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Technical overview – Kinnex kits for single-cell RNA, full-length RNA and 16S rRNA sequencing

Sequel II and IIe systems ICS v11.0 Revio system ICS v13.0+ SMRT Link v13.0+

Kinnex kits for single-cell RNA, full-length RNA and 16S rRNA sequencing

Technical overview

- 1. Kinnex product introduction
- 2. Kinnex product configuration overview
- 3. Kinnex library preparation & sequencing workflow overview
- 4. Kinnex single-cell RNA workflow key highlights
- 5. Kinnex full-length RNA workflow key highlights
- 6. Kinnex 16S rRNA workflow key highlights
- 7. Technical documentation & applications support resources

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Kinnex product introduction

Kinnex kits for single-cell RNA, full-length RNA, and 16S rRNA sequencing

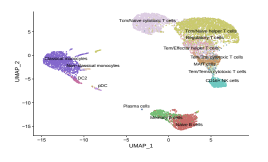
Kinnex kits offer scalable, cost-effective RNA sequencing solutions



Single-cell RNA sequencing [(10x Chromium Single Cell 3' or 5' cDNA input)]

> Up to 4-plex Kinnex library [16-fold concatenation]

80M reads (Revio SMRT Cell) [40M reads (Sequel II/IIe SMRT Cell 8M)]



Identify cell type-specific isoform expression

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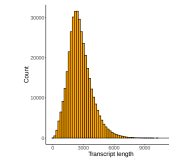


Kinnex full-length RNA kit²

Full-length RNA sequencing

Up to 48-plex Kinnex library [8-fold concatenation]

40M reads (Revio SMRT Cell) [15M reads (Sequel II/IIe SMRT Cell)]



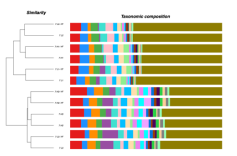
Full-length transcripts from 1–10 kb for bulk RNA samples



Full-length 16S rRNA for species identification

Up to 1,536-plex Kinnex library [12-fold concatenation]

60M (Revio SMRT Cell) [25M reads (Sequel II/IIe SMRT Cell)]

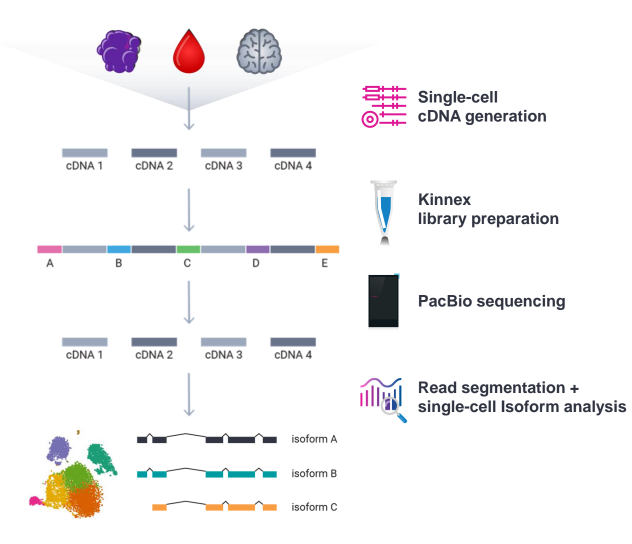


Taxonomic profiling for human, animal, and environmental samples

¹ Note: Kinnex single-cell RNA kit requires SMRT Link v13.1 or higher.

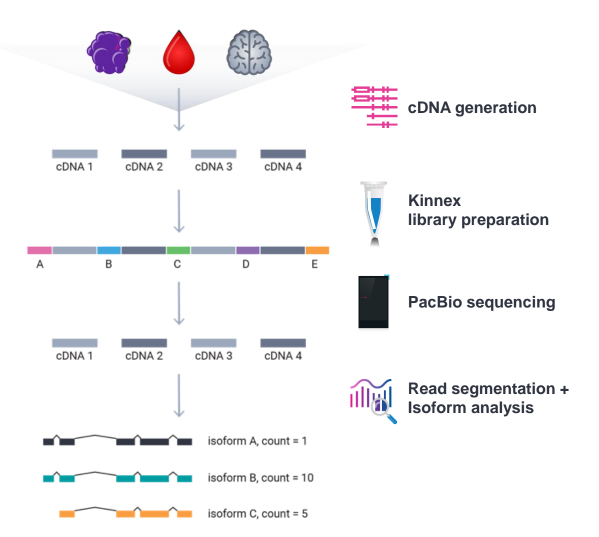
² Kinnex full-length RNA kit and Kinnex 16S rRNA kit requires SMRT Link v13.0 or higher.

Kinnex single-cell RNA kit for single-cell isoform sequencing



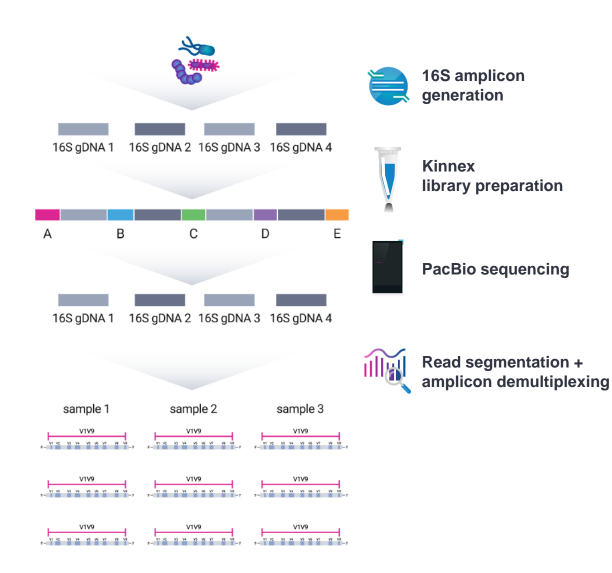
- 10x Chromium Single Cell 3' kit (v3.1) and 5' kit (v2)
- 15–75 ng cDNA input
- 3,000 to 10,000 target cell recovery
- 2-day Kinnex library preparation using Kinnex single-cell RNA kit
- Barcoded Kinnex adapters support up to 4-plex multiplexing
- SMRT Link Run Design support for 'Kinnex single-cell RNA' application type option with auto-analysis (read segmentation + single-cell isoform analysis)¹
- SMRT Link single-cell Iso-Seq isoform-classification software to identify novel genes and isoforms
- Output compatible with tertiary single-cell analysis tools (e.g., *Seurat, Scanpy, Kana*)

Kinnex full-length RNA kit for high-accuracy, full-length isoform sequencing



- Input 300 ng total RNA, RIN ≥7
- Generate up to 12-plex barcoded cDNA using Iso-Seq express 2.0 kit (103-071-500)
- 2-day Kinnex library preparation using Kinnex full-length RNA kit (103-072-000)
- SMRT Link Run Design support for 'Kinnex full-length RNA' application type with auto-analysis (read segmentation + isoform analysis)¹
- SMRT Link Iso-Seq isoform-classification software to identify novel genes and isoforms with abundance information

Kinnex 16S rRNA kit for full-length 16S sequencing



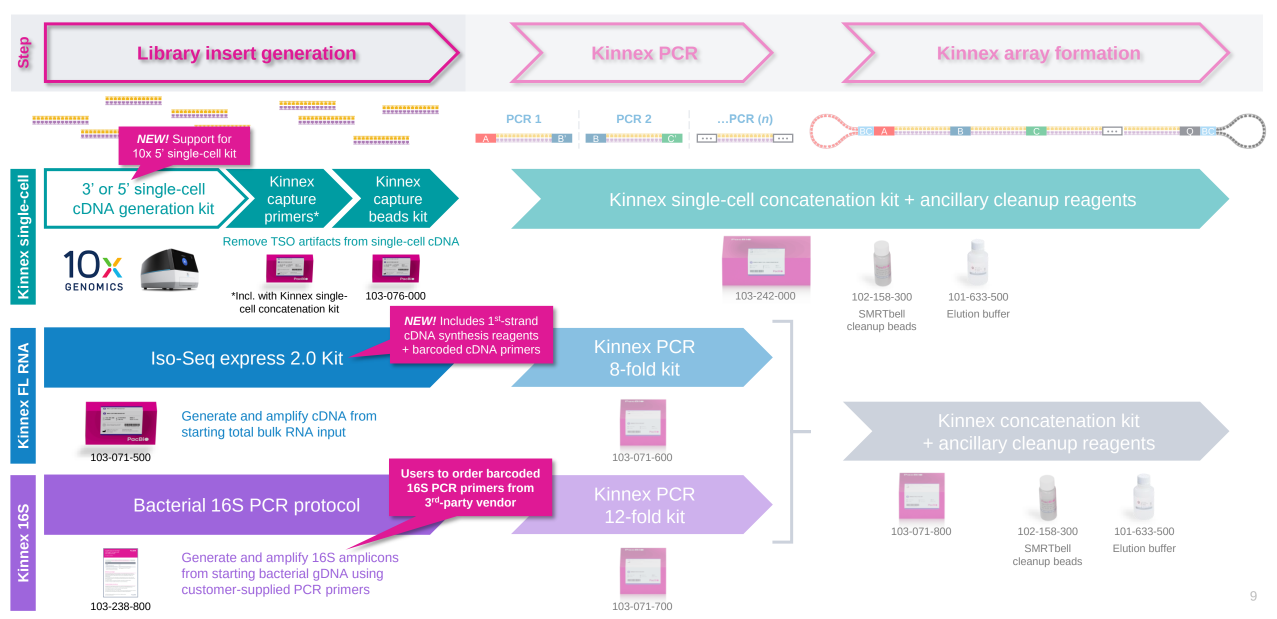
- Official protocol to generate barcoded 16S amplicons compatible with Kinnex 16S rRNA kit
- Protocol supports up to 384-plex multiplexing
- 2-day Kinnex library preparation using Kinnex 16S rRNA kit (103-072-100)
- SMRT Link Run Design support for 'Kinnex 16S rRNA' application type option with auto-analysis (read segmentation only)¹
- Demultiplex 16S amplicon barcodes in SMRT Link to generate per-sample read BAM files
- Analyze per-sample BAM files using GitHub tools or other custom 16S analysis pipeline

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Kinnex product configuration overview

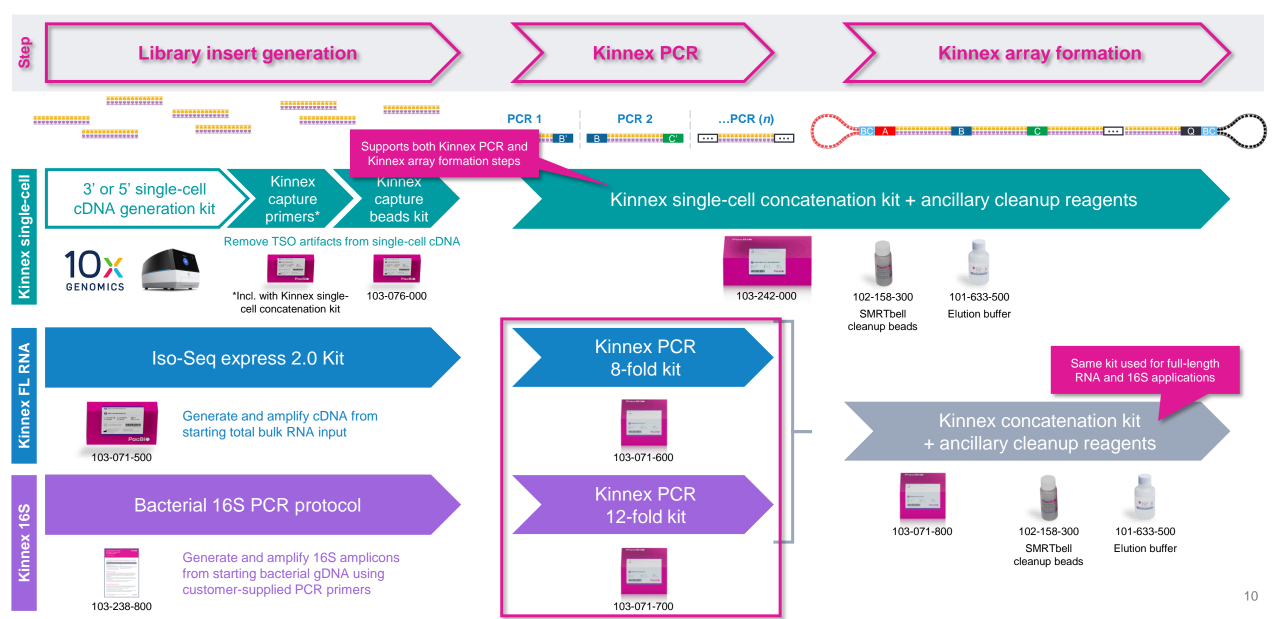
Kinnex product design overview and supported applications

Kinnex kits utilize the MAS-Seq concatenation method to increase throughput on Sequel II/IIe & Revio systems



Kinnex product design overview and supported applications

Kinnex kits utilize the MAS-Seq concatenation method to increase throughput on Sequel II/IIe & Revio systems



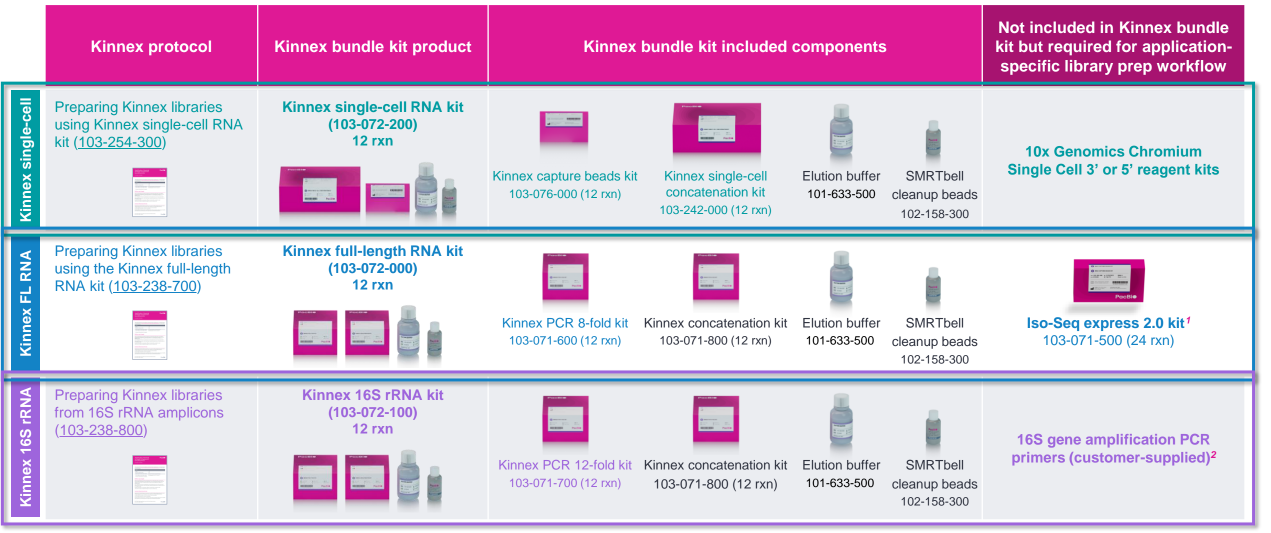
Kinnex bundle kit product features

All three Kinnex bundle kit products are compatible with Sequel II/IIe and Revio systems

			Concatenation .	Samp	le multiplexing	capacity	Throughput	SMRT Link
	Kinnex protocol	Kinnex bundle kit product	factor	Amplicon multiplexing	Library multiplexing	Total multiplexing capacity	per SMRT Cell	support
Kinnex single-cell	Preparing Kinnex libraries using Kinnex single-cell RNA kit (<u>103-254-300</u>)	Kinnex single-cell RNA kit (103-072-200) 12 rxn	16-fold	None	4-plex	4-plex	~30-40 M (Sequel II/IIe) ~80-100 M (Revio)	Full Demux BC Read seg. Sc-Iso-Seq
Kinnex FL RNA	Preparing Kinnex libraries using the Kinnex full-length RNA kit (<u>103-238-700</u>)	Kinnex full-length RNA kit (103-072-000) 12 rxn	8-fold	12-plex (Using Iso-Seq express 2.0 kit ¹) S1 S2 S3	4-plex	48-plex	~15 M (Sequel Ⅱ/Ⅱe) ~40 M (Revio)	Full Demux BC Read seg. Iso-Seq
Kinnex 16S rRNA	Preparing Kinnex libraries from 16S rRNA amplicons (103-238-800)	Kinnex 16S rRNA kit (103-072-100) 12 rxn	12-fold	384-plex (Using customer- supplied 16S PCR primers) S1F ### S1R S2F ### S2R	4-plex	1,536-plex	~25 M (Sequel Ⅱ/Ⅱe) ~60 M (Revio)	Partial Demux BC Read seg.

Kinnex bundle kit product components

Kinnex full-length RNA and 16S rRNA kits share a common workflow; Kinnex single-cell RNA kit uses existing MAS-Seq single-cell workflow with additional 10x 5' single-cell cDNA and sample multiplexing support



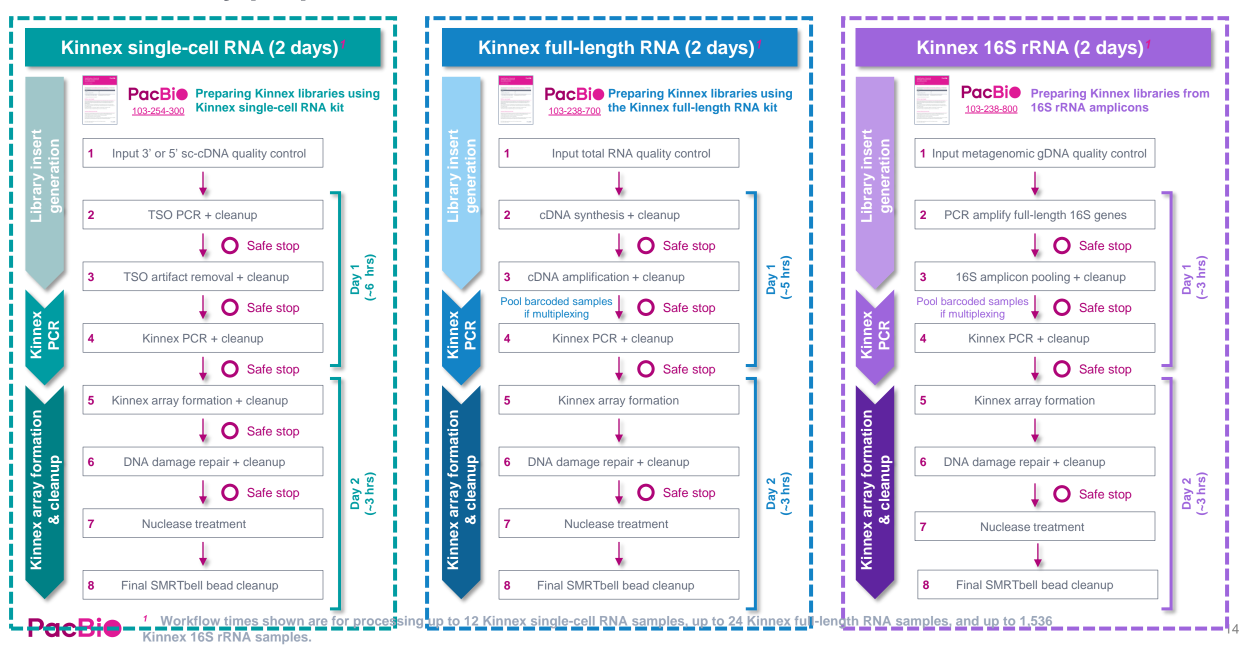
¹ Note: Iso-Seq express 2.0 kit (103-071-500) is not included in Kinnex full-length RNA bundle kit and must be purchased separately from PacBio.

PacBio ² Refer to *Procedure & checklist – Preparing Kinnex libraries from 16S rRNA amplicons (103-238-800)* for recommended 16S gene-specific forward and reverse PCR primer sequences to order.

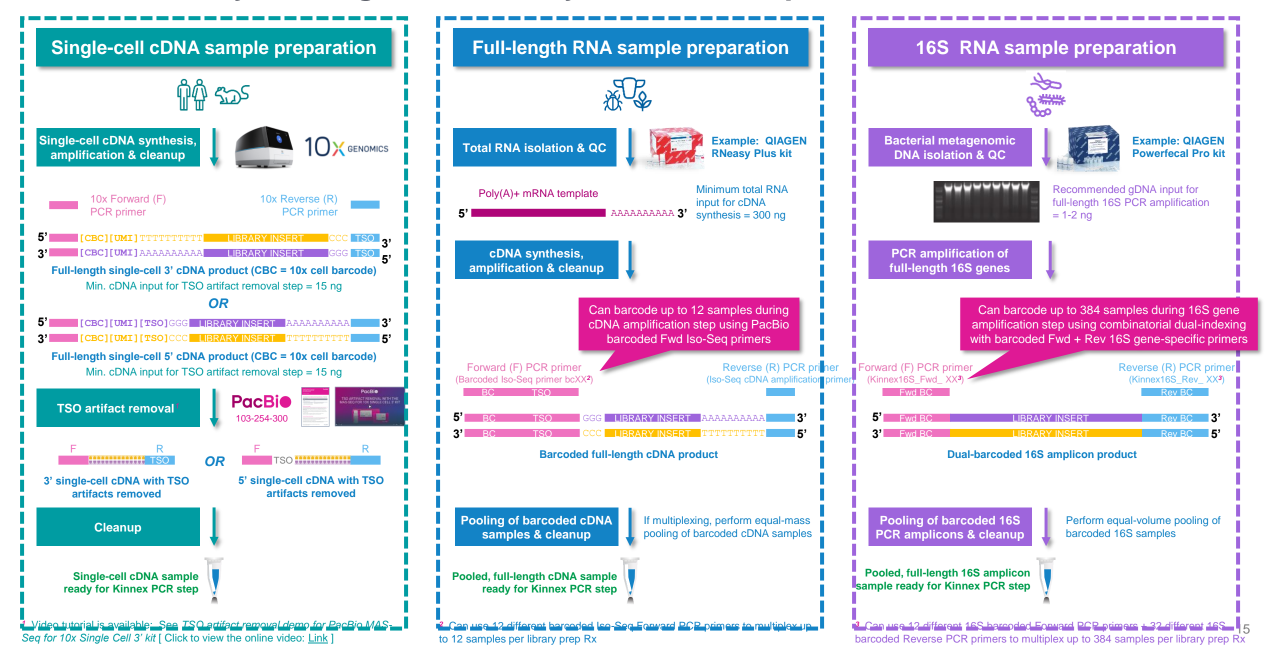
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Kinnex library preparation & sequencing workflow overview

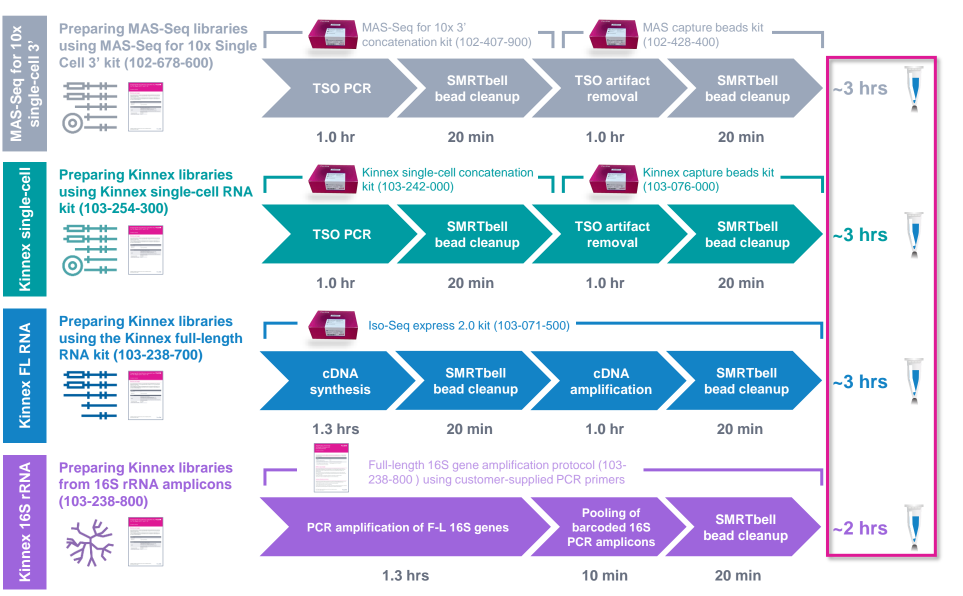
Kinnex library preparation workflow overview



Kinnex library insert generation key workflow steps

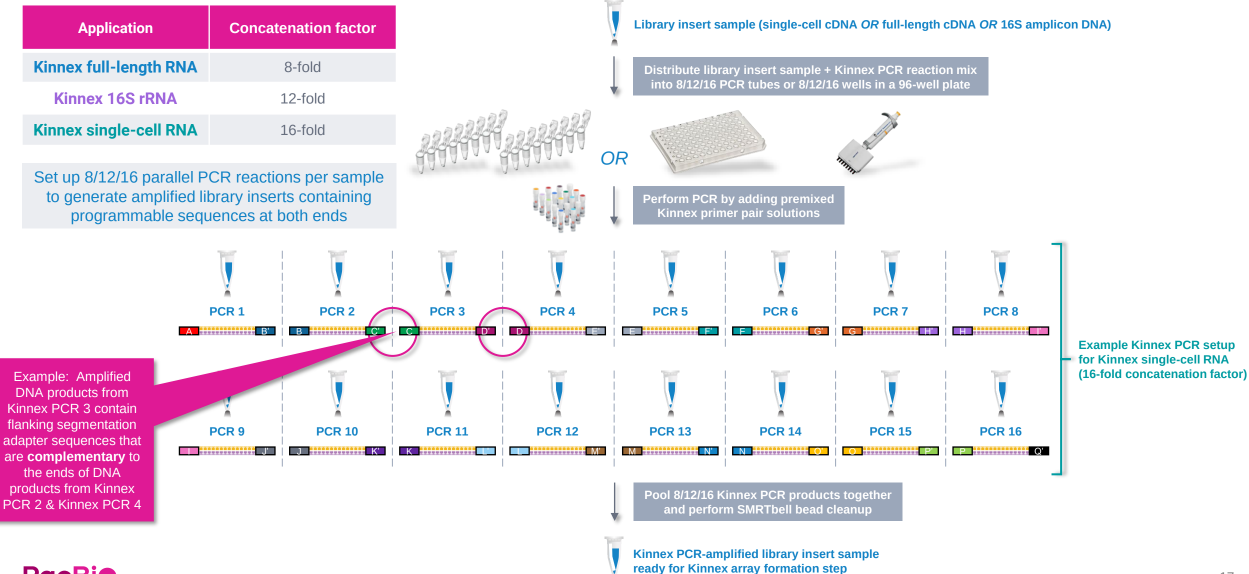


Application-specific Kinnex library insert generation workflow comparison



Kinnex PCR key workflow steps

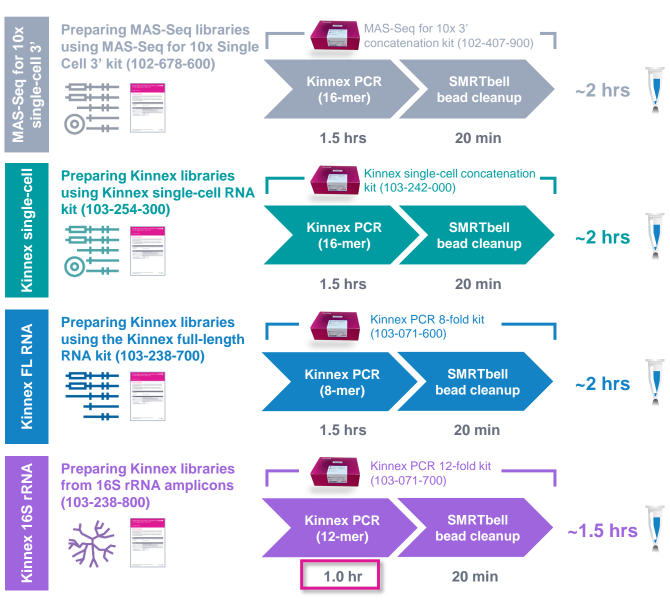
Kinnex PCR step incorporates programmable segmentation adapter sequences into library insert samples



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Application-specific Kinnex PCR workflow comparison



Step	Temperature	Duration	Cycles
Initial Denaturation	98°C	3 min	1
Denaturation	98°C	20 s	
Annealing	68°C	30 s	9
Extension	72°C	4 min	
Final Extension	72°C	5 min	1
Hold	4°C	Hold	

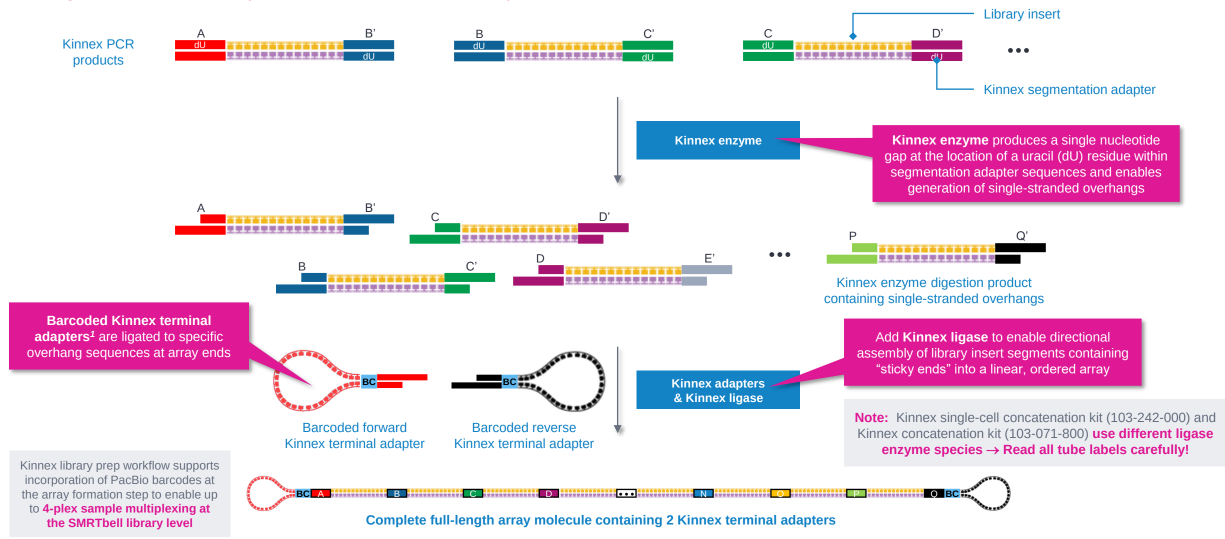
Step	Temperature	Duration	Cycles
Initial Denaturation	98°C	3 min	1
Denaturation	98°C	20 s	
Annealing	68°C	30 s	9
Extension	72°C	4 min	
Final Extension	72°C	5 min	1
Hold	4°C	Hold	

Step	Temperature	Duration	Cycles
Initial Denaturation	98°C	3 min	1
Denaturation	98°C	20 s	
Annealing	68°C	30 s	9
Extension	72°C	4 min	
Final Extension	72°C	5 min	1
Hold	4°C	Hold	

Step	Temperature	Duration	Cycles
Initial Denaturation	98°C	3 min	1
Denaturation	98°C	20 s	
Annealing	68°C	30 s	9
Extension	72°C	90 s	
Final Extension	72°C	5 min	1
Hold	4°C	Hold	

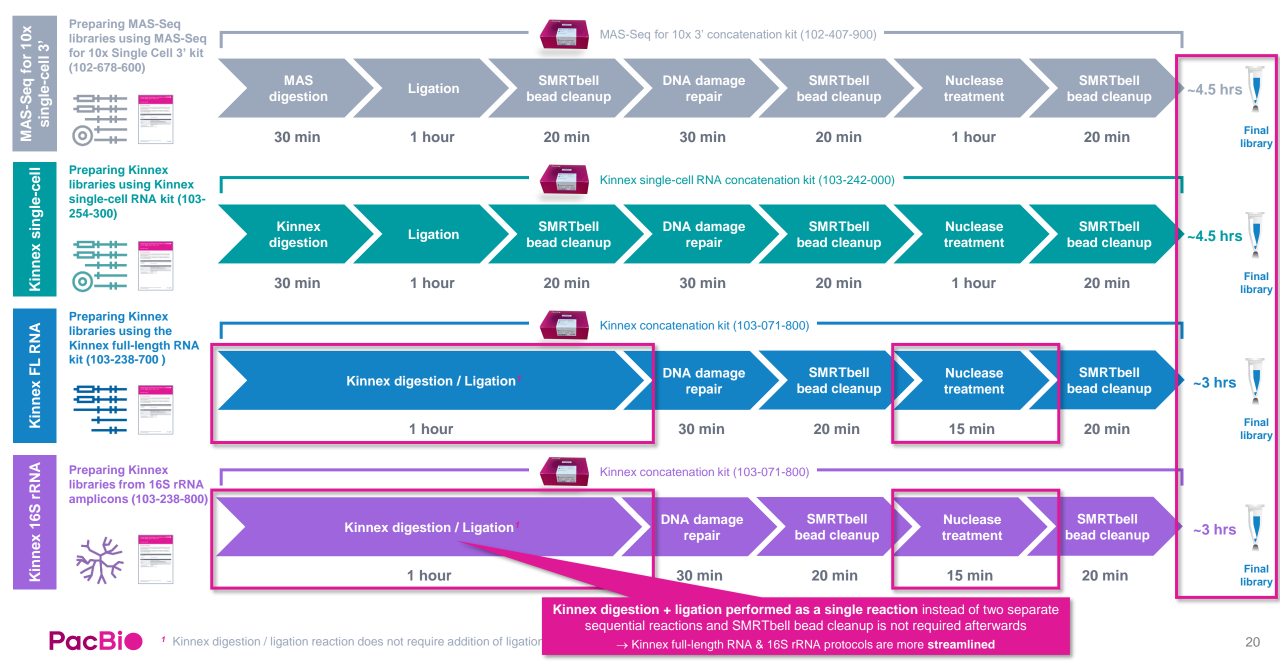
Kinnex array formation key workflow steps

Kinnex array formation step involves assembly of Kinnex PCR products ("segments") containing a library insert + programmable "sticky ends" into a linear array



Pace ¹ Four barcoded terminal Kinnex adapters (Kinnex adapter bcM0001-bcM0004) are available for Kinnex array formation step. Note: Kinnex concatenation workflow is not compatible with standard SMRTbell adapters from SMRTbell prep kit 3.0 and is also not compatible with SMRTbell barcoded adapter plate 3.0.

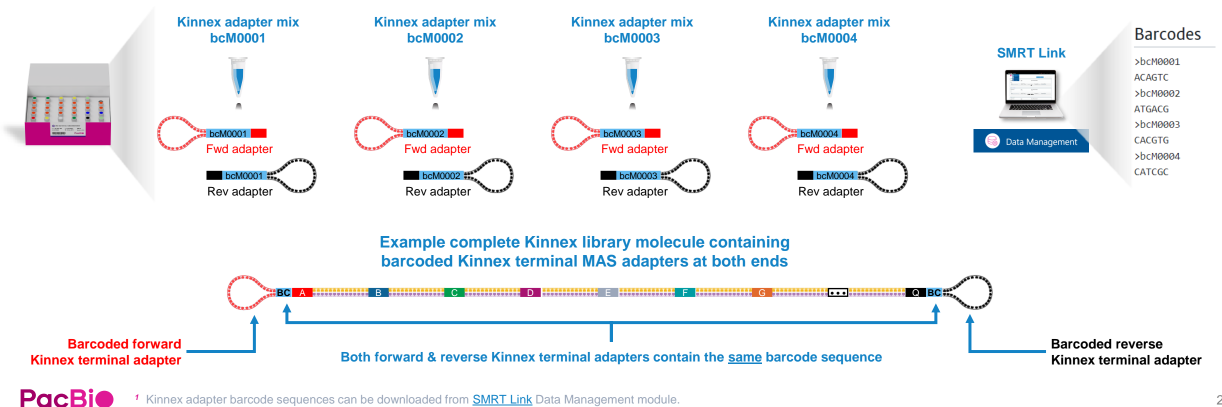
Application-specific Kinnex array formation workflow comparison



New Kinnex SMRTbell adapter design

Kinnex terminal adapters incorporate barcode sequences to enable up to 4-plex sample multiplexing at the library level

- New Kinnex adapters contain barcode sequences¹ to enable (optional) sample multiplexing at the SMRTbell library level (up to 4-plex)
 - Forward and reverse Kinnex adapter pairs are pre-mixed in Kinnex concatenation kits
 - Kinnex concatenation kits contain a total of 4 barcoded Kinnex adapter mixes (bcM0001-bcM0004) to enable multiplexing of up to 4 samples per SMRT Cell

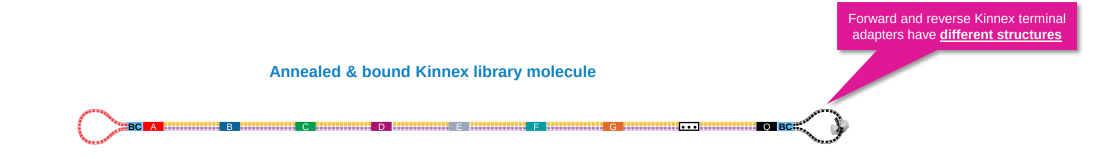


¹ Kinnex adapter barcode sequences can be downloaded from SMRT Link Data Management module.

New Kinnex SMRTbell adapter design (cont.)

Kinnex terminal adapters use a new design that enables improved SMRT sequencing performance

- Kinnex adapters enable:
 - Longer polymerase read length → Improved HiFi conversion rate (HiFi reads/Total P1 reads)
 - Improved P1 loading efficiency

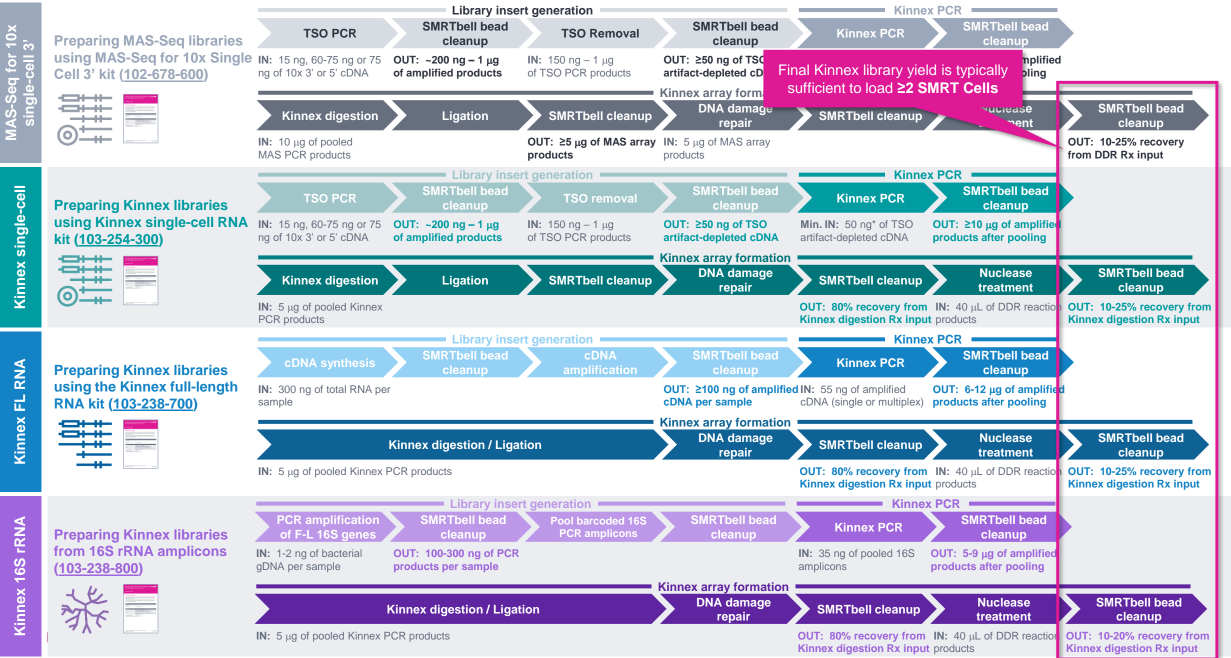


• New Kinnex adapter design requires a different sequencing primer (Kinnex sequencing primer 103-179-000)

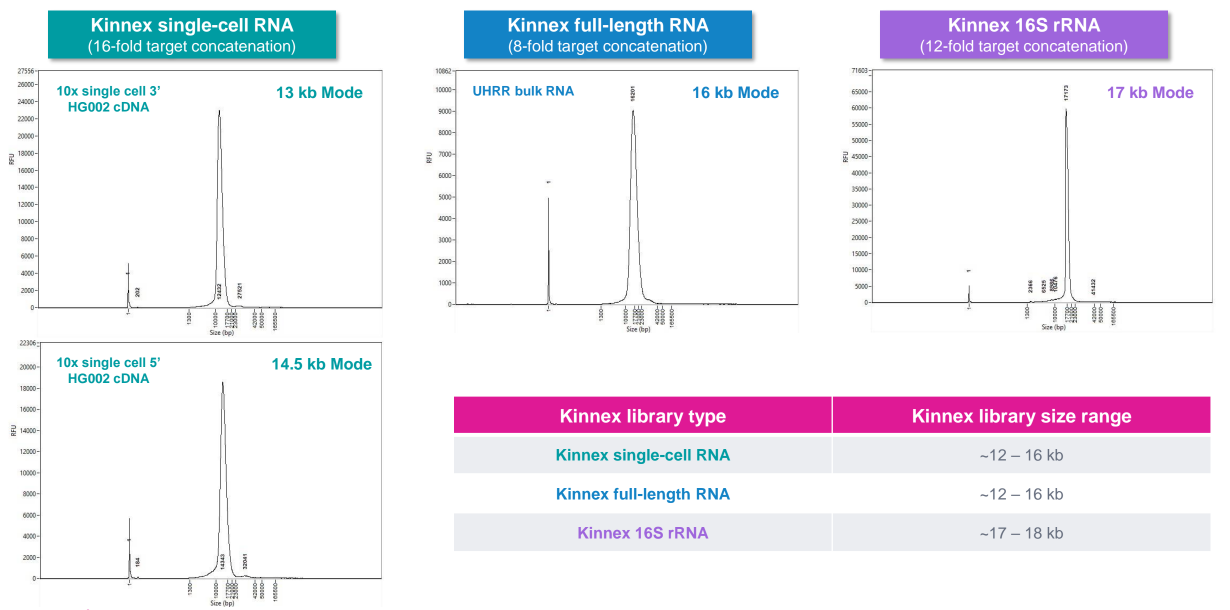




Application-specific Kinnex library prep input & expected step yield comparison



Example Femto Pulse DNA sizing QC results for final Kinnex SMRTbell libraries



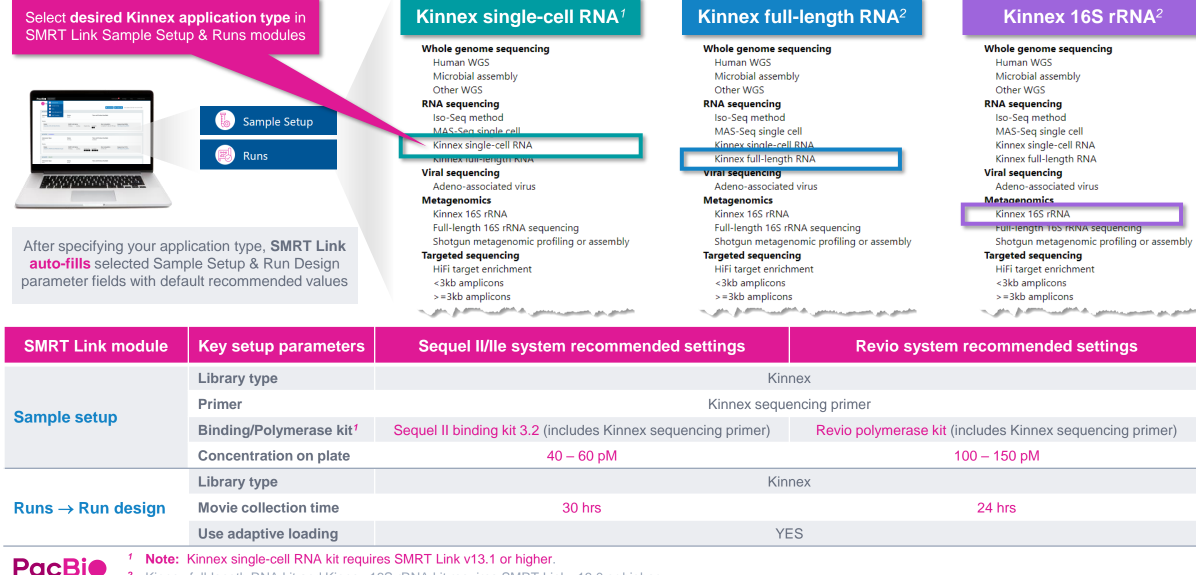
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Kinnex sequencing preparation workflow overview

Sample Setup & Run Design recommendations for Kinnex libraries

Follow SMRT Link instructions to prepare Kinnex libraries for sequencing^{1,2}



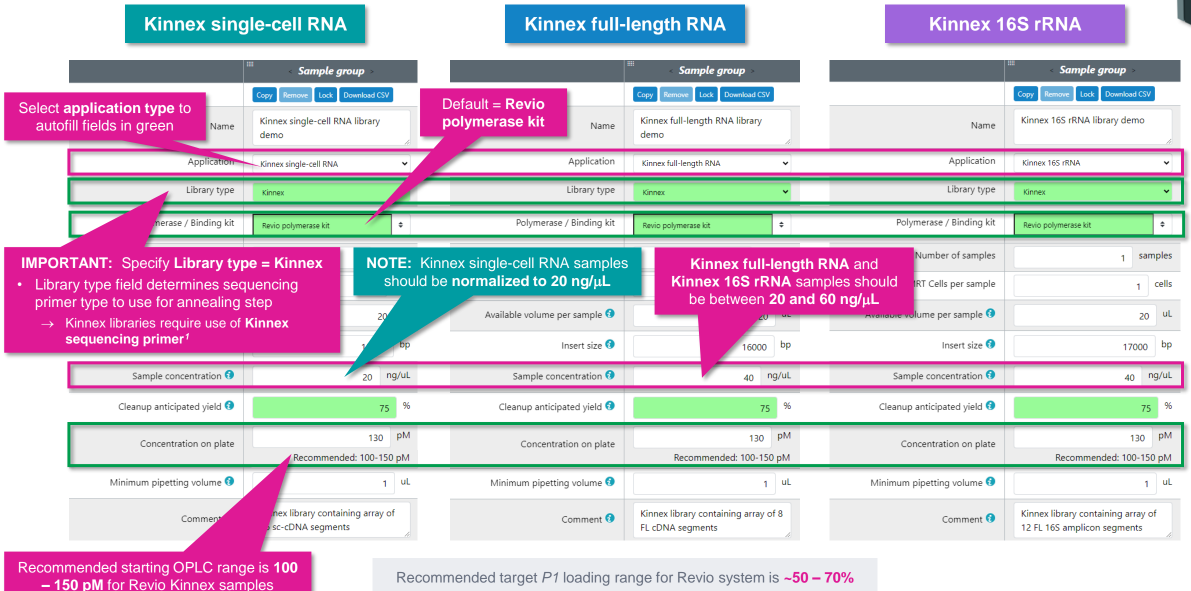
² Kinnex full-length RNA kit and Kinnex 16S rRNA kit requires SMRT Link v13.0 or higher.

SMRT Link Sample Setup and Run Design for Kinnex kits video demonstration

Video demonstration of SMRT Link Sample Setup and Run Design setup procedure for Kinnex kits supporting full-length RNA sequencing, single-cell RNA sequencing and full-length 16S rRNA sequencing

Sample Setup / Sample Calculation Sequel II binding kit 3.1/3.2, Revio polymerase kit	t week		Conversion Calculator
	+ Add Sam	npie Group	
		 Sample group 	
		Copy Remove Lock Download CSV	
	Name	My Batch of Samples	
	Application	Kinnex full-length RNA V	
	Library type	Kinnex	<u>Demo video</u> for Sample Setup and Run Design for Kinnex kits
	Polymerase / Binding kit	Revio polymerase kit \$	(SMRT Link v13.0+)
	Number of samples	1 samples	 Demo video for Sample Setup and Run Design for Kinnex kits in SMRT Link v13.0+
	SMRT Cells per sample	1 cells	-
	Available volume per sample 🜖	20 UL	 Kinnex kits support full-length RNA sequencing (Kinnex full-length RNA kit), full-
	Insert size 🔞	16000 bp	length 16S rRNA sequencing (Kinnex 16S
	Sample concentration 📀	40 ng/uL	rRNA kit) and full-length single-cell RNA sequencing (Kinnex single-cell RNA kit)
	Cleanup anticipated yield 🔞	75 %	
	Concentration on plate	130 pM	
		Recommended: 100-150 pM	
YouTube	Minimum pipetting volume 🜖	1 4	
	Comment 🚯		

SMRT Link Sample Setup procedure for Revio system

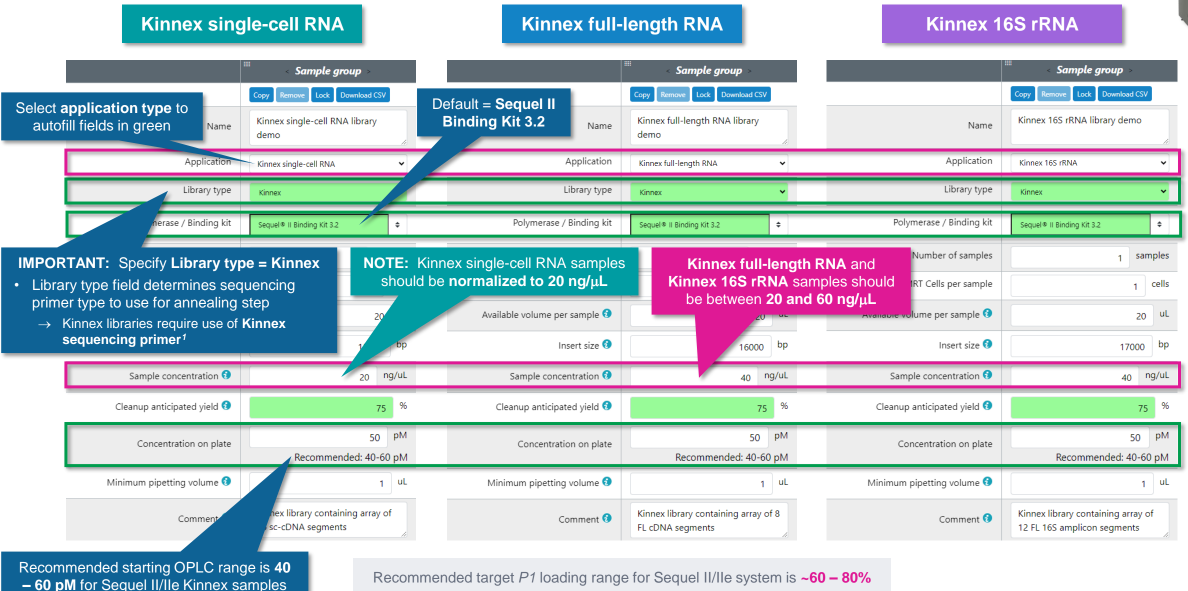


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¹ Sample Setup Library type field specifies structure of SMRTbell library and determines sequencing primer type to use for annealing step. For Kinnex libraries, the forward and reverse Kinnex terminal SMRTbell adapters have different structures and require use of Kinnex sequencing primer for primer annealing step.

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SMRT Link Sample Setup procedure for Sequel II/IIe systems



Recommended target P1 loading range for Sequel II/IIe system is ~60 – 80%

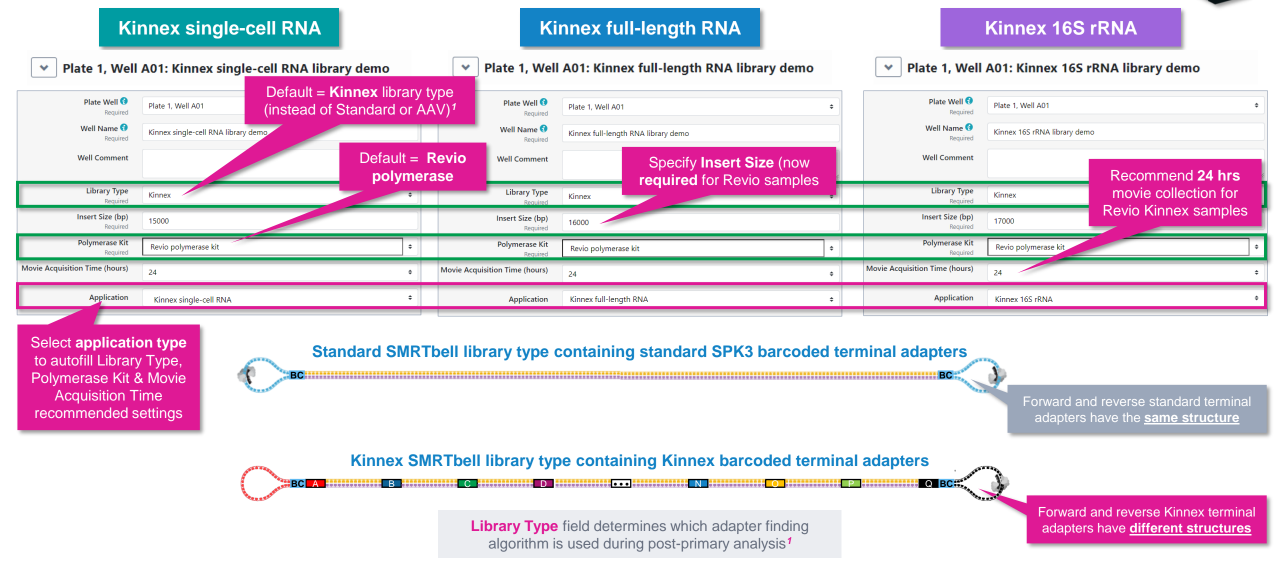
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Sample Setup Library type field specifies structure of SMRTbell library and determines sequencing primer type to use for annealing step. For Kinnex libraries, the forward and reverse Kinnex terminal SMRTbell adapters have different structures and require use of Kinnex sequencing primer for primer annealing step.

SMRT Link Run Design procedure for Revio system

Sample and run information

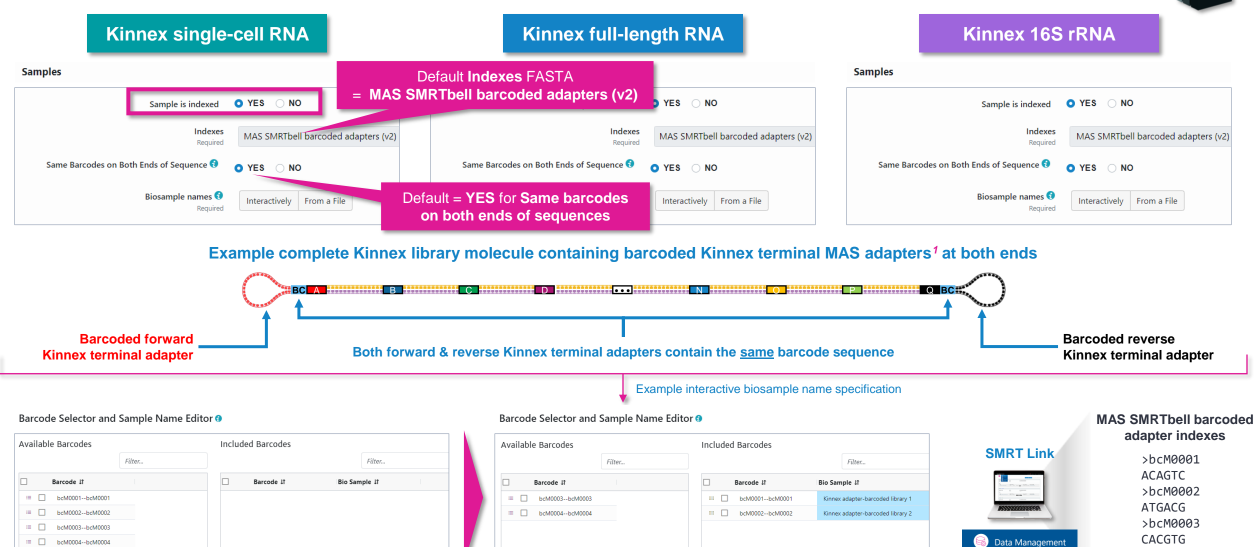
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1 Note: When sequencing a Kinnex library sample, if 'Standard' library type is mistakenly selected instead of 'Kinnex' then a higher missing adapter rate (> 95%) and a slight degradation in barcode demultiplexing performance (~93-96% barcoded HiFi read yield) will be observed.

SMRT Link Run Design procedure for Revio system (cont.)

Sample indexing (barcoding) information



¹ Four barcoded terminal Kinnex adapters (Kinnex adapter bcM0001-bcM0004) are available for Kinnex array formation step. Kinnex adapter barcode sequences can be downloaded from <u>SMRT Link</u> Data Management module.

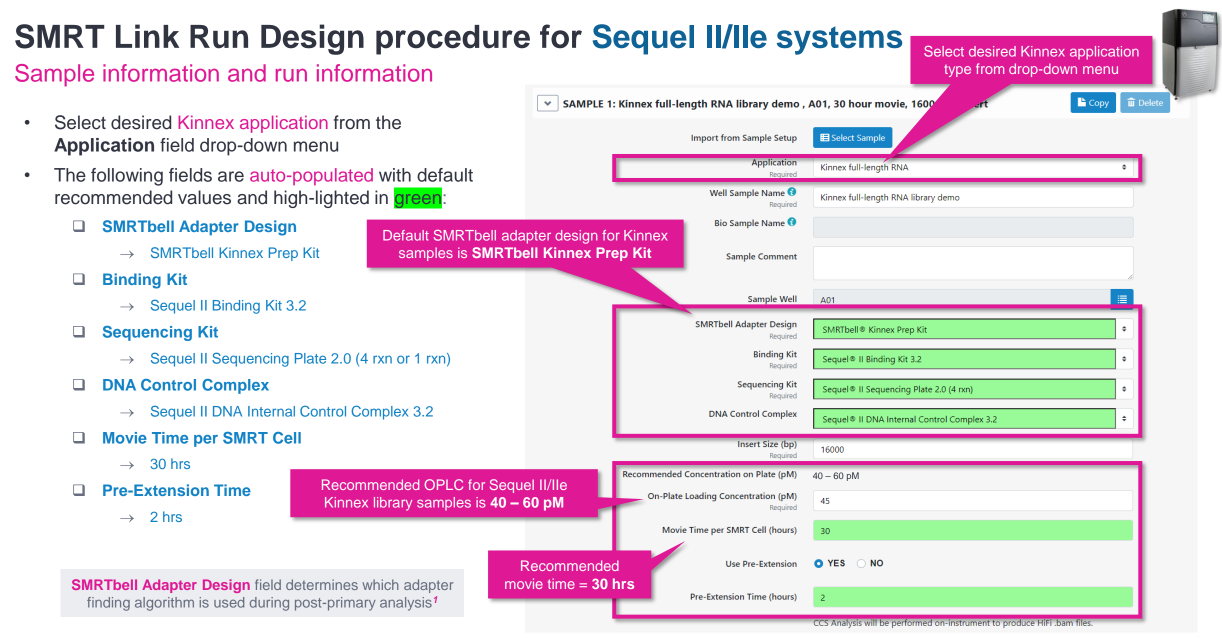
>bcM0004 CATCGC

SMRT Link Run Design procedure for Revio system (cont.)

Run options and data options

Kinnex single-cell RNA	Kinnex full-length RNA	Kinnex 16S rRNA
Run Options OPLC now required for Revio samples	v Run Options	✓ Run Options
Library Concentration (pM) Required	Library Concentration (pM) Required	Library Concentration (pM) 130 Required
Use Adaptive Loading O YES O NO	Use Adaptive Loading 🔹 YES 🔷 NO	Use Adaptive Loading O YES O NO
Default = YES for Use Adaptive Loading Default = NO Include Base King		V Data Options
Include Base Kinetics YES ONO	Include Base Kinetics VES ONO	Include Base Kinetics VES • NO
Consensus Mode O MOLECULE O STRAND	Consensus Mode O MOLECULE O STRAND	Consensus Mode MOLECULE STRAND
Assign Data Torget 3 General Project	Assign Data To Project 📀 General Project	Assign Data To Project 📀 General Project
Default Consensus Mode = MOLECULE ¹	Can leave Include Base Kinetics and Consensus Mode fields at their default settings for Kinnex library samples	





Example sample information entered into a Sequel IIe system run design worksheet for a Kinnex fulllength RNA library sample.

Pacbio ¹ Note: When sequencing a Kinnex library sample, if 'Overhand-SMRTbell Prep Kit 3.0' is mistakenly selected instead of 'SMRTbell Kinnex Prep kit' in the SMRTbell Adapter Design field, then a higher missing adapter rate (> 95%) and a slight degradation in barcode demultiplexing performance (~93-96% barcoded HiFi read yield) will be observed.

SMRT Link Run Design procedure for Sequel II/IIe systems (cont.)

Advanced options

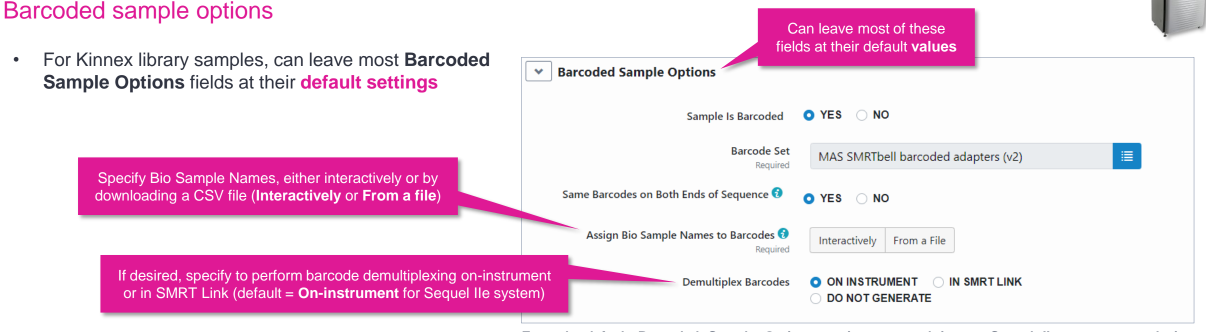
- For all Kinnex library samples, leave the following Advanced Options fields at their default settings
 - Use Adaptive Loading
 - \rightarrow YES
 - □ Loading Target (P1 + P2)
 - \rightarrow 0.85
 - Maximum Loading Time
 - \rightarrow 2 hours
 - **CCS Analysis Output Include Low Quality Reads**
 - \rightarrow NO
 - **CCS Analysis Output Include Kinetics Information**
 - \rightarrow NO
 - Pre-Extension Time
 - \rightarrow 2 hrs
- If desired, specify to use an alternative project folder for the Add Data to Project field

Advanced Options		Leave these Advanced Options fields at their default values
Use Adaptive Loading	• YES ONO	
Loading Target (P1 + P2)	0.85	
Maximum Loading Time (hours)	2	
CS Analysis Output - Include Low Quality Reads 🕄	OYES ONO	
CCS Analysis Output - Include Kinetics Information	🔵 YES 💿 NO	
Add Data to Project 🕄	General Project	÷

Can specify to use a different Project folder



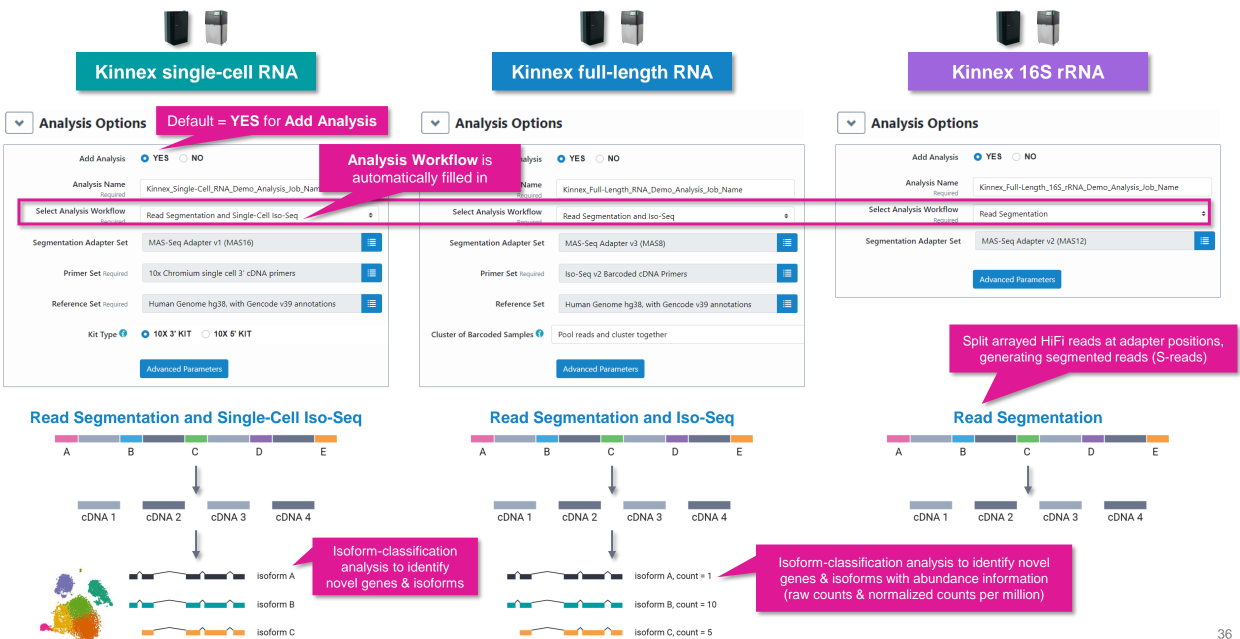
SMRT Link Run Design procedure for Sequel II/IIe systems (cont.)



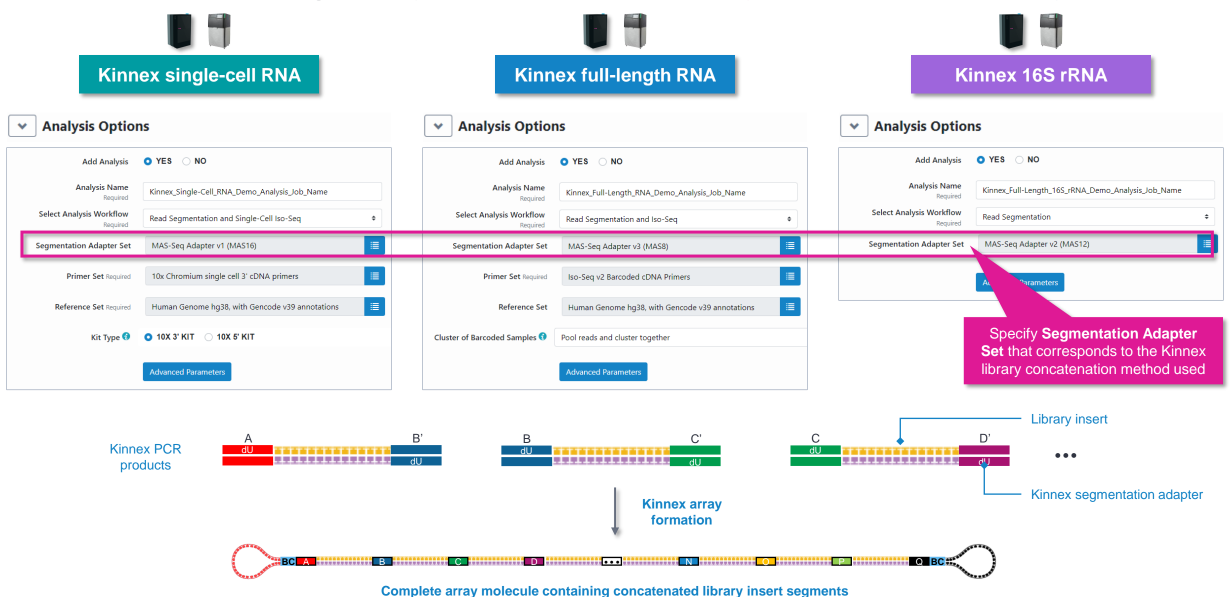
Example default Barcoded Sample Options settings entered into a Sequel lle system run design worksheet for a Kinnex full-length RNA library sample.

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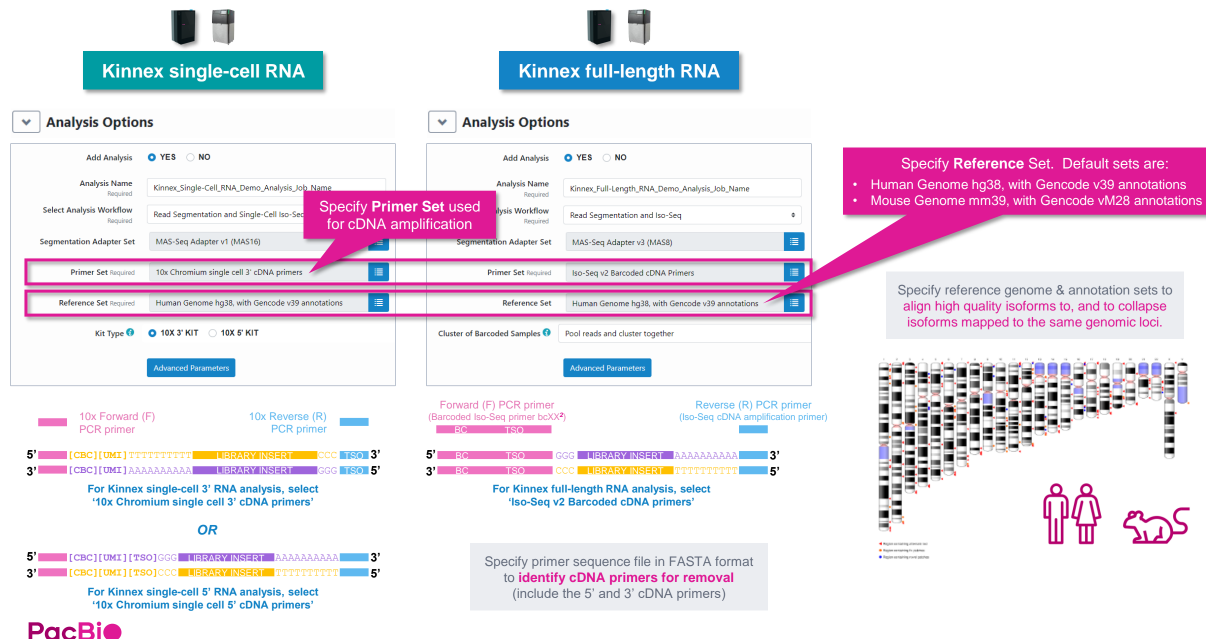
SMRT Link Run Design analysis options for Revio system & Sequel II/IIe systems



SMRT Link Run Design analysis options for Revio system & Sequel II/IIe systems (cont.)

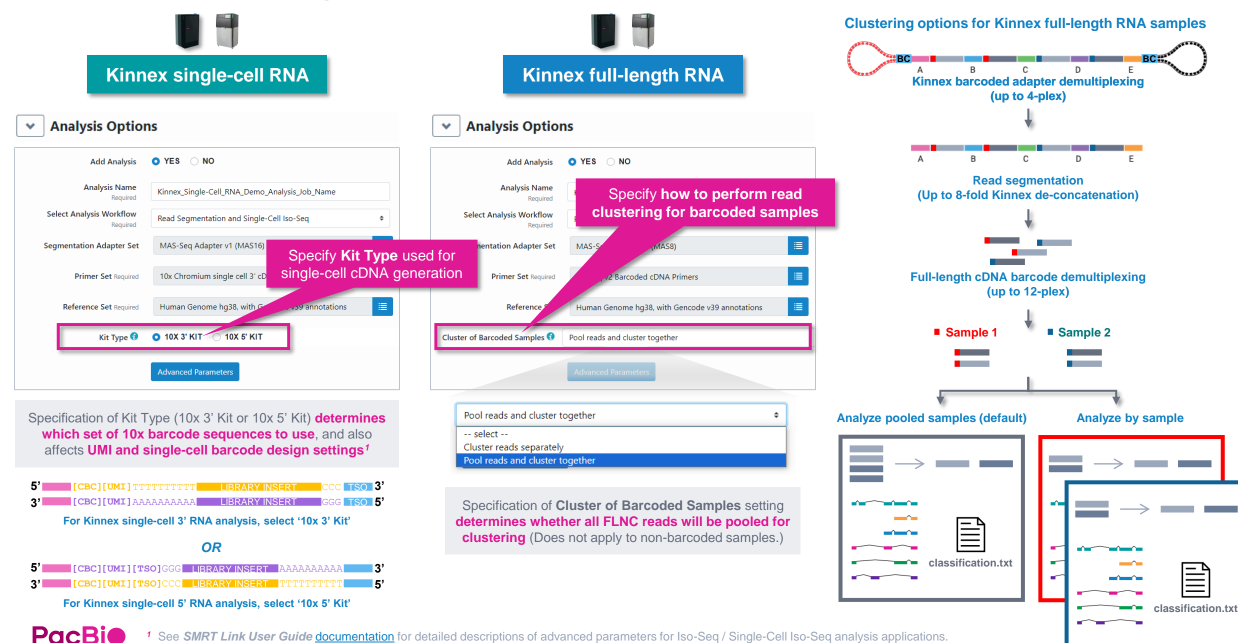


SMRT Link Run Design analysis options for Revio system & Sequel II/IIe systems (cont.)



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SMRT Link Run Design analysis options for Revio system & Sequel II/IIe systems (cont.)



¹ See SMRT Link User Guide documentation for detailed descriptions of advanced parameters for Iso-Seq / Single-Cell Iso-Seq analysis applications.

SMRT Link Read Segmentation and Iso-Seq analysis video demonstration

Video demonstration of SMRT Link Read Segmentation and Iso-Seq application workflow for analysis of Kinnex full-length RNA samples

Analysis Application Required	Analysis Name	
Read Segmentation and Iso-Seq \$	test	
Import Analysis Settings	Analysis Datasets	
Associated Inputs	Displaying rows 1 to 1 out of 1	
Segmentation Adapter Set	ID IT Name IT	
MAS-Seq Adapter v3 (MAS8)	21 3230211_KPoS_64007	
Primer Set Required		
Iso-Seq v2 Barcoded cDNA Primers		
Reference Set		
Human Genome hg38, with Gencode v39 annotations	Domo video for Bood Segmentation and Ico Seg workflow	
Cluster of Barcoded Samples 🜖	Demo video for Read Segmentation and Iso-Seq workflow	
Pool reads and cluster together ÷	(SMRT Link v13.0+)	
Advanced Parameters	 Workflow supports full-length isoform analysis for data generated on PacBio Sequel II/IIe and Revio systems using Kinnex full-length RNA kit 	
Auvanceu Parameters	End-to-end workflow begins with HiFi reads and outputs full-length isoform classifications with supporting read count information	
YouTube		

SMRT Link Read Segmentation and Demultiplex Barcodes video demonstration

Video demonstration of SMRT Link Read Segmentation and Demultiplex Barcodes workflow for analysis of Kinnex 16S rRNA samples

Data Utility Required	Analysis Name
Read Segmentation \$	test-ReadSeg
The Import Analysis Settings	Analysis Datasets
Associated Inputs	Displaying rows 1 to 1 out of 1
Segmentation Adapter Set	ID If Name If
MAS-Seq Adapter v2 (MAS12)	21 20231020-4_84028_13
Advanced Parameters	
	Demo video for Read Segmentation and Demultiplex Barcodes workflow (SMRT Link v13.0+)
	 Demo video for analyzing Kinnex 16S rRNA data generated using Kinnex 16S rRNA kit in SMRT Link v13.0 and up
YouTube	

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Kinnex single-cell RNA library preparation & sequencing workflow key highlights

Kinnex single-cell RNA library preparation procedure description

Procedure & checklist – Preparing Kinnex libraries using Kinnex single-cell RNA kit (<u>103-254-300</u>) describes the workflow for constructing single-cell Kinnex libraries from 10x Chromium 3' or 5' single cell cDNA using the **Kinnex single-cell RNA kit**¹ for library preparation and sequencing on PacBio Sequel II, Sequel IIe & Revio systems.

OverviewSamples per kit12Workflow time3	days for up to 12 samples	Preparing Kinnex [™] libraries using Kinnex single-cell RNA kit Procedure & checklist
cDNA input	>15 ng per library	Before you begin This procedure describes the workflow for constructing single-cell Kinnex libraries from 10x Chromium 3 or 5 cDNA using the Kinnex single-cell RNA kt (103-072-200) for library prep and sequencing on PacBio* Sequel® IL Sequel ILe, and Revio* systems. This kit is intended for use with single-cell CDNA generated using the 10x Chromium Next GEM Single Cell 3' kit v3.1 or 10x Chromium Next GEM Single Cell 5' kit v2, standard throughput. It has not been tested for use on low throughput (LT) which are currently unsupported.
Quantity	cDNA concentration should be >1 ng/ μ L with up to 15 μ L in volume. See step 2.1 for 10x cDNA input requirement.	Overview Samples per kit 12 Workflow time 3 days for up to 12 samples
Average segment lengths	500-1,000 bp	eDNA input >15 ng per library
Average 16-segment array lengt	hs 10–15 kb Concatenation factor	Quantity cDNA concentration shiuld be >Ing/µL with up to 15 µL in volume. See <u>step 21</u> for 10x cDNA input require to 1 Average segment lengths 500-1,000 bp Average 10-segment array/lengths 10-15 kb
Kinnex single-cell RNA kit	16-fold A B C C C C C C C C C C C C C C C C C C	Kinnex single-cell RNA library prep protocol u Kinnex single-cell RNA kit <u>Do not use</u> SMRTbell prep kit 3.0 with this protocol
(103-072-200)		© 2024 PacBio All rights reserved. Research use only. Not for use in diagnostic procedures. PacBi● PacBio Documentation (103-254-300)

Comparison of Kinnex single-cell RNA kit vs. MAS-Seq for 10x Single Cell 3' kit

	Kinnex single-cell RNA kit (103-072-200)	MAS-Seq for 10x Single Cell 3' kit (102-659-600)
# reactions	12	8
Concatenation	16-fold	
Compatibility	10x Chromium Single Cell 3' kit (v3.1) 10x Chromium Single Cell 5' kit (v2)	10x Chromium Single Cell 3' kit (v3.1)
Sample multiplexing support	4-plex using Kinnex barcoded adapters	None
Workflow time	2 days	
SMRT Link support	Yes (since SMRT Link v13.1)	Yes (since SMRT Link v11.1)
Sequencing primer	Kinnex sequencing primer	Standard sequencing primer
Sequencing OPLC	50 pM (Sequel II/IIe) 130 pM (Revio)	85 pM (Sequel II/IIe) 225 pM (Revio)
Run time	Sequel II/IIe: 30hr movie with adaptive loading Revio: 24hr movie with adaptive loading	
S-read yield	~30-40 M reads (Sequel II/IIe) ~80-100 M reads (Revio)	

Kinnex single-cell RNA experimental design considerations

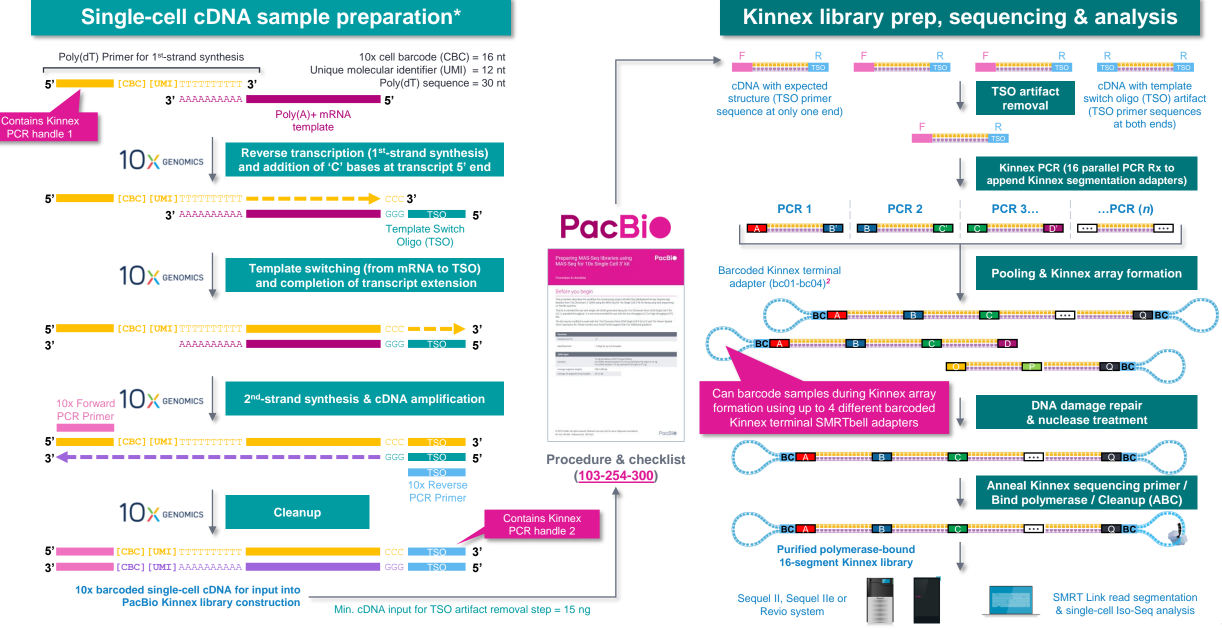
Kinnex single-cell RNA application use case recommendations for PacBio systems

	Sequel II and IIe systems	Revio system
Experimental goal	Characterize alternative splicing in single cells / cell types	
Sample multiplexing ¹	Not recommended	Up to 2 samples per Revio SMRT Cell (2-plex)
Cell input into 10x Chromium single cell 3' or 5'	3,000 – 10,000 cells for running a single (non-multiplexed) sample on one Sequel II SMRT Cell 8M)	3,000 – 6,000 cells per sample if multiplexing 2 samples per Revio SMRT Cell (2-plex)
		8,000 – 10,000 cells per sample if running a single (non- multiplexed) sample on one Revio SMRT Cell
Expected coverage	Obtain ≥3,000 – 10,000 unique reads/single cell	Obtain up to ~10,000 unique reads/single cell
Kinnex library prep protocol	Procedure & checklist – Preparing Kinnex libraries using Kinnex single-cell RNA kit (103-254-300)	
Single-cell cDNA input into Kinnex library prep workflow	15-75 ng of 10x Chromium 3' or 5' single cell cDNA	
SMRT Link data analysis workflows	Read Segmentation and Single-cell Iso-Seq Analysis	
	Annotation & quantification: SQANTI3	
Community data analysis tools	Differential analysis: TappAS	
	Fusion calling: pbfusion	
	Visualization: SWAN	



¹ Kinnex single-cell concatenation kit (103-242-000) can support up to 4-plex sample multiplexing per SMRT Cell through the use of four different barcoded Kinnex terminal SMRTbell adapters during Kinnex single-cell RNA library construction.

Kinnex single-cell RNA method overview



* Refer to 10x Genomics Support website to download 10x Chromium single cell 3'/5' reagent kit user guides.

² Kinnex adapter barcode sequences can be downloaded from <u>SMRT Link</u> Data Management module.

Kinnex single-cell RNA library preparation procedural notes



TSO PCR

- Set up TSO PCR reactions **ON ICE** using the **CORRECT** TSO PCR primer set (3' 10x cDNA \rightarrow 3' TSO PCR primer; 5' 10x cDNA \rightarrow 5' TSO PCR primer)
 - PCR polymerase 3'→5' exonuclease activity negatively impacts amplification yield if prepared at room temp.

TSO removal [Video demonstration]

- **IMPORTANT:** For bead capture steps, allow enough time for beads to magnetize as binding buffer is highly viscous; pipette mix with care and **avoid generating bubbles**
 - Wide-bore pipette tips are recommended to help minimize foaming (specifically when resuspending Kinnex capture beads)
- Fully resuspend beads during all wash steps to remove artifact cDNA effectively
- Fully resuspend beads before DNA quantification using Qubit dsDNA HS assay

SMRTbell bead cleanup

- Prior to Kinnex array formation, perform a **1.5X** SMRTbell bead cleanup
 - If the cDNA contains smaller fragments <200 bp, it is recommended to increase the SMRTbell cleanup bead ratio to 1.8 – 2.0X

Kinnex array formation



Kinnex digestion

- Recommended input amount to proceed with Kinnex array formation is $5\,\mu g$ of Kinnex PCR amplicons
 - Proceeding with <3 μg is not recommended since lower input amounts may lead

PacBito insufficient final library yields to enable optimal sequencing results

Kinnex PCR



Kinnex PCR

- Set up Kinnex PCR reactions ON ICE
 - PCR polymerase 3'→5' exonuclease activity negatively impacts amplification yield if prepared at room temp.
- Perform Kinnex PCR using 25 ng of cDNA after TSO removal step

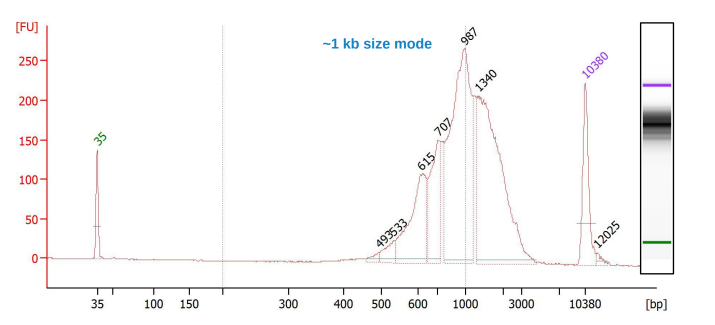
Final SMRTbell bead cleanup

Perform 1.0X SMRTbell bead cleanup on final library

Example Kinnex single-cell RNA library preparation QC results

Kinnex single-cell 3' RNA library prepared with human cDNA

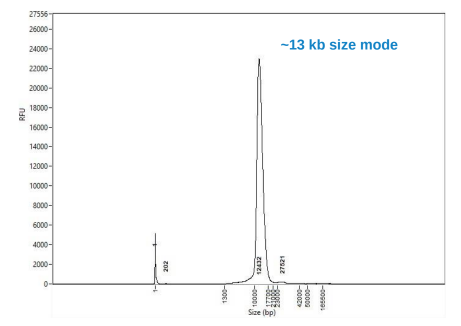
Kinnex PCR DNA sizing QC (Single-cell 3' cDNA)



Example Bioanalyzer DNA sizing QC analysis results for Kinnex PCR products generated for a 10x Chromium single-cell 3' cDNA samples prepared from a human cell line (HG002).

Final Kinnex library yield is typically sufficient to load ≥2 SMRT Cells

Final Kinnex single-cell RNA library QC



Example Femto Pulse DNA sizing QC analysis results for final Kinnex single-cell RNA library.

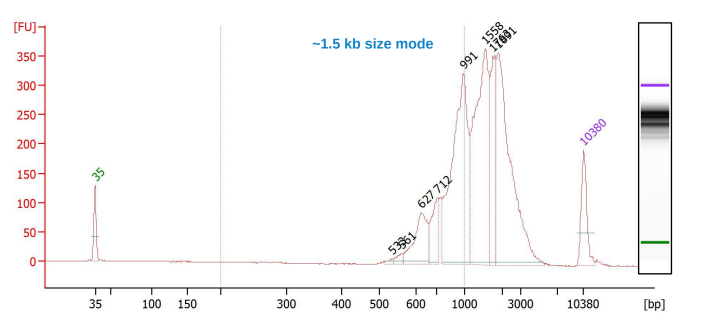
10x single cell 3' cDNA input	15 ng
cDNA input for Kinnex array formation	5000 ng
Post-nuclease treatment & final library cleanup yield (%) ¹	1100 ng (22.0%)

¹ Post-nuclease treatment & final cleanup yields typically ranged from ~10% to ~25% when using single-cell 3' cDNA samples for Kinnex single-cell RNA library construction.

Example Kinnex single-cell RNA library preparation QC results (cont.)

Kinnex single-cell 5' RNA library prepared with human cDNA

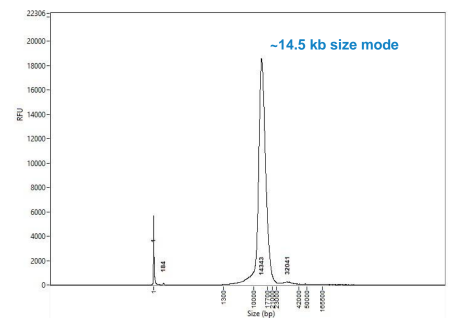
Kinnex PCR DNA sizing QC (Single-cell 5' cDNA)



Example Bioanalyzer DNA sizing QC analysis results for Kinnex PCR products generated for a 10x Chromium single-cell 5' cDNA samples prepared from a human cell line (HG002).

Final Kinnex library yield is typically sufficient to load ≥2 SMRT Cells

Final Kinnex single-cell RNA library QC



Example Femto Pulse DNA sizing QC analysis results for final Kinnex single-cell RNA library.

10x single cell 5' cDNA input	15 ng
cDNA input for Kinnex array formation	5000 ng
Post-nuclease treatment & final library cleanup yield (%) ¹	1008 ng (20.2%)

¹ Post-nuclease treatment & final cleanup yields typically ranged from ~10% to ~25% when using single-cell 5' cDNA samples for Kinnex single-cell RNA library construction.

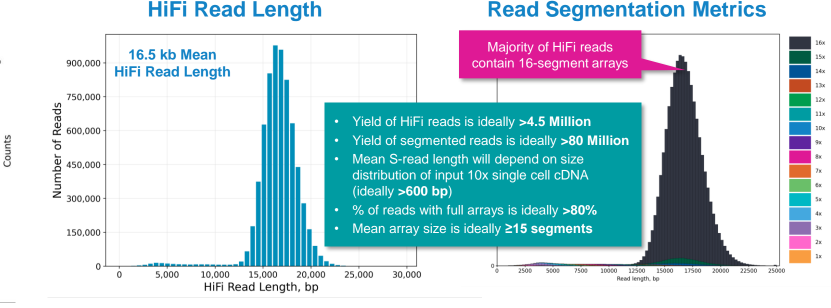
Example sequencing performance for Kinnex single-cell RNA libraries prepared with human cDNA

Revio system example data¹ – Kinnex single-cell RNA 3' library sample

Raw Data Report

Raw Base Yield	1,289 Gb
Mean Polymerase Read Length	73.16 kb
P0	27%
P1	70%
P2	3%

Example sequencing metrics for a human Kinnex single-cell RNA 3' library sample run on a Revio system with Revio polymerase kit / 130 pM on-plate loading concentration (OPLC) / 24-hrs movie time.



HiFi Reads	6.7 M
HiFi Base Yield	111.24 Gb
Mean HiFi Read Length	16.55 kb
Median HiFi Read Quality	Q28
HiFi Read Mean # of Passes	8

For human Kinnex single-cell RNA libraries, per-Revio SMRT Cell HiFi read counts were typically \sim 4 – 7 Million depending on the final library insert size and *P1* loading performance.

Input HiFi Reads	6,673,602
Segmented reads (S-reads)	104,869,257
Mean length of S-reads	1,031 bp
Percent of reads with full arrays	93.89%
Mean array size (concentration factor)	15.71

For Kinnex single-cell RNA libraries, per-Revio SMRT Cell segmentation read counts were typically >80 Million.

PacBi

¹ HiFi read lengths, reads/data per SMRT Cell and other sequencing performance results can vary depending on DNA sample quality, insert size, *P1* loading performance & movie time. Note: Shorter library insert sizes (<15 kb), lower DNA quality samples, and suboptimal *P1* loading performance may result in HiFi data yields <90 Gb per Revio SMRT Cell.

Example sequencing performance for Kinnex single-cell RNA libraries prepared with human cDNA

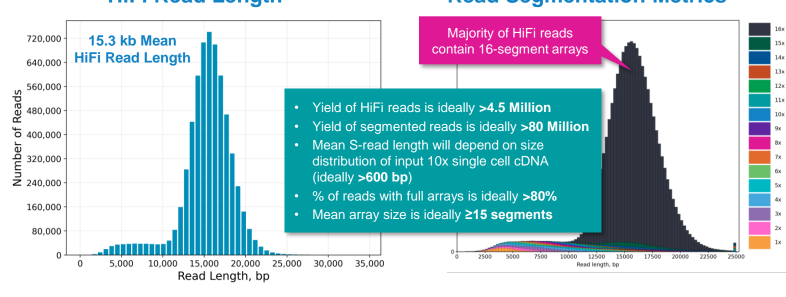
Revio system example data¹ – Kinnex single-cell RNA 5' library sample

How the set of the set

Raw Data Report

Raw Base Yield	1,116 Gb
Mean Polymerase Read Length	74,7 kb
P0	40%
P1	59%
P2	1%

Example sequencing metrics for a human Kinnex single-cell RNA 5' library sample run on a Revio system with Revio polymerase kit / 130 pM on-plate loading concentration (OPLC) / 24-hrs movie time.



HiFi Reads	6.1 M
HiFi Base Yield	93.7 Gb
Mean HiFi Read Length	15.3 kb
Median HiFi Read Quality	Q30
HiFi Read Mean # of Passes	9

For human Kinnex single-cell RNA libraries, per-Revio SMRT Cell HiFi read counts were typically \sim 4 – 7 Million depending on the final library insert size and *P1* loading performance.

Input HiFi Reads	6,104,086
Segmented reads (S-reads)	91,323,803
Mean length of S-reads	980 bp
Percent of reads with full arrays	87.46%
Mean array size (concentration factor)	14.96

For Kinnex single-cell RNA libraries, per-Revio SMRT Cell segmentation read counts were typically >80 Million.

PacBi

¹ HiFi read lengths, reads/data per SMRT Cell and other sequencing performance results can vary depending on DNA sample quality, insert size, *P1* loading performance & movie time. Note: Shorter library insert sizes (<15 kb), lower DNA quality samples, and suboptimal *P1* loading performance may result in HiFi data yields <90 Gb per Revio SMRT Cell.

HiFi Read Length

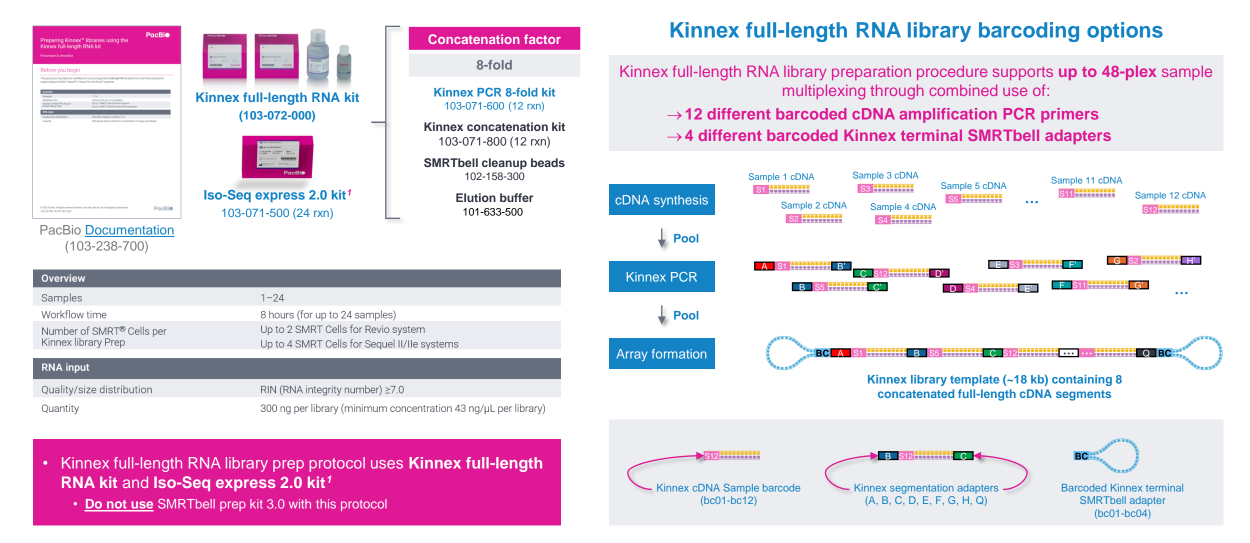
Read Segmentation Metrics

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Kinnex full-length RNA library preparation & sequencing workflow key highlights

Kinnex full-length RNA library preparation procedure description

Procedure & checklist – Preparing Kinnex libraries using the Kinnex full-length RNA kit (103-238-700) describes the workflow for constructing Kinnex libraries from total RNA samples using the Kinnex full-length RNA kit for sequencing on PacBio Sequel II, Sequel IIe & Revio systems.



Comparison of Iso-Seq express 2.0 kit vs. Iso-Seq express oligo kit





	Iso-Seq express 2.0 kit (103-071-500)		Iso-Seq express oligo kit (101-737-500)
Description	Includes Iso-Seq Express template switching oligo, barcoded cDNA PCR Primers, and other reagents needed for performing 1 st -strand cDNA synthesis and PCR amplification of cDNA products		Includes Iso-Seq express template switching oligo and non-barcoded cDNA PCR Primer to be used in conjunction with third-party reagents for performing 1 st -strand cDNA synthesis and PCR amplification of cDNA products
# reactions		2	24
Storage	Refer to product insert for storage instructions		-70°C to -80°C
Compatible SMRTbell library types	Standard (non-concatenated) library Kinnex (concatenated) library		Standard (non-concatenated) library
Sample multiplexing support	Includes 12 barcoded Iso-Seq PCR primers (bc01 – bc12) for up to 12- plex sample multiplexing		Requires additional purchase of barcoded PCR primers for cDNA amplification from a third-party vendor
Kit contents	Iso-Seq RT buffer Iso-Seq RT primer mix Iso-Seq RT enzyme mix Iso-Seq cDNA PCR mix	Iso-Seq template switch oligo Iso-Seq cDNA amplification primer Iso-Seq primers (bc01 – bc12)	Iso-Seq Express template switching oligo Iso-Seq Express cDNA PCR Primer
SMRT Link support for barcoded cDNA primer Iso- Seq libraries	YES (select 'Iso-Seq v2 Barcoded cDNA Primers' for primer set to use for Iso-Seq analysis application)		YES (select 'Iso-Seq 12 Barcoded cDNA Primers' or 'Custom cDNA Primers' for primer set to use for Iso-Seq analysis application)

Kinnex full-length RNA experimental design considerations

Kinnex full-length RNA application use case recommendations for PacBio systems

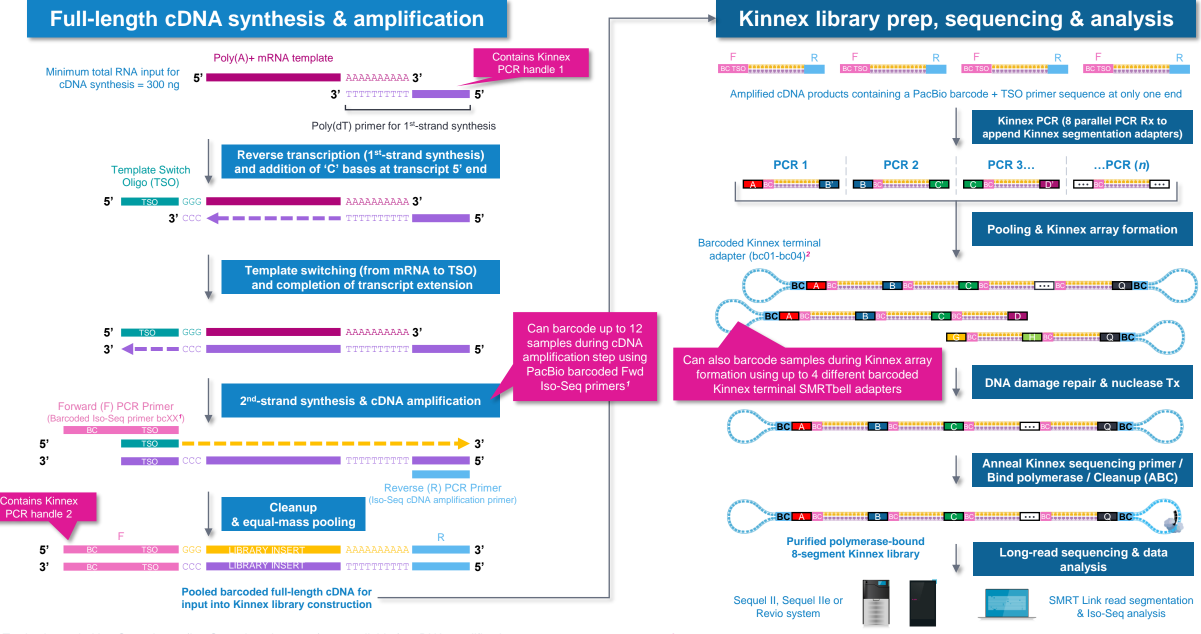
Example application	Human genetics disease studies	Biopharma for identifying highly expressed targets	Plant & animal whole genome annotation	
Experimental goal	Isoform discovery and quantification of moderate-to-rare transcripts			
Example study design	Disease vs. normal tissues with multiple replicates			
Target depth of coverage per sample	10 M reads per sample	5 M reads per sample	≤5 M reads per sample	
Comple multiplevip a1	Sequel II/IIe system: Up to 2 samples per SMRT Cell 8M (2-plex)	Sequel II/IIe system: Up to 3 samples per SMRT Cell 8M (3-plex)	Sequel II/IIe system: Up to 3 tissue types per SMRT Cell 8M (3-plex)	
Sample multiplexing ¹	Revio system: Up to 4 samples per Revio SMRT Cell (4-plex)	Revio system: Up to 8 samples per Revio SMRT Cell (8-plex)	Revio system: Up to 8 tissue types per Revio SMRT Cell (8-plex) ²	
Expected data throughput (per SMRT	Sequel II/IIe s	ystem: 15 M reads per SMRT Cell 8M divided	by <i>N</i> samples	
Cell)	Revio system: 40 M reads divided by N samples			
Kinnex library prep protocol	Procedure & checklist – Preparing Kinnex libraries using the Kinnex full-length RNA kit (103-238-700)			
Total RNA input into Kinnex library prep workflow	300 ng total RNA (RIN ≥7) for 1 st -strand cDNA synthesis			
SMRT Link data analysis workflows	Read Segmentation and Iso-Seq analysis application with option to "pool reads and cluster together" to get a master isoform classification file with per-sample full-length read counts			
Community data analysis tools	Community data analysis tools Annotation & quantification: PIGEON, SQANTI3, Differential analysis: TappAS, Fusion calling: pbfusion, Visualization: SWAN			

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¹ Kinnex concatenation kit (103-071-800) can support up to 48-plex sample multiplexing through the combined use of 12 different barcoded cDNA amplification primers and 4 different barcoded Kinnex terminal SMRTbell adapters during Kinnex full-length RNA library construction.

² If targeting <5 M transcripts reads per sample \rightarrow can multiplex up to 12 tissues types per Revio SMRT Cell.

Kinnex full-length RNA method overview



¹ Twelve barcoded Iso-Seq primers (Iso-Seq primer bc01–12) are available for cDNA amplification step.

² Kinnex adapter barcode sequences can be downloaded from <u>SMRT Link</u> Data Management module.

Kinnex full-length RNA library preparation procedural notes

Library insert generation

cDNA amplification

SMRTbell bead **cDNA** amplification

Set up on ice and add PCR reaction to thermal cycler after the lid has preheated to

SMRTbell bead

Pool + SMRTbell bead cleanup

Kinnex PCR

Kinnex PCR

Kinnex PCR

(8-mer)

- Can transfer entire volume of primers to PCR tubes for ease of use with multi-channel pipettes (8 primer mix tubes)
- Set up on ice and add PCR reaction to thermal cycler after lid has preheated to ٠ 105°C to avoid digestion of primers by polymerase exonuclease activity

Pooling of 8 Kinnex PCR products + SMRTbell bead cleanup

- Pool exactly 23 µL from each Kinnex PCR reaction for a total combined volume of 184 μL
- Add exactly **193** µL of SMRTbell cleanup beads (**1.05X**) •
- Kinnex PCR mix significantly increases stringency of SMRTbell clean up beads, so accurate pipetting is critical

105°C to avoid digestion of primers by polymerase exonuclease activity Barcoded primers are used during cDNA amplification .

12 barcoded forward primers (bc01-bc12) available for use in combination with • Iso-Seq cDNA amplification primer

SMRTbell bead cleanup

Previous non-Kinnex full-length RNA (bulk monomer) Iso-Seg protocol (Procedure & checklist – Preparing Iso-Seg libraries using SMRTbell prep kit 3.0 [102-396-000]) offered 3 options for post-cDNA amplification SMRTbell bead cleanup: 0.82X, 0.86X, and $0.95X \rightarrow$ For simplification. Kinnex full-length RNA protocol now only specifies to use 0.9X

Kinnex array formation



Kinnex digestion

- Recommended input amount to proceed with Kinnex array formation is 5 µg of Kinnex PCR amplicons
 - Proceeding with <3 µg is not recommended since lower input amounts may lead to ٠ insufficient final library yields to enable optimal sequencing results

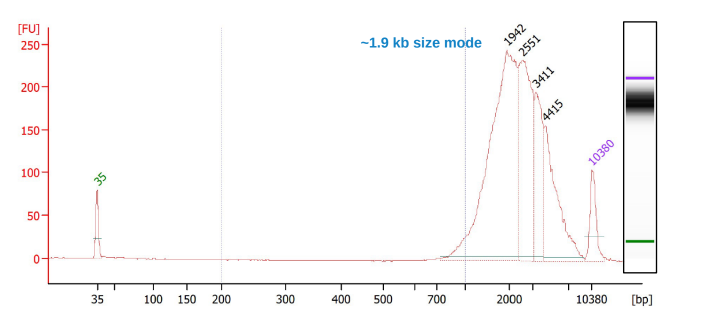
Final SMRTbell bead cleanup

Perform 1.0X SMRTbell bead cleanup on final library

Example Kinnex full-length RNA library preparation QC results

Kinnex full-length RNA library prepared with human UHRR total RNA

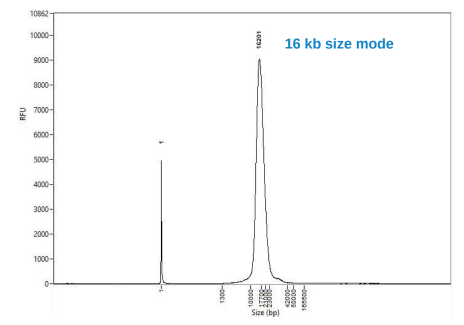
Amplified full-length cDNA QC



Example Bioanalyzer DNA sizing QC analysis results for amplified full-length cDNA generated from a universal human RNA reference (UHRR) total RNA sample.

Final Kinnex library yield is typically sufficient to load ≥2 SMRT Cells

Final Kinnex full-length RNA library QC



Example Femto Pulse DNA sizing QC analysis results for final Kinnex full-length RNA library.

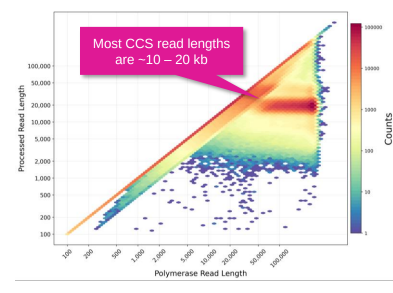
Total RNA input for cDNA synthesis	300 ng
cDNA input for Kinnex array formation	5900 ng
Post-nuclease treatment & final library cleanup yield (%) ¹	1460 ng (24.7%)

¹ Post-nuclease treatment & final cleanup yields typically ranged from ~10% to ~25% when using UHRR total RNA samples for Kinnex full-length RNA library construction.

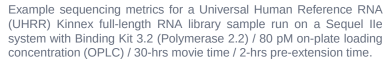
Example sequencing performance for Kinnex full-length RNA libraries prepared with human cDNA

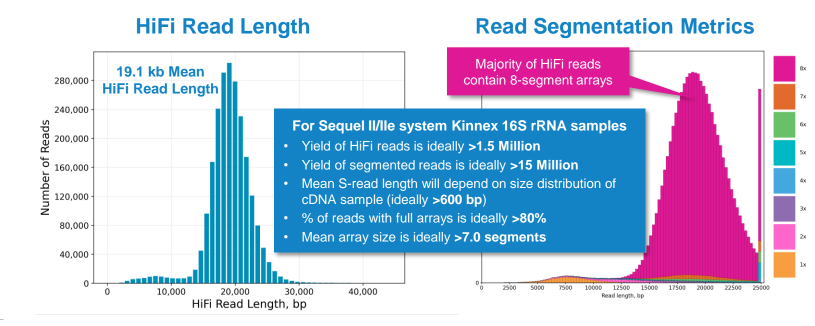
Sequel IIe system example data¹

Raw Data Report



Raw Base Yield	652 Gb
Mean Polymerase Read Length	115.36 kb
P0	28%
P1	71%
P2	1%





HiFi Reads	2.3 M
HiFi Base Yield	43.2 Gb
Mean HiFi Read Length	19.1 kb
Median HiFi Read Quality	Q32
HiFi Read Mean # of Passes	10

For UHRR Kinnex full-length RNA libraries, per-SMRT Cell 8M HiFi read counts typically ranged from ~2 - 3 Million depending on the final library insert size.

Reads	2,260,039
Segmented reads (S-reads)	17,213,165
Mean length of S-reads	2,420 bp
Percent of reads with full arrays	91.07%
Mean array size (concentration factor)	7.62

For UHRR Kinnex libraries, per-SMRT Cell 8M segmentation read counts were typically ~15 - 20 Million.

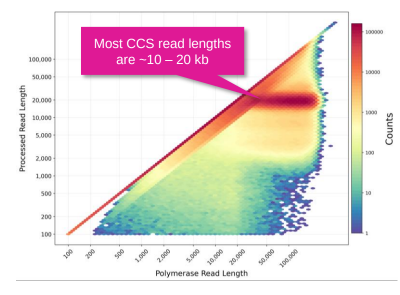


¹ HiFi read lengths, reads/data per SMRT Cell and other sequencing performance results can vary depending on DNA sample quality, insert size, P1 loading performance & movie time. Note: Shorter library insert sizes (<15 kb), lower DNA quality samples, and suboptimal P1 loading performance may result in HiFi data yields <30 Gb per Sequel II SMRT Cell 8M.

Example sequencing performance for Kinnex full-length RNA libraries prepared with human cDNA

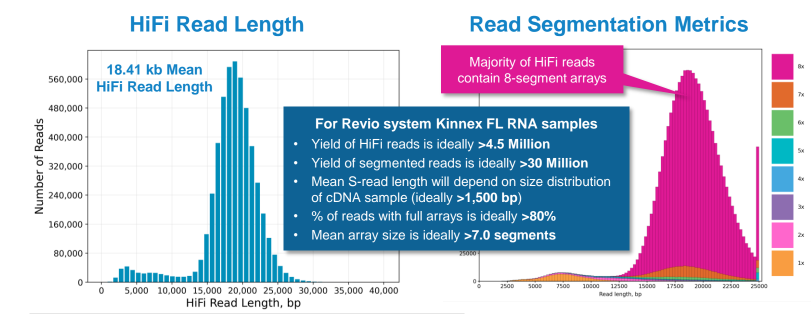
Revio system example data¹

Raw Data Report



Raw Base Yield	1,168 Gb
Mean Polymerase Read Length	58.6 kb
P0	16%
P1	79%
P2	5%

Example sequencing metrics for a Universal Human Reference RNA (UHRR) Kinnex full-length RNA library sample run on a Revio system with Revio polymerase kit / 130 pM on-plate loading concentration (OPLC) / 24-hrs movie time.



HiFi Reads	5.1 M
HiFi Base Yield	93.47 Gb
Mean HiFi Read Length	18.41 kb
Median HiFi Read Quality	Q28
HiFi Read Mean # of Passes	7

For UHRR Kinnex full-length RNA libraries, per-Revio SMRT Cell HiFi read counts were typically ~5 - 6 Million depending on the final library insert size and P1 loading performance.

Input HiFi Reads	5,027,154
Segmented reads (S-reads)	37,216,151
Mean length of S-reads	2,393 bp
Percent of reads with full arrays	85.84%
Mean array size (concentration factor)	7.40

For UHRR Kinnex libraries, per-Revio SMRT Cell segmentation read counts were typically ~30 - 45 Million.



¹ HiFi read lengths, reads/data per SMRT Cell and other sequencing performance results can vary depending on DNA sample quality, insert size, P1 loading performance & movie time. Note: Shorter library insert sizes (<15 kb), lower DNA quality samples, and suboptimal P1 loading performance may result in HiFi data yields <90 Gb per Revio SMRT Cell.

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Kinnex 16S rRNA library preparation & sequencing workflow key highlights

Kinnex 16S rRNA library preparation procedure description

Procedure & checklist – Preparing Kinnex libraries from 16S rRNA amplicons (103-238-800) describes the workflow for constructing Kinnex libraries from full-length 16S amplicons using the Kinnex 16S rRNA kit* for sequencing on PacBio Sequel II, Sequel IIe & Revio systems.



This procedure provides instructions for generating Kinnex libraries from full-length 16S amplicons for sequencing on PacBio® Sequel® II, Sequel IIe, and Revio™ systems.

- 1. Amplification of full-length 16S genes (V1-V9 regions) from metagenomic samples using barcoded Forward and Reverse 16S primers
- Concatenation of 16S amplicons to ~19 kb 2.
- 3. Multiplexed sequencing on the Sequel II/IIe and Revio systems

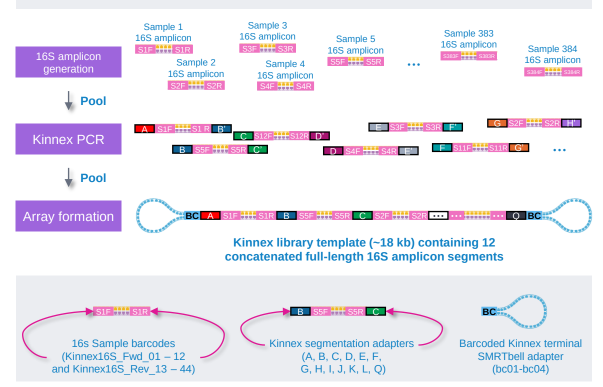
Barcoded 16S-specific primers (12 forward and 32 reverse) can be used in different combinations allowing for the multiplexing of up to 384 samples on one SMRT® Cell. If combined with barcoded Kinnex adapters (4-plex), a total of 1536 samples can be sequenced.

- Kinnex full-length RNA library prep protocol uses Kinnex 16S rRNA kit
 - Do not use SMRTbell prep kit 3.0 with this protocol

Kinnex full-length 16s rRNA library barcoding options

Kinnex 16S rRNA library preparation procedure supports up to 1,536-plex sample multiplexing through combined use of:

- \rightarrow 12 different 16S barcoded Forward PCR primers¹
- \rightarrow 32 different 16S barcoded Reverse PCR primers¹
- \rightarrow 4 different barcoded Kinnex terminal SMRTbell adapters



PacBi 16S barcoded Forward and Reverse PCR primer oligos (HPLC purified) are not included in Kinnex 16S rRNA kit and must be supplied by users through a third-party oligo vendor.

Kinnex 16S rRNA experimental design considerations

Kinnex 16S rRNA application use case recommendations for PacBio systems

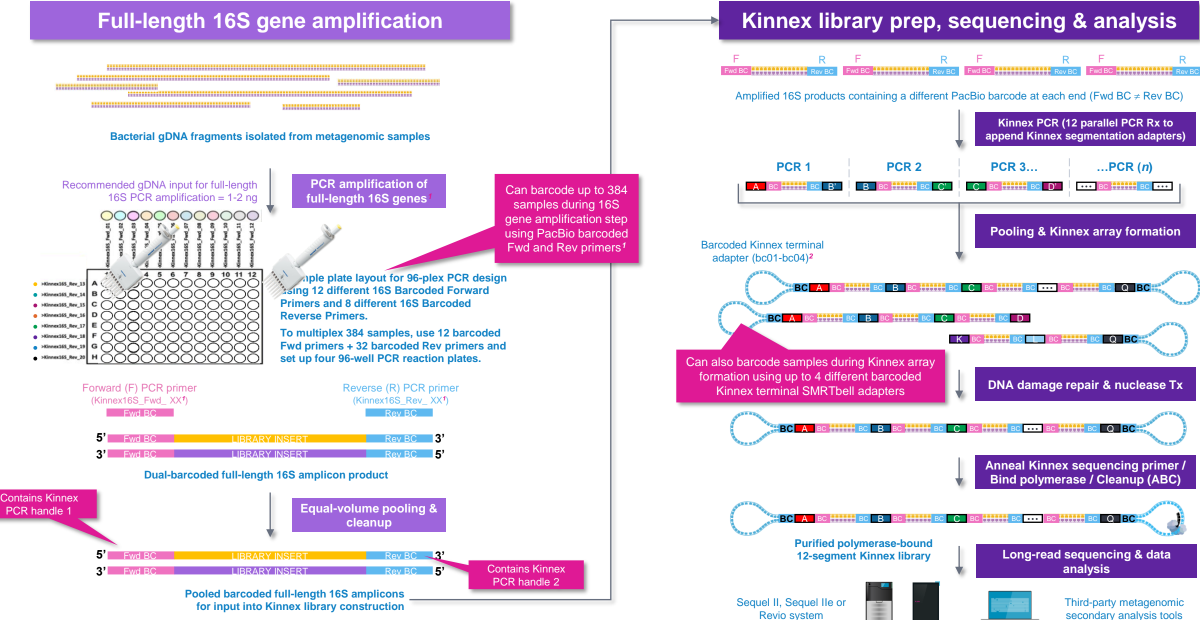
	Sequel II and IIe systems		Revio	system
Experimental goal	Determine the microbial diversity (phylogeny and taxonomy) of bacteria in a metagenomic sample			
Sample multiplexing ¹	Up to 384 samples per SMRT Cell 8M (384-plex)		Up to 1,536 samples per Revio	SMRT Cell (1536-plex)
Expected coverage per sample ²	96-plex	260 K	96-plex	625 K
	192-plex	130 K	192-plex	313 K
	384-plex	65 K	384-plex	156 K
	768-plex	33 K	768-plex	78 K
	1,536-plex	16 K	1,536-plex	39 K
Kinnex library prep protocol	Procedure & checklist – Preparing Kinnex libraries from 16S rRNA amplicons (103-238-800)			
Metagenomic DNA input amount input into 16S gene amplification	1-2 ng of input gDNA per metagenomic sample			
16S amplicon DNA input into Kinnex library prep workflow	35 ng of purified pooled 16S amplicon DNA			
SMRT Link data analysis workflows	Read Segmentation			
Community data analysis tools	pb-16S-nf			

¹ Kinnex concatenation kit (103-071-800) can support up to 1,536-plex sample multiplexing through the combined use of 12 different 16S barcoded Forward PCR primers + 32 different 16S barcoded Reverse PCR primers and 4 different barcoded Kinnex terminal SMRTbell adapters during Kinnex 16s rRNA library construction.



² With proper full array formation and adequate sequencing, one SMRT Cell on the Sequel II, IIe, and Revio systems are expected to achieve 20–25 million and 50–60 million 16S sequences, respectively. For most 16S analysis applications, typically aim for ~30-50 K reads/sample.

Kinnex 16S rRNA method overview



¹ 12 different 16S barcoded Forward PCR primers + 32 different 16S barcoded Reverse PCR primers are available for 16S gene amplification step to multiplex up to 384 samples.

² Kinnex adapter barcode sequences can be downloaded from <u>SMRT Link</u> Data Management module.

Kinnex 16S rRNA library preparation procedural notes

Library insert generation

PCR amplification of full-length 16S genes

PCR amplification of F-L 16S genes

٠

٠

Pool barcoded 16S SMRTbell bead PCR amplicons

Set up on ice and add PCR reaction to thermal cycler after the lid has preheated to

Customer supplies Kapa PCR mix (HiFi HotStart ReadyMix) and oligos

Up to 384-plex can be done at this point using combinatorial indexing

105°C to avoid digestion of primers by polymerase exonuclease activity

SMRTbell bead

SMRTbell bead **cleanup**

Kinnex PCR

Kinnex PCR

Kinnex PCR

(12-mer)

- Can transfer entire volume of primers to PCR tubes for ease of use with multi-channel pipettes (12 primer mix tubes)
- Set up on ice and add PCR reaction to thermal cycler after lid has preheated to **105°C** to avoid digestion of primers by polymerase exonuclease activity

SMRTbell bead cleanup

- Pool exactly 23 µL from each Kinnex PCR reaction for a total combined volume of 276 μL
- Add exactly 304 µL of SMRTbell clean up beads (1.1X)
- Kinnex PCR mix significantly increases stringency of SMRTbell clean up beads, so accurate pipetting is critical

Kinnex array formation

DNA damage SMRTbell bead Nuclease SMRTbell bead Kinnex digestion / Ligation cleanup treatment cleanup repair

Kinnex digestion

- Recommended input amount to proceed with Kinnex array formation is 5 µg of Kinnex ٠ PCR amplicons
 - Proceeding with <3 µg is not recommended since lower input amounts may lead to ٠ insufficient final library yields to enable optimal sequencing results

Final SMRTbell bead cleanup

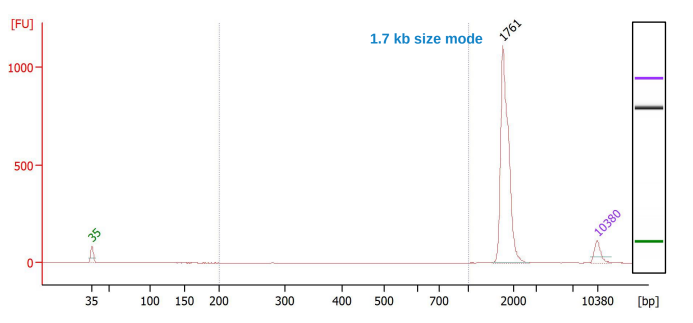
Perform 1.0X SMRTbell bead cleanup on final library

Example Kinnex 16S rRNA library preparation QC results

Kinnex full-length 16S RNA library prepared from mock microbial community genomic DNA

Final Kinnex library yield is typically sufficient to load ≥2 SMRT Cells

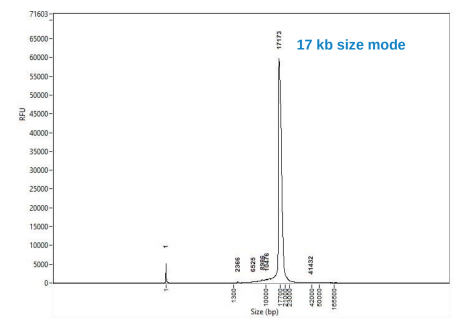
16S amplicon DNA QC



Example Bioanalyzer DNA sizing QC analysis results for pooled 16S amplicon DNA samples generated from mock microbial community genomic DNA (ATCC MSA-1003 20 Strain Staggered Mix).



Final Kinnex 16S rRNA library QC



Example Femto Pulse DNA sizing QC analysis results for final Kinnex 16S rRNA library.

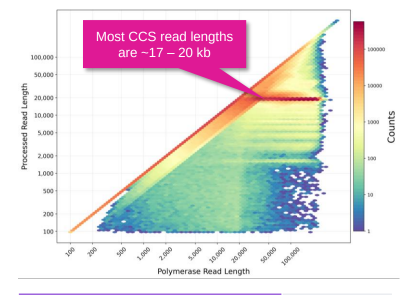
gDNA input for 16S PCR	1.1 ng
16S amplicon DNA input Kinnex PCR products for Kinnex array formation	6000 ng
Post-nuclease treatment & final library cleanup yield (%) ¹	1080 ng (18%)

¹ Post-nuclease treatment & final cleanup yields typically ranged from ~10% to ~20% when using mock microbial community genomic DNA for Kinnex full-length 16S rRNA library construction.

Example sequencing performance for Kinnex 16S rRNA libraries prepared from mock microbial community genomic DNA

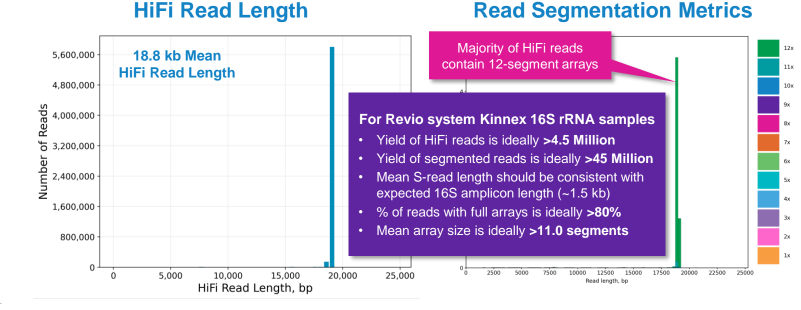
Revio system example data¹ (1,536-plex data set)

Raw Data Report



Raw Base Yield	1,222 Gb
Mean Polymerase Read Length	69.0
P0	27%
P1	70%
P2	3%

Example sequencing metrics for a Kinnex 16S rRNA library sample run on a Revio system with Revio polymerase kit / 130 pM on-plate loading concentration (OPLC) / 24-hrs movie time.



HiFi Reads	6.1 M
HiFi Base Yield	114.21 Gb
Mean HiFi Read Length	18.78 kb
Median HiFi Read Quality	Q32
HiFi Read Mean # of Passes	7

For Kinnex 16S rRNA libraries, per-Revio SMRT Cell HiFi read counts were typically \sim 4 – 6 Million depending on the final library insert size and *P1* loading performance.

Input HiFi Reads	6,050,730
Segmented reads (S-reads)	71,720,714
Mean length of S-reads	1,560 bp
Percent of reads with full arrays	95.03%
Mean array size (concentration factor)	11.85

For Kinnex 16S rRNA libraries, per-Revio SMRT Cell segmentation read counts were typically ~45 – 60 Million.



¹ HiFi read lengths, reads/data per SMRT Cell and other sequencing performance results can vary depending on DNA sample quality, insert size, *P1* loading performance & movie time. Note: Shorter library insert sizes (<15 kb), lower DNA quality samples, and suboptimal *P1* loading performance may result in HiFi data yields <90 Gb per Revio SMRT Cell.

Improving sequencing performance of "difficult" 16S samples

Performing AMPure PB bead size-selection on Kinnex full-length 16S rRNA libraries can help improve *P1* loading of challenging metagenomic samples

Sample Name	P1 %	Gb Yield	Mean Length	Mean QV
16S_collaborator_SOP	26	46	18,813 bp	Q29
16S_collaborator_3.1X AMPure	80	87	18,851 bp	Q28

Some bacterial 16S samples may have carry-over contaminants present leading to low *P1* loading on Revio and Sequel II/IIe systems

 \rightarrow Using AMPure size-selection (3.1X 35% AMPure PB beads) can help mitigate this issue

PacBi

Technical documentation & applications support resources

Technical resources for Kinnex library preparation, sequencing & data analysis

Single-cell cDNA sample preparation literature & other resources

- 10x Genomics Chromium Next GEM Single Cell 3' v3.1 (Single Index) How-to Video [Link]
- 10x Genomics Chromium Single Cell 3' Reagent Kits User Guide v3.1 (CG000204)
- 10x Genomics Chromium Single Cell 5' Reagent Kits User Guide v2 Chemistry Dual Index (CG000331)

Kinnex library preparation literature & other resources

- Application note Kinnex 16S rRNA kit for full-length 16S sequencing (<u>102-326-601</u>)
- Application note Kinnex full-length RNA kit for isoform sequencing (<u>102-326-591</u>)
- Application note Kinnex single-cell RNA kit for single-cell isoform sequencing (<u>102-326-549</u>)
- Brochure Scalable, cost-effective RNA sequencing with PacBio Kinnex kits (<u>102-326-597</u>)
- Procedure & checklist Preparing Kinnex libraries using Kinnex single-cell RNA kit (<u>103-254-300</u>)
- Procedure & checklist Preparing Kinnex libraries using Kinnex full-length RNA kit (<u>103-238-700</u>)
- Procedure & checklist Preparing Kinnex libraries from 16s rRNA amplicons (<u>103-238-800</u>)
- Technical overview Kinnex kits for single-cell RNA, full-length RNA and 16S rRNA sequencing (103-343-700)
- Technical overview Kinnex library preparation for full-length 16S rRNA gene sequencing (<u>103-344-800</u>)
- Technical overview Kinnex library preparation using Kinnex full-length RNA kit (<u>103-344-700</u>)
- Technical overview Kinnex library preparation using Kinnex single-cell RNA kit (<u>103-344-600</u>)
- Video tutorial PacBio Kinnex single-cell RNA TSO artifact removal demo for Kinnex single-cell RNA kit [Link]
- Video tutorial SMRT Link Sample Setup and Run Design setup procedure for Kinnex kits [Link]
- Whitepaper Bulk and single-cell isoform sequencing for human disease research (<u>102-326-576</u>)

Technical resources for Kinnex library preparation, sequencing & data analysis (cont.)

Data analysis resources

- Application note Bioinformatics tools for full length isoform sequencing (<u>102-326-593</u>)
- SMRT Link MAS-Seq troubleshooting guide (<u>102-994-400</u>)
- SMRT Link software installation guide [Link]
- SMRT Link user guide [Link]
- SMRT Tools reference guide [Link]
- Video tutorial Analyzing Kinnex 16S rRNA data in SMRT Link: [Link]
- Video tutorial Read Segmentation and Iso-Seq workflow in SMRT Link: [Link]

Publications and posters

- Schertzer, M.D. et al. (2023) Cas13d-mediated isoform-specific RNA knockdown with a unified computational and experimental toolbox. BioRxiv preprint [Link]
- Al'Khafaji, A.M. et al. (2023) High-throughput RNA isoform sequencing using programmable cDNA concatenation. Nature biotechnology. [Link]
- ASM Microbe Poster (2023) Increasing throughput of full-length 16S sequencing using concatenation [Link]

Webinars

- PacBio webinar (2023) Understanding clonal evolution using game theory and single-cell long-read isoform analysis [Link]
- PacBio Iso-Seq social club webinar (2022) Introduction to Iso-Seq method [Link]
- PacBio Iso-Seq social club webinar (2022) SQANTI3 for isoform classification and annotation [Link]
- PacBio Iso-Seq social club webinar (2022) TappAS for isoform differential expression analysis [Link]
- PacBio Iso-Seq social club webinar (2022) Single-cell Iso-Seq applications in cancer and neurological disorders [Link]

Technical resources for Kinnex library preparation, sequencing & data analysis (cont.)

Example PacBio data sets

Application	Dataset	Data type	PacBio system
Kinnex single-cell RNA sequencing	Homo sapiens - PBMC 10x Chromium Single Cell 5' and 3' libraries [Link]	HiFi long read	Sequel II & Revio systems
	Homo sapiens - HG002 (10x 5') [Link]	HiFi long read	Revio system
Kinnex full-length RNA sequencing	Homo sapiens – universal human reference RNA (UHRR) [<u>Link</u>]	HiFi long read	Sequel II & Revio systems
	Homo sapiens – HG002 [<u>Link</u>]	HiFi long read	Revio system
	Homo sapiens – Heart [Link]	HiFi long read	Revio system
	Homo sapiens – Cerebellum [Link]	HiFi long read	Revio system
Kinnex 16S rRNA sequencing	ZymoBIOMICS Fecal Reference with TruMatrix Technology (human) [Link]	HiFi long read	Sequel II & Revio systems
	Mixture: ZymoBIOMICS Gut Microbiome Standard, ZymoBIOMICS Fecal Reference with TruMatrix™ Technology, ATCC 20 Strain Even Mix Genomic Material, ATCC 20 Strain Staggered Mix Genomic Material [Link]	HiFi long read	Sequel II & Revio systems

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