filters with the **same** name are ORed together, duplicating existing requirements. The filter string should be enclosed in **single quotes**.

dataset filter [-h] infile outfile filters [filters ...]

Required	Description
infile	The name of the XML file to filter.
outfile	The name of the output filtered XML file.
filters	The values to filter on. (Example: rq>0.85)

Examples

Filter on read quality >0.99 (Q20):

% dataset filter in.consensusreadset.xml hifi.consensusreadset.xml 'rq >= 0.99'

Filter on read quality and length:

 $\$ dataset filter in.consensus readset.xml filtered.consensus readset.xml 'rq >= 0.99 AND length >= 10000'

Filter for very long and very short reads:

\$ dataset filter in.consensusreadset.xml filtered.consensusreadset.xml 'length >= 40000; length <= 400'

Filter for specific high-quality barcodes:

% dataset filter mixed.consensusreadset.xml samples1-3.consensusreadset.xml 'bc = [0,1,2] AND bq >= 26'

merge command: Combine XML files.

dataset merge [-h] outfile infiles [infiles ...]

Required	Description
infiles	The names of the XML files to merge.
outfile	The name of the output XML file.

split command: Split a Data Set XML file.

Required	Description
infile	The name of the XML file to split.

Options	Description
outfiles	The names of the resulting XML files.
contigs	Splits the XML file based on contigs. (Default = FALSE)
barcodes	Splits the XML file based on barcodes. (Default = FALSE)
zmws	Splits the XML file based on ZMWs. (Default = FALSE)
byRefLength	Splits contigs by contig length. (Default = TRUE)
noCounts	Updates the Data Set counts after the split. (Default = FALSE)
chunks x	Splits contigs into x total windows. (Default = 0)
maxChunks x	Splits the contig list into at most x groups. (Default = 0)
targetSize x	Specifies the minimum number of records per chunk. (Default = 5000)
breakContigs	Breaks contigs to get closer to maxCounts. (Default = False)
subdatasets	Splits the XML file based on sub-datasets. (Default = False)
outdir OUTDIR	Specifies an output directory for the resulting XML files. (Default = <in-place>, not the current working directory.)</in-place>

 ${\tt validate}$ command: Validate XML and Resourceld files. (This is an internal testing functionality that may be useful.)

Note: This command requires that pyxb (**not** distributed with SMRT Link) be installed. If **not** installed, validate simply checks that the files pointed to in ResourceIds exist.

dataset validate [-h] [--skipFiles] infile

Required	Description
infile	The name of the XML file to validate.

Options	Description
skipFiles	Skips validating external resources. (Default = False)

summarize command: Summarize a Data Set XML file.

dataset summarize [-h] infile

Required	Description
infile	The name of the XML file to summarize.

consolidate command: Consolidate XML files.

Required	Description
infile	The name of the XML file to consolidate.
datafile	The name of the resulting data file.
xmlfile	The name of the resulting XML file.

Options	Description
numFiles x	Specifies the number of data files to produce. (Default = 1)
noTmp	Do not copy to a temporary location to ensure local disk use. (Default = False)

loadstats command: Load an ${\tt sts.xml}$ file containing pipeline statistics into a Data Set XML file.

dataset loadstats [-h] [--outfile OUTFILE] infile statsfile

Required	Description
infile	The name of the Data Set XML file to modify.
statsfile	The name of the .sts.xml file to load.

Options	Description	
outfile OUTFILE	The name of the XML file to output. (Default = None)	

newuuid command: Refresh a Data Set's Unique ID.

dataset newuuid [-h] [--random] infile

Required	Description
infile	The name of the XML file to refresh.

Options	Description
random	Generates a random UUID, instead of a hash. (Default = False)

loadmetadata command: Load a .metadata .xml file into a Data Set XML file.

dataset loadmetadata [-h] [--outfile OUTFILE] infile metadata

Required	Description
infile	The name of the Data Set XML file to modify.
metadata	The .metadata.xml file to load, or Data Set to borrow from.

Options	Description
outfile OUTFILE	Specifies the XML file to output. (Default = None)

copyto command: Copy a Data Set and resources to a new location.

dataset copyto [-h] [--relative] infile outdir

Required	Description
infile	The name of the XML file to copy.
outdir	The directory to copy to.

Options	Description	
relative	Makes the included paths relative instead of absolute. (Default = False)	

absolutize command: Make the paths in an XML file absolute.

dataset absolutize [-h] [--outdir OUTDIR] infile

Required	Description	
infile	The name of the XML file whose paths should be absolute.	

Options	Description	
outdir OUTDIR	Specifies an optional output directory. (Default = None)	

relativize command: Make the paths in an XML file relative.

dataset relativize [-h] infile

Required	Description	
infile	The name of the XML file whose paths should be relative.	

Examples - Filter reads

To filter one or more BAM file's worth of subreads, aligned or otherwise, and then place them into a single BAM file:

```
# usage: dataset filter <in_fn.xml> <out_fn.xml> <filters>
dataset filter in_fn.subreadset.xml filtered_fn.subreadset.xml 'rq>0.85'
# usage: dataset consolidate <in_fn.xml> <out_data_fn.bam> <out_fn.xml>
dataset consolidate filtered_fn.subreadset.xml consolidate.subreads.bam
out fn.subreadset.xml
```

The filtered Data Set and the consolidated Data Set should be read-forread equivalent when used with SMRT® Analysis software.

Example - Resequencing pipeline

- Align two movie's worth of subreads in two SubreadSets to a reference
- · Merge the subreads together.
- Split the subreads into Data Set chunks by contig.
- Process using gcpp on a chunkwise basis (in parallel).
- 1. Align each movie to the reference, producing a Data Set with one BAM file for each execution:

```
pbalign movie1.subreadset.xml referenceset.xml movie1.alignmentset.xml
pbalign movie2.subreadset.xml referenceset.xml movie2.alignmentset.xml
```

2. Merge the files into a FOFN-like Data Set; BAMs are not touched:

```
# dataset merge <out_fn> <in_fn> [<in_fn> <in_fn> ...]
dataset merge merged.alignmentset.xml movie1.alignmentset.xml movie2.alignmentset.xml
```

- 3. Split the Data Set into chunks by contig name; BAMs are not touched:
 - Note that supplying output files splits the Data Set into that many output files (up to the number of contigs), with multiple contigs per file.
 - Not supplying output files splits the Data Set into one output file per contig, named automatically.
 - Specifying a number of chunks instead will produce that many files, with contig or even subcontig (reference window) splitting.

dataset split --contigs --chunks 8 merged.alignmentset.xml

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4. Process the chunks:

gcpp --reference referenceset.xml --output
chunklconsensus.fasta,chunklconsensus.fastq,chunklconsensus.vcf,chunklconsensus.gff
chunklcontigs.alignmentset.xml

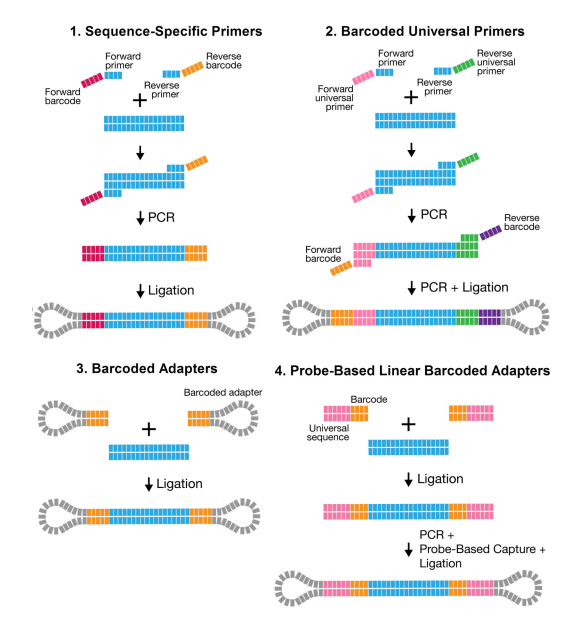
The chunking works by duplicating the original merged Data Set (no BAM duplication) and adding filters to each duplicate such that only reads belonging to the appropriate contigs are emitted. The contigs are distributed among the output files in such a way that the total number of records per chunk is about even.

Demultiplex Barcodes

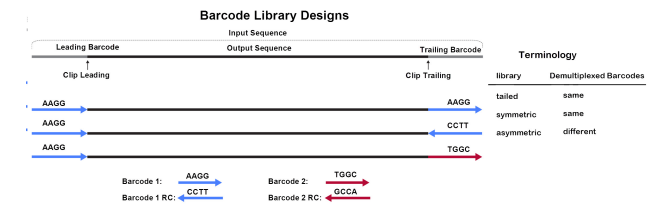
The **Demultiplex Barcodes** application identifies barcode sequences in PacBio single-molecule sequencing data.

Demultiplex Barcodes can demultiplex samples that have a unique persample barcode pair and were pooled and sequenced on the same SMRT® Cell. There are four different methods for barcoding samples with PacBio technology:

- 1. Sequence-specific primers
- 2. Barcoded universal primers
- 3. Barcoded adapters
- 4. Linear Barcoded Adapters for Probe-based Captures



In addition, there are three different barcode library designs. **Demultiplex Barcodes** supports raw subread and CCS reads demultiplexing.



In the overview above, the input sequence is flanked by adapters on both sides. The bases adjacent to an adapter are **barcode regions**. A read can have up to two barcode regions, leading and trailing. Either or both adapters can be missing and consequently the leading and/or trailing region is not being identified.

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.

For the **asymmetric** design, **different** barcodes are attached to the sides of the insert sequence of interest. To identify the different barcodes, a read with leading and trailing barcode regions is required.

Output barcode pairs are generated from the identified barcodes. The barcode names are combined using "--", for example bc1002--bc1054. The sort order is defined by the barcode indices, starting with the lowest.

Workflow

By default, **Demultiplex Barcodes** processes input reads grouped by ZMW, **except** if the <code>--per-read</code> option is used. All barcode regions along the read are processed individually. The final per-ZMW result is a summary over all barcode regions. Each ZMW is assigned to a pair of selected barcodes from the provided set of candidate barcodes. Subreads from the same ZMW will have the same barcode and barcode quality. For a particular target barcode region, every barcode sequence gets aligned as given and as reverse-complement, and higher scoring orientation is chosen. This results in a list of scores over all candidate barcodes.

If only same barcode pairs are of interest (symmetric/tailed), use the
 --same option to filter out different barcode pairs.

If only different barcode pairs are of interest (asymmetric), use the

 -different option to require at least two barcodes to be read, and remove pairs with the same barcode.

Parameter presets

Recommended parameter combinations are available using --preset for HiFi input:

```
    HIFI-SYMMETRIC
```

```
--ccs --min-score 80 --min-end-score 50 --min-ref-span 0.75 --same
```

HIFI-ASYMMETRIC

```
--ccs --min-score 80 --min-end-score 50 --min-ref-span 0.75 --different --min-scoring-regions 2
```

NONE (Default)

Half adapters

For an adapter call with only one barcode region, the high-quality region finder cuts right through the adapter. The preceding or succeeding subread was too short and was removed, or the sequencing reaction started/stopped there. This is called a **half adapter**. Thus, there are also 1.5, 2.5, N+0.5 adapter calls.

ZMWs with half or only one adapter can be used to identify the same barcode pairs; positive-predictive value might be reduced compared to high adapter calls. For asymmetric designs with different barcodes in a pair, at least a single full-pass read is required; this can be two adapters, two half adapters, or a combination.

Usage

- Any existing output files are overwritten after execution.
- Always use --peek-guess to remove spurious barcode hits.

Analysis of subread data

```
lima movie.subreads.bam barcodes.fasta prefix.bam
lima movie.subreadset.xml barcodes.barcodeset.xml prefix.subreadset.xml
```

Analysis of CCS reads

```
lima --css movie.ccs.bam barcodes.fasta prefix.bam
lima --ccs movie.consensusreadset.xml barcodes.barcodeset.xml
prefix.consensusreadset.xml
```

If you do not need to import the demultiplexed data into SMRT Link, use the -no-pbi option to minimize memory consumption and run time.

Symmetric or tailed options

```
Raw: --same
CCS read: --same --ccs
```

Asymmetric options

Raw: --different
CCS reads: --different --ccs

Example execution

lima m54317_180718_075644.subreadset.xml \ Sequel_RSII_384_barcodes_v1.barcodeset.xml \ m54317_180718_075644.demux.subreadset.xml \ --different --peek-guess

Options	Description			
same	Retains only reads with the same barcodes on both ends of the insert sequence, such as symmetric and tailed designs.			
different	Retains only reads with different barcodes on both ends of the insert sequence, asymmetric designs. Enforcesmin-passes ≥ 1 .			
min-length n	Omits reads with lengths below $\tt n$ base pairs after demultiplexing. ZMWs with no reads passing are omitted. (Default = 50)			
max-input-length n	Omits reads with lengths above ${\tt n}$ base pairs for scoring in the demultiplexing step. (Default = 0, deactivated)			
min-score n	Omits ZMWs with average barcode scores below n. A barcode score measures the alignment between a barcode attached to a read and an ideal barcode sequence, and is an indicator how well the chosen barcode pair matches. It is normalized to a range between 0 (no hit) and 100 (a perfect match). (Default = 0, PacBio recommends setting it to 26.)			
min-end-score n	Specifies the minimum end barcode score threshold applied to the individual leading and trailing ends. (Default = 0)			
min-passes n	Omits ZMWs with less than n full passes, a read with a leading and trailing adapter. (Default = 0, no full-pass needed) Example: 0 pass : insert - adapter - insert 1 pass : insert - adapter - INSERT - adapter - insert 2 passes: insert - adapter - INSERT - adapter - INSERT - adapter - insert			
score-full-pass	Uses only reads flanked by adapters on both sides (full-pass reads) for barcode identification.			
min-ref-span	Specifies the minimum reference span relative to the barcode length. (Default = 0.5)			
per-read	Scores and tags per subread, instead of per ZMW.			
ccs	Sets defaults to -A 1 -B 4 -D 3 -I 3 -X 1.			
peek n	Looks at the first n ZMWs of the input and return the mean. This lets you test multiple test barcode.fasta files and see which set of barcodes was used.			
guess n	This performs demultiplexing twice. In the first iteration, all barcodes are tested per ZMW. Afterwards, the barcode occurrences are counted and their mean is tested against the threshold n; only those barcode pairs that pass this threshold are used in the second iteration to produce the final demultiplexed output. A prefix.lima.guess file shows the decision process;same is being respected.			
guess-min-count	Specifies the minimum ZMW count to whitelist a barcode. This filter is ANDed with the minimum barcode score specified byguess. (Default = 0)			

Options	Description		
peek-guess	Sets the following options: peek 50000guess 45guess-min-count 10. Demultiplex Barcodes will run twice on the input data. For the first 50,000 ZMWs, it will guess the barcodes and store the mask of identified barcodes. In the second run, the barcode mask is used to demultiplex all ZMWs.		
	If combined withccs then the barcode score threshold is increased byguess 75.		
single-side	Identifies barcodes in molecules that only have barcodes adjacent to one adapter.		
window-size-mult window-size-bp	The candidate region size multiplier: barcode_length * multiplier. (Default = 3) Optionally, you can specify the region size in base pairs using		
	window-size-bp. If set,window-size-mult is ignored.		
num-threads n	Spawns $\bf n$ threads; 0 means use all available cores. This option also controls the number of threads used for BAM and PBI compression. (Default = 0)		
chunk-size n	Specifies that each thread consumes n ZMWs per chunk for processing. (Default = 10).		
no-bam	Does not produce BAM output. Useful if only reports are of interest, as run time is shorter.		
no-pbi	Does not produce a .bam.pbi index file. The on-the-fly .bam.pbi file generation buffers the output data. If you do not need a .bam.pbi index file for SMRT Link import, use this option to decrease memory usage to a minimum and shorten the run time.		
no-reports	Does not produce any reports. Useful if only demultiplexed BAM files are needed.		
dump-clips	Outputs all clipped barcode regions generated to the <pre><pre><pre><pre>fix>.lima.clips file.</pre></pre></pre></pre>		
dump-removed	Outputs all records that did not pass the specified thresholds, or are without barcodes, to the <pre><pre><pre></pre></pre></pre>		
split-bamsplit-bam-named	Specifies that each barcode has its own BAM file called prefix.idxBest-idxCombined.bam, such as prefix.0-0.bam. Optionally,split-bam-named names the files by their barcode names instead of their barcode indices.		
isoseq	Removes primers as part of the Iso-Seq® pipeline. See "Demultiplexing Iso-Seq® data" on page 33 for details.		
bad-adapter-ratio n	Specifies the maximum ratio of bad adapters. (Default = 0).		

Input files

Input data in PacBio-enhanced BAM format is either:

- Sequence data Unaligned subreads, directly from Sequel II systems and Sequel IIe systems.
- Unaligned CCS reads, generated by CCS analysis.

Barcodes are provided as a FASTA file or BarcodeSet file:

- One entry per barcode sequence.
- No duplicate sequences.

- · All bases must be in upper-case.
- Orientation-agnostic (forward or reverse-complement, but not reversed.)

Example

```
>bc1000
CTCTACTTACTTACTG
>bc1001
GTCGTATCATCATGTA
>bc1002
AATATACCTATCATTA
```

Note: Name barcodes using an alphabetic character prefix to avoid later barcode name/index confusion.

Output files

Demultiplex Barcodes generates multiple output files by default, all starting with the same prefix as the output file, using the suffixes .bam, .subreadset.xml, and .consensusreadset.xml. The report prefix is lima. **Example**:

```
lima m54007\_170702\_064558.subreads.bam barcode.fasta /my/path/ m54007\_170702\_064558 demux.subreadset.xml
```

For all output files, the prefix is

```
/my/path/m54007_170702_064558_demux.
```

- contains clipped records, annotated with barcode tags, that passed filters and respect the --same option.
- fix>.lima.report: A tab-separated file describing each ZMW,
 unfiltered. This is useful information for investigating the
 demultiplexing process and the underlying data. A single row contains
 all reads from a single ZMW. For --per-read, each row contains one
 subread, and ZMWs might span multiple rows.
- prefix>.lima.summary: Lists how many ZMWs were filtered, how
 many ZMWs are the same or different, and how many reads were
 filtered.

```
Undesired hybrids : xxx (xx%) <- Only with --peek-guess
Undesired same barcode pairs : xxx (xx%) <- Only with --different
Undesired diff barcode pairs : xxx (xx%) <- Only with --same
Undesired 5p--5p pairs : xxx (xx%) <- Only with --isoseq Undesired 3p--3p pairs : xxx (xx%) <- Only with --isoseq Undesired single side : xxx (xx%) <- Only with --isoseq Undesired no hit : xxx (xx%) <- Only with --isoseq
ZMWs for (B):
With same barcode : 162244 (92%) With different barcodes : 14112 (8%)
Coefficient of correlation : 32.79%
(4)
ZMWs for (A):
Allow diff barcode pair : 157264 (74%)
Allow same barcode pair : 188026 (88%)

Bad adapter yield loss : 10112 (5%) <- Only with --bad-adapter-ratio

Bad adapter impurity : 10348 (5%) <- Only without --bad-adapter-ratio
Reads for (B):
Above length
                                    : 1278461 (100%)
Below length
                                        : 2787 (0%)
```

Explanation of each block:

- 1. Number of ZMWs that went into lima, how many ZMWs were passed to the output file, and how many did not qualify.
- 2. For those ZMWs that did not qualify: The marginal counts of each filter. (Filter are described in the Options table.)

 When running with --peek-guess or similar manual option combination and different barcode pairs are found during peek, the full SMRT Cell may contain low-abundant different barcode pairs that were identified during peek individually, but not as a pair. Those unwanted barcode pairs are called hybrids.
- 3. For those ZMWs that passed: How many were flagged as having the same or different barcode pair, as well as the coefficient of variation for the barcode ZMW yield distribution in percent.
- 4. For all input ZMWs: How many allow calling the same or different barcode pair. This is a simplified version of how many ZMW have at least one full pass to allow a different barcode pair call and how many ZMWs have at least half an adapter, allowing the same barcode pair call.
- 5. For those ZMWs that qualified: The number of reads that are above and below the specified --min-length threshold.
- counts: A .tsv file listing the counts of each
 observed barcode pair. Only passing ZMWs are counted.

Example: column -t prefix.lima.count

IdxFirst	IdxCombined	IdxFirstNamed	IdxCombinedNamed	Counts	MeanScore
0	0	bc1001	bc1001	1145	68
1	1	bc1002	bc1002	974	69
2	2	bc1003	bc1003	1087	68

• clips: Contains clipped barcode regions generated
using the --dump-clips option. Example:

```
head -n 6 prefix.lima.clips

>m54007 170702 064558/4850602/6488_6512 bq:34 bc:11

CATGTCCCCTCAGTTAAGTTACAA

>m54007 170702 064558/4850602/6582_6605 bq:37 bc:11

TTTTGACTAACTGATACCAATAG

>m54007 170702 064558/4916040/4801 4816 bq:93 bc:10
```

- contains records that did not pass the
 specified thresholds, or are without barcodes, using the option
 --dump-removed.lima does not generate a .pbi, nor Data Set for this
 file. This option cannot be used with any splitting option.
- fix>.lima.guess: A.tsv file that describes the barcode
 subsetting process activated using the --peek and --guess options.

IdxFirst	IdxCombined	IdxFirstNamed	IdxCombinedNamed	NumZMWs	MeanScore	Picked
0	0	bc1001t	bc1001t	1008	50	1
1	1	bc1002t	bc1002t	1005	60	1
2	2	bc1003t	bc1003t	5	24	0
3	3	bc1004t	bc1004t	555	61	1

- One DataSet, subreadset.xml, or .consensusreadset.xml file is generated per output BAM file.
- .pbi: One PBI file is generated per output BAM file.

What is a universal spacer sequence and how does it affect demultiplexing?

For library designs that include an identical sequence between adapter and barcode, such as probe-based linear barcoded adapters samples, Demultiplex Barcodes offers a special mode that is activated if it finds a shared prefix sequence among all provided barcode sequences.

Example

>custombc1
ACATGACTGTGACTATCTCACACATATCAGAGTGCG
>custombc2
ACATGACTGTGACTATCTCAACACACAGACTGTGAG

In this case, Demultiplex Barcodes detects the shared prefix

ACATGACTGTGACTATCTCA and removes it internally from all barcodes. Subsequently, it increases the window size by the length ${\tt L}$ of the prefix sequence.

- If -window-size-bp N is used, the actual window size is L + N.
- If --window-size-mult M is used, the actual window size is (L + |bc|) * M.

Because the alignment is semi-global, a leading reference gap can be added without any penalty to the barcode score.

What are bad adapters?

In the <code>subreads.bam</code> file, each subread has a context flag <code>cx</code>. The flag specifies, among other things, whether a subread has flanking adapters, before and/or after. Adapter-finding was improved and can also find molecularly-missing adapters, or those obscured by a local decrease in accuracy. This may lead to missing or obscured bases in the flanking barcode. Such adapters are labelled "bad", as they don't align with the adapter reference sequence(s). Regions flanking those bad adapters are problematic, because they can fully or partially miss the barcode bases, leading to wrong classification of the molecule. <code>lima</code> can handle those adapters by <code>ignoring</code> regions flanking bad adapters. For this, <code>lima</code> computes the ratio of number of bad adapters divided by number of all adapters.

By default, --bad-adapter-ratio is set to 0 and does not perform any filtering. In this mode, bad adapters are handled just like good adapters.

But the *.lima.summary file contains one row with the number of ZMWs that have at least 25% bad adapters, but otherwise pass all other filters. This metric can be used as a diagnostic to assess library preparation.

If --bad-adapter-ratio is set to non-zero positive (0,1), bad adapter flanking barcode regions are treated as missing. If a ZMW has a higher ratio of bad adapters than provided, the ZMW is filtered and consequently removed from the output. The *.lima.summary file contains two additional rows.

```
With bad adapter : 10349 (28%)
Bad adapter yield loss : 10112 (5%)
```

The first row counts the number of ZMWs that have bad adapter ratios that are too high; the percentage is with respect to the number of all ZMW not passing. The second row counts the number of ZMWs that are removed solely due to bad adapter ratios that are too high; the percentage is with respect the number of all input ZMWs and consequently is the effective yield loss caused by bad adapters.

If a ZMW has ~50% bad adapters, one side of the molecule is molecularly-missing an adapter. For 100% bad adapter, **both** sides are missing adapters. A lower than ~40% percentage indicates decreased local accuracy during sequencing leading to adapter sequences not being found. If a high percentage of ZMWs is molecularly-missing adapters, you should improve library preparation.

Demultiplexing Iso-Seq® data

Demultiplex Barcodes is used to identify and remove Iso-Seq cDNA primers. If the Iso-Seq sample is barcoded, the barcodes should be included as part of the primer. Only by using the command-line can users use lima with the --isoseq option for demultiplexing Iso-Seq data.

The input Iso-Seq data format for demultiplexing is .ccs.bam. Users must first generate a CCS reads BAM file for an Iso-Seq Data Set before running lima. The recommended parameters for running CCS analysis for Iso-Seq are min-pass=1, min accuracy=0.9, and turning Polish to OFF.

- 1. Primer IDs must be specified using the suffix _5p to indicate 5' cDNA primers and the suffix _3p to indicate 3' cDNA primers. The 3' cDNA primer should not include the Ts and is written in reverse complement.
- 2. Below are four example primer sets. The first is unbarcoded, the second has barcodes (shown in lower case) adjacent to the 3' primer.

Example 1: The Iso-Seq cDNA Primer primer set, included with the SMRT Link installation.

Users following the standard Iso-Seq Express protocol **without** multiplexing, or running a Data Set that has **already** been demultiplexed (either using Run Design or the SMRT® Analysis application) should use this default option.

```
>IsoSeq_5p
GCAATGAAGTCGCAGGGTTGGG
>IsoSeq_3p
GTACTCTGCGTTGATACCACTGCTT
```

Example 2: The Iso-Seq 12 Barcoded cDNA Primers set, included with the SMRT Link installation.

Users using barcoded cDNA primers listed in the Appendix 3 Recommended barcoded NEBNext single cell cDNA PCR primer and IsoSeg Express cDNA PCR primer sequences section of the document

Procedure & checklist - Preparing Iso-Seq® libraries using SMRTbell Prep Kit 3.0, should select this option.

```
>bc1001_5p
CACATATCAGAGTGCGGCAATGAAGTCGCAGGGTTGGGG
>bc1002_5p
ACACACAGACTGTGAGGCAATGAAGTCGCAGGGTTGGGG
```

(There are a total of 24 sequence records, representing 12 pairs of F/R barcoded cDNA primers.)

Example 3: An example of a custom cDNA primer set. 4 tissues were multiplexed using barcodes on the 3' end only.

```
>IsoSeq_5p
GCAATGAAGTCGCAGGGTTGGG
>dT_BC1001_3p
AAGCAGTGGTATCAACGCAGAGTACCACATATCAGAGTGCG
>dT_BC1002_3p
AAGCAGTGGTATCAACGCAGAGTACACACAGACTGTGAG
>dT_BC1003_3p
AAGCAGTGGTATCAACGCAGAGTACACACATCTCGTGAGAG
>dT_BC1004_3p
AAGCAGTGGTATCAACGCAGAGTACCACGCACACACGCGCG
```

Example 4: Special Handling for the TeloPrime cDNA Kit

The Lexogen TeloPrime cDNA kit contains As in the 3' primer that **cannot** be differentiated from the polyA tail. For best results, remove the As from the 3' end as shown below:

```
>TeloPrimeModified_5p
TGGATTGATATGTAATACGACTCACTATAG
>TeloPrimeModified_3p
CGCCTGAGA
```

- 3. Use the --isoseq mode. Note that this **cannot** be combined with the --guess **option**.
- 4. The output will be only different pairs with a 5p and 3p combination:

```
demux.5p--tissue1_3p.bam
demux.5p--tissue2_3p.bam
```

The --isoseq parameter set is very conservative for removing any spurious and ambiguous calls, and guarantees that only proper asymmetric (barcoded) primer are used in downstream analyses. Good libraries reach >75% CCS reads passing the Demultiplex Barcodes filters.

BAM tags

In SMRT Link v12.0, LB and SM tags are set by the user in Run Design. The SM tag can also be set in Demultiplex Barcodes in SMRT Analysis.

Non-demultiplex case:

- LB: Well Sample Name.
- SM: Bio Sample Name.

Multiplexed case, BAM pre-demultiplexing:

- LB: Well Sample Name.
- SM: Tag removed.

Multiplexed case, BAMs post-demultiplexing:

- LB: Well Sample Name for all child barcode BAMs.
- SM: Each individual Bio Sample Name for the specific barcode.
- BC: Barcode sequence or hyphenated barcode sequences of the pair.
- DS: Appends barcode information used in demultiplexing: BarcodeFile, BarcodeHash, BarcodeCount, BarcodeMode, BarcodeQuality.
- · Example read group header after demultiplexing:

```
ID:66d5a6af/3--3
PL:PACBIO
DS:READTYPE=SUBREAD;
   Ipd:CodecV1=ip;
   PulseWidth:CodecV1=pw;
  BINDINGKIT=101-500-400;
   SEQUENCINGKIT=101-427-800;
   BASECALLERVERSION=5.0.0;
   FRAMERATEHZ=100.000000;
   BarcodeFile=Sequel 16 barcodes v3.barcodeset.xml;
   BarcodeHash=f2b1fa0b43eb6ccbb30749883bb550e3;
   BarcodeCount=16;
   BarcodeMode=Symmetric;
   BarcodeQuality=Score
PU:m54010 200212 162236
SM:MySampleName
PM:SEOUEL
BC: ACAGTCGAGCGCTGCGT
```

export-datasets

The export-datasets tool takes one or more PacBio Data Set XML files and packages all contents (including index files and supplemental Data Sets) into a single ZIP archive. Data Set resources, such as BAM files, are reorganized and renamed to flatten the directory structure, avoid redundant file writes, and convert all resource paths from absolute paths to relative paths. Where multiple Data Sets are provided, the contents of each is nested in a directory named after the UniqueId attribute in the XML.

The resulting archive is primarily intended to be directly imported into SMRT Link using the Data Management interface, but it may also be unpacked manually and used on the command line.

Usage

export-datasets [options] <dataset>...

Options	Description
-o,output	Name of output ZIP file. (Default = datasets_ <timestamp>.zip)</timestamp>
keep-parent-ref	Keeps the reference to the parent Data Set when archiving a demultiplexed child Data Set.
no-scraps	Excludes the scraps.bam file if present in the XML file.
-h,help	Displays help information and exits.
log-file	Writes the log to a file. (Default = stderr)
log-level	Specifies the log level; values are [ERROR, DEBUG, INFO, WARN]. (Default = WARN)
logback	Override all logger configuration using a specified logback.xml file.
log2stdout	If True, log output is displayed to the console. (Default = False)
debug	Alias for setting the log level to DEBUG. (Default = False)
quiet	Alias for setting the log level to ERROR. (Default = False)
verbose	Alias for setting the log level to INFO. (Default = False)

Input files

· One or more PacBio Dataset XML files.

Output file

One output ZIP file.

Examples

mapped.alignmentset.xml

```
export-datasets m64001_200704_012345.subreadset.xml
export-datasets sample1.consensusreadset.xml sample2.consensusreadset.xml
\sample3.consensusreadset.xml -o barcoded_ccs.zip
export-datasets /opt/smrtlink/jobs/0000/0000001/0000001234/outputs/
```

export-job

The export-job tool packages a SMRT Link Analysis job for export to another system, usually for reimportation into another SMRT Link instance. All internal paths in job output files are converted from absolute to relative paths, and many of the internal details of the Cromwell workflows are omitted. The export is **not** a complete record of the job, but rather a collection of job output files and metadata.

Note that export-job will include any external Data Sets referenced in output Data Sets inside the job, for example ReferenceSets associated with mapped Data Sets, or BarcodeSets associated with demultiplexed Data Sets. However, these Data Sets will **not** be imported along with the job. The exported job does **not** include the input reads used to run the job; these may be exported separately using the export-datasets tool.

Important: Only SMRT Link v10.0 or later generates the necessary metadata files for export-job to save a full record of job execution. Jobs created with older versions of SMRT Link will still be archived, but the metadata will be empty and/or incorrect.

Usage

export-job [options] <job_dir>

Options	Description	
<job_dir></job_dir>	Path to a SMRT Link job directory.	
-o,output	Name of output ZIP file. (Default = job_ <timestamp>.zip)</timestamp>	
-h,help	Displays help information and exits.	
log-file	Writes the log to a file. (Default = stderr)	
log-level	Specifies the log level; values are [ERROR, DEBUG, INFO, WARN]. (Default = WARN)	
logback	Override all logger configuration using a specified logback.xml file.	
log2stdout	If True, log output is displayed to the console. (Default = False)	
debug	Alias for setting the log level to DEBUG. (Default = False)	
quiet	Alias for setting the log level to ERROR. (Default = False)	
verbose	Alias for setting the log level to INFO. (Default = False)	

Input

A path to a job directory.

Output file

One output ZIP file.

Examples

export-job /path/to/smrtlink/jobs-root/0000/0000000/000000860 -o job860.zip

To reimport on another system:

pbservice import-job job860.zip

gcpp

gcpp is a variant-calling tool provided by the GCpp package which provides several variant-calling algorithms for PacBio sequencing data.

Usage

```
gcpp -j8 --algorithm=arrow \
    -r lambdaNEB.fa \
    -o variants.gff \
    aligned subreads.bam
```

This example requests variant-calling, using 8 worker processes and the Arrow algorithm, taking input from the file aligned_subreads.bam, using the FASTA file lambdaNEB.fa as the reference, and writing output to variants.gff.

A particularly useful option is --referenceWindow/-w; which allows the variant-calling to be performed exclusively on a **window** of the reference genome.

Input files

- A sorted file of reference-aligned reads in PacBio's standard BAM format.
- A FASTA file that follows the PacBio FASTA file convention. If specifying an input FASTA file, a FASTA index file (.fai) with the same name and path is **required**. If the .fai file is not supplied, gcpp exits and displays an error message.

Note: The --algorithm=arrow option requires that certain metrics be in place in the input BAM file. It requires per-read SNR metrics, and the per-base PulseWidth metric for Sequel data.

The selected algorithm will stop with an error message if any features that it requires are unavailable.

Output files

Output files are specified as comma-separated arguments to the $-\circ$ flag. The file name extension provided to the $-\circ$ flag is meaningful, as it determines the output file format. For example:

```
gcpp aligned subreads.bam -r lambda.fa -o myVariants.gff,myConsensus.fasta
```

will read input from aligned_subreads.bam, using the reference lambda.fa, and send variant call output to the file myVariants.gff, and consensus output to myConsensus.fasta.

The file formats currently supported (using extensions) are:

- .gff: PacBio GFFv3 variants format; convertible to BED.
- .vcf: VCF 4.2 variants format (that is compatible with v4.3.)
- .fasta: FASTA file recording the consensus sequence calculated for each reference contig.

• .fastq: FASTQ file recording the consensus sequence calculated for each reference contig, as well as per-base confidence scores.

Options	Description
-j	Specifies the number of worker processes to use.
algorithm=	Specifies the variant-calling algorithm to use; values are plurality, arrow and poa. (Default = arrow)
-r	Specifies the FASTA reference file to use.
-0	Specifies the output file format; values are .gff, .vcf, .fasta, and .fastq.
maskRadius	When using the <code>arrow</code> algorithm, setting this option to a value N greater than 0 causes <code>gcpp</code> to pass over the data a second time after masking out regions of reads that have >70% errors in $2*N+1$ bases. This setting has little to no effect at low coverage, but for high-coverage datasets (>50X), setting this parameter to 3 may improve final consensus accuracy. In rare circumstances, such as misassembly or mapping to the wrong reference, enabling this parameter may cause worse performance.
minConfidence MINCONFIDENCE -q MINCONFIDENCE	Specifies the minimum confidence for a variant call to be output to variants.{gff,vcf} (Default = 40)
minCoverage MINCOVERAGE -x MINCOVERAGE	Specifies the minimum site coverage for variant calls and consensus to be calculated for a site. (Default = 5)

Available algorithms

At this time there are three algorithms available for variant calling: plurality, poa and arrow.

- plurality is a simple and very fast procedure that merely tallies the most frequent read base or bases found in alignment with each reference base, and reports deviations from the reference as potential variants. This approach is prone to insertion and deletion errors.
- poa uses the partial order alignment algorithm to determine the consensus sequence. It is a heuristic algorithm that approximates a multiple sequence alignment by progressively aligning sequences to an existing set of alignments.
- arrow uses the per-read SNR metric and the per-pulse pulsewidth metric as part of its likelihood model.

Confidence values

The ${\tt arrow}$ and ${\tt plurality}$ algorithms make a confidence metric available for every position of the consensus sequence. The confidence should be interpreted as a phred-transformed posterior probability that the consensus call is incorrect; such as:

$$QV = -10\log_{10}(p_{err})$$

 ${\tt gcpp}$ clips reported QV values at 93; larger values **cannot** be encoded in a standard FASTQ file.

Chemistry specificity

The --algorithm=arrow parameter is trained per-chemistry. arrow identifies the sequencing chemistry used for each run by looking at metadata contained in the input BAM data file. This behavior can be overridden by a command-line option.

When multiple chemistries are represented in the reads in the input file, the Arrow will model reads appropriately using the parameter set for its chemistry, thus yielding optimal results.

Genome Assembly

The Genome Assembly application generates *de novo* assemblies using HiFi reads. The application is fast, produces contiguous assemblies, and is suitable for genomes of any size.

The Genome Assembly application is powered by the IPA HiFi genome assembler and includes the following features:

- Separates haplotypes during assembly using a novel phasing stage (Nighthawk).
- Polishes the contigs with phased reads using Racon.
- Improves haplotype separation using the purge dups tool.

Workflow of the Genome Assembly application

Analysis steps are highly optimized to produce assemblies of large genomes efficiently.















The workflow consists of seven stages:

- 1. Sequence database construction.
- 2. Fast overlap computation using the Pancake tool.
- 3. A dedicated phasing stage using the Nighthawk tool.
- 4. Filtering chimeras and residual repeats.
- 5. Layout based on the string graph.
- 6. Polishing using the Racon tool.
- 7. Purging haplotype duplicates from the primary assembly using the third-party tool purge dups.

The workflow accepts HiFi XML Data Sets as input.

IPA HiFi genome assembler

- · Scales well on a cluster.
- The workflow has an embedded downsampling feature:
 - If the genome size and the desired coverage are specified, the initial stage (sequence database construction) downsamples the input Data Set to the desired coverage.
 - Otherwise, the full coverage is used.

Usage

The Genome Assembly application is run using the <code>pbcromwell run</code> command, with the <code>pb_assembly_hifi</code> parameter to specify the application. See "pbcromwell" on page 81 for details.

To view information on the available Genome Assembly options, enter:

```
pbcromwell show-workflow-details pb assembly hifi
```

The **minimum** command needed to run the workflow requires the input and the number of threads. The following example uses 16 threads:

```
pbcromwell run pb_assembly_hifi -e <input.xml> --nproc 16
```

The following example performs an assembly using an input XML Data Set, and uses all default settings, including 1 CPU:

```
pbcromwell run pb assembly hifi -e <input.consensusreadset.xml>
```

Note: The default options for this workflow are equivalent to the following command:

```
pbcromwell run pb_assembly_hifi \
-e <input.consensusreadset.xml> \
--task-option reads=None \
--task-option ipa2_genome_size=0 \
--task-option ipa2_downsampled_coverage=0 \
--task-option ipa2_advanced_options="" \
--task-option ipa2_run_polishing=True \
--task-option ipa2_run_phasing=True \
--task-option ipa2_run_purge_dups=True \
--task-option ipa2_ctg_prefix="ctg." \
--task-option ipa2_reads_db_prefix="reads" \
--task-option ipa2_cleanup_intermediate_files=True \
--task-option dataset_filters="" \
--task-option filter_min_qv=20 \
--nproc 8
```

The default options for this workflow should work well for any genome types.

If the assembly is run on a single local node with high CPU count, such as 64 cores, we recommend that the job submission for pbcromwell is configured so that it uses 4 concurrent jobs and 16 threads per job.

We found this to be more efficient than using 64 threads and 1 concurrent job, as many steps are very data I/O-dependent.

You can apply a similar principle for compute environments with more or fewer cores. For example, for a machine with 80 cores, one can use 20 threads and 4 concurrent jobs.

Genome Assembly parameters input files

Option	Default value	Description
-e,eid_ccs	NONE	Optional parameter, required iftask-option reads <input/> is not specified. This is a SMRT Link-specific input parameter and supports only PacBio Consensusreadset XML files as input.
task-option reads	NONE	Optional parameter, required if <code>-e <input/></code> is not specified. Supports multiple input formats: FASTA, FASTQ, BAM, XML, FOFN and gzipped versions of FASTA/FASTQ.
task-option ipa2_genome_size	0	The approximate number of base pairs expected in the genome. This is used only for downsampling; if the value is \leq 0, downsampling is disabled. Note : It is better to slightly overestimate rather than underestimate the genome length to ensure good coverage across the genome.
task-option ipa2_downsampled_coverage	0	The input Data Set can be downsampled to a desired coverage, provided that both the ipa2_downsampled_coverage and ipa2_genome_size options are specified and >0.
		Downsampling applies to the entire assembly process, including polishing.
		This parameter selects reads randomly, using a fixed random seed for reproducibility.
task-option ipa2_advanced_options	NONE	A semicolon-separated list of KEY=VALUE pairs. New line characters are not accepted. (These are described later in this document.)
task-option ipa2_run_polishing	TRUE	Enables or disables the polishing stage of the workflow. Polishing can be disabled to perform fast draft assemblies.
task-option ipa2_run_phasing	TRUE	Enables or disables the phasing stage of the workflow. Phasing can be disabled to assemble haploid genomes, or to perform fast draft assemblies.
task-option ipa2_run_purge_dups	TRUE	Enables or disables identification of "duplicate" alternate haplotype contigs which may be assembled in the primary contig file, and moves them to the associate contig (haplotig) file.
task-option ipa2_ctg_prefix	.ctg	The prefix used to label the output generated contigs.
task-option ipa2_reads_db_prefix	reads	The prefix of the sequence and seed databases which will be used internally for assembly.
task-option ipa2_cleanup_intermediate _files	TRUE	Removes intermediate files from the run directory to save space.
task-option dataset_filters	NONE	(General pbcromwell option) A semicolon-separated (not comma-separated) list of other filters to add to the Data Set.
task-option filter_min_qv	20	(General pbcromwell option) Phred-scale integer QV cutoff for filtering HiFi reads. The default for all applications is 20 (QV 20), or 99% predicted accuracy.
task-option downsample_factor	0	Downsampling factor applied directly to the input Data Set. This parameter is not related to ipa2_downsampled_coverage.
task-option mem_scale_factor	8	Controls the amount of requested memory for individual Cromwell tasks in the workflow. The default value of 8 is good for larger genomes, but it may be too much memory for smaller genomes.

Option	Default value	Description
config	NONE	(General pbcromwell option) Java configuration file for running Cromwell.
nproc	1	(General pbcromwell option) Number of processors, (except per task).

- *.bam file containing PacBio data.
- *.fasta or *.fastq file containing PacBio data.
- *.xml file containing PacBio data.
- *.fofn files with file names of files containing PacBio data.

Output files

- final_purged_primary.fasta file containing assembled primary contigs.
- final_purged_haplotigs.fasta file containing assembled haplotigs.

Advanced parameters

Advanced parameters should be rarely modified. For the special cases when that is required, advanced parameters are documented below.

Advanced parameters specified on the command line:

- Are in the form of key = value pairs.
- Each pair is separated by a semicolon (;) character.
- The full set of advanced parameters is surrounded by one set of double quotes.
- The specified value of a parameter overwrites the default options for that key. All desired options of that parameter must be explicitly listed, not just the ones which should change from the default.
- Setting an empty value **clears** the parameter; it does **not** reset the value back to default.

Example

--task-option ipa2_advanced_options="config_seeddb_opt=-k 28;config_block_size=2048"

Complete list of available advanced parameters and default values:

Advanced parameters	Default value	Description
config_genome_size	0	The approximate number of base pairs expected in the genome, used to determine the coverage cutoff. This is only used for downsampling; 0 turns downsampling off.
		Note : It is better to slightly overestimate rather than underestimate the genome length to ensure good coverage across the genome.
config_coverage	0	The input Data Set can be downsampled to a desired coverage, provided that both the Downsampled coverage and Genome Length parameters are specified and above 0.
		Downsampling applies to the entire assembly process, including polishing. This feature selects reads randomly, using a fixed random seed for reproducibility.
config_polish_run	1	Enables or disables the polishing stage of the workflow. Polishing can be disabled to perform fast draft assemblies. 0 disables this feature; 1 enables it.
config_phase_run	1	Enables or disables the phasing stage of the workflow. Phasing can be disabled to assemble haploid genomes, or to perform fast draft assemblies. 0 disables this feature; 1 enables it.
config_purge_dups_run	1	Enables or disables the purge_dups stage of the workflow. 0 disables this feature; 1 enables it.
config_autocomp_max_cov	1	If enabled, the maximum allowed overlap coverage at either the 5' or the 3' end of every read is automatically determined based on the statistics computed from the overlap piles. This value is appended to the config ovl_filter opt value internally, and supersedes the manually specifiedmax-cov andmax-diff values of that parameter. These options are used to determine potential repeats and filter out those reads before the string graph is constructed. 0 disables this feature; 1 enables it.
config_block_size	4096	The overlapping process is performed on pairs of blocks of input sequences, where each block contains the number of sequences which crop up to this size (in Mbp). Note : The number of pairwise comparisons grows quadratically with the number of blocks (meaning more cluster jobs), but also the larger the block size the more resources are required to execute each pairwise comparison.
config_existing_db_prefix	NONE	Allows injection of an existing SeqDB, so that one doesn't have to be built from scratch. The provided existing DB is symbolically linked and used for assembly. (This option is intended for debugging purposes.)

Advanced parameters	Default value	Description
config_ovl_filter_opt	max-diff 80max-cov 100min-cov 2bestn 10min-len 4000gapFiltminDepth 4	Overlap filter options. gapFilt - Enables the chimera filter, which analyzes each overlap pile, and determines whether a pread is chimeric based on the local coverage across the pread. minDepth - Option for the chimera filter. The chimera filter is ignored when a local region of a read has
	idt-stage2 98	coverage lower than this value. The other parameters are:
		min-cov - Minimum allowed coverage at either the 5' or the 3' end of a read. If the coverage is below this value, the read is blacklisted and all of the overlaps it is incident with are ignored. This helps remove potentially chimeric reads.
		max-cov - Maximum allowed coverage at either the 5' or the 3' end of a read. If the coverage is above this value, the read is blacklisted and all of the overlaps it is incident with are ignored. This helps remove repetitive reads which can make tangles in the string graph. Note that this value is a heuristic which works well for ∼30x seed length cutoff. If the cutoff is set higher, we advise that this value be also increased. Alternatively, using the autocompute max cov option can automatically estimate the value of this parameter, which can improve contiguity (for example, in cases when the input genome size or the seed coverage were overestimated).
		max-diff - Maximum allowed difference between the coverages at the 5' and 3' ends of any particular read. If the coverage is above this value, the read is blacklisted and all of the overlaps it is incident with are ignored. If the autocompute _max_cov option is used, then the same computed value is supplied to this parameter as well.
		bestn - Keep at most this many overlaps on the 5' and the 3' side of any particular read.
		min-len - Filter overlaps where either A-read or the B-read are shorter than this value.
		idt-stage2 - Filter overlaps with identity below 98%high-copy-sample-rate - Controls the downsampling of reads from high copy elements to the expected coverage determined by maxCov*rate, where rate is the value of this parameter. If rate is 0, then these high coverage reads are discarded.
config_ovl_min_idt	98	The final overlap identity threshold. Applied during the final filtering stage, right before the overlaps are passed to the layout stage.
config_ovl_min_len	1000	The minimum length of either A-read or a B-read to keep the overlap. Applied during the final filtering stage, right before the overlaps are passed to the layout stage.
config_ovl_opt	one-hit-per- target min-idt 96	Overlapping options for the pancake overlapping tool. The options set by this parameter here are passed directly to pancake. For details on pancake options, use pancake -h. The defaults used here are:one-hit-per-target which keeps only the best hit in case there are multiple possible overlaps between a pair of reads (tandem repeats); andmin-idt 96 which will filter out any overlap with identity lower than 96%.
config_phasing_opt	NONE	Options for the phasing tool nighthawk. The options set by this parameter are passed directly to nighthawk. For details on nighthawk options, use nighthawk -h.

Advanced parameters	Default value	Description
config_phasing_split_opt	split-type noverlaps limit 3000000	Options that control the chunking of the phasing jobs, and through that regulate the time and memory consumption of each individual chunk.
		The defaults are:split-type noverlaps which splits the chunks by the number of overlaps; andlimit 3000000 which allow at most approximately 3 million overlaps per chunk.
		Empirically, the current defaults keep the maximum memory consumption (RSS) of the phasing jobs under 4 GB per chunk.
config_seeddb_opt	-k 28 -w 120	Options to control the seed computation. These options are passed directly to the pancake seeddb command.
	space 1	Defaults: -k 28 is the k-mer size of 28 bp; -w 120 is the minimizer window size of 120 bp; andspace 1 specifies the spacing for spaced seed construction, with 1 gap in between every two bases of the seed.
		For more details on these and other options, use pancake seeddb -h.
config_seqdb_opt	compression 1	Options to control the construction of the sequence database. These options are passed directly to the pancake seqdb command.
		Current default iscompression 1 which turns on the 2-bit encoding compression of the sequences.
		For more details on these and other options, use pancake seqdb -h.
config_use_hpc	0	This parameter enables (1) or disables (0) an experimental Homopolymer Compression feature.
		If this feature is enabled, the overlaps are computed from homopolymer-compressed sequences. The layout stage is somewhat slower because the sequences have to be aligned to determine the correct homopolymer-expanded coordinates.
config_use_seq_ids	1	This feature is mostly useful for debugging purposes. If 0 is specified, then the overlaps contain original sequence names instead of their numerical IDs.
		The default of ${\tt 1}$ uses the numerical IDs to represent reads, which uses memory much more efficiently.
config_purge_map_opt	min-map-len 1000 min-idt 98.0 bestn 5	This option is used to control the mapping of the reads to contigs for the purge dups tool. The mapper used is pancake, and the options set by this parameter are used directly by pancake. For details on pancake options, use pancake -h.
		Optionmin-map-len 1000 removes any alignment which did not span more than 1000 bp during the mapping process;min-idt 98.0 removes any alignment with identity below 98.0%, andbestn 5 keeps at most 5 top scoring alignments for each query read (one primary alignment and at most 4 secondary alignments).

Advanced parameters	Default value	Description
config_purge_dups_calcuts	NONE	The third-party tool purge_dups can accept user-defined cutoffs for purging. On some genomes, the automated computation of the cutoffs in purge_dups can result in suboptimal values, and in this case a user can specify them manually.
		This option is passed directly to the purge_dups_calcuts tool. For details on the possible values that can be passed to this tool, use ipa_purge_dups_calcuts without parameters.
		Relevant parameters include:
		-1 INT Lower bound for read depth.
		-m INT Transition between haploid and diploid.-u INT Upper bound for read depth.
<pre>config_purge_dups_get_seq s</pre>	NONE	User-specified options for the third party purge_dups_get_seqs tool used as part of the purge_dups_workflow. This option is passed directly to the purge_dups_get_seqs tool.
		For details on the possible values that can be passed to this tool, use <code>ipa_purge_dups_get_seqs</code> without parameters.
		Relevant parameters include:
		<pre>-e BOOL Only remove sequences at the ends of a contig (Default = FALSE).</pre>
		-s BOOL Split contigs (Default = FALSE) .
		-p STR Prefix of output files (Default = NULL).
		-c BOOL Keep high coverage contigs in the primary contig set (Default = FALSE).
		-a BOOL Do not add prefix to haplotigs (Default = FALSE).
		-g INT Maximum gap size between duplications (Default = 10K).
		-1 INT Minimum primary contig length (Default = 10K).
		-m INT Minimum ratio of remaining primary contig length to the original contig length (Default = 0.05).
<pre>config_m4filt_high_copy_s ample_rate</pre>	1.0	This option is passed to the <code>high-copy-sample-rate</code> parameter of the overlap filter, which controls the downsampling of reads from high copy elements to the expected coverage determined by <code>maxCov*rate</code> , where <code>rate</code> is the value of this parameter. If <code>0</code> , then these high coverage reads are discarded.
		Note : This parameter supersedes the config_ovl_filter_opt options.
config_max_polish_block_m b	100	During the polishing stage, contigs are grouped into chunks of approximate size specified by this parameter (in megabases). Each chunk is processed separately and in parallel (depending on the system configuration).
config_layout_opt	NONE	This value is passed directly to the legacy assembly layout stage, if the legacy layout stage is used. To see a list of valid options for this parameter, enter <code>ipa2_ovlp_to_graph</code> -h on the command line.
		This parameter is not currently passed to the new layout implementation.

Advanced parameters	Default value	Description
config_use_legacy_layout	0	Legacy layout implementation is used if this value is set to 1. Otherwise, the new layout implementation is used by default.
		Important note: The legacy implementation requires the Networkx Python package to be installed, which is no longer a necessary prerequisite of the Genome Assembly application.

HiFiViral SARS-CoV-2 Analysis

Use this application to analyze multiplexed samples sequenced with the HiFiViral SARS-CoV-2 Kit. For **each** sample, this analysis provides:

- Consensus sequence (FASTA).
- · Variant calls (VCF).
- HiFi reads aligned to the reference (BAM).
- Plot of HiFi read coverage depth across the SARS-CoV-2 genome.

Across all samples, this analysis provides:

- Job summary table including passing sample count at 90 and 95% genome coverage.
- Sample summary table including, for each sample: Count of variable sites, genome coverage, read coverage, and probability of multiple strains, and other metrics.
- Plate QC graphical summary of performance across samples in assay plate layout.
- Plot of HiFi read depth of coverage for all samples.

Notes:

- The application accepts HiFi reads (BAM format) as input. HiFi reads are reads generated with CCS analysis that have a quality value equal to or greater than Phred-scaled Q20.
- This application is for SARS-CoV-2 analysis only and is not recommended for other viral studies. The Wuhan reference genome is included with SMRT Link and used by default, but advanced users may specify other reference genomes. We have not tested the application with reference genomes other than the Wuhan reference genome.
- The application is intended to identify variable sites and call a single consensus sequence per sample. The output consensus sequence is produced based on the dominant variant observed. Minor variant information that passes through a default threshold may be encoded in the raw VCF, but does **not** get propagated into the consensus sequence FASTA.
- The HiFiViral SARS-CoV-2 Analysis application can be run using the Auto Analysis feature available in Run Design. This feature allows users to complete all necessary analysis steps immediately after sequencing without manual intervention. The Auto Analysis workflow includes CCS, Demultiplex Barcodes, and HiFiViral SARS-CoV-2 Analysis.

HiFiViral SARS-CoV-2 application workflow

- 1. Process the reads using the \min tool to trim the probe arm sequences.
- 2. Align the reads to the reference genome using pbmm2.
- 3. Call and filter variants using bcftools, generating the raw variant calls in VCF file format. Filtering in this step removes low-quality calls (less than Q20), and normalizes indels.

4. Filter low-frequency variants using vcfcons and generate a consensus sequence by injecting variants into the reference genome. At each position, a variant is called only if both the base coverage exceeds the minimum base coverage threshold (Default = 4) and the fraction of reads that support this variant is above the minimum variant frequency threshold (Default = 0.5). See here for details.

Preparing input data for the HiFiViral SARS-CoV-2 Analysis application

- Run the Demultiplex Barcodes cromwell workflow, where the input to that application are HiFi reads, and the primers are multiplexed barcode primers. See "Demultiplex Barcodes" on page 24 for details. If HiFi reads have not been generated on the instrument, run CCS analysis first. See "ccs" on page 8 for details.
 - Provide the proper barcode sequences:
 Barcoded M13 Primer Plate.
 - Use the task option lima_symmetric_barcodes=false. (The barcode pairs are asymmetric.)
 - Provide the correctly-formatted barcode pair-to-Bio Sample CSV file.
 For details, see the --task-option sample_wells_csv option in the table further down this page.

Input files

- movie.consensusreadset.xml: Previously-demultiplexed HiFi reads, packaged as separate BAM files wrapped in an XML Data Set. (See "Preparing input data for the HiFiViral SARS-CoV-2 Analysis application" on page 52 for details.)
- sars cov2.referenceset.xml: The Wuhan reference genome.
- HiFiViral_SARS-CoV-2_Enrichment_Probes.barcodeset.xml: Dataset XML specifying the probe sequence file in FASTA format.
- [Optional] Plate QC csv: Four-column CSV file that includes barcode, biosample, plateID and wellID.
 To specify this optional file, add the following option to pbservice:

 -task-option sample wells csv=<path to CSV file>

Output files

- pb_sars_cov2_kit.probe_counts_zip: Zipped TSV files with probe counts, per sample.
- pb sars cov2 kit.variants csv: CSV variant calls for all samples.
- pb_sars_cov2_kit.vcf_zip: Zipped VCF files containing the final variant calls, per sample.
- pb_sars_cov2_kit.raw_vcf_zip: Zipped VCF files containing the raw variant calls, per sample.
- pb_sars_cov2_kit.fasta_zip: Zipped final consensus sequences, by sample, in FASTA format. This is a single consensus sequence with Ns for each sample.
- pb_sars_cov2_kit.frag_fasta_zip: Zipped file of consensus sequences, split on Ns, in FASTA format, by sample.

- pb_sars_cov2_kit.mapped_zip: Zipped BAM files containing the output from mapping HiFi reads to the reference genome, by sample.
- samples.consensus_mapped.bam.zip: Zipped BAM files containing the output from mapping consensus FASTA files to the reference genome, by sample.
- samples.coverage.png.zip: Zipped per-sample coverage graphs in png format.
- hifi_reads.fastq.zip: Zipped file of per-sample trimmed HiFi reads in FASTQ format.
- sample_summary.csv: Sample Summary file in CSV format, with one row per sample.

Options	Description
task-option sample_wells_csv	Specifies a 4-column CSV file used to generate the Plate QC Report, which displays analysis results for each sample in the assay plate. The CSV file must contain barcode pairs, Bio Sample name, Plate IDs, and Well IDs. The report is useful for diagnosing sample issues based on plate location. (Default = None)
task-option min_coverage	Specifies the minimum read coverage. Below this value, the consensus sequence will be set to Ns and no variants are called. (Default = 4)
task-option min_alt_freq	Specifies that only variants whose frequency is greater than this value are reported. This frequency is determined based on the read depth (DP) and allele read count (AD) information in the VCF output file. We recommend using the default value to properly call the dominant alternative variant while also filtering out potential artifacts. (Default = 0.5)
task-option min_bq	Specifies that reads with barcode scores below this minimum value are not included in analysis. (Default = 80)
task-option mimux_overrides	Specifies additional options to pass to the mimux preprocessing tool for trimming and filtering reads by probe sequences. Options should be entered in space-separated format.
	Available options include:
	max-len: Specifies the maximum sequence length. (Default = 800)
	same: Specifies that only reads with arms sequences from the same probe are used.
task-option probes_fasta	Specifies a FASTA file containing probes sequences if using probes other than those supplied with the HiFiViral SARS-CoV-2 kit.

Running the SARS-CoV-2 Analysis application

```
pbcromwell run pb_sars_cov2_kit \
-e <movie.consensusreadset.xml> \
-e $SMRT_ROOT/current/bundles/smrtinub/current/private/pacbio/barcodes/HiFiViral_SARS-CoV-2_Enrichment_Probes.barcodeset.xml \
-e eid_ref_dataset_2:$SMRT_ROOT/current/bundles/smrtinub/current/private/pacbio/canneddata/referenceset/SARS-CoV-2/sars_cov2.referenceset.xml
--task-option min_alt_freq=0.5 \
--task-option min_bq=80 \
--task-option mimux_overrides="--max-len=800 --same" \
--task-option sample_wells_csv=None \
--config cromwell.conf \
--nproc 8
```

ipdSummary

The ipdsummary tool detects DNA base-modifications from kinetic signatures. It is part of the kineticsTool package.

kineticsTool loads IPDs observed at each position in the genome, compares those IPDs to value expected for unmodified DNA, and outputs the result of this statistical test. The expected IPD value for unmodified DNA can come from either an in-silico control or an amplified control. The in-silico control is trained by PacBio and shipped with the package. It predicts the IPD using the local sequence context around the current position. An amplified control Data Set is generated by sequencing unmodified DNA with the same sequence as the test sample. An amplified control sample is usually generated by whole-genome amplification of the original sample.

Modification detection

The basic mode of kineticsTool does an independent comparison of IPDs at each position on the genome, for each strand, and outputs various statistics to CSV and GFF files (after applying a significance filter).

Modifications identification

kineticsTool also has a Modification Identification mode that can decode multi-site IPD "fingerprints" into a reduced set of calls of specific modifications. This feature has the following benefits:

- Different modifications occurring on the same base can be distinguished; for example, 6mA and 4mC.
- The signal from one modification is combined into one statistic, improving sensitivity, removing extra peaks, and correctly centering the call.

Algorithm: Synthetic control

Studies of the relationship between IPD and sequence context reveal that most of the variation in mean IPD across a genome can be predicted from a 12-base sequence context surrounding the active site of the DNA polymerase. The bounds of the relevant context window correspond to the window of DNA in contact with the polymerase, as seen in DNA/polymerase crystal structures. To simplify the process of finding DNA modifications with PacBio data, the tool includes a pre-trained lookup table mapping 12-mer DNA sequences to mean IPDs observed in C2 chemistry.

Algorithm: Filtering and trimming

kineticsTool uses the Mapping QV generated by pbmm2 and stored in the cmp.h5 or BAM file (or AlignmentSet) to **ignore** reads that are not confidently mapped. The default minimum Mapping QV required is 10, implying that pbmm2 has 90% confidence that the read is correctly mapped. Because of the range of read lengths inherent in PacBio data, this can be changed using the --mapQvThreshold option.

There are a few features of PacBio data that require special attention to achieve good modification detection performance. kineticsTool inspects the alignment between the observed bases and the reference sequence for an IPD measurement to be included in the analysis. The PacBio read sequence **must** match the reference sequence for k around the cognate base. In the current module, k=1. The IPD distribution at some locus can be thought of as a mixture between the "normal" incorporation process IPD, which is sensitive to the local sequence context and DNA modifications, and a contaminating "pause" process IPD, which has a much longer duration (mean >10 times longer than normal), but happen rarely (~1% of IPDs).

Note: Our current understanding is that pauses do **not** carry useful information about the methylation state of the DNA; however a more careful analysis may be warranted. Also note that modifications that drastically increase the roughly 1% of observed IPDs are generated by pause events. Capping observed IPDs at the global 99th percentile is motivated by theory from robust hypothesis testing. Some sequence contexts may have naturally longer IPDs; to avoid capping too much data at those contexts, the cap threshold is adjusted per context as follows:

```
capThreshold = max(global99, 5*modelPrediction,
percentile(ipdObservations, 75))
```

Algorithm: Statistical testing

We test the hypothesis that IPDs observed at a particular locus in the sample have longer means than IPDs observed at the same locus in unmodified DNA. If we have generated a Whole Genome Amplified Data Set, which removes DNA modifications, we use a case-control, two-sample t-test. This tool also provides a pre-calibrated "synthetic control" model which predicts the unmodified IPD, given a 12-base sequence context. In the synthetic control case we use a one-sample t-test, with an adjustment to account for error in the synthetic control model.

Usage

To run using a BAM input, and output GFF and HDF5 files:

```
ipdSummary aligned.bam --reference ref.fasta m6A,m4C --gff basemods.gff \ --csv h5 kinetics.h5
```

To run using <code>cmp.h5</code> input, perform methyl fraction calculation, and output GFF and CSV files:

```
ipdSummary aligned.cmp.h5 --reference ref.fasta m6A,m4C --methylFraction \ --gff basemods.gff --csv kinetics.csv
```

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Output options	Description
gff FILENAME	GFF format.
csv FILENAME	Comma-separated value format.
bigwig FILENAME	BigWig file format.

Input files

- A standard PacBio alignment file either AlignmentSet XML, BAM, or cmp.h5 - containing alignments and IPD information.
- Reference sequence used to perform alignments. This can be either a FASTA file or a ReferenceSet XML.

Output files

The tool provides results in a variety of formats suitable for in-depth statistical analysis, quick reference, and consumption by visualization tools. Results are generally indexed by reference position and reference strand. In all cases the strand value refers to the strand carrying the modification in the DNA sample. Remember that the kinetic effect of the modification is observed in read sequences aligning to the opposite strand. So reads aligning to the positive strand carry information about modification on the negative strand and vice versa, but the strand containing the putative modification is always reported.

 modifications.gff: Compliant with the GFF Version 3 specification. Each template position/strand pair whose probability value exceeds the probability value threshold appears as a row. The template position is 1-based, per the GFF specifications. The strand column refers to the strand carrying the detected modification, which is the opposite strand from those used to detect the modification. The GFF confidence column is a Phred-transformed probability value of detection.

The auxiliary data column of the GFF file contains other statistics which may be useful for downstream analysis or filtering. These include the coverage level of the reads used to make the call, and +/-20 bp sequence context surrounding the site.

modifications.csv: Contains one row for each (reference position, strand) pair that appeared in the Data Set with coverage at least x.
 x defaults to 3, but is configurable with the --minCoverage option. The reference position index is 1-based for compatibility with the GFF file in the R environment. Note that this output type scales poorly and is not recommended for large genomes; the HDF5 output should perform much better in these cases.

Output columns: In-silico control mode

Column	Description
refId	Reference sequence ID of this observation.
tpl	1-based template position.
strand	Native sample strand where kinetics were generated. 0 is the strand of the original FASTA, 1 is opposite strand from FASTA.
base	The cognate base at this position in the reference.
score	Phred-transformed probability value that a kinetic deviation exists at this position.
tMean	Capped mean of normalized IPDs observed at this position.
tErr	Capped standard error of normalized IPDs observed at this position (standard deviation/sqrt(coverage)).
modelPrediction	Normalized mean IPD predicted by the synthetic control model for this sequence context.
ipdRatio	tMean/modelPrediction.
coverage	Count of valid IPDs at this position.
frac	Estimate of the fraction of molecules that carry the modification.
fracLow	2.5% confidence bound of the frac estimate.
fracUpp	97.5% confidence bound of the frac estimate.

Output columns: Case control mode

Column	Description
refId	Reference sequence ID of this observation.
tpl	1-based template position.
strand	Native sample strand where kinetics were generated. 0 is the strand of the original FASTA, 1 is opposite strand from FASTA.
base	The cognate base at this position in the reference.
score	Phred-transformed probability value that a kinetic deviation exists at this position.
caseMean	Mean of normalized case IPDs observed at this position.
controlMean	Mean of normalized control IPDs observed at this position.
caseStd	Standard deviation of case IPDs observed at this position.
controlStd	Standard deviation of control IPDs observed at this position.
ipdRatio	tMean/modelPrediction.
testStatistic	T-test statistic.
coverage	Mean of case and control coverage.
controlCoverage	Count of valid control IPDs at this position.
caseCoverage	Count of valid case IPDs at this position.

isoseq3

The isoseq3 tool enables analysis and functional characterization of transcript isoforms for bulk and single-cell sequencing data generated on PacBio instruments. The analysis can be performed *de novo*, without a reference genome. If a reference genome is available, an optional collapse step to produce unique isoform files based on genomic coordinates is available. The input to isoseq3 tools should be HiFi (CCS) reads in BAM format.

Recommended commands for bulk Iso-Seq® Analysis

Command	Description	Output format	Recommended next step
lima	Remove cDNA primers.	fl.bam	isoseq3 refine
isoseq3 refine	Remove polyA tail and artificial concatemers.	flnc.bam	isoseq3 cluster
isoseq3 cluster	De novo isoform-level clustering.	unpolished.bam	pbmm2
pbmm2	Align to the genome.	aligned.sorted.bam	isoseq3 collapse
isoseq3 collapse	Collapse redundant transcripts based on exonic structures.	collapsed.gff	pigeon

Recommended commands for single-cell Iso-Seq® Analysis

Command	Description	Output format	Recommended next step
lima	Remove cDNA primers.	fl.bam	isoseq3 tag
isoseq3 tag	Extract UMIs and cell barcodes.	fl.tagged.bam	isoseq3 refine
isoseq3 refine	Remove polyA tail and artificial concatemers.	flnc.bam	isoseq3 correct
isoseq3 correct	Correct cell barcodes and tag reads that are real cells.	flnc.corrected.bam	isoseq3 bcstats
isoseq3 bcstats	Summarize barcode statistics for real/non-real cells.	bcstats.reports.tsv	isoseq3 dedup or isoseq3 groupdedup
isoseq3 dedup or isoseq3 groupdedup	Deduplicate UMIs.	dedup.bam	pbmm2
pbmm2	Align to the genome.	aligned.sorted.bam	isoseq3 collapse
isoseq3 collapse	Collapse redundant transcripts based on exonic structures.	collapsed.gff	pigeon

Usage

isoseq3 <tool>

Options	Description
-h,help	Displays help information and exits.
version	Displays program version number and exits

Typical bulk Iso-Seq workflow

1. Visualize primers, then remove primers and demultiplex:

```
cat primers.fasta
>5p
GCAATGAAGTCGCAGGGTTGGGG
>3p
GTACTCTGCGTTGATACCACTGCTT
```

lima movie.ccs.bam primers.fasta demux.bam --isoseq

See "Demultiplex Barcodes" on page 24 for details on the lima tool.

2. Identify and remove polyA tails; also remove artificial concatemers. The output are full-length, non-concatemer (FLNC) reads:

isoseq3 refine demux.5p--3p.bam primers.fasta flnc.bam --require-polya

3. Cluster FLNC reads at the isoform levels to generate consensus transcript isoform sequences. This generates unpolished.hq.bam and unpolished.hq.fasta.gz files, which are the high-quality (HQ) transcripts that should be analyzed further.

isoseq3 cluster flnc.bam unpolished.bam --use-qvs

4. (Optional) Map transcripts to the genome and collapse HQ transcripts based on genomic mapping:

```
pbmm2 align unpolished.bam reference.fasta aligned.sorted.bam --preset ISOSEQ --sort isoseq3 collapse --do-not-collapse-extra-5exons aligned.sorted.bam out.gff Or isoseq3 collapse --do-not-collapse-extra-5exons aligned.sorted.bam movie.ccs.bam out.gff
```

See "pbmm2" on page 90 for details.

refine command: Remove polyA and concatemers from full-length (FL) reads and generate full-length non-concatemer (FLNC) transcripts (FL to FLNC).

Usage

isoseq refine [options] <ccs.demux.bam|xml> <primer.fasta|xml> <flnc.bam|xml>

Inputs/outputs	Description
ccs.demux.bam xml	Input demultiplexed CCS reads BAM or ConsensusReadSet XML file. This is usually the output from running lima with theisoseq option, such as demux.5p3p.bam.
primer.fasta xml	Input primer FASTA or BarcodeSet XML file.
flnc.bam xml	Output FLNC BAM or ConsensusReadSet XML file.

Preprocessing	Description
min-polya-length	Specifies the minimum poly(A) tail length. (Default = 20)
require-polya	Requires reads to have a poly(A) tail and remove it.
min-rq	Specifies the minimum CCS read quality. (Default = -1, deactivated)

Options	Description
help, -h	Displays help information and exits.
version	Displays program version number and exits.
verbose, -v	Sets the verbosity level.
-j,num-threads	Specifies the number of threads to use when processing; 0 means autodetection. (Default = 0)
log-file	Writes the log to a file. (Default = stderr)
log-level	Specifies the log level; values are [DEBUG, INFO, WARN, TRACE, FATAL]. (Default = WARN)

cluster command: Cluster FLNC reads and generate transcripts.

Usage

isoseq3 cluster [options] input output

Example

isoseq3 cluster flnc.bam unpolished.bam --use-gvs

Custom BAM tags

isoseq3 cluster adds the following custom PacBio tags to the output BAM file:

- ib: Barcode summary: triplets delimited by semicolons, each triplet contains two barcode indices and the ZMW counts, delimited by commas. **Example**: 0,1,20;0,3,5
- im: ZMW names associated with this isoform.
- is: Number of ZMWs associated with this isoform.

Inputs/outputs	Description
input	flnc.bam file or movie.consensusreadset.xml file.
output	unpolished.bam file prefix (the actual output files will be unpolished.hq.bam and unpolished.lq-bam) or unpolished.transcriptset.xml file.

Options	Description
s1	Specifies the number of seeds for minimer-only clustering. (Default = 1000)
s2	Specifies the number of seeds for DP clustering. (Default = 1000)

Options	Description
poa-cov	Specifies the maximum number of CCS reads used for POA consensus. (Default = 10)
use-qvs	Use CCS analysis Quality Values; setspoa-cov to 100.
split-bam	Splits BAM output files into a maximum of N files; 0 means no splitting. (Default = 0)
min-subreads-split	Subread threshold for High-Quality/Low-Quality split; only works withuse-qvs. (Default = 7)
log-level	Specifies the log level; values are [DEBUG, INFO, WARN, ERROR, CRITICAL]. (Default = WARN)
-v,verbose	Uses verbose output.
-j,num-threads	Specifies the number of threads to use; 0 means autodetection. (Default = 0)
log-file	Writes the log to a file. (Default = stdout)

 ${\tt summarize}\ command:\ Create\ a\ .csv-format\ barcode\ overview\ from\ transcripts.$

Usage

isoseq3 summarize [options] input output

Example

isoseq3 summarize unpolished.bam summary.csv

Inputs/outputs	Description
input	unpolished.bam file. (The output from isoseq3 cluster.)
output	summary.csv file.

Options	Description
log-level	Specifies the log level; values are [DEBUG, INFO, WARN, ERROR, CRITICAL]. (Default = WARN)
-v,verbose	Uses verbose output.
log-file	Writes the log to file. (Default = stdout)

collapse command: Collapse transcripts based on genomic mapping.

Usage

isoseq3 collapse [options] <alignments.bam|xml> <ccs.bam|xml> <out.fastq>

Example - Bulk Iso-Seq

```
isoseq3 collapse --do-not-collapse-extra-5exons aligned.sorted.bam out.gff Or isoseq3 collapse --do-not-collapse-extra-5exons aligned.sorted.bam ccs.bam out.gff
```

Example - Single-Cell Iso-Seq

isoseq3 collapse aligned.sorted.bam out.gff \mathbf{Or} isoseq3 collapse aligned.sorted.bam ccs.bam out.gff

Inputs/outputs	Description
alignments	Alignments mapping transcripts to the reference genome. (BAM or XML file).
ccs.bam	Optional input BAM file containing CCS reads.
out.gff	Collapsed transcripts in GFF format.

Options	Description
min-aln-coverage	Ignores alignments with less than the Minimum Query Coverage. (Default = 0 . 95)
min-aln-identity	Ignores alignments with less than the Minimum Alignment Identity. (Default = 0.50)
max-fuzzy-junction	Ignores mismatches or indels shorter than or equal to N. (Default = 5)
max-5p-diff	Specifies the maximum allowed 5' difference if on same exon. (Default = 1000)
max-3p-diff	Specifies the maximum allowed 3' difference if on same exon. (Default = 100)
do-not-collapse-extra- 5exons	Do not collapse 5' shorter transcripts which miss one or multiple 5' exons to a longer transcript.
max-batch-mem	Specifies the maximum memory for batch loading, in megabytes (MB). Batches can be slightly larger than this value. Value ≤0 loads all data in memory at once. (Default = 4096)
split-group-size	Specifies that groups larger than this will be linearly split for parallel processing. (Default = 100)
keep-non-real-cells	Do not skip reads with non-real cells.
version	Displays program version number and exits.
log-file	Writes the log to file. (Default = stderr)
log-level	Specifies the log level; values are [DEBUG, INFO, WARN, ERROR, CRITICAL]. (Default = WARN)
-j,num-threads	Specifies the number of threads to use; 0 means autodetection. (Default = 0)

correct command: Correct group barcodes (xc:z) given a set of known correct barcodes. Additionally, real (as opposed to background RNA) cells are annotated with the rc tag after this step. In subsequent steps, such as isoseq3 groupdedup, only reads from real cells as indicated by the rc tag will be used.

Usage

isoseq3 correct [options] <flnc.bam> <flnc.corrected.bam>

Inputs/outputs	Description
flnc.bam	Input BAM file containing full-length non-concatemer reads, tagged with UMIs and cell barcodes.
flnc.corrected.bam	Output BAM file containing barcode-corrected full-length non-concatemer reads.

Options	Description
barcodes,-B	Specifies a plain text file containing known "true" barcodes, one per line. This include list specifies the barcode-set to which raw cell barcodes are remapped by minimum edit distance. May be gzip-compressed.
max-edit-distance,-M	Specifies the maximum edit distance for mapping barcodes to those specified by the -barcodes option. Increasing this value increases yield, but potentially introduce errors. (Default = 2)
filter,-F	Specifies the filtering mode.
	missing removes reads which could not be corrected.
	• failing removes reads which could not be corrected as well as reads failing the -max-edit-distance threshold.
	(Default = none)
help, -h	Displays help information and exits.
version	Displays program version number and exits.
verbose, -v	Sets the verbosity level.
-j,num-threads	Specifies the number of threads to use when processing; 0 means autodetection. (Default = 0)
log-file	Writes the log to a file. (Default = stderr)
log-level	Specifies the log level; values are [DEBUG, INFO, WARN, TRACE, FATAL]. (Default = WARN)

 ${\tt tag}$ command: Remove cell barcodes and unique molecular identifiers (UMIs) from full-length reads and generate tagged full-length transcripts (FL to FLT).

Usage

isoseq3 tag [options] <fl.bam> <fl.tagged.bam>

Inputs/outputs	Description
fl.bam	Input BAM file containing full-length reads.
fl.tagged.bam	Output BAM file containing full-length reads, with UMI and cell barcodes extracted and tagged.

Options	Description
design	Specifies which bases to use as cell/molecular barcodes as part of the barcoding design. (Default = $T-8U-10B$)
min-read-length	Specifies the minimum read length after trimming. (Default = 50)
help, -h	Displays help information and exits.
version	Displays program version number and exits.

Options	Description
verbose, -v	Sets the verbosity level.
-j,num-threads	Specifies the number of threads to use when processing; 0 means autodetection. (Default = 0)
log-file	Writes the log to a file. (Default = stderr)
log-level	Specifies the log level; values are [DEBUG, INFO, WARN, TRACE, FATAL]. (Default = WARN)

<code>groupdedup</code> command: Deduplicate reads grouped by UMIs (barcodesorted FLTNC to DEDUP). Note that after barcode correction, the BAM file **must** be sorted by the corrected barcode BAM tag (CB) first.

Usage

samtools sort -t CB corrected.bam -o corrected.sorted.bam
isoseq3 groupdedup [options] <fltnc.bam|xml (one or more).>

Inputs/outputs	Description
fltnc.bam xml (one or more)	Input BAM file(s) containing full-length non-concatemer transcripts, sorted by corrected barcode BAM tag (CB).
dedup.bam	Output BAM file(s) containing deduplicated reads.

Options	Description
poa-cov	Specifies the maximum number of CCS reads used to determine poa consensus, using the poa variant-calling algorithm. (Default = 10)
max-tag-mismatches	Specifies the maximum number of mismatches between tags. (Default = 1)
max-tag-shift	Specifies that tags may be shifted by at maximum of ${\tt N}$ bases. (Default = 1)
max-length-difference	Specifies the maximum insert lengths difference. (Default = 50)
max-insert-pad	Specifies the maximum number of missing flanking bases on either insert side. (Default = 5)
min-concordance-perc	Specifies the minimum insert alignment concordance, in %. (Default = 97)
max-insert-gaps	Specifies the maximum number of insert gaps per 20 bp window. (Default = 5)
barcode-tag,-C	Specifies the BAM cell/group tag by which to group. If not specified, checks the ${\tt CB}$ and ${\tt XC}$ tags.
no-poa	Select the highest-pass read in the data set instead of generating a consensus sequence using the poa variant-calling algorithm.
keep-non-real-cells	Specifies not to skip reads with non-real cells, as indicated by the rc tag.
batch-size	Specifies the number of BAM records to load per batch. (Default = 500000)
help, -h	Displays help information and exits.
version	Displays program version number and exits.

Options	Description
verbose, -v	Sets the verbosity level.
-j,num-threads	Specifies the number of threads to use when processing; 0 means autodetection. (Default = 0)
log-file	Writes the log to a file. (Default = stderr)
log-level	Specifies the log level; values are [DEBUG, INFO, WARN, TRACE, FATAL]. (Default = WARN)

bcstats command: Generates statistics for group barcodes (CB: Z,XC: Z, or CR: Z) and (optionally) molecular barcodes (XM: Z, UR: Z, or UB: Z). This tool should be run after isoseq3 correct to get the rc tag, which indicates which reads come from real cells (as opposed to ambient RNA).

Usage

isoseq3 bcstats [options] <Input BAMs>

Inputs	Description
input.bam (one or more)	Input BAM file(s) containing full-length non-concatemer transcripts.

Options	Description
molecular,-U	Specifies statistics for molecular barcodes (UMIs) as well as cell barcodes; this produces a .tsv file with one line for each cell barcode and one line for each molecular barcode.
-o,output	Specifies an output .tsv file of statistics for input BAM files. (Default = /dev/stdout)
-R,json	Specifies the path to an output JSON report. (Default = /dev/stderr)
deduplicated,-D	Specifies that the output is deduplicated. This allows for faster analysis and sanity checks across versions.
batch-size,-Z	Specifies the batch size for processing. (Default = 65536)
percentile,-P	Specifies the percentile to use when calculating real vs non-real cells. This option is relevant only when RealCellCalculationMethod (method) is set to percentile. (Default = 99)
-T,target	Specifies whether to determine real vs non-real cells by read count (readcount) or UMI count (umicount). (Default = umicount)
-M,method	Specifies whether to determine real vs non-real cells using the Knee-finding (knee) or the percentile-based method (percentile). (Default = knee)
help, -h	Displays help information and exits.
version	Displays program version number and exits.
verbose, -v	Sets the verbosity level.
-j,num-threads	Specifies the number of threads to use when processing; 0 means autodetection. (Default = 0)
log-file	Writes the log to a file. (Default = stderr)
log-level	Specifies the log level; values are [DEBUG, INFO, WARN, TRACE, FATAL]. (Default = WARN)

 ${\tt dedup}\ command:\ Deduplicate\ reads\ grouped\ by\ UMIs.\ \textbf{Note}:\ This\ tool\ has\ been\ superceded\ by\ the\ {\tt groupdedup}\ tool.$

Usage

isoseq3 dedup [options] <fltnc.bam|xml> <dedup.bam|xml>

Inputs/outputs	Description
fltnc.bam xml (one or more)	Input BAM file(s) containing full-length non-concatemer transcripts, sorted by corrected barcode BAM tag (CB).
dedup.bam xml	Output BAM file(s) or XML file(s) containing deduplicated reads.

Options	Description
poa-cov	Specifies the maximum number of CCS reads used to determine poa consensus, using the poa variant-calling algorithm. (Default = 10)
max-tag-mismatches	Specifies the maximum number of mismatches between tags. (Default = 1)
max-tag-shift	Specifies that tags may be shifted by at maximum of ${\tt N}$ bases. (Default = 1)
max-insert-length-diff	Specifies the maximum insert lengths difference. (Default = 50)
max-insert-pad	Specifies the maximum number of missing flanking bases on either insert side. (Default = 5)
min-concordance-perc	Specifies the minimum insert alignment concordance, in %. (Default = 97)
max-insert-gaps	Specifies the maximum number of insert gaps per 20 bp window. (Default = 5)
keep-non-real-cells	Specifies not to skip reads with non-real cells, as indicated by the rc tag.
help, -h	Displays help information and exits.
version	Displays program version number and exits.
verbose, -v	Sets the verbosity level.
-j,num-threads	Specifies the number of threads to use when processing; 0 means autodetection. (Default = 0)
log-file	Writes the log to a file. (Default = stderr)
log-level	Specifies the log level; values are [DEBUG, INFO, WARN, TRACE, FATAL]. (Default = WARN)

juliet

juliet is a general-purpose minor variant caller that identifies and phases minor single nucleotide substitution variants in complex populations. It identifies codon-wise variants in coding regions, performs a reference-guided *de novo* variant discovery, and annotates known drug-resistance mutations. Insertion and deletion variants are currently ignored; support will be added in a future version. There is no technical limitation with respect to the target organism or gene.

The underlying model is a statistical test, the Bonferroni-corrected Fisher's Exact test. It compares the number of observed mutated codons to the number of expected mutations at a given position.

<code>juliet</code> uses JSON target configuration files to define different genes in longer reference sequences, such as overlapping open reading frames in HIV. These predefined configurations ease batch applications and allow immediate reproducibility. A target configuration may contain multiple coding regions within one reference sequence and optional drug resistance mutation positions.

Notes:

- The preinstalled target configurations are meant for a quick start. It is the user's responsibility to ensure that the target configurations used are correct and up-to-date.
- If the target configuration none was specified, the provided reference is assumed to be in-frame.

Performance

At a coverage of 6,000 CCS reads with a predicted accuracy (RQ) of ≥ 0.99 , the false positive and false negative rates are below 1% and 0.001% (10^{-5}), respectively.

Usage

juliet --config "HIV" data.align.bam patientZero.html

Required	Description
input_file.bam	Input aligned BAM file containing CCS reads, which must be PacBiocompliant, that is, cigar M is forbidden.
output_file.html	Output report HTML file.

Configuration	Description
config,-c	Path to the target configuration JSON file, predefined target configuration tag, or the JSON string.
mode-phasing,-p	Phase variants and cluster haplotypes.

Restrictions	Description
region,-r	Specifies the genomic region of interest; reads are clipped to that region. Empty means all reads.
drm-only,-k	Only reports DRM positions specified in the target configuration. Can be used to filter for drug-resistance mutations - only known variants from the target configuration are called.
min-perc,-m	Specifies the minimum variant percentage to report. Example :min-perc 1 will only show variant calls with an observed abundance of more than 1%. (Default = 0)
max-perc,-n	Specifies the maximum variant percentage to report. Example :max-perc 95 will only show variant calls with an observed abundance of less than 95%. (Default = 100)

Chemistry override (specify both)	Description
sub,-s	Specifies the substitution rate. Use to override the learned rate. (Default = 0)
del,-d	Specifies the deletion rate. Use to override the learned rate. (Default = 0)

Options	Description
help, -h	Displays help information and exits.
verbose, -v	Sets the verbosity level.
version	Displays program version number and exits.
debug	Returns all amino acids, irrespective of their relevance.
mode-phasing,-p	Phases variants and cluster haplotypes.

Input files

- BAM-format files containing CCS reads. These must be PacBiocompliant, that is, cigar M is forbidden.
- Input CCS reads should have a minimal predicted accuracy of 0.99.
- Reads should be created with CCS analysis using the --richQVs option. Without the --richQVs information, the number of false positive calls might be higher, as juliet is missing information to filter actual heteroduplexes in the sample provided.
- juliet currently does **not** demultiplex barcoded data; you must provide one BAM file per barcode.

Output files

A JSON and/or HTML file:

```
juliet data.align.bam patientZero.html
juliet data.align.bam patientZero.json
juliet data.align.bam patientZero.html patientZero.json
```

The HTML file includes the same content as the JSON file, but in more human-readable format. The HTML file contains four sections:

1. Input data

Summarizes the data provided, the exact call for juliet, and juliet version for traceability purposes.

2. Target config

Predefined v1.1, PacBio internal

Summarizes details of the provided target configuration for traceability. This includes the configuration version, reference name and length, and annotated genes. Each gene name (in bold) is followed by the reference start, end positions, and possibly known drug resistance mutations.

▼ Target config

Config Version:

```
Reference Name: HIV HXB2
Reference Length: 9719
Genes:
  • 5'LTR (1-634)
  • p17 (790-1186)
  • p24 (1186-1879)
  • p2 (1879-1921)
  • p7 (1921-2086)
  • p1 (2086-2134)
  • p6 (2134-2292)
  • Protease (2253-2550)
       o atv/r: V321 L33F M46I M46L 147V G48V G48M I50L I54V I54T I54A I54L I54M V82A V82T V82F V82S I84V N88S L90M
       o DRV/r: V32I L33F I47V I47A I50V I54L I54M L76V V8F I84V
       o fpv/r: v32I L33F M46I M46L I47V I47A I50V I54V I54T I54A I54L I54M L76V V82A V82T V82F V82S I84V L90M
       o IDV/r: V32I M46I M46L I47V I54V I54T I54A I54L I54M L76V V82A V82T V82F V82S I84V N88S L90M
       O NFV: D30N L33F M46I M46L I47V G48V G48M I54V I54T I54A I54L I54M V82A V82T V82F V82S I84V N88D N88S L90M
       o SOV/r: G48V G48M I54V I54T I54A I54L I54M V82A V82T I84V N88S L90M
```

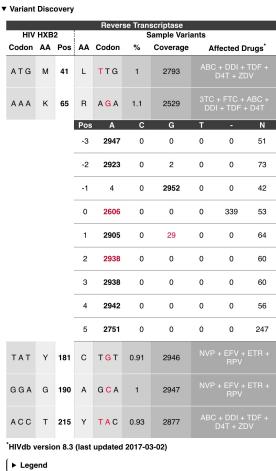
3. Variant discovery

For each gene/open reading frame, there is one overview table.

Each row represents a variant position.

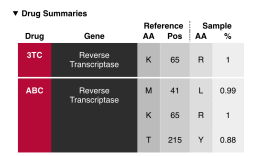
o TPV/r: V32I L33F M46I M46L I47V I47A I54V I54A I54M V82T V82L I84V

- Each variant position consists of the reference codon, reference amino acid, relative amino acid position in the gene, mutated codon, percentage, mutated amino acid, coverage, and possible affected drugs.
- Clicking the row displays counts of the multiple-sequence alignment counts of the -3 to +3 context positions.



4. Drug summaries

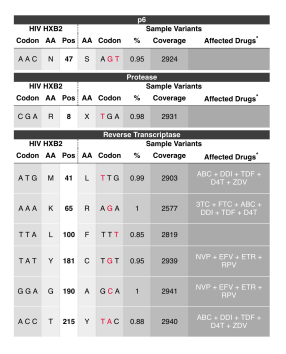
Summarizes the variants grouped by annotated drug mutations:



Predefined target configuration

juliet ships with one predefined target configuration, for HIV. Following is the command syntax for running that predefined target configuration:

juliet --config "HIV" data.align.bam patientZero.html



• **Note**: For the predefined configuration HIV, use the HIV HXB2 complete genome for alignment.

Customized target configuration

To define your own target configuration, create a JSON file. The root child genes contains a list of coding regions, with begin and end, the name of the gene, and a list of drug resistant mutations. Each DRM consists of its name and the positions it targets. The drms field is optional. If provided, the referenceSequence is used to call mutations, otherwise it will be tested against the major codon. All indices are with respect to the provided alignment space, 1-based, begin-inclusive and end-exclusive [).

Target configuration: Example 1- A customized json target configuration file named my customized hiv.json:

```
"version": "Free text to version your config files"
  "databaseVersion": "DrugDB version x.y.z (last updated YYYY-MM-DD)"
}
```

Run with a customized target configuration using the --config option:

juliet --config my customized hiv.json data.align.bam patientZero.html

Valid formats for DRMs/positions

```
Only the reference position.

M130 Reference amino acid and reference position.

M103L Reference aa, reference position, mutated aa.

M103LKA Reference aa, reference position, list of possible mutated aas.

Reference position and mutated aa.

Reference position and list mutated aas.
```

Missing amino acids are processed as wildcard (*).

Example

```
{ "name": "ATV/r", "positions": [ "V32I", "L33", "46IL", "I54VTALM", "V82ATFS", "84" ] }
```

Target configuration: Example 2 - BCR-ABL:

For BCR-ABL, using the ABL1 gene with the following reference NM_005157.5, a typical target configuration looks like this:

```
"genes": [
     {
          "name": "ABL1",
          "begin": 193,
          "end": 3585,
          "drms": [
                  "name": "imatinib",
                  "positions": [
                    "T315AI", "Y253H", "E255KV", "V299L", "F317AICLV", "F359CIV" ]
              },
                  "name": "dasatinib",
                  "positions": [ "T315AI", "V299L", "F317AICLV" ]
              },
                  "name": "nilotinib",
                  "positions": [ "T315AI", "Y253H", "E255KV", "F359CIV" ]
              },
                  "name": "bosutinib",
                  "positions": [ "T315AI" ]
          ]
     }
 ],
"referenceName": "NM 005157.5",
 "referenceSequence": "TTAACAGGCGCGTCCC..."
```

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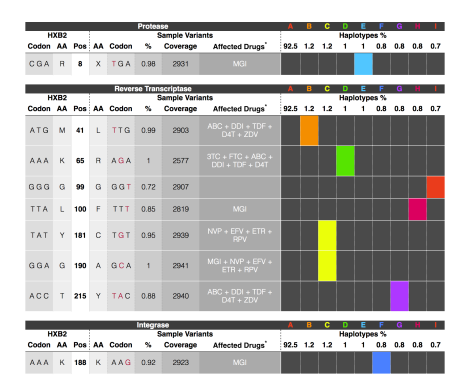
No target configuration

If **no** target configuration is specified, either make sure that the sequence is in-frame, or specify the region of interest to mark the correct reading frame, so that amino acids are correctly translated. The output is labeled with unknown as the gene name:

juliet data.align.bam patientZero.html

Phasing

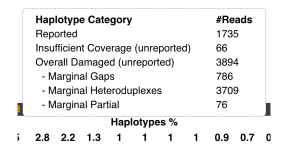
The default mode is to call amino-acid/codon variants independently. Using the <code>--mode-phasing</code> option, variant calls from distinct haplotypes are clustered and visualized in the HTML output.



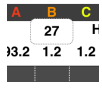
- The row-wise variant calls are "transposed" onto per-column haplotypes. Each haplotype has an ID: [A-Z] {1} [a-z]?.
- For each variant, colored boxes in this row mark haplotypes that contain this variant.
- Colored boxes per haplotype/column indicate variants that co-occur.
 Wild type (no variant) is represented by plain dark gray. A color palette helps to distinguish between columns.
- The JSON variant positions has an additional haplotype_hit boolean array with the length equal to the number of haplotypes. Each entry indicates if that variant is present in the haplotype. A haplotype block under the root of the JSON file contains counts and read names. The order of those haplotypes matches the order of all haplotype_hit arrays.

There are two types of tooltips in the haplotype section of the table.

The first tooltip is for the **Haplotypes** % and shows the number of reads that count towards (A) Actually reported haplotypes, (B) Haplotypes that have less than 10 reads and are not being reported, and (C) Haplotypes that are not suitable for phasing. Those first three categories are mutually exclusive and their sum is the total number of reads going into juliet. For (C), the three different marginals provide insights into the sample quality; as they are marginals, they are not exclusive and can overlap. The following image shows a sample with bad PCR conditions:



The second type of tooltip is for each haplotype percentage and shows the number of reads contributing to this haplotype:



Microbial Genome Analysis

The Microbial Genome Analysis application is powered by the IPA HiFi genome assembler and includes a base modification detection feature performed after the assembly.

Workflow of the Microbial Genome Analysis application

The workflow consists of several steps around 2 main stages:

- 1. **Chromosomal stage**: Assemble large contigs using IPA, the HiFi genome assembly tool.
- 2. Separate reads that were used for accurate large contigs from all other reads.
- 3. **Plasmid stage**: Assemble plasmids (using IPA) from the reads separated in the previous step.
- 4. De-duplicate plasmids.
- 5. Collect all contigs into a single FASTA file.
- 6. Rotate circular contigs.
- 7. Align the input Data Set to the assembled contigs.
- 8. Polish assembled contigs using Racon.
- 9. Perform base modification detection.

The application accepts HiFi XML Data Sets as input, and has an embedded downsampling feature:

- If the genome size and the desired coverage are specified, both stages
 of assembly are downsampled, as with the Genome Assembly
 application.
- Otherwise, the full coverage is used.

The embedded downsampling feature is **not** applied to the alignment stage; **all** input reads will be aligned against the assembled contigs.

Usage

The Microbial Genome Analysis application is run using the pbcromwell run command, with the pb_microbial_analysis parameter to specify the application. See "pbcromwell" on page 81 for details.

To view information on the available Microbial Genome Analysis options, enter:

pbcromwell show-workflow-details pb microbial analysis

The minimum command needed to run the workflow requires the input Data Set.

The following example performs assembly and base modification detection using an input XML Data Set, and uses all default settings, including 1 CPU:

pbcromwell run pb_microbial_analysis -e <input.consensusreadset.xml>

Note: To specify different task options on the command line, consider the following example:

```
pbcromwell run pb_microbial_analysis \
-e <input.consensusreadset.xml> \
--task-option ipa2_genome_size=0 \
--task-option ipa2_downsampled_coverage=0 \
--task-option microasm_plasmid_contig_len_max=300000
--task-option ipa2_cleanup_intermediate_files=True \
--task-option dataset_filters="" \
--task-option filter_min_qv=20 \
--nproc 8
```

The default options for this application should work well for any genome type.

As microbes are relatively small, we rarely find much advantage to using more than 4 threads, or more than 2 concurrent jobs.

Microbial Genome Analysis parameters

Option	Default value	Description
-e,eid_ccs	NONE	This is a SMRT Link-specific input parameter and supports only PacBio Consensusreadset XML files as input.
task-option reads	NONE	Optional parameter, required if -e <input/> is not specified. Supports multiple input formats: FASTA, FASTQ, BAM, XML, FOFN and gzipped versions of FASTA/FASTQ.
task-option ipa2_genome_size	10M	The approximate number of base-pairs expected in the chromosomal genome. Used only for downsampling in the assembly stages (that is, not in polishing). If value <=0, then downsampling is off.
		Default: 10M (10 Mega basepairs).
		Note: <code>ipa2_genome_size</code> is currently the only option that accepts a metric suffix. Commas and decimals are not accepted anywhere. <code>ipa2_genome_size</code> actually accepts a String, which can be an integer followed by one of these metric suffixes: <code>k/M/G</code> . For example: <code>4500k</code> means "4,500 kilobases" or "4,500,000". <code>M</code> stands for Mega and <code>G</code> stands for Giga.
task-option ipa2_downsampled_coverage	100	The maximum coverage after downsampling with respect to the estimated <code>genome_size</code> . (The default <code>genome_size</code> was chosen to be larger than most realistic microbes.) If value <=0, then downsampling is off.
		Default: 100
		Example : If your genome is 1G in length, and you specify ipa2_genome_size=2G, you have over-estimated by 2x. If you also specify ipa2_downsampled_coverage=100, your data will be downsampled to 200x coverage, simply because of the over-estimate. The cost of extra coverage is greater runtime. The defaults are usually fine.

Option	Default value	Description
task-option ipa2_advanced_options_chro m	See Description column	A semicolon-separated list of KEY=VALUE pairs. New line characters are not accepted. (These are described later in this document.)
		<pre>config_block_size = 100;</pre>
		<pre>config_seeddb_opt = -k 28 -w 20space 0use- hpc-seeds-only;</pre>
		<pre>config_ovl_opt =one-hit-per-targetmin-idt 98tracebackmask-hpmask-repeatstrim trim-window-size 30trim-match-frac 0.75</pre>
task-option ipa2_advanced_options_plas mid	See Description column	A semicolon-separated list of KEY=VALUE pairs. New line characters are not accepted. (These are described later in this document.)
		<pre>con-fig_block_size = 100; con-fig_ovl_filter_opt =max-diff 80max-cov 100min-cov 2bestn 10min-len 500</pre>
		gapFiltminDepth 4idt-stage2 98;
		<pre>con-fig_ovl_min_len = 500; con-fig_seeddb_opt = -k 28 -w 20space 0use- hpc-seeds-only;</pre>
		config ovl opt =one-hit-per-targetmin-idt
		98min-map-len 500min-anchor-span 500 tracebackmask-hpmask-repeatstrimtrim- window-size 30trim-match-frac 0.75smart- hit-per-targetsecondary-min-ovl-frac 0.05; con-fig_layout_opt =allow-circular;
task-option ipa2_cleanup_intermediate_ files	TRUE	Removes intermediate files from the run directory to save space.
task-option microasm_plasmid_contig_le n_max	300000	After the chromosomal stage, in task filter_draft_contigs, separates long contigs (presumed to be chromosomal) from shorter contigs (to be re-assembled in the plasmid stage). Then, after the plasmid stage, in dedup_plasmids, contigs less than this value are ignored. The default value is usually fine.
task-option microasm_run_secondary_pol ish	TRUE	Specifies that an additional round of polishing will be applied after the two stages of assembly have completed.
task-option run_basemods	TRUE	Specifies that base modification analysis be performed.
task-option kineticstools_identify_mod s	m4C,m6A	Specify the base modifications to identify, in a commaseparated list.
task-option kineticstools_p_value	0.001	Specifies the probability value cutoff for detecting base modifications.
task-option motif_min_score	35	Specifies the minimum QMod score used to identify a motif.
task-option motif_min_fraction	0.30	Specifies the minimum methylated fraction to identify a motif.
task- optionrun_find_motifs	TRUE	Specifies that motif-finding be performed.
task-option dataset_filters	NONE	(General pbcromwell option) A semicolon-separated (not comma-separated) list of other filters to add to the Data Set.
task-option filter_min_qv	20	(General pbcromwell option) Phred-scale integer QV cutoff for filtering HiFi reads. The default for all applications is 20 (QV 20), or 99% predicted accuracy.

Option	Default value	Description
task-option log_level	INFO	Specifies the logging (verbosity) level for tools which support this feature. Values are: [TRACE, DEBUG, INFO, WARN, FATAL]
task-option nproc	1	Specified the number of CPU threads to use.
task-option max_nchunks	40	Specifies the maximum number of chunks per task.
task-option tmp_dir	/tmp	Specifies an optional temporary directory for used by several subtools, such as sort.

Input files

- *.bam file containing PacBio data.
- *.xml file containing PacBio data.

Output files

- final_assembly.fasta: File containing all assembled contigs, rotated.
- assembly.rotated.polished.renamed.fsa: File for NCBI, but otherwise identical to final_assembly.fasta; only the headers are changed.
- pb_microbial_analysis.motifs_csv: File containing motifs and modifications.

Advanced assembly options

Note: Advanced parameters should be rarely modified. Advanced parameters for the Microbial Genome Assembly workflow are the **same** as those for the Genome Assembly workflow. See "Advanced parameters" on page 45 for details.

Advanced parameters specified on the command line:

- Are in the form of key = value pairs.
- Each pair is separated by a semicolon (;) character.
- The full set of advanced parameters is surrounded by one set of double quotes.
- The specified value of a parameter overwrites the default options for that key. All desired options of that parameter must be explicitly listed, not just the ones which should change from the default.
- Setting an empty value clears the parameter; it does not reset the value back to default.

Example

--task-option ipa2_advanced_options_chrom="config_seeddb_opt=-k
28;config block size=2048"

motifMaker

The motifMaker tool identifies motifs associated with DNA modifications in prokaryotic genomes. Modified DNA in prokaryotes commonly arises from restriction-modification systems that methylate a specific base in a specific sequence motif. The canonical example is the m6A methylation of adenine in GATC contexts in *E. coli*. Prokaryotes may have a very large number of active restriction-modification systems present, leading to a complicated mixture of sequence motifs.

PacBio SMRT sequencing is sensitive to the presence of methylated DNA at single base resolution, via shifts in the polymerase kinetics observed in the real-time sequencing traces. For more background on modification detection, see here.

Algorithm

Existing motif-finding algorithms such as MEME-chip and YMF are suboptimal for this case for the following reasons:

- They search for a **single** motif, rather than attempting to identify a complicated mixture of motifs.
- They generally don't accept the notion of aligned motifs the input to
 the tools is a window into the reference sequence which can contain
 the motif at any offset, rather than a single center position that is
 available with kinetic modification detection.
- Implementations generally either use a Markov model of the reference (MEME-chip), or do exact counting on the reference, but place restrictions on the size and complexity of the motifs that can be discovered.

Following is a rough overview of the algorithm used by motifMaker: Define a motif as a set of tuples: (position relative to methylation, required base). Positions not listed in the motif are implicitly degenerate. Given a list of modification detections and a genome sequence, define the following objective function on motifs:

Motif score(motif) = (# of detections matching motif) / (# of genome sites matching motif) * (Sum of log-pvalue of detections matching motif) = (fraction methylated) * (sum of log-pvalues of matches)

Then, search (close to exhaustively) through the space of all possible motifs, progressively testing longer motifs using a branch-and-bound search. The "fraction methylated" term must be less than 1, so the maximum achievable score of a child node is the sum of scores of modification hits in the current node, allowing pruning of all search paths whose maximum achievable score is less than the best score discovered so far.

Usage

Run the find command, and pass the reference FASTA and the modifications.gff (.gz) file output by the PacBio modification detection workflow.

The reprocess subcommand annotates the GFF file with motif information for better genome browsing.

MotifMaker [options] [command] [command options]

find command: Run motif-finding.

find [options]

Options	Description
-h,help	Displays help information and exits.
* -f,fasta	Reference FASTA file.
* -g,gff	Modifications.gff or .gff.gz file.
-m,minScore	Specifies the minimum Qmod score to use in motif finding. (Default = 40.0)
* -o,output	Outputs motifs.csv file.
-x,xml	Outputs motifs XML file.

reprocess command: Update a modifications.gff file with motifinformation based on new Modification QV thresholds.

reprocess [options]

Options	Description
-c,csv	Raw modifications.csv file.
* -f,fasta	Reference FASTA file.
* -g,gff	Modifications.gff or .gff.gz file.
-m,minFraction	Specifies that only motifs above this methylated fraction are used. (Default = 0.75)
-m,motifs	Motifs.csv file.
* -o,output	Reprocessed modifications.gff file.

Output files

Using the find command:

• Output CSV file: Contains motif information.

Using the reprocess command:

 Output GFF file: Contains added motif information based on new Modification QV thresholds. The format of the output file is the same as the input file.

pbcromwell

The pbcromwell tool is PacBio's wrapper for the cromwell scientific workflow engine used to power SMRT Link. pbcromwell includes advanced utilities for interacting directly with a Cromwell server.

pbcromwell is designed primarily for running workflows distributed and supported by PacBio, but it is written to handle any valid WDL source (version 1.0), and is very flexible in how it takes input. PacBio workflows are expected to be found in the directory defined by the SMRT_PIPELINE_BUNDLE_DIR environment variable, which is automatically defined by the SMRT Link distribution.

Note that pbcromwell does **not** interact with SMRT Link services; to run a Cromwell workflow as a SMRT Link job, use pbservice. (For details, see "pbservice" on page 97.)

Note: Interaction with the Cromwell server is primarily intended for developers and power users.

Usage

Options	Description
output-dir OUTPUT_DIR	Specifies the output directory for cromwell output. (Default = cromwell_out)
overwrite	Overwrites the output directory, if it exists. (Default = False)
-i INPUTS,inputs INPUTS	Specifies cromwell inputs and settings as JSON files. (Default = None)
-e ENTRY_POINTS,entry ENTRY_POINTS	Specifies the entry point Data Set; may be repeated for workflows that take more than one input Data Set. Note that all PacBio workflows require at least one such entry point.
-n NPROC,nproc NPROC	Specifies the number of processors per task. (Default = 1)
-c MAX_NCHUNKS,max-nchunks MAX_NCHUNKS	Specifies the maximum number of chunks per task. (Default = None)
target-size TARGET_SIZE	Specifies the target chunk size. (Default = None)
queue QUEUE	Specifies the cluster queue to use. (Default = None)
-o OPTIONS,options OPTIONS	Specifies additional cromwell engine options, as a JSON file. (Default = None)
-t TASK_OPTIONS,task-option TASK_OPTIONS	Specifies workflow- or task-level option as $key=value$ strings, specific to the application. May be specified multiple times for multiple options. (Default = [])
-b BACKEND,backend BACKEND	Specifies the backend to use for running tasks. (Default = None)
-r MAX_RETRIES,maxRetries MAX_RETRIES	Specifies the maximum number of times to retry a failing task. (Default = 1)

Options	Description
tmp-dir TMP_DIR	Specifies an optional temporary directory for cromwell tasks, which must exist on all compute hosts. (Default = None)
config CONFIG	Specifies a Java configuration file for running cromwell. (Default = None)
dry-run	Specifies that cromwell is not executed, but instead writes out final inputs and then exits. (Default = True)
workflow	Specifies the workflow ID (such as pb_ccs or cromwell.workflows.pb_ccs for PacBio workflows only) or a path to a Workflow Description Language (WDL) source file.

Enter pbcromwell {command} -h for a command's options.

Examples

Show available PacBio-developed workflows:

pbcromwell show-workflows

Show details for a PacBio workflow:

pbcromwell show-workflow-details pb_ccs

Generate cromwell.conf with HPC settings:

pbcromwell configure --default-backend SGE --output-file cromwell.conf

Launch a PacBio workflow:

pbcromwell run pb_ccs -e /path/to/movie.subreadset.xml --nproc 8 --config /full/ path/ to/cromwell.conf

Options	Description
-h,help	Displays help information and exits.
version	Displays program version number and exits.
log-file LOG_FILE	Writes the log to file. (Default = None, writes to stdout.)
log-level=INFO	Specifies the log level; values are [DEBUG, INFO, WARNING, ERROR, CRITICAL.] (Default = INFO)
debug	Alias for setting the log level to DEBUG. (Default = False)
verbose, -v	Sets the verbosity level. (Default = None)

pbcromwell run command: Run a Cromwell workflow. Multiple input modes are supported, including a pbsmrtpipe-like set of arguments (for PacBio workflows **only**), and JSON files already in the native Cromwell format.

All PacBio workflows have similar requirements to the pbsmrtpipe pipelines in previous SMRT Link versions:

- 1. One or more PacBio dataset XML entry points, usually a SubreadSet or ConsensusReadSet (--entry-point <FILE>.)
- 2. Any number of workflow-specific task options (--task-option <OPTION>.)
- 3. Engine options independent of the workflow, such as number of processors per task (--nproc), or compute backend (--backend).

Output is directed to a new directory: --output-dir, which defaults to cromwell_out. This includes final inputs for the Cromwell CLI, and subdirectories for logs (workflow and task outputs), links to output files, and the Cromwell execution itself, which has a complex nested directory structure. Detailed information about the workflow execution can be found in the file metadata.json, in the native Cromwell format.

Note that output file links do **not** include the individual resource files of Data Sets and reports (BAM files, index files, plot PNGs, and so on.) Follow the symbolic links to their real path (for example using readlink -f) to find report plots.

For further information about Cromwell, see the official documentation here.

Workflow examples

Run the CCS analysis:

pbcromwell run pb ccs -e <SUBREADS> --nproc 8 --config /full/path/to/cromwell.conf

Run the Iso-Seq workflow, including mapping to a reference, and execute on SGE:

pbcromwell run pb_isoseq3 -e <SUBREADS> -e <PRIMERS> -e <REFERENCE> --nproc 8 -- config
/full/path/to/cromwell.conf

Run a user-defined workflow:

pbcromwell run my_workflow.wdl -i inputs.json -o options.json --config /full/path/to/ cromwell.conf

Set up input files but exit before starting Cromwell:

pbcromwell run pb ccs -e <SUBREADS> --nproc 8 --dry-run

Print details about the named PacBio workflow, including input files and task options. **Note**: The prefix cromwell.workflows. is optional.

```
pbcromwell show-workflow-details pb_ccs
pbcromwell show-workflow-details cromwell.workflows.pb ccs
```

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pbcromwell show-workflow-details command: Display details about a specified PacBio workflow, including input files and task options.

Usage

Options	Description
workflow_id	Specifies the workflow ID, such as pb_ccs or cromwell.workflows.pb_ccs for PacBio workflows only.
inputs-json INPUTS_JSON	Writes a JSON template containing workflow inputs to the specified file. (Default = None)

pbcromwell configure command: Generate the Java configuration file used by cromwell to define backends and other important engine options that **cannot** be set at runtime. You can pass this to pbcromwell run using the --config argument.

Usage

Options	Description
port PORT	Specifies the port that cromwell should listen to. (Default = 8000)
local-job-limit LOCAL_JOB_LIMIT	Specifies the maximum number of local jobs/tasks that can be run at once. (Default = 10)
jms-job-limit JMS_JOB_LIMIT	Specifies the maximum number of jobs/tasks that can be submitted to the queueing system at one time. (Default = 500)
db-port DB_PORT	Specifies the database port for cromwell to use; if undefined, database configuration is omitted. (Default = None)
db-user DB_USER	Specifies the user name used to connect to Postgres. (Default = smrtlink_user)
db-password DB_PASSWORD	Specifies the password used to connect to Postgres.
default-backend DEFAULT_BACKEND	Specifies the default job execution backend. Choices are: Local, SGE, Slurm, PBS, or LSF. (Default = Local)
max-workflows MAX_WORKFLOWS	Specifies the maximum number of workflows that cromwell can run at once. (Default = 100)
output-file OUTPUT_FILE	Specifies the name of the output configuration file. (Default = cromwell.conf)
timeout TIMEOUT	Specifies the time to wait for task completion before checking the cluster job status. (Default = 600)

Options	Description
cache	Enables call caching. (Default = True)
no-cache	Disables call caching. (Default = True)

pbindex

The ${\tt pbindex}$ tool creates an index file that enables random access to PacBio-specific data in BAM files.

Usage

pbindex <input>

Options	Description
-h,help	Displays help information and exits.
version	Displays program version number and exits.

Input file

• *.bam file containing PacBio data.

Output file

- * . pbi index file, with the same prefix as the input file name.

pbmarkdup

The pbmarkdup tool marks PCR duplicates in CCS reads Data Sets from amplified libraries. PCR duplicates are different reads that arose from amplifying a single-source molecule. pbmarkdup can also optionally remove the duplicate reads.

Note: pbmarkdup only works with CCS reads, not with Subreads.

pbmarkdup uses a reference-free comparison method. Duplicates are identified as pairs of reads that:

- 1. Have the same length within 10 bp, and
- 2. Have high percent identity alignments at the molecule ends at >98% identity of the first and last 250 bp.

Clusters are formed from sets of two or more duplicate reads, and a single read is selected as the representative of each cluster. Other reads in the cluster are considered duplicates.

How are duplicates marked?

In FASTA and FASTQ formats, reads from duplicate clusters have annotated names. The following is a FASTA example:

>m64013 191117 050515/67104/ccs DUPLICATE=m64013 191117 050515/3802014/ccs DS=2

This shows a marked duplicate read $m64013_191117_050515/67104/ccs$ that is a duplicate of $m64013_191117_050515/3802014/ccs$ in a cluster with 2 reads (DS value). Accordingly, the following is the read selected as the representative of the molecule:

 $>m64013_191117_050515/3802014/ccs$ DS=2

In BAM format, duplicate reads are flagged with 0×400 . The read-level tag ds provides the number of reads in a cluster (like the DS attribute described above for FASTA/FASTQ), and the du tag provides the name of the representative read (like the DUPLICATE attribute described above for FASTA/FASTQ).

Usage

pbmarkdup [options] <INFILE.bam|xml|fa|fq|fofn> <OUTFILE.bam|xml|fa.gz|fq.gz>

Options Description	
-h,help	Displays help information and exits.
version	Displays program version number and exits.
log-file	Logs to a file, instead of stderr.

Options	Description				
log-level	Specifies the log level; values are [TRACE, DEBUG, INFO, WARN, FATAL] (Default = WARN)				
	log-le	vel IN	FO produces a summ	ary report such as:	
	LIBRARY	READS	UNIQUE MOLECULES	DUPLICATE READS	
	<unnamed></unnamed>	25000	24948 (99.8%)	52 (0.2%)	
	SS-lib	496	493 (99.4%)	3 (0.6%)	
	TOTAL	25496	25441 (99.8%)	55 (0.2%)	
-j,num-threads	Specifies th (Default = 0		er of threads to use, (means autodetection	۱.

Duplicate marking options	Description
cross-library, -x	Identifies duplicate reads across sequencing libraries. Libraries are specified in the BAM read group ${\tt LB}$ tag.

Output options	Description
-rmdup, -r	Excludes duplicates from OUTFILE. (This is redundant whendup-file is specified.)
dup-file FILE	Stores duplicate reads in an extra file other than OUTFILE. The format of this file can be different from the output file.
clobber, -f	Overwrites OUTFILE if it exists.

Input files

CCS reads from one or multiple movies in any of the following formats:

- PacBio BAM (.ccs.bam)
- PacBio dataset (.dataset.xml)
- File of File Names (.fofn)
- FASTA (.fasta,.fasta.gz)
- FASTQ (.fastq,.fastq.gz)

Output files

CCS reads with duplicates marked in a format inferred from the file extension:

- PacBio BAM (.ccs.bam)
- PacBio dataset (.dataset.xml), which also generates a corresponding BAM file.
- FASTA (.fasta.gz)
- FASTQ (.fastq.gz)

Allowed input/output combinations

Input File	Output BAM	Output Dataset	Output FASTQ	Output FASTA
Input BAM	Allowed	Allowed	Allowed	Allowed
Input Dataset	Allowed	Allowed	Allowed	Allowed
Input FASTQ	Not Allowed	Not Allowed	Allowed	Allowed
Input FASTA	Not Allowed	Not Allowed	Not Allowed	Allowed

Examples

Run on a single movie:

pbmarkdup movie.ccs.bam output.bam

Run on multiple movies:

pbmarkdup movie1.fasta movie2.fasta output.fasta

Run on multiple movies and output duplicates in separate file:

pbmarkdup moviel.ccs.bam movie2.fastq uniq.fastq --dup-file dups.fasta

pbmm2

The pbmm2 tool aligns native PacBio data, outputs PacBio BAM files, and is a SMRT minimap2 wrapper for PacBio data.

pbmm2 supports native PacBio input and output, provides sets of recommended parameters, generates sorted output on-the-fly, and post-processes alignments. Sorted output can be used directly for polishing using GenomicConsensus, if BAM has been used as input to pbmm2.

pbmm2 adds the following SAM tag to each aligned record:

 mg, stores gap-compressed alignment identity, defined as nM/(nM + nMM + nInsEvents + nDelEvents).

Usage

pbmm2 <tool>

Options	Description	
-h,help	Displays help information and exits.	
version	Displays program version number and exits.	

index command: Indexes references and stores them as .mmi files. Indexing is optional, but recommended if you use the same reference with the same --preset multiple times.

Usage

```
pbmm2 index [options] <ref.fa|xml> <out.mmi>
```

Input file

 *.fasta, *.fa file containing reference contigs or *.referenceset.xml.

Output file

• out.mmi (minimap2 index file.)

Notes:

- You can use existing minimap2 .mmi files with pbmm2 align.
- If you use an index file, you cannot override parameters -k, -w, nor -u
 in pbmm2 align.
- The minimap2 parameter -H (homopolymer-compressed k-mer) is always on for SUBREAD and UNROLLED presets, and can be disabled using -u.

Options	Description
preset	Specifies the alignment mode:
	• "SUBREAD" -k 19 -w 10
	• "CCS" -k 19 -w 10 -u
	• "ISOSEQ" -k 15 -w 5 -u
	• "UNROLLED" -k 15 -w 15
	The option is not case-sensitive. (Default = SUBREAD)
-k	Specifies the k-mer size, which cannot be larger than 28. (Default = -1)
-w	Specifies the Minimizer window size. (Default = -1)
-u,no-kmer-compression	Disables homopolymer-compressed k-mer. (Compression is on by default for the SUBREAD and UNROLLED presets.)
short-sa-cigar	Populates the SA tag with the short cigar representation.

align command: Aligns PacBio reads to reference sequences. The output argument is optional; if not provided, the BAM output is streamed to stdout.

Usage

pbmm2 align [options] <ref.fa|xml|mmi> <in.bam|xml|fa|fq> [out.aligned.bam|xml]

Input files

- *.fasta file containing reference contigs, or *.referenceset.xml, or
 *.mmi index file.
- *.bam, *.subreadset.xml, *.consensusreadset.xml,
 *.transcriptset.xml, *.fasta, *.fa, *.fastq, or *.fastq file containing PacBio data.

Output files

- *.bam aligned reads in BAM format.
- *.alignmentset, *.consensusalignmentset.xml, Or
 - *.transcriptalignmentset.xml if XML output was chosen.

The following Data Set Input/output combinations are allowed:

SubreadSet > AlignmentSet

pbmm2 align hg38.referenceset.xml movie.subreadset.xml hg38.movie.alignmentset.xml

ConsensusReadSet > ConsensusAlignmentSet

pbmm2 align hg38.referenceset.xml movie.consensusreadset.xml
hg38.movie.consensusalignmentset.xml --preset CCS

TranscriptSet > TranscriptAlignmentSet

pbmm2 align hg38.referenceset.xml movie.transcriptset.xml
hg38.movie.transcriptalignmentset.xml --preset ISOSEQ

FASTA/FASTQ input

In addition to native PacBio BAM input, reads can also be provided in FASTA and FASTQ formats.

Attention: The resulting output BAM file **cannot** be used as input into GenomicConsensus!

With FASTA/FASTQ input, the --rg option sets the read group. **Example**:

pbmm2 align hg38.fasta movie.Q20.fastq hg38.movie.bam --preset CCS --rg
'@RG\tID:myid\tSM:mysample'

All three reference file formats .fasta, .referenceset.xml, and .mmi can be combined with FASTA/FASTQ input.

Options	Description
-h,help	Displays help information and exits.
chunk-size	Processes N records per chunk. (Default = 100)
sort	Generates a sorted BAM file.
-m,sort-memory	Specifies the memory per thread for sorting. (Default = 768M)
-j,alignment-threads	Specifies the number of threads used for alignment. 0 means autodetection. (Default = 0)
-J,sort-threads	Specifies the number of threads used for sorting. 0 means 25% of $-j$, with a maximum of 8. (Default = 0)
sample	Specifies the sample name for all read groups. Defaults, in order of precedence: A) SM field in the input read group B) Biosample name C) Well sample name D) "UnnamedSample".
rg	Specifies the read group header line such as '@RG\tID:xyz\tSM:abc'. Only for FASTA/Q inputs.
-y,min-gap-comp-id-perc	Specifies the minimum gap-compressed sequence identity, in percent. (Default = 70)
-1,min-length	Specifies the minimum mapped read length, in base pairs. (Default = 50)
-N,best-n	Specifies the output at maximum ${\tt N}$ alignments for each read. 0 means no maximum. (Default = 0)
strip	Removes all kinetic and extra QV tags. The output cannot be polished.
split-by-sample	Specifies one output BAM file per sample.
no-bai	Omits BAI index file generation for sorted output.
unmapped	Specifies that unmapped records be included in the output.
median-filter	Picks one read per ZMW of median length.
zmw	Processes ZMW Reads; subreadset.xml input is required. This activates the UNROLLED preset.

Options	Description
hqregion	Processes the HQ region of each ZMW; subreadset.xml input is required. This activates the UNROLLED preset.

Parameter set options and overrides	Description
preset	Specifies the alignment mode:
	• "SUBREAD" -k 19 -w 10 -o 5 -O 56 -e 4 -E 1 -A 2 -B 5 -z 400 -Z 50 -r 2000 -L 0.5
	• "CCS" -k 19 -w 10 -u -o 5 -O 56 -e 4 -E 1 -A 2 -B 5 - z 400 -Z 50 -r 2000 -L 0.5
	• "ISOSEQ" -k 15 -w 5 -u -o 2 -O 32 -e 1 -E 0 -A 1 -B 2 -z 200 -z 100 -C 5 -r 200000 -G 200000 -L 0.5
	• "UNROLLED" -k 15 -w 15 -o 2 -O 32 -e 1 -E 0 -A 1 -B 2 -z 200 -z 100 -r 2000 -L 0.5
	(Default = SUBREAD)
-k	Specifies the k-mer size, which cannot be larger than 28. (Default = -1)
-w	Specifies the Minimizer window size. (Default = −1)
-u,no-kmer-compression	Disables homopolymer-compressed k-mer. (Compression is on by default for the SUBREAD and UNROLLED presets.)
-A	Specifies the matching score. (Default = -1)
-В	Specifies the mismatch penalty. (Default = -1)
-z	Specifies the Z-drop score. (Default = −1)
-Z	Specifies the Z-drop inversion score. (Default = −1)
-r	Specifies the bandwidth used in chaining and DP-based alignment. (Default = -1)
-o,gap-open-1	Specifies the gap open penalty 1. (Default = -1)
-0,gap-open-2	Specifies the gap open penalty 2. (Default = −1)
-e,gap-extend-1	Specifies the gap extension penalty 1. (Default = −1)
-E,gap-extend-2	Specifies the gap extension penalty 2. (Default = −1)
-L,lj-min-ratio	Specifies the long join flank ratio. (Default = -1)
-G	Specifies the maximum intron length; this changes -r. (Default = -1)
-C	Specifies the cost for a non-canonical GT-AG splicing. (Default = −1)
no-splice-flank	Specifies that you do not prefer splicing flanks GT-AG.

Examples

Generate an index file for reference and reuse it to align reads:

```
pbmm2 index ref.fasta ref.mmi
pbmm2 align ref.mmi movie.subreads.bam ref.movie.bam
```

Align reads and sort on-the-fly, with 4 alignment and 2 sort threads:

pbmm2 align ref.fasta movie.subreads.bam ref.movie.bam --sort -j 4 -J 2 $\,$

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Align reads, sort on-the-fly, and create a PBI:

pbmm2 align ref.fasta movie.subreadset.xml ref.movie.alignmentset.xml --sort

Omit the output file and stream the BAM output to stdout:

pbmm2 align hg38.mmi movie1.subreadset.xml | samtools sort > hg38.movie1.sorted.bam

Align the CCS reads fastq input and sort the output:

pbmm2 align ref.fasta movie.Q20.fastq ref.movie.bam --preset CCS --sort --rg
'@RG\tID:myid\tSM:mysample'

Alignment parallelization

The number of alignment threads can be specified using the -j, --alignment-threads option. If **not** specified, the maximum number of threads are used, minus one thread for BAM I/O and minus the number of threads specified for sorting.

Sorting

Sorted output can be generated using the --sort option.

- By default, 25% of threads specified with the -j option (Maximum = 8) are used for sorting.
- To override the default percentage, the -J, --sort-threads option
 defines the explicit number of threads used for on-the-fly sorting. The
 memory allocated per sort thread is defined using the -m, --sortmemory option, accepting suffixes M,G.

Benchmarks on human data show that 4 sort threads are recommended, but that no more than 8 threads can be effectively leveraged, even with 70 cores used for alignment. We recommend that you provide more memory to **each** of a **few** sort threads to avoid disk I/O pressure, rather than providing less memory to each of many sort threads.

What are parameter sets and how can I override them?

Per default, pbmm2 uses recommended parameter sets to simplify the multitudes of possible combinations. See the available parameter sets in the option table shown earlier.

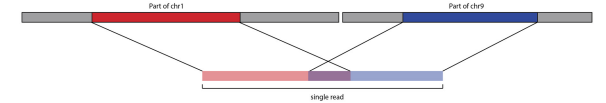
What other special parameters are used implicitly?

We implicitly use the following minimap2 parameters:

- Soft clipping with -Y.
- Long cigars for tag CG with -L.
- X/= cigars instead of M with --eqx.
- No overlapping guery intervals with repeated matches trimming.
- No secondary alignments are produced using the --secondary=no option.

What is repeated matches trimming?

A repeated match occurs when the same query interval is shared between a primary and supplementary alignment. This can happen for translocations, where breakends share the same flanking sequence:



And sometimes, when a LINE gets inserted, the flanks are duplicated, leading to complicated alignments, where we see a split read sharing a duplication. The inserted region itself, mapping to a random other LINE in the reference genome, may also share sequence similarity to the flanks:



To get the best alignments, minimap2 decides that two alignments may use up to 50% (default) of the same query bases. This does **not** work for PacBio, as pbmm2 requires that a single base may never be aligned twice. Minimap2 offers a feature to enforce a query interval overlap to 0%. If a query interval gets used in two alignments, one or both get flagged as secondary and get filtered. This leads to yield loss, and more importantly, missing SVs in the alignment.

Papers (such as this) present dynamic programming approaches to find the optimal split to uniquely map query intervals, while maximizing alignment scores. We don't have per base alignment scores available, thus our approach is much simpler. We align the read, find overlapping query intervals, determine one alignment to be maximal reference-spanning, then trim all others. By trimming, pbmm2 rewrites the cigar and the reference coordinates on-the-fly. This allows us to increase the number of mapped bases, which slightly reduces mapped concordance, but boosts SV recall rate.

How can I set the sample name?

You can override the sample name (SM field in the RG tag) for **all** read groups using the --sample option. If not provided, sample names derive from the Data Set input using the following order of precedence: A) SM field in the input read group B) Biosample name C) Well sample name D) UnnamedSample. If the input is a BAM file and the --sample option was **not** used, the SM field is populated with UnnamedSample.

Can I split output by sample name?

Yes, the --split-by-sample option generates one output BAM file per sample name, with the sample name as the file name prefix, if there is more than one aligned sample name.

Can I remove all those extra per-base and per-pulse tags?

Yes, the --strip option removes the following extraneous tags if the input is BAM: dq, dt, ip, iq, mq, pa, pc, pd, pe, pg, pm, pq, pt, pv, pw, px, sf, sq, st. Note that the resulting output BAM file cannot be used as input into GenomicConsensus.

Where are the unmapped reads?

Per default, unmapped reads are omitted. You can add them to the output BAM file using the --unmapped option.

Can I output at maximum the N best alignments per read?

Use the option -N, --best-n. If set to 0, (the default), maximum filtering is disabled.

Is there a way to only align one subread per ZMW?

Using the --median-filter option, only the subread closest to the median subread length per ZMW is aligned. Preferably, full-length subreads flanked by adapters are chosen.

pbservice

The pbservice tool performs a variety of useful tasks within SMRT Link.

- To get help for pbservice, use pbservice -h.
- To get help for a specific pbservice command, use pbservice <command> -h.

Note: pbservice requires authentication when run from a remote host, using the same credentials used to log in to the SMRT Link GUI. (This also routes all requests through HTTPS port 8243, so the services port is **not** required if authentication is used.) Access to services running on localhost will continue to work without authentication.

All pbservice commands include the following optional parameters:

Options	Description
host=http://localhost	Specifies the server host. Override the default with the environmental variable PB_SERVICE_HOST.
port=8070	Specifies the server port. Override the default with the environmental variable PB_SERVICE_PORT.
log-file LOG_FILE	Writes the log to file. (Default = None, writes to stdout.)
log-level=INFO	Specifies the log level; values are [DEBUG, INFO, WARNING, ERROR, CRITICAL.] (Default = INFO)
debug=False	Alias for setting the log level to DEBUG. (Default = False)
quiet=False	Alias for setting the log level to CRITICAL to suppress output. (Default = False)
user USERNAME	Specifies the user to authenticate as; this is required if the target host is anything other than localhost.
ask-pass	Prompts the user to enter a password.
password PASSWORD	Supplies the password directly. This exposes the password in the shell history (or log files), so this option is not recommended unless you are using a limited account without Unix login access.

status command: Use to get system status.

import-dataset command: Import Local Data Set XML. The file location **must** be accessible from the host where the services are running; often on a shared file system.

Required	Description
xml_or_dir	Specifies a directory or XML file for the Data Set.

import-run command: Create a SMRT Link Run Design and optional Auto Analysis jobs from a CSV file. This is equivalent to the Run Design CSV import feature in the SMRT Link UI; the resulting runs can then be edited in the UI or executed on-instrument.

Required	Description
csv_file	Specifies a Run Design CSV-format file.

Option	Description
dry-run	Prints the generated run XML without POSTing to the server.

import-fasta command: Import a FASTA file and convert to a ReferenceSet file. The file location **must** be accessible from the host where the services are running; often on a shared file system.

Required	Description
fasta_path	Path to the FASTA file to import.

Options	Description
name	Specifies the name of the ReferenceSet to convert the FASTA file to.
organism	Specifies the name of the organism.
ploidy	Ploidy.
block=False	Blocks during importing process.

run-analysis command: Run a secondary analysis pipeline using an analysis.json file.

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```
[--debug] [--quiet] [--block]
json path
```

Required	Description
json_path	Path to the analysis.json file.

Options	Description
block=False	Blocks during importing process.

emit-analysis-template command: Output an analysis.json
template to stdout that can be run using the run-analysis command.

get-job command: Get a job summary by Job Id.

Required	Description
job_id	Job id or UUID.

get-jobs command: Get job summaries by Job Id.

Options	Description
-m=25,max-items=25	Specifies the maximum number of jobs to get.

get-dataset command: Get a Data Set summary by Data Set Id or UUID.

Required	Description
id_or_uuid	Data Set Id or UUID.

get-datasets command: Get a Data Set list summary by Data Set type.

```
[-t DATASET TYPE]
```

Required	Description
-t=subreads,dataset- type=subreads	Specifies the type of Data Set to retrieve: subreads, alignments, references, barcodes.

delete-dataset command: Delete a specified Data Set.

Note: This is a "soft" delete - the database record is tagged as inactive so it won't display in any lists, but the files will **not** be removed.

Required	Description
ID	A valid Data Set ID, either UUID or integer ID, for the Data Set to delete.

Examples

To obtain system status, the Data Set summary, and the job summary:

```
pbservice status --host smrtlink-release --port 9091
```

To import a Data Set XML:

```
pbservice import-dataset --host smrtlink-release --port 9091 \
path/to/subreadset.xml
```

To obtain a job summary using the Job Id:

```
pbservice get-job --host smrtlink-release --port 9091 \
--log-level CRITICAL 1
```

To obtain Data Sets by using the Data Set type subreads:

```
pbservice get-datasets --host smrtlink-alpha --port 8081 \
   --quiet --max-items 1 -t subreads
```

To obtain Data Sets by using the Data Set type alignments:

```
pbservice get-datasets --host smrtlink-alpha --port 8081 \backslash --quiet --max-items 1 -t alignments
```

To obtain Data Sets by using the Data Set type references:

```
pbservice get-datasets --host smrtlink-alpha --port 8081 \ --quiet --max-items 1 -t references
```

To obtain Data Sets by using the Data Set type barcodes:

```
pbservice get-datasets --host smrtlink-alpha --port 8081 \backslash --quiet --max-items 1 -t barcodes
```

To obtain Data Sets by using the Data Set UUID:

pbservice get-dataset --host smrtlink-alpha --port 8081 \ --quiet 43156b3a-3974-4ddb-2548-bb0ec95270ee

pbsv

pbsv is a structural variant caller for PacBio reads. It identifies structural variants and large indels (Default: ≥20 bp) in a sample or set of samples relative to a reference. pbsv identifies the following types of variants: Insertions, deletions, duplications, copy number variants, inversions, and translocations.

pbsv takes as input read alignments (BAM) and a reference genome (FASTA); it outputs structural variant calls (VCF).

Usage

Options	Description
-h,help	Displays help information and exits.
version	Displays program version number and exits.
log-file	Logs to a file, instead of stdout.
log-level	Specifies the log level; values are [TRACE, DEBUG, INFO, WARN, FATAL.] (Default = WARN)
discover	Finds structural variant signatures in read alignments (BAM to SVSIG).
call	Calls structural variants from SV signatures and assign genotypes (SVSIG to VCF).

pbsv discover command: Finds structural variant (SV) signatures in read alignments. The input read alignments must be sorted by chromosome position. Alignments are typically generated with pbmm2. The output SVSIG file contains SV signatures.

Usage

pbsv discover [options] <ref.in.bam|xml> <ref.out.svsig.gz>

Required	Description
ref.in.bam xml	Coordinate-sorted aligned reads in which to identify SV signatures.
ref.out.svsig.gz	Structural variant signatures output.

Options	Description
-h,help	Displays help information and exits.
-s,sample	Overrides sample name tag from BAM read group.
-q,min-mapq	Ignores alignments with mapping quality < N. (Default = 20)
-m,min-ref-span	Ignores alignments with reference length < N bp. (Default = 100)
-w,downsample-window-length	Specifies a window in which to limit coverage, in base pairs. (Default = $10K$)
-a,downsample-max- alignments	Considers up to N alignments in a window; 0 means disabled. (Default = 100)

Options	Description
-r,region	Limits discovery to this reference region: CHR CHR : START-END.
-1,min-svsig-length	Ignores SV signatures with length < ${\tt N}$ bp. (Default = 7)
-b,tandem-repeats	Specifies tandem repeat intervals for indel clustering, as an input BED file.
-k,max-skip-split	Ignores alignment pairs separated by > $\rm N$ bp of a read or reference. (Default = 100)
-y,min-gap-comp-id-perc	Ignores alignments with gap-compressed sequence identity < N%. (Default = 70)

 ${\tt pbsv}$ call command: Calls structural variants from SV signatures and assigns genotypes. The input SVSIG file is generated using ${\tt pbsv}$ discover. The output is structural variants in VCF format.

Usage

pbsv call [options] <ref.fa|xml> <ref.in.svsig.gz|fofn>
<ref.out.vcf>

Required	Description
ref.fa xml	Reference FASTA file or ReferenceSet XML file against which to call variants.
ref.in.svsig.gz fofn	SV signatures from one or more samples. This can be either an SV signature SVSIG file generated by pbsv discover, or a FOFN of SVSIG files.
ref.out.vcf	Variant call format (VCF) output file.

Options	Description
-h,help	Displays help information and exits.
-j,num-threads	Specifies the number of threads to use, 0 means autodetection. (Default = 0)
-r,region	Limits discovery to this reference region: CHR CHR : START-END
-t,types	Calls these SV types: "DEL", "INS", "INV", "DUP", "BND", "CNV". (Default = "DEL, INS, INV, DUP, BND")
-m,min-sv-length	Ignores variants with length < N bp. (Default = 20)
max-ins-length	Ignores insertions with length >N bp. (Default = 15K)
max-dup-length	Ignores duplications with length >N bp. (Default = 1M)
cluster-max-length-perc- diff	Does not cluster signatures with difference in length >₽%. (Default = 25)
cluster-max-ref-pos-diff	Does not cluster signatures > N bp apart in the reference. (Default = 200)
cluster-min-basepair-perc- id	Does not cluster signatures with base pair identity <p%. (default="10)</td"></p%.>
-x,max-consensus-coverage	Limits to N reads for variant consensus. (Default = 20)
-s,poa-scores	Scores POA alignment with triplet match, mismatch, gap. (Default = "1, -2, -2")
min-realign-length	Considers segments with >N length for realignment. (Default = 100)

Options	Description
-A,call-min-reads-all-samples	Ignores calls supported by < N reads total across samples. (Default = 3)
-O,call-min-reads-one-sample	Ignores calls supported by $< N$ reads in every sample. (Default = 3)
-S,call-min-reads-per- strand-all-samples	Ignores calls supported by < $\tt N$ reads per strand total across samples. (Default = 1)
-B,call-min-bnd-reads-all-samples	Ignores BND calls supported by < $\tt N$ reads total across samples. (Default = 2)
-P,call-min-read-perc-one-sample	Ignores calls supported by < P% of reads in every sample. (Default = 20)
preserve-non-acgt	Preserves non-ACGT in REF allele instead of replacing with ${\tt N.}$ (Default = ${\tt false})$
hifi,ccs	Uses options optimized for HiFi reads: -S 0 -P 10. (Default = False)
gt-min-reads	Specifies the minimum supporting reads to assign a sample a non-reference genotype. (Default = 1)
annotations	Annotates variants by comparing with sequences in FASTA. (Default annotations are ALU, L1, and SVA.)
annotation-min-perc-sim	Annotates variant if sequence similarity >P%. (Default = 60)
min-N-in-gap	Considers ≥ N consecutive "N" bp as a reference gap. (Default = 50)
filter-near-reference-gap	Flags variants < N bp from a gap as "NearReferenceGap". (Default = 1000)
filter-near-contig-end	Flags variants < N bp from a contig end as "NearContigEnd". (Default = 1K)

Following is a typical SV analysis workflow starting from subreads:

1. Align PacBio reads to a reference genome, per movie:

Subreads BAM input:

pbmm2 align ref.fa moviel.subreads.bam ref.moviel.bam --sort --median-filter --sample sample1

CCS reads BAM input:

pbmm2 align ref.fa movie1.ccs.bam ref.movie1.bam --sort --preset CCS --sample sample1

CCS reads FASTQ input:

pbmm2 align ref.fa movie1.Q20.fastq ref.movie1.bam --sort --preset CCS --sample sample1
--rg '@RG\tID:movie1'

2. Discover the signatures of structural variation, per movie or per sample:

```
pbsv discover ref.moviel.bam ref.samplel.svsig.gz
pbsv discover ref.movie2.bam ref.sample2.svsig.gz
```

3. Call structural variants and assign genotypes (all samples); for CCS analysis input append --ccs:

```
pbsv call ref.fa ref.sample1.svsig.gz ref.sample2.svsig.gz
ref.var.vcf
```

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Launching a multi-sample pbsv analysis - requirements

- 1. Merge multiple Bio Sample SMRT Cells to one Data Set with the Bio Samples specified.
 - Each SMRT Cell must have exactly one Bio Sample name multiple Bio Sample names cannot be assigned to one SMRT Cell.
 - Multiple SMRT Cells can have the same Bio Sample name.
 - All of the inputs need to already have the appropriate Bio Sample records in their CollectionMetadata. If they don't, they are treated as a single sample.
- 2. Create a ReferenceSet from a FASTA file.
 - The ReferenceSet is often already generated and registered in SMRT Link.
 - If the ReferenceSet doesn't exist, use the dataset create command to create one:

dataset create --type ReferenceSet --name reference_name reference.fasta

Launching a multi-sample analysis

```
# Set subreads and ref FASTA
sample1=sample1.subreadset.xml sample2=sample2.subreadset.xml
ref=reference.fasta

pbmm2 align ${ref} ${sample1} sample1.bam --sort --median-filter --sample Sample1
pbmm2 align ${ref} ${sample2} sample2.bam --sort --median-filter --sample Sample2
samtools index sample1.bam
samtools index sample2.bam
pbindex sample1.bam
pbindex sample2.bam
pbindex sample2.bam
pbsv discover sample1.bam sample1.svsig.gz
pbsv discover sample2.bam sample2.svsig.gz
pbsv call ${ref} sample1.svsig.gz sample2.svsig.gz out.vcf
```

out.vcf: A pbsv VCF output file, where columns starting from column 10 represent structural variants of Sample 1 and Sample 2:

```
#CHROM POS ID REF ALT QUAL FILTER INFO FORMAT Sample1 Sample2 chr01 222737 pbsv.INS.1 T TTGGTGTTTGTTTTTTTTTTTTTTTTTTT . PASS SVTYPE=INS;END=222737;SVLEN=21;SVANN=TANDEM GT:AD:DP 0/1:6,4:10 0/1:6,5:11
```

pbvalidate

The phyalidate tool validates that files produced by PacBio software are compliant with PacBio's own internal specifications.

Input files

phyalidate supports the following input formats:

- BAM
- FASTA
- · Data Set XML

See here for further information about each format's requirements.

Usage

Required	Description
file	Input BAM, FASTA, or Data Set XML file to validate.

Options	Description
-h,help	Displays help information and exits.
version	Displays program version number and exits.
log-file LOG_FILE	Writes the log to file. Default (None) will write to stdout.
log-level	Specifies the log level; values are [DEBUG, INFO, WARNING, ERROR, CRITICAL.] (Default = CRITICAL)
debug=False	Alias for setting the log level to DEBUG. (Default = False)
quiet	Alias for setting the log level to CRITICAL to suppress output. (Default = False)
verbose, -v	Sets the verbosity level. (Default = None)
quick	Limits validation to the first 100 records (plus file header); equivalent tomax-records=100. (Default = False)
max MAX_ERRORS	Exits after MAX_ERRORS were recorded. (Default = None; checks the entire file.)
max-records MAX_RECORDS	Exits after MAX_RECORDS were inspected. (Default = None; checks the entire file.)
type	Uses the specified file type instead of guessing. [BAM, Fasta, AlignmentSet, ConsensusSet, ConsensusAlignmentSet, SubreadSet, BarcodeSet, ContigSet, ReferenceSet, HdfSubreadSet] (Default = None)

Options	Description
index	Requires index files:.fai or .pbi. (Default = False)
strict	Turns on additional validation, primarily for Data Set XML. (Default = False)

BAM options	Description
unaligned	Specifies that the file should contain only unmapped alignments. (Default = None, no requirement.)
unmapped	Alias forunaligned. (Default = None)
aligned	Specifies that the file should contain only mapped alignments. (Default = None, no requirement.)
mapped	Alias foraligned. (Default = None)
contents	Enforces the read type: [SUBREAD, CCS] (Default = None)
reference REFERENCE	Specifies the path to an optional reference FASTA file, used for additional validation of mapped BAM records. (Default = None)

Examples

To validate a BAM file:

pbvalidate in.subreads.bam

To validate a FASTA file:

pbvalidate in.fasta

To validate a Data Set XML file:

pbvalidate in.subreadset.xml

To validate a BAM file and its index file (.pbi):

pbvalidate --index in.subreads.bam

To validate a BAM file and exit after 10 errors are detected:

pbvalidate --max 10 in.subreads.bam

To validate up to 100 records in a BAM file:

pbvalidate --max-records 100 in.subreads.bam

To validate up to 100 records in a BAM file (equivalent to --max-records=100):

pbvalidate --quick in.subreads.bam

To validate a BAM file, using a specified log level:

pbvalidate --log-level=INFO in.subreads.bam

To validate a BAM file and write log messages to a file rather than to ${\tt stdout}$:

pbvalidate --log-file validation_results.log in.subreads.bam

pigeon

The pigeon tool is a PacBio transcript toolkit that contains tools to classify and filter full-length transcript isoforms into categories (see table below) against a reference annotation.

pigeon is based on the SQANTI3 software, and pigeon output is compatible with downstream analysis with the Seurat software.

pigeon characterizes isoforms obtained by full-length transcript sequencing. pigeon combines the long read-defined transcripts with the reference annotation as well as with other orthogonal data to provide a wide range of descriptors of transcript quality. pigeon then generates a comprehensive report to facilitate quality control and filtering of the isoform models.

Pigeon isoform categories

Category	Description
FSM (Full Splice Match)	The reference and query isoform have the same number of exons and each internal junction matches the positions of the reference. The exact 5' start and 3' end can differ within the first/last exons.
ISM (Incomplete Splice Match)	The query isoform has fewer external exons than the reference, but each internal junction matches the positions of the reference. The exact 5' start and 3' end can differ within the first/last exons.
NIC (Novel In Catalog)	The query isoform does not have a FSM or ISM match, but is using a combination of known donor/acceptor sites.
NNC (Novel Not in Catalog)	The query isoform does not have a FSM or ISM match, and has at least one donor or acceptor site that is not annotated.
Antisense	The query isoform does not have overlap with a same-strand reference gene, but is anti-sense to an annotated gene.
Fusion	The query isoform overlaps two or more reference genes.
More junctions	The query isoform overlaps two or more reference genes, however some junctions are shared by multiple genes.
Genic Intron	The query isoform is completely contained within a reference intron.
Genic Genomic	The query isoform overlaps with introns and exons.
Intergenic	The query isoform is in the intergenic region.

Usage

pigeon [options] <index,classify,filter,report,make-seurat>

Options	Description
-h,help	Displays help information and exits.
version	Displays program version number and exits.
log-file	Logs to a file, instead of stdout.
log-level	Specifies the log level; values are [TRACE, DEBUG, INFO, WARN, FATAL.] (Default = WARN)

index command: Creates indices for sorted genomic annotation files. (This is **required** for input to the classification stage.) Acceptable annotation files include GTF for annotations and CAGE peak BED files.

Usage

pigeon index [options] <input file>

Examples

pigeon index gencode.v25.annotation.gtf
pigeon index refTSS_v3.1_human_coordinate.hg38.bed

Required	Description
Input file	Annotation text file. Annotations must be grouped by reference, with the name in the first column, such as BED or GTF.

Options	Description
-h,help	Displays help information and exits.
version	Displays program version number and exits.
log-file	Logs to a file, instead of stdout.
log-level	Specifies the log level; values are [TRACE, DEBUG, INFO, WARN, FATAL.] (Default = WARN)

sort command: Sorts the mapped collapsed GFF file. (This is **required** for input to the classification stage.)

Usage

pigeon sort [options] <Input file>

Example

pigeon sort collapsed.gff -o sorted.gf

Required	Description
Input file	Mapped collapsed isoform GFF file.

Options	Description
-0	Specifies the output file name. If not specified, the output file name is the input file name with the suffix . sorted.
-h,help	Displays help information and exits.
version	Displays program version number and exits.
log-file	Logs to a file, instead of stdout.
log-level	Specifies the log level; values are [TRACE, DEBUG, INFO, WARN, FATAL.] (Default = WARN)

classify command: Classifies isoforms according to their structural categories and detects novel isoforms, as compared to known genome annotations.

Usage

pigeon classify [options] <isoforms.gff> <annotations.gtf> <reference.fa>

Required	Description
isoforms.gff	Sorted collapsed GFF file containing the Isoforms to classify.
annotations.gtf	GTF file containing reference annotations. Note: The associated .pgi file is required . (Use the pigeon index command to generate one.)
reference.fa	FASTA file containing references; this may be gzipped [.gz]. Note: The associated .fai file is required. (Use the samtools faidx command to generate one.)

Input/output options	Description
-r,ref	Specifies a ReferenceSet dataset XML file. May be used in place of the reference annotation and sequence positional arguments, but the dataset must provide these inputs. May (optionally) provide supplemental reference resources such as the polyA motif and CAGE peak input files.
-d,out-dir	Specifies the destination directory for all output files. (Default = [.])
-o,out-prefix	Specifies a prefix for all non-matrix output files. If not specified, the primary input isoform file is used to determine the prefix. Example : An input file of my_isoforms.gff yields the output file name prefix my_isoform.
fl,flnc	Specifies the name of a full-length PacBio abundance file.
cage-peak	Specifies the name of a CAGE (Cap Analysis of Gene Expression) data file in BED format. Note : The associated .pgi file is required . (Use the pigeon index command to generate one.) See here for more information on CAGE.
poly-a	Specifies the name of a text file containing a ranked list of polyA motifs, one per line.
-c,coverage	Specifies junction coverage data. Associated .pgi file is required. Use pigeon index to generate one.
gene-id	Specifies the use of the gene ID instead of the gene name, such as ENSG00000206195.11 rather than DUXAP8.

Algorithm setting	Description
sites	Specifies a comma-separated list of splice sites to be considered as canonical. (Default = GTAG, GCAG, ATAC)
window	Specifies the size of the window (in base pairs) in the genomic DNA screened for Adenine content downstream of TTS. (Default = 20)
min-ref-length	Specifies the minimum reference transcript length, in base pairs. (Default = 200)
is-fusion	Specifies that the inputs are fusion isoforms.
polya-search-distance	Specifies the distance (in base pairs) to look upstream of the 3' end for the highest ranking polyA motif signal. (Default = 50)

Options	Description
-h,help	Displays help information and exits.
version	Displays program version number and exits.
-j,num-threads	Specifies the number of threads to use when processing; 0 means autodetection. (Default = 0)
log-file	Logs to a file, instead of stdout.
log-level	Specifies the log level; values are [TRACE, DEBUG, INFO, WARN, FATAL.] (Default = WARN)

filter command: Removes isoform classifications that indicate amplification artifacts such as intrapriming or template switching.

Note: pigeon filter can only be called after pigeon classify is first called.

Usage

pigeon filter [options] <classification_file>

Examples

pigeon filter classification.txt
pigeon filter classification.txt --isoforms sorted.gff

Required	Description
classification file	Classification.txt output file from pigeon filter.

Options	Description
-h,help	Displays help information and exits.
version	Displays program version number and exits.
log-file	Logs to a file, instead of stdout.
log-level	Specifies the log level; values are [TRACE, DEBUG, INFO, WARN, FATAL.] (Default = WARN)
isoforms	Specifies the generation of a filtered GFF. This requires that you specify the GFF file that was used as input to pigeon classify.

report command: Reports gene saturation. Gene saturation can be determined by subsampling the classification output and determining the number of unique genes at each subsample size.

Usage

pigeon report [options] <classification.txt> <subsample.txt>

Example

pigeon report <classification.filtered_lite_classification.txt> <saturation.txt>

Required	Description
classification.txt	Input classification text file. Could be after pigeon classify or pigeon filter.
subsample.txt	Output subsampling text report file.

Options	Description
-s,sub-sample-increment	Specifies the number of reads between subsampling data points. (Default = 10000)
-h,help	Displays help information and exits.
version	Displays program version number and exits.
log-file	Logs to a file, instead of stdout.
log-level	Specifies the log level; values are [TRACE, DEBUG, INFO, WARN, FATAL] (Default = WARN)

make-seurat command: Generates the following output files required to run tertiary analysis using the Seurat software:

- barcodes.tsv
- genes.tsv
- matrix.mtx
- info.csv
- annotated.info.csv

Usage

pigeon make-seurat [options] <classification.txt> -a <annotation GTF|XML> --dedup <dedup fasta> --group <collapse group TXT>

Example

pigeon make-seurat --dedup dedup.fasta --group
scisoseq.mapped_transcripts.collapse.group.txt
scisoseq_classification.filtered_lite_classification.txt -d output -a
absolutized.referenceset.xml

Required	Description
classification.txt	Text file containing transcript classifications. Could be generated by pigeon classify or pigeon filter.
dedup	Specifies a molecule FASTA file generated by isoseq3 groupdedup or isoseq3 dedup.
-g,group	Specifies an output groups text file generated by isoseq collapse.
-a,annotation	Specifies reference annotations for Seurat gene matrix. May be a GTF/GFF or Reference-Set XML file.

Input/output options	Description
-o,out-prefix	Specifies a prefix for all non-matrix files. If not specified, the primary input classification file is used to determine the prefix. Example:
	my_isoforms_classification.txt input yields an output prefix of my_isoforms.

Input/output options	Description
-d,out-dir	Specifies the destination directory for all output files. (Default = [.])

Matrix options	Description
keep-novel-genes	Specifies that novel genes be kept in the matrix output.
keep-ribo-mito-genes	Specifies that ribosomal and mitochondrial genes be kept in the matrix output.

Options	Description
-h,help	Displays help information and exits.
version	Displays program version number and exits.
log-file	Logs to a file, instead of stdout.
log-level	Specifies the log level; values are [TRACE, DEBUG, INFO, WARN, FATAL.] (Default = WARN)

primrose

The primrose tool analyzes the kinetic signatures of cytosine bases in CpG motifs to identify the presence of 5mC.

primrose uses a convolution neural network (CNN) to predict the methylation state (5mC) of each CpG in a HiFi read. Methylation is assumed to be symmetric between strands with output reported in the forward direction with respect to the HiFi read sequence. The output uses the Mm and M1 tags as defined in the SAM Format Optional Fields Specification. (See here for details.)

Algorithm

The CNN is trained using invitro modified controls of methylated and unmethylated human DNA. The unmethylated control comprises a human shotgun library that has undergone whole genome amplification (WGA), a process that includes PCR, and therefore removes all modifications. The methylated control is generated by subjecting a human WGS library to invitro methylation using a CpG methyltransferase enzyme (M. SSSI). Kinetic data, pulse width and inter-pulse distance, over a 16 base pair window for both the forward and reverse strand, is used as input to the CNN. The output of the CNN is a probability scale measure of whether the CpG is symmetrically 5mC-modified.

Usage

primrose [options] <HiFi INPUT> <HiFi OUTPUT>

Required	Description
HiFi INPUT	Input BAM file or ConsensusReadSet XML file.
HiFi OUPUT	Output BAM file or ConsensusReadSet XML file.

Options	Description
keep-kinetics	Specifies that the kinetics tracks (IPD and PulseWidth records) fi , fp , fn , ri , rp and rn are included in the output BAM file.
min-passes	Specifies the minimum number of passes. (Default = 3)
model	Specifies a path to a trained TensorFlow model directory, or to an exported ONNX model file.
-h,help	Displays help information and exits.
version	Displays program version number and exits.
-j,num-threads	Specifies the number of threads to use when processing; 0 means autodetection. (Default = 0)
log-file LOG_FILE	Writes the log to file. Default (None) will write to stderr.
log-level	Specifies the log level; values are [TRACE, DEBUG, INFO, WARNING, , FATAL.] (Default = WARNING)

Example

primrose --log-level INFO movie.hifi.bam movie.hifi_5mc.bam

runqc-reports

The <code>runqc-reports</code> tool generates up to five different Run QC reports, depending on Data Set type: Raw Data, Adapters, Loading, Control, and CCS reads. Generating a complete set of reports requires the presence of an sts.xml resource in the Data Set, but either the CCS Analysis report (or a fallback Subreads report) will **always** be generated. All report JSON and plot PNG files are written to the current working directory, unless otherwise specified.

Usage

Required	Description
dataset_xml	Input SubreadSet or ConsensusReadSet XML, which must contain an sts.xml resource for the full Run QC report set to be generated.
-o OUTPUT_DIR	Output report directory. (Default = Current working directory)

Options	Description
-h,help	Displays help information and exits.
version	Displays program version number and exits.
log-file LOG_FILE	Writes the log to file. Default (None) will write to stdout.
log-level	Specifies the log level; values are [DEBUG, INFO, WARNING, ERROR, CRITICAL.] (Default = WARNING)
debug	Alias for setting the log level to DEBUG. (Default = False)
quiet	Alias for setting the log level to CRITICAL to suppress output. (Default = False)
verbose, -v	Sets the verbosity level. (Default = None)

Examples

```
runqc-reports moviename.subreadset.xml
runqc-reports moviename.consensusreadset.xml
```

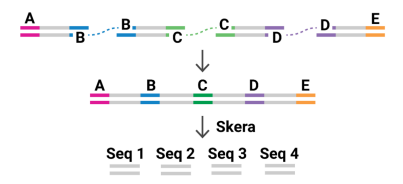
skera

The skera tool splits arrayed HiFi reads at adapter positions, generating segmented reads (S-reads) which are the individual fragments. For each input HiFi read, the tool creates multiple BAM records, one for each fragment. An arrayed HiFi read can contain many fragments (or S-reads).

The skera tool also includes a command to reconstitute the original parent reads from input fragments.

Segmented reads

An arrayed HiFi read is comprised of multiple fragments. The figure below displays an arrayed HiFi read with four fragments (Seq 1 through Seq 4).



Segmented read BAM header

Arrayed HiFi reads have a READTYPE of CCS in the DS (Description) tag in the BAM header. After running skera split, segmented reads (S-reads) will have a READTYPE of SEGMENT. Additionally, a SOURCE key is added with a value of CCS.

Header DS tag for arrayed HiFi read:

DS:READTYPE=CCS;

Header DS tag for segmented read (S-Read):

DS:READTYPE=SEGMENT;SOURCE=CCS;

Segmented read names

For segmented reads (S-reads), the base QNAME follows the CCS read conventions, while also appending the 0-based coordinate interval (qStart, qEnd) that represents its span within the arrayed HiFi read.

Default:

{movieName}/{holeNumber}/ccs/{qStart} {qEnd}

If the CCS read is split by strand:

{movieName}/{holeNumber}/ccs/fwd/{qStart}_{qEnd}
{movieName}/{holeNumber}/ccs/rev/{qStart} {qEnd}

BAM tag	Туре	Description	Example
di	Integer	Index of the read segment.	di:i:0
qs	Integer	qStart of the segment in the parent read.	qs:i:16
qe	Integer	qEnd of the segment in the parent read.	qe:i:450
dl	Integer	Leading segmentation adapter index.	dl:i:0
dr	Integer	Trailing segmentation adapter index.	dr:i:1
ds	Binary	Binary JSON file.	ds:b:10,21,23

Notes:

- All tags of type Integer are 0-based.
- qs and qe tags are left open, right closed.
- ${\tt dl}$ and ${\tt dr}$ tags match the ordering of adapter sequences in the user input Adapter FASTA.
- The ds tag contains a number of fields used for supporting the skera undo command, which are **not** intended for general use.

skera split command: Splits sequences at adapter locations.

Usage

skera split [options] <Input dataset> <ADAPTERS.fasta> <Output dataset>

Required	Description
Input dataset	Input HiFi reads in BAM/ConsensusReadSet XML format.
ADAPTERS.fasta	Segmentation adapters FASTA or AdapterSet XML file containing segmentation adapters. If not specified, the adapters specified in the XML metadata are used. Segmentation adapters must be ordered in the expected order of adapters in the reads. There should be one entry per adapter (forward or reverse-complement orientation) with no overlapping adapter sequences. Duplicate names or sequences are not allowed.
	Example:
	>A
	AGCTTACTTGTGAAGA
	>B
	ACTTGTAAGCTGTCTA
	>C
	ACTCTGTCAGGTCCGA
	>D
	ACCTCCTCCAGAA
	>E
	>AACCGGACACTTAG
Output dataset	Output S-reads in BAM/ConsensusReadSet XML format.

Options	Description
-h,help	Displays help information and exits.
version	Displays program version number and exits.
-j,num-threads	Specifies the number of threads to use when processing; 0 means autodetection. (Default = 0)
log-file LOG_FILE	Writes the log to file. Default (None) will write to stderr.
log-level	Specifies the log level; values are [TRACE, DEBUG, INFO, WARN, FATAL]. (Default = WARN)

skera undo command: Reconstitutes the original parent reads from input fragments (S-reads). **Note**: To run skera undo, **both** the passing and non passing BAM files **must** be present in the same directory.

Usage

skera undo [options] <Input dataset> <Output dataset>

Required	Description
Input dataset	skera split reads in BAM/ConsensusReadSet XML format.
Output dataset	Arrayed HiFi reads in BAM/ConsensusReadSet XML format.

Options	Description
-h,help	Displays help information and exits.
version	Displays program version number and exits.
-j,num-threads	Specifies the number of threads to use when processing; 0 means autodetection. (Default = 0)
log-file LOG_FILE	Writes the log to file. Default (None) will write to stderr.
log-level	Specifies the log level; values are [TRACE, DEBUG, INFO, WARN, FATAL]. (Default = WARN)

Input files for skera split

- *.bam: Arrayed HiFi reads.
- <adapter>.fasta: A FASTA file containing segmentation adapters.

Output files for skera split

- * .bam: BAM file containing S-reads that were segmented from arrayed HiFi reads.
- *.non_passing.bam: BAM file containing HiFi reads that failed segmentation. Any arrayed HiFi read in which at least one segment is flanked by adapter sequences that are out of order (as specified by the segmentation adapter FASTA file) is output to the *non_passing.bam file to enable the skera undo command.

Note: To run skera undo, both the passing and non_passing BAM files must be present in the same directory.

 *.summary.json, *summary.csv: Contains the following summary statistics:

- **Reads**: The number of input arrayed HiFi reads.
- **Segmented Reads (S-reads)**: The number of generated S-reads.
- **Mean Length of S-reads**: The mean read length of the generated S-reads.
- Percentage of Reads with Full Array: The percentage of input reads containing all adapter sequences in the order listed in the segmentation adapter FASTA file.
- Mean Array Size: The mean number of fragments (or S-reads) found in the input reads.
- *.ligations.csv: Contains counts of adapter ligations found in arrayed HiFi reads. For each sequence found between two adapters, a ligation between two adapters is counted. The ligations are counted in the order of the adapters in the arrayed HiFi read (adapter_1, adapter 2).
- *.read_lengths.csv: Contains the HiFi read length, S-Read length, and array size for each HiFi read.

Examples

skera split <movie>.hifi_reads.bam adapters.fasta <movie>.skera.bam

skera undo <movie>.skera.bam <movie>.undo.bam

summarize Modifications

The summarizeModifications tool generates a GFF summary file (alignment_summary.gff) from the output of base modification analysis (for example, ipdSummary) combined with the coverage summary GFF generated by resequencing pipelines. This is useful for power users running custom workflows.

Usage

Input files

- modifications: Base Modification GFF file.
- alignmentSummary: Alignment Summary GFF file.

Output files

• gff_out: Coverage summary for regions (bins) spanning the reference with Base Modification results for each region.

Options	Description
-h,help	Displays help information and exits.
version	Displays program version number and exits.
log-file LOG_FILE	Writes the log to file. Default (None) will write to stdout.
log-level	Specifies the log level; values are [DEBUG, INFO, WARNING, ERROR, CRITICAL] (Default = INFO)
debug	Alias for setting the log level to DEBUG. (Default = False)
quiet	Alias for setting the log level to CRITICAL to suppress output. (Default = False)
verbose, -v	Sets the verbosity level. (Default = None)

Appendix A - Application entry points and output files

Note: To print information about a specific PacBio workflow, including input files and task options, use the pbcromwell show-workflow-details command, which is available for **all** applications.

Examples

```
pbcromwell show-workflow-details pb_demux_ccs
pbcromwell show-workflow-details cromwell.workflows.pb export ccs
```

(The prefix cromwell.workflows is optional.)

CCS Analysis

Analysis application name: cromwell.workflows.pb ccs

Entry point

```
:id: eid_subread
:name: Entry eid_subread
:fileTypeId: PacBio.DataSet.SubreadSet
```

Key output files

File name	Datastore Sourceld
FASTQ file	ccs_fastq_out
FASTA file	ccs_fasta_out
<moviename>.hifi.reads.bam file</moviename>	ccs_bam_out
Consensus Sequences	pb_ccs.ccsxml
CCS Analysis Statistics	pb_ccs.report_ccs
All Reads (BAM)	reads_bam
<moviename>.hifi.reads.fasta</moviename>	ccs_fasta_out
<moviename>.hifi.reads.fastq</moviename>	ccs_fastq_out

Demultiplex Barcodes

Analysis application name: cromwell.workflows.pb demux ccs

Entry points

```
:id: eid_ccs
:name: Entry eid_ccs
:fileTypeId: PacBio.DataSet.ConsensusReadSet
:id: eid_barcode
:name: Entry eid_barcode
:fileTypeId: PacBio.DataSet.BarcodeSet
```

File name	Datastore Sourceld
Barcode Report Details	pb_demux_ccs.summary_csv
Demultiplexed Datasets	pb_demux_ccs.demuxed_files_datastore
Unbarcoded Reads	pb_demux_ccs.unbarcoded

Export Reads

Analysis application name: cromwell.workflows.pb_export_ccs

Entry point

:id: eid_ccs
:name: Entry eid_ccs
:fileTypeId: PacBio.DataSet.ConsensusReadSet

Key output files

File name	Datastore Sourceld
<pre><moviename>.hifi_reads.fastq.gz</moviename></pre>	ccs_fastq_out
<pre><moviename>.hifi_reads.fasta.gz</moviename></pre>	ccs_fasta_out
<pre><moviename>.hifi_reads.bam.gz</moviename></pre>	ccs_bam_out

Note: If users select a lower cutoff Phred QV value, the string hifi is replaced by the QV value in the file names.

Example: <moviename>.q10.fastq.gz.

Genome Assembly

Analysis application name: cromwell.workflows.pb_assembly_hifi

Entry point

:id: eid_ccs
:name: Entry eid_ccs
:fileTypeId: PacBio.DataSet.ConsensusReadSet

Key output files

File name	Datastore Sourceld
Final Polished Assembly, Primary Contigs	pb_assembly_hifi.final_primary_contigs_fasta
Final Polished Assembly, Haplotigs	pb_assembly_hifi.final_haplotigs_fasta
List of Circular Contigs	pb_assembly_hifi.circular_contigs
Summary Report	pb_assembly_hifi.report_polished_assembly

HiFi Mapping

Analysis application name: cromwell.workflows.pb ccs subreads

Entry points

:id: eid_ccs
:name: Entry eid_ccs
:fileTypeId: PacBio.DataSet.ConsensusReadSet
:id: eid_ref_dataset
:name: Entry eid_ref_dataset
:fileTypeId: PacBio.DataSet.ReferenceSet

File name	Datastore Sourceld
Mapped reads	pb_align_ccs.mapped
Coverage summary	pb_align_ccs.coverage_gff

HiFiViral SARS-CoV-2 Analysis

Analysis application name: cromwell.workflows.pb_sars_cov2_kit

Entry points

```
:id: eid_ccs (HiFi reads, demultiplexed as separate BAM files)
:name: Entry eid_ccs
:fileTypeId: PacBio.DataSet.ConsensusReadSet
:id: eid_ref_dataset_2 (Reference Genome)
:name: Entry eid_ref_dataset
:fileTypeId: PacBio.DataSet.ReferenceSet
```

Key output files

File name	Datastore Sourceld
All Samples, Probe Counts TSV	pb_sars_cov2_kit.probe_counts_zip
All Samples, Raw Variant Calls VCF	pb_sars_cov2_kit.raw_vcf_zip
All Samples, Variant Call VCF	pb_sars_cov2_kit.vcf_zip
All Samples, Variant Calls CSV	pb_sars_cov2_kit.variants_csv
All Samples, Consensus Sequence FASTA	pb_sars_cov2_kit.fasta_zip
All Samples, Consensus Sequence By Fragments FASTA	pb_sars_cov2_kit.frag_fasta_zip
All Samples, Aligned Reads BAM	pb_sars_cov2_kit.mapped_zip
All Samples, Consensus Sequence Aligned BAM	pb_sars_cov2_kit.aligned_frag_zip
All Samples, Trimmed HiFi reads FASTQ	pb_sars_cov2_kit.trimmed_zip
Failed Sample Info	pb_sars_cov2_kit.sample_failures_csv
Failed Sample Analysis Logs	pb_sars_cov2_kit.errors_zip
Demultiplex Summaries	pb_sars_cov2_kit.lima_summary_zip
Sample Summary Table CSV	pb_sars_cov2_kit.summary_csv
All Samples, Genome Coverage Plots	pb_sars_cov2_kit.coverage_png_zip

Iso-Seq Analysis

Analysis application name: cromwell.workflows.pb_isoseq3_ccsonly

Entry points

```
:id: eid_ccs
:name: Reads
:fileTypeId: PacBio.DataSet.ConsensusReadSet

:id: eid_barcode
:name: Primers
:fileTypeId: PacBio.DataSet.BarcodeSet

:id: eid_ref_dataset
:name: Reference (Optional)
:fileTypeId: PacBio.DataSet.ReferenceSet
```

Key output files

File name	Datastore Sourceld
Collapsed Filtered Isoforms FASTQ	pb_isoseq3_ccsonly.collapse_fastq
Collapsed Filtered Isoforms GFF	pb_isoseq3_ccsonly.collapse_gff
Group TXT	pb_isoseq3_ccsonly.collapse_group
Abundance TXT	pb_isoseq3_ccsonly.collapse_abundance
Read Stat TXT	pb_isoseq3_ccsonly.collapse_readstat
Collapsed Isoforms Abundance TXT (Files are numbered consecutively, 1 for each barcoded sample.)	pb_isoseq3_ccsonly.collapse_abundance Separate clusters, one per barcoded sample.
Collapsed Isoforms Abundance TXT (Files are numbered consecutively, 1 for each barcoded sample.)	pb_isoseq3_ccsonly.collapse_abundance Pooled clusters, one per barcoded sample.
High-Quality Transcripts	pb_isoseq3_ccsonly.hq_fastq
Low-Quality Transcripts	pb_isoseq3_ccsonly.lq_fastq
High-Quality Transcripts Counts (Files are numbered consecutively, 1 for each barcoded sample.)	pb_isoseq3_ccsonly.barcode_overview_report Separate clusters, one per barcoded sample.
High-Quality Transcripts Counts (Files are numbered consecutively, 1 for each barcoded sample.)	pb_isoseq3_ccsonly.barcode_overview_report Pooled clusters, one per barcoded sample.
CCS reads FASTQ	pb_isoseq3_ccsonly.ccs_fastq_zip
Full-length CCS reads	pb_isoseq3ccsonly.flnc_bam
Polished Report	pb_isoseq3ccsonly.polish_report_csv
Cluster Report	pb_isoseq3ccsonly.report_isoseq

Mark PCR **Duplicates**

Analysis application name: cromwell.workflows.pb_mark_duplicates

Entry points

:id: eid_ccs
:name: Entry eid_ccs
:fileTypeId: PacBio.DataSet.ConsensusReadSet

File name	Datastore Sourceld
PCR Duplicates	pb_mark_duplicates.duplicates
Deduplicated reads	<pre>pb_mark_duplicates.deduplicated In the SMRT Link UI, this displays as <original_dataset_name> (deduplicated).</original_dataset_name></pre>

Microbial Genome **Analysis**

Analysis application name: cromwell.workflows.pb_microbial_analysis

Entry point

:id: eid_ccs

:name: Entry eid_ccs
:fileTypeId: PacBio.DataSet.ConsensusreadSet

Key output files

File name	Datastore Sourceld
Final Polished Assembly	pb_microbial_analysis.assembly_fasta
Final Polished Assembly Index	pb_microbial_analysis.assembly_fasta.fai
Final Polished Assembly for NCBI	pb_microbial_analysis.ncbi_fasta
Mapped BAM	pb_microbial_analysis.mapped
Target sequences used to produce the Mapped BAM	pb_microbial_analysis.mapped_target_fasta
Target sequence Index used to produce the Mapped BAM	pb_microbial_analysis.mapped_target_fasta_fai
List of Circular Contigs	pb_microbial_analysis.circular_list
Coverage Summary	pb_microbial_analysis.coverage_gff
Coverage Report	pb_microbial_analysis.report_coverage
Mapping Statistics Report	pb_microbial_analysis.report_mapping_stats
Mapped BAM Datastore	pb_microbial_analysis.mapped_bam_datastore
Polished Assembly Report	pb_microbial_analysis.report_polished_assembly
Full Kinetics Summary	pb_microbial_analysis.basemods_gff
IPD Ratios	pb_microbial_analysis.basemods_csv
Motifs and Modifications	pb_microbial_analysis.motifs_csv
Motifs and Modifications	pb_microbial_analysis.motifs_gff

Read Segmentation

Analysis application name: cromwell.workflows.pb segment reads

Entry points

:id: eid_ccs :name: HiFi Reads

:fileTypeId: PacBio.DataSet.ConsensusReadSet

:id: Barcode

:name: MAS-Seq Adapters

:fileTypeId: PacBio.DataSet.BarcodeSet

File name	Datastore Sourceld
Input Adapters	pb_segment_reads.adapters_fasta
Report Read Segmentation	pb_segment_reads.report_read_segmentation
Non-Passing Reads	pb_segment_reads.non_passing_bam

File name	Datastore Sourceld
Segmented Reads	pb_segment_reads.segmented_reads
Segmented Reads BAM	pb_segment_reads.segmented_reads_bam

Read Segmentation and Single-Cell Iso-Seq Analysis

Analysis application name:

cromwell.workflows.pb_segment_reads_and_sc_isoseq

Entry points

```
:id: eid_ccs
:name: HiFi Reads
:fileTypeId: PacBio.DataSet.ConsensusReadSet
:id: eid_ref_dataset
:name: Reference Genome
:fileTypeId: PacBio.DataSet.ReferenceSet
:id: eid_barcode
:name: MAS-Seq Adapters
:fileTypeId: PacBio.DataSet.BarcodeSet
:id: eid_barcode_2
:name: Single-CeIl Iso-Seq Primers
:fileTypeId: PacBio.DataSet.BarcodeSet
```

File name	Datastore Sourceld
Input Adapters	pb_segment_reads_and_sc_isoseq.adapters_fasta
Report Read Segmentation	pb_segment_reads_and_sc_isoseq.report_read_segmentation
Barcode statistics	scisoseq.bcstats_report_tsv
Transcript classifications	scisoseq.classification_txt
Transcript classification file, after SQANTI filtering criteria	scisoseq_classification.filtered_lite_classification.txt
Transcript junction file	scisoseq_junctions.txt
Transcript junction file, after SQANTI filtering criteria	scisoseq_classification.filtered_lite_junctions.txt
Unique transcripts, sorted GFF file	scisoseq_transcripts.sorted.gff
Unique transcripts, after SQANTI filtering criteria, GFF file	scisoseq_transcripts.sorted.filtered_lite.gff
Segmented reads, unmapped BAM file	segmented.bam
Reads that did not pass segmentation, unmapped BAM file	segmented.non_passing.bam
Read segmentation report	read_segmentation.report.json
Deduplicated reads mapped BAM file and index file	scisoseq.mapped.bam, scisoseq.mapped.bam.bai
Deduplicated reads, unmapped BAM file	scisoseq.5p3p.tagged.refined.corrected.sorted.dedup.bam

File name	Datastore Sourceld
Single-cell isoform and gene matrix, gzipped	scisoseq.seurat_info.tar.gz

Single-Cell Iso-Seq Analysis

Analysis application name: cromwell.workflows.pb_sc_isoseq

Entry points

:id: eid_ccs :name: HiFi Reads

:fileTypeId: PacBio.DataSet.ConsensusReadSet

:id: eid ref dataset :name: Reference Genome

:fileTypeId: PacBio.DataSet.ReferenceSet

:id: eid_barcode

:name: Single-Cell Iso-Seq Primers :fileTypeId: PacBio.DataSet.BarcodeSet

File name	Datastore Sourceld
Barcode statistics	scisoseq.bcstats_report_tsv
Transcript classifications	scisoseq.classification_txt
Transcript classification file, after SQANTI filtering criteria	scisoseq_classification.filtered_lite_classification.txt
Transcript junction file	scisoseq_junctions.txt
Transcript junction file, after SQANTI filtering criteria	scisoseq_classification.filtered_lite_junctions.txt
Unique transcripts, sorted GFF file	scisoseq_transcripts.sorted.gff
Unique transcripts, after SQANTI filtering criteria, GFF file	scisoseq_transcripts.sorted.filtered_lite.gff
Segmented reads, unmapped BAM file	segmented.bam
Reads that did not pass segmentation, unmapped BAM file	segmented.non_passing.bam
Read segmentation report	read_segmentation.report.json
Deduplicated reads mapped BAM file and index file	scisoseq.mapped.bam, scisoseq.mapped.bam.bai
Deduplicated reads, unmapped BAM file	scisoseq.5p3p.tagged.refined.corrected.sorted.dedup.bam
Single-cell isoform and gene matrix, gzipped	scisoseq.seurat_info.tar.gz

Structural Variant Calling

Analysis application name: cromwell.workflows.pb_sv_ccs

Entry points

```
:id: eid_ccs
:name: Entry eid_ccs
:fileTypeId: PacBio.DataSet.ConsensusReadSet
:id: eid_ref_dataset
:name: Entry eid_ref_dataset
:fileTypeId: PacBio.DataSet.ReferenceSet
```

Key output files

File name	Datastore Sourceld
Structural Variants	pb_sv_ccs.variants
Aligned reads (Bio Sample Name)	pb_sv_ccs.alignments_by_sample_datastore

Trim Ultra-Low Adapters

Analysis application name: cromwell.workflows.pb_trim_adapters

Entry points

```
:id: eid_ccs
:name: Entry eid_ccs
:fileTypeId: PacBio.DataSet.ConsensusReadSet
:id: eid_barcode
:name: Entry eid_barcode
:fileTypeId: PacBio.DataSet.BarcodeSet
```

Note: The barcodes need to be a single primer sequence.

Key output files

File name	Datastore Sourceld
Reads Missing Adapters	pb_trim_adapters.unbarcoded
PCR Adapter Data CSV	pb_trim_adapters.summary_csv
Trimmed reads	<pre>pb_trim_adapters.trimmed In the SMRT Link UI, this displays as <original_dataset_name> (trimmed).</original_dataset_name></pre>

Variant Calling

Analysis application name: cromwell.workflows.pb variant calling

Entry points

```
:id: eid_ccs
:name: Entry eid_ccs
:fileTypeId: PacBio.DataSet.ConsensusReadSet
:id: eid_ref_dataset
:name: Entry eid_ref_dataset
:fileTypeId: PacBio.DataSet.ReferenceSet
```

Key output files

File name	Datastore Sourceld
Structural Variants VCF	pb_variant_calling.structural_variants_vcf
DeepVariant gVCF	pb_variant_calling.variants_gvcf
DeepVariant Phased VCF	pb_variant_calling.phased_variants_vcf

5mC CpG Detection

Analysis application name: cromwell.workflows.pb_detect_methyl

Entry points

:id: eid_ccs
:name: Entry eid_ccs
:fileTypeId: PacBio.DataSet.ConsensusReadSet

:id: eid_barcode

:name: Entry eid barcode :fileTypeId: PacBio.DataSet.BarcodeSet

File name	Datastore Sourceld
HiFi reads with 5mC Calls	pb_detect_methyl.ccsbam

Appendix B - Third party command-line tools

Following is information on the third-party command-line tools included in the smrtcmds/bin subdirectory.

bamtools

- A C++ API and toolkit for reading, writing, and manipulating BAM files.
- See https://sourceforge.net/projects/bamtools/ for details.

cromwell

- · Scientific workflow engine used to power SMRT Link.
- See https://cromwell.readthedocs.io/en/stable/ for details.

ipython

- · An interactive shell for using the PacBio API.
- See https://ipython.org/ for details.

purge_dups

- Removes haplotigs and contig overlaps in a de novo assembly based on read depth.
- See https://github.com/dfguan/purge_dups for details.

python

- · An object-oriented programming language.
- See https://www.python.org/ for details.

samtools, BCFtools

- A set of programs for interacting with high-throughput sequencing data in SAM/BAM/CRAM/VCF/BCF2 formats.
- See http://www.htslib.org/ for details.

Appendix C - Microbial Genome Analysis advanced options (HiFi reads only)

Use this application to generate de novo assemblies of small prokaryotic genomes between 1.9-10 Mb and companion plasmids between 2 – 220 kb from Hifi (CCS) reads.

The workflow also performs base modification analysis of the assembled genome.

The Microbial Genome Analysis application:

- Includes chromosomal- and plasmid-level de novo genome assembly, circularization, polishing, and rotation of the origin of replication for each circular contig.
- Performs base modification detection (m4C and m6A).
- · Facilitates assembly of larger genomes (yeast) as well.
- · Accepts HiFi read data (BAM format) as input.

The workflow consists of two assembly stages: **chromosomal** and **plasmid**.

Chromosonal stage: Intended for contig assembly of large sequences. This stage uses more stringent filtering (using advanced options) to produce contiguous assemblies of complex regions, but it may miss small sequences in the input sample (such as plasmids.)

Plasmid stage: Intended for a fine-grained assembly. This stage assembles **only** the unmapped and poorly mapped reads. It also relaxes the overlapping parameters, using advanced options.

Both stages use an automated random subsampling process to reduce the input Data Set for assembly (by default to 100x). Note that the subsampling is **only** applied to the contig construction process, while the alignment and reports stages of the workflow still uses the **full** input Data Set.

Available options for the two assembly stages are **identical**. The only differences are:

- 1. Chromosonal stage parameters are prefixed with:

 ipa2_advanced_options_chrom. In the SMRT Link interface for the

 Microbial Genome Analysis application, this corresponds to the

 Advanced Assembly Options for chromosomal stage field.
- 2. Plasmid stage parameters are prefixed with:

 ipa2_advanced_options_plasmid. In the SMRT Link interface for the

 Microbial Genome Analysis application, this corresponds to the

 Advanced Assembly Options for plasmid stage field.

The same sub-options are available to each stage, but the defaults are very different. The current defaults are:

```
ipa2_advanced_options_chrom =
"config_block_size = 100; config_seeddb_opt = -k 28 -w 20 --space 0 --use-hpc-seeds-
only; config_ovl_opt = --one-hit-per-target --min-idt 98 --traceback --mask-hp --mask-
repeats --trim --trim-window-size 30 --trim-match-frac 0.75;"

ipa2_advanced_options_plasmid =

"config_block_size = 100; config_ovl_filter_opt = --max-diff 80 --max-cov 100 --min-cov
2 --bestn 10 --min-len 500 --gapFilt --minDepth 4 --idt-stage2 98; config_ovl_min_len =
500; config_seeddb_opt = -k 28 -w 20 --space 0 --use-hpc-seeds-only; config_ovl_opt = -
one-hit-per-target --min-idt 98 --min-map-len 500 --min-anchor-span 500 --traceback --
mask-hp --mask-repeats --trim --trim-window-size 30 --trim-match-frac 0.75 --smart-hit-
per-target --secondary-min-ovl-frac 0.05; config_layout_opt = --allow-circular;"
```

Options are separated by **semicolons**; within each option, parameters are separated by **spaces**.

Users should not need to modify any of default options. If the defaults are modified, workflow behavior could be very different.

Note: The options available in <code>ipa2_advanced_options_*</code> are exactly the same as the <code>config_*</code> options available for the Genome Assembly tool. See "Genome Assembly parameters input files" on page 44 for details.