# PacBi•

# COMPLETE BIOLOGICAL INSIGHTS FOR CONFIDENT DECISIONS

Unlock greater value with HiFi long-read sequencing systems













#### **Genomes**

Complete + accurate genomes

#### **Transcriptomes**

Ful-length RNA isoforms

#### **Epigenomes**

Directly phased methylomes

#### **Metagenomics**

Species-level resolution for profiling and assembly

## Targeted sequencing

Comprehensive variant calling for regions of interest

HiFi sequencing is SMRT® business

#### The HiFi accuracy you love, at the price and throughput you need



HiFi sequencing is the only known long-read technology to provide 99.9% or better accuracy in every run. HiFi reads eliminate the tradeoff between read length and accuracy, allowing you to address your most complex biological questions.

With over 1,000 peer-reviewed publications in 2023 alone and over 1,200 HiFi sequencers globally, the PacBio portfolio of reliable and established sequencing systems includes both the production-scale Revio® and the Vega™ benchtop system.

#### The benefits of HiFi reads



Long read lengths



Supported analysis tools consolidated into a single pipeline



Easy library preparation



High read accuracy



Low coverage requirements



Compatible ecosystem partners to enable an end-to-end workflow



Small file sizes to minimize compute time



A single technology solution

# 99.9 99.9 98.2 98.3 96.9 97.0 % Indels SVs ■ Revio system with SPRQ™ ■ Vega system

Data shown is for a single Revio or Vega SMRT® Cell

# Two systems, one gold standard

Vega and Revio systems deliver the same high-standard HiFi sequencing, ensuring consistent, accurate data. HiFi reads cover all regions of the genome, providing the most complete and precise assemblies and variant calls. Unlike short reads, which can miss complex regions, HiFi captures full-length transcripts and difficult areas. With either HiFi system, you get the best possible genome with low coverage requirements.







Built on Revio technology, the Vega **benchtop system** brings HiFi sequencing into any lab, giving you full control to produce complete genomes, full-length isoforms, and directly-phased methylomes.





#### HiFi at scale

The Revio system revolutionized HiFi sequencing with **production-scale power**. Now with SPRQ chemistry, Revio requires less DNA input, expands multiomic applications, decreases costs, and includes on-board 6mA methylation calling.

1 SMRT® Cell per run	Up to 4 SMRT Cells per run
60 Gb per run	480 Gb per run (4 SMRT Cells)
2 μg DNA input	500 ng DNA input
12 or 24-hour run times	12, 24, or 30-hour run times
5mC and 6mA methylation calling	5mC and 6mA methylation calling
Vega price: \$169k system at \$1,100 per run Vega Access: \$79k system at \$1,750 per run*	\$599k system at \$995 per SMRT Cell

Prices shown in US dollar and at list price, may differ by region. Your local sales representative can provide detailed pricing in your currency.

 $<sup>{\</sup>rm *Vega\ Access\ offers\ the\ Vega\ system\ at\ \$79k\ and\ \$1,750\ per\ run\ for\ the\ first\ 152\ sequencing\ reagents\ and\ \$1100\ thereafter.}$ 

#### Seamless workflow

Revio and Vega systems feature powerful NVIDIA GPUs for accurate, ready-to-use data directly from the sequencer. With streamlined consumables and flexible software, both systems simplify the sequencing workflow – for any genome center, service provider, or research lab.



#### Boost your efficiency

Fully automated sequencing with loading under a minute

Run preview and the capability to pre-stage consumables for the next run keeps you one step ahead

On-board primary analysis including Google DeepConsensus, methylation calling, and demultiplexing in every run







#### Sequencing, simplified

Universal HiFi workflow from extraction to analysis across all HiFi systems

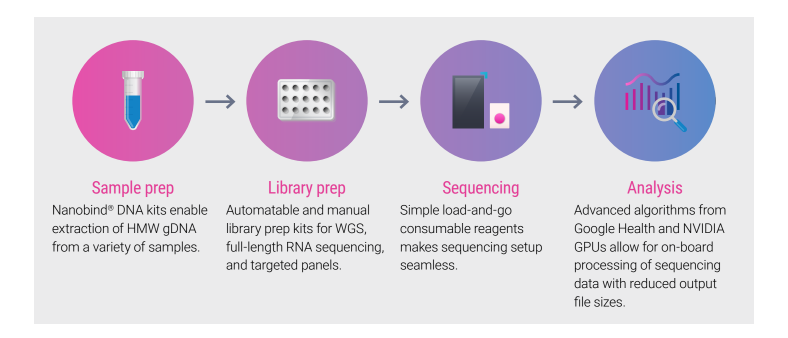
Intuitive user interface with simple, NFC tagged consumables

Standard BAM file format with direct-to-cloud upload

Web-based **SMRT**<sup>®</sup> **Link** software supports sequencing prep, run design, and data analysis in a single interface across a wide range of applications.

**SMRT® Analysis** provides fast, easy, and powerful computational tools and workflows to achieve a complete view of the genome or region of interest.

#### Universal HiFi workflow from sample prep to analysis





Our 30+ **PacBio Compatible partners** integrate products optimized for HiFi sequencing at every step of the workflow, from sample prep to analysis.



Nanobind extraction kits employ proprietary Nanobind disks for the extraction of DNA from diverse sample types, including saliva, blood, cell, and tissue samples. With Nanobind, researchers can extract longer fragments from high-quality DNA in less than two hours, with minimal sample input and without the use of harmful chemicals or mechanical homogenization.















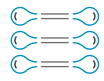




Whether you are doing full-length RNA sequencing, single-cell RNA sequencing, or 16S rRNA, **Kinnex™** offers full-length isoform discovery and abundance information, cell type-specific isoform characterization, and species-level metagenomics community identification that are difficult to capture with short reads.



**PureTarget™** generates targeted native DNA libraries with an amplification-free approach which retains epigenetic signals and has no PCR artifacts or errors. Now you can screen the most challenging genomic regions at scale. The PureTarget product line includes a panel of the most important repeat expansions for human health in genes like *FMR1*, *C9orf72*, and *RFC1*.



SMRTbell® library generation using **HiFi prep kit 96 and HiFi plex prep kit 96** offers automation-friendly and scalable library prep solutions for projects and genomes sizes from microbial to human.

#### What can you do with 1 SMRT Cell?

PacBio is the only sequencing technology to offer HiFi reads that provide accuracy of 99.9%. With one SMRT Cell on either the Vega or Revio system, you can sequence a human genome at 20x coverage, multiplex up to 384 microbial genomes, or analyze full-length RNA isoforms—all with the accuracy and completeness of HiFi sequencing. This flexibility allows you to scale your research across whole genomes, targeted regions, or transcriptomes with ease.



Vega™ system HiFi within reach

1 SMRT® Cell 1 SMRT Cell 4 SMRT Cells

HiFi at scale

Whole genome sequencing           Human genome (20x coverage)         1         2         8           Human methylation profiling (5x)         4         8         32           De novo assembly (1 Gb genome)         2         4         16           Microbial de novo assembly (2 Gb total)         384         384         1,536           Targeted panels           Amplicon sequencing         >1,000         >1,000         >4,000           Target enrichment           20 Mb panel         12         16         64           2 Mb panel         72         96         384           100 kb panel         288         384         1,536           PureTarget™ repeat expansion panel         48         48         192           RNA sequencing           Kinnex™ single-cell RNA sequencing         1 (3,000-6,000 cells)         1 (6,000-10,000 cells)         4 (6,000-10,000 cells)           Kinnex full-length RNA sequencing         6         12         48           10M reads         3         6         24           Microbial           Shotgun metagenomic profiling         64 communities         128 communities         512 communities				
Human genome (20x coverage)       1       2       8         Human methylation profiling (5x)       4       8       32         De novo assembly (1 Gb genome)       2       4       16         Microbial de novo assembly (2 Gb total)       384       384       1,536         Targeted panels         Amplicon sequencing       >1,000       >1,000       >4,000         Target enrichment         20 Mb panel       12       16       64         2 Mb panel       72       96       384         100 kb panel       288       384       1,536         PureTarget™ repeat expansion panel       48       48       192         RNA sequencing         Kinnex™ single-cell RNA sequencing       1 (3,000−6,000 cells)       1 (6,000−10,000 cells)       4 (6,000−10,000 cells)         Kinnex full-length RNA sequencing       6       12       48         10M reads       3       6       24         Microbial         Shotgun metagenomic profiling       64 communities       128 communities       512 communities         Shotgun metagenomic assembly       8 communities       16 communities       64 communities	Application	Samples per run		
Human methylation profiling (5x)       4       8       32         De novo assembly (1 Gb genome)       2       4       16         Microbial de novo assembly (2 Gb total)       384       384       1,536         Targeted panels         Amplicon sequencing       >1,000       >1,000       >4,000         Target enrichment         20 Mb panel       12       16       64         2 Mb panel       72       96       384         100 kb panel       288       384       1,536         PureTarget™ repeat expansion panel       48       48       192         RNA sequencing         Kinnex™ single-cell RNA sequencing       1 (3,000−6,000 cells)       1 (6,000−10,000 cells)       4 (6,000−10,000 cells)         Kinnex full-length RNA sequencing       5M reads       6       12       48         10M reads       3       6       24         Microbial         Shotgun metagenomic profiling       64 communities       128 communities       512 communities         Shotgun metagenomic assembly       8 communities       16 communities       64 communities	Whole genome sequencing			
De novo assembly (1 Gb genome)         2         4         16           Microbial de novo assembly (2 Gb total)         384         384         1,536           Targeted panels           Amplicon sequencing         >1,000         >1,000         >4,000           Target enrichment           20 Mb panel         12         16         64           2 Mb panel         72         96         384           100 kb panel         288         384         1,536           PureTarget™ repeat expansion panel         48         48         192           RNA sequencing           Kinnex™ single-cell RNA sequencing         1 (3,000-6,000 cells)         1 (6,000-10,000 cells)         4 (6,000-10,000 cells)           Kinnex full-length RNA sequencing         1 (3,000-6,000 cells)         1 (6,000-10,000 cells)         4 (6,000-10,000 cells)           Kinnex full-length RNA sequencing         6         12         48           10M reads         3         6         24           Microbial           Shotgun metagenomic assembly         8 communities         128 communities         512 communities	Human genome (20x coverage)	1	2	8
Microbial de novo assembly (2 Gb total)         384         384         1,536           Targeted panels           Amplicon sequencing         >1,000         >1,000         >4,000           Target enrichment	Human methylation profiling (5x)	4	8	32
Targeted panels         Amplicon sequencing       >1,000       >1,000       >4,000         Target enrichment	De novo assembly (1 Gb genome)	2	4	16
Amplicon sequencing       >1,000       >1,000       >4,000         Target enrichment       12       16       64         2 Mb panel       72       96       384         100 kb panel       288       384       1,536         PureTarget™ repeat expansion panel       48       48       192         RNA sequencing       1 (3,000−6,000 cells)       1 (6,000−10,000 cells)       4 (6,000−10,000 cells)         Kinnex full-length RNA sequencing       6       12       48         10M reads       3       6       24         Microbial         Shotgun metagenomic profiling       64 communities       128 communities       512 communities         Shotgun metagenomic assembly       8 communities       16 communities       64 communities	Microbial de novo assembly (2 Gb total)	384	384	1,536
Target enrichment20 Mb panel1216642 Mb panel7296384100 kb panel2883841,536PureTarget™ repeat expansion panel4848192RNA sequencingKinnex™ single-cell RNA sequencing1 (3,000-6,000 cells)1 (6,000-10,000 cells)4 (6,000-10,000 cells)Kinnex full-length RNA sequencing5M reads6124810M reads3624Microbial44 communities128 communities512 communitiesShotgun metagenomic profiling64 communities128 communities512 communitiesShotgun metagenomic assembly8 communities16 communities64 communities	Targeted panels			
20 Mb panel       12       16       64         2 Mb panel       72       96       384         100 kb panel       288       384       1,536         PureTarget™ repeat expansion panel       48       48       192         RNA sequencing         Kinnex™ single-cell RNA sequencing       1 (3,000-6,000 cells)       1 (6,000-10,000 cells)       4 (6,000-10,000 cells)         Kinnex full-length RNA sequencing         5M reads       6       12       48         10M reads       3       6       24         Microbial         Shotgun metagenomic profiling       64 communities       128 communities       512 communities         Shotgun metagenomic assembly       8 communities       16 communities       64 communities	Amplicon sequencing	>1,000	>1,000	>4,000
2 Mb panel       72       96       384         100 kb panel       288       384       1,536         PureTarget™ repeat expansion panel       48       48       192         RNA sequencing         Kinnex™ single-cell RNA sequencing       1 (3,000−6,000 cells)       1 (6,000−10,000 cells)       4 (6,000−10,000 cells)         Kinnex full-length RNA sequencing         5M reads       6       12       48         10M reads       3       6       24         Microbial         Shotgun metagenomic profiling       64 communities       128 communities       512 communities         Shotgun metagenomic assembly       8 communities       16 communities       64 communities	Target enrichment			
100 kb panel2883841,536PureTarget™ repeat expansion panel4848192RNA sequencingKinnex™ single-cell RNA sequencing1 (3,000−6,000 cells)1 (6,000−10,000 cells)4 (6,000−10,000 cells)Kinnex full-length RNA sequencing5M reads6124810M reads3624MicrobialShotgun metagenomic profiling64 communities128 communities512 communitiesShotgun metagenomic assembly8 communities16 communities64 communities	20 Mb panel	12	16	64
PureTarget™ repeat expansion panel4848192RNA sequencingI (3,000-6,000 cells)1 (6,000-10,000 cells)4 (6,000-10,000 cells)Kinnex full-length RNA sequencingImage: Company of the	2 Mb panel	72	96	384
RNA sequencing  Kinnex™ single-cell RNA sequencing  I (3,000-6,000 cells)  I (6,000-10,000 cells)  Kinnex full-length RNA sequencing  5M reads  6  12  48  10M reads  3  6  24  Microbial  Shotgun metagenomic profiling  64 communities  128 communities  512 communities  512 communities  64 communities  16 communities  64 communities	100 kb panel	288	384	1,536
Kinnex™ single-cell RNA sequencing1 (3,000−6,000 cells)1 (6,000−10,000 cells)4 (6,000−10,000 cells)Kinnex full-length RNA sequencing5M reads6124810M reads3624MicrobialShotgun metagenomic profiling64 communities128 communities512 communitiesShotgun metagenomic assembly8 communities16 communities64 communities	PureTarget™ repeat expansion panel	48	48	192
Kinnex full-length RNA sequencing  5M reads 6 12 48 10M reads 3 6 24  Microbial  Shotgun metagenomic profiling 64 communities 128 communities 512 communities Shotgun metagenomic assembly 8 communities 16 communities 64 communities	RNA sequencing			
5M reads 6 12 48 10M reads 3 6 24  Microbial  Shotgun metagenomic profiling 64 communities 128 communities 512 communities Shotgun metagenomic assembly 8 communities 16 communities 64 communities	Kinnex™ single-cell RNA sequencing	1 (3,000-6,000 cells)	1 (6,000-10,000 cells)	4 (6,000-10,000 cells)
10M reads 3 6 24  Microbial  Shotgun metagenomic profiling 64 communities 128 communities 512 communities  Shotgun metagenomic assembly 8 communities 16 communities 64 communities	Kinnex full-length RNA sequencing			
MicrobialShotgun metagenomic profiling64 communities128 communities512 communitiesShotgun metagenomic assembly8 communities16 communities64 communities	5M reads	6	12	48
Shotgun metagenomic profiling64 communities128 communities512 communitiesShotgun metagenomic assembly8 communities16 communities64 communities	10M reads	3	6	24
Shotgun metagenomic assembly 8 communities 16 communities 64 communities	Microbial			
	Shotgun metagenomic profiling	64 communities	128 communities	512 communities
Kinnex 16S rRNA 1.024 communities 1.536 communities 6.144 communities	Shotgun metagenomic assembly	8 communities	16 communities	64 communities
1,000 0011114111111111111111111111111111	Kinnex 16S rRNA	1,024 communities	1,536 communities	6,144 communities

All sample throughputs are estimates for either the Vega system with 1 SMRT Cell or the Revio system using SPRQ chemistry with both 1 or 4 SMRT Cells. Coverage may vary based on sample quality, library quality, and fragment lengths. Currently available SMRTbell® adapter index plates 96A-96D contain a total of 384 SMRTbell barcoded adapters. Microbial de novo assembly assumes microbes with 2 Gb of total genome size at 30x per sample. Single-cell transcriptomics assumes ≥80 million reads per library on the Revio system and ~50-60 million reads per library on the Vega system. Full-length RNA sequencing assumes a total of 60M reads for Revio SPRQ and 30M reads for Vega, regardless of plexity. Amplicon sequencing assumes a 12-hour movie time for 1–5 kb, 24-hour movie time for 5+ kb, and >50x per sample. Target enrichment assumes >50x per sample.

#### Go further with HiFi long-reads





#### Annual throughput 200 Vega SMRT Cells 1,250 Revio SMRT Cells

	Est samples per year	Est library prep + sequencing cost per sample	Est samples per year	Est library prep + sequencing cost per sample
Whole genomes				
Human genome (20x)	200	\$1,156	2,500	\$554
Human methylation profiling (5x)	800	\$330	10,000	\$180
de novo assembly (1 Gb genome)	400	\$613	5,000	\$312
Microbial <i>de novo</i> assembly (2 Gb total)	76,800	\$42	480,000	\$42
Targeted panels				
Amplicons	>200,000	<\$5	1.2M	<\$5
PureTarget repeat expansion panel	9,600	\$240	60,000	\$240
RNA sequencing				
Kinnex single-cell RNA sequencing	200 (3,000-6,000 cells each)	\$1,575	1,250 (6,000–10,000 cells each)	\$1,470
Kinnex full-length RNA sequencing	<b>5M reads</b> : 1,200 <b>10M reads</b> : 600	<b>5M reads</b> : \$420 <b>10M reads</b> : \$685	<b>5M reads</b> : 15,000 <b>10M reads</b> : 7,500	<b>5M reads</b> : \$200 <b>10M reads</b> : \$325
Microbial				
Shotgun metagenomic profiling	12,800 communities	\$50	160,000 communities	\$50
Shotgun metagenomic assembly	1,600 communities	\$200	20,000 communities	\$120
Kinnex 16S rRNA	200,000 communities	\$4	1.9M communities	\$4

Assume U.S. list pricing for PacBio products. All sample throughputs are estimates for either the Vega system with 1 SMRT Cell or the Revio system using SPRQ chemistry with 4 SMRT Cells. Coverage may vary based on sample quality, library quality, and fragment lengths. Currently available SMRTbell® adapter index plates 96A-96D contain a total of 384 SMRTbell barcoded adapters. Microbial de novo assembly assumes microbes with 2 Gb of total genome size at 30x per sample. Single-cell transcriptomics assumes  $\approx 80$  million reads per library on the Revio system and  $\approx 50$ -60 million reads per library on the Vega system. Full-length RNA sequencing assumes a total of 60M reads for Revio SPRQ and 30M reads for Vega, regardless of plexity. Amplicon sequencing assumes a 12-hour movie time for 1–5 kb, 24-hour movie time for 5+ kb, and  $\approx 50$  per sample.





### World-class service + support teams

With our dedicated support teams by your side and the **Customer Hub** as your one-stop online solution, managing orders and getting assistance has never been easier.



#### Financing available

PacBio Capital enhances accessibility by eliminating the up-front capital expenditure barrier. This financing program offers highly competitive rates, six-month same-as-cash financing option, and a streamlined credit and funding process to fast-track your purchase.



#### An ecosystem to support you

#### The PacBio Compatible program

integrates products optimized for HiFi sequencing at every step of the workflow for a highly efficient experience. This includes seamless sample prep and library prep workflows and integrated automation products.

#### **Ordering information**

Part number	Product
103-525-500	Vega system
103-517-600	Vega polymerase kit
103-274-300	Vega sequencing plate
103-406-700	Vega SMRT Cell tray
102-090-600	Revio system
103-520-100	Revio SPRQ polymerase kit
103-504-900	Revio SPRQ sequencing plate
102-202-200	Revio SMRT Cell tray

#### Ready to start HiFi sequencing?

#### Talk to a sales rep today.

North America: nasales@pacb.com South America: sasales@pacb.com EMEA: emea@pacb.com

Asia Pacific: apsales@pacb.com



#### Learn more about applications

pacb.com/applications



#### Learn more about the Vega system

pacb.com/vega



#### Learn more about the Revio system

pacb.com/revio

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