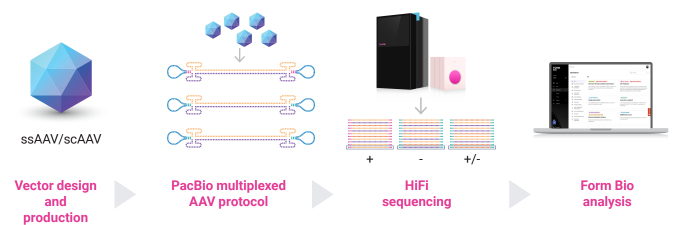
## AAV SEQUENCING — APPLICATION BRIEF BEST PRACTICES

# Accelerate discovery and comprehensively assess adeno-associated virus (AAV) vector quality with highly accurate long reads (HiFi reads)

With HiFi sequencing on the Revio® and Vega™ systems, you can sequence AAV genome populations to identify truncation, mutation, and host integration events. The AAV workflow from PacBio® accommodates both scAAV and ssAAV constructs with easy on-instrument HiFi read generation.

#### From AAV to SMRTbell® libraries with data analysis solutions



Use the PacBio AAV protocol<sup>1</sup> to:

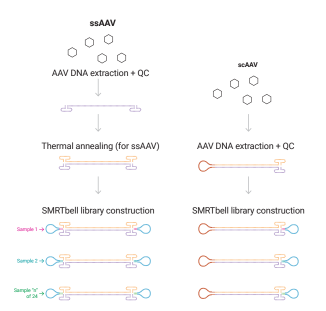
- Convert ssAAV DNA to double-stranded DNA by thermal annealing
- Multiplex up to 24 samples using barcoded SMRTbell adapters

Use the on-instrument AAV mode<sup>2</sup> to:

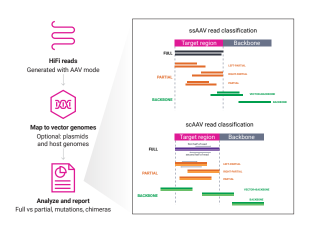
 Generate HiFi reads for proper handling of scAAV and ssAAV structures Analyze your AAV PacBio data with:

- $\bullet$  GitHub command-line analysis,  $^{\!3}$  or
- Form Bio's cloud-based analytical solution<sup>4</sup>

#### HiFi library workflow for AAV



#### **Bioinformatics workflow for AAV**

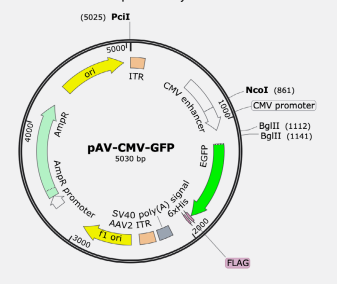


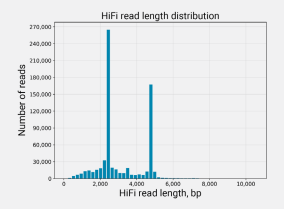
HiFi reads generated with on-instrument AAV mode are aligned to a reference vector genome and optionally other helper plasmids and host genomes. Reads are characterized as ssAAV or scAAV based on the presence or absence of self-complementary alignments, with further characterization as full or partial vectors, backbone or other contaminations, etc. For more information, visit the AAV wiki³ or the AAV bioinformatics webinar.<sup>5</sup>



#### **AAV** sample characterization

**Example 1.** HiFi sequencing identifies unexpected self-complementary structures in ssAAV



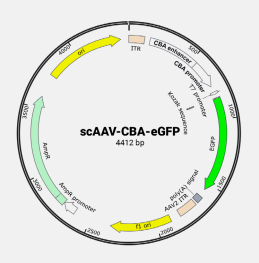


#### ssAAV pAV-CMV-GFP

ssAAV	Full	35.8%
	Partial	10.4%
	Backbone	0.0%
scAAV	Full	29.0%
	Partial	19.4%
	Backbone	0.8%

**Table 1. Characterization of ssAAV using HiFi sequencing.** Expected full-length ssAAV reads ( $\sim$ 2.5 kb) make up only 35.8% of reads. Unexpected self-complementary structures (scAAV) of the full vector ( $\sim$ 5 kb) make up 29% of the reads.

### **Example 2.** HiFi sequencing confirms expected scAAV vector genomes



#### scAAV CBA-eGFP

ssAAV	Full	0.4%
	Partial	0.5%
	Backbone	0.0%
scAAV	Full	95.1%
	Partial	3.6%
	Backbone	0.1%

**Table 2. Characterization of scAAV using HiFi sequencing.** Percentage of reads characterized as full, partial, or backbone, based on alignment to reference vector sequences. Reads mapping with complementary structures will be characterized as scAAV.

#### KEY REFERENCES

- Procedure & checklist Preparing multiplexed AAV SMRTbell libraries using SMRTbell prep kit 3.0
- 2. SMRT Link user guide (v11.0)
- 3. https://github.com/Magdoll/AAV
- 4. https://info.formbio.com/pacbio
- Bioinformatics webinar series on-demand: Optimizing and confirming AAV designs with PacBio sequencing



Learn about AAV gene therapy: **pacb.com/gene-therapy** 

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