

# Pippin™

Automated  
DNA Size  
Selection

NGS Library  
Construction



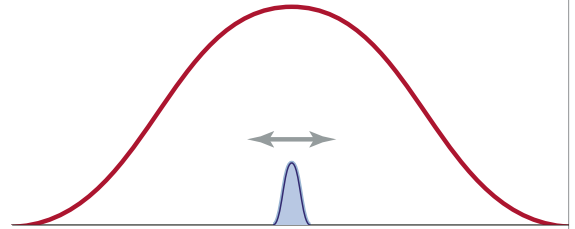
# Flexible and Versatile

## Size Selection modes

### Tight:

Narrowest fragment range based on a target size

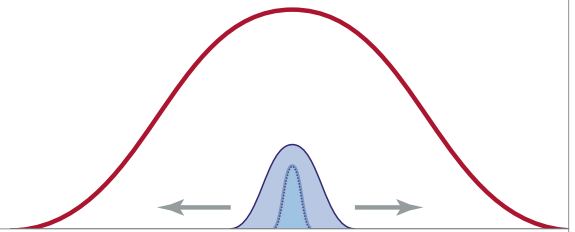
- Paired-end sequencing
- Emulsion prep



### Range:

Set with start and end base pair values

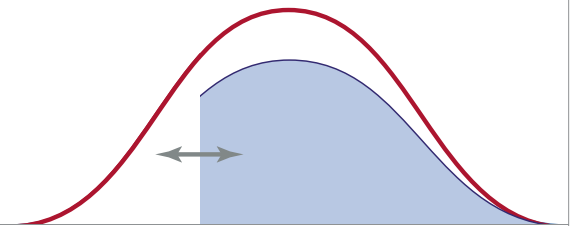
- RNA-seq
- Genotyping by sequencing
- ChIP-seq
- MicroRNA



### High-Pass:

Collects sizes above a base pair threshold

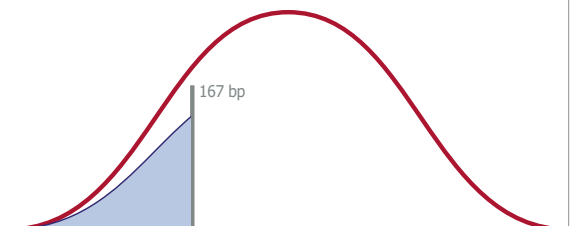
- Long-range genomics
- Long-read sequencing



### Low-Pass:

Collects pre-library nucleic acids below a threshold

- Cell free DNA analysis



Pippin Prep™

Blue Pippin™

Pippin HT

Unlike any other method, automated DNA size selection allows users to select the optimal median fragment size, or adjust the base-pair range of selection. For any NGS platform, and particularly for multi-platform labs, the Pippin system's flexibility streamlines workflows and improves data quality.

# A High Value Step

## For Short-Read Sequencing...

### Key Applications and Benefits

#### Paired-end Sequencing

Narrow size distributions improve assembly

#### RNA-seq

Removes adapter dimers from libraries

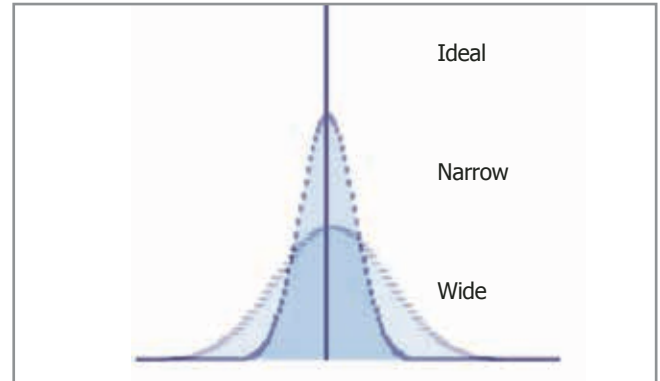
#### ddRAD-seq

Precisely collects fragments generated between restriction sites

#### ChIP-seq

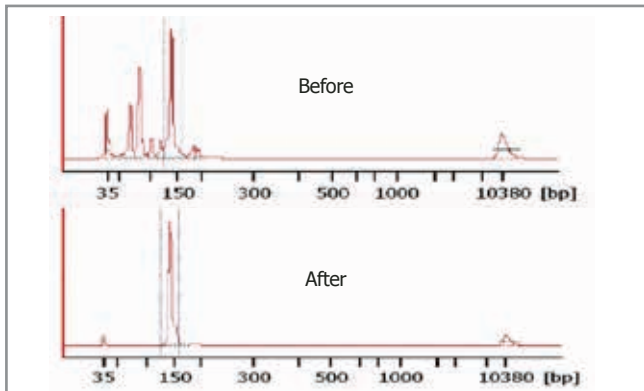
Improves identification of binding sites

### Optimize Size Selection for NGS Platforms



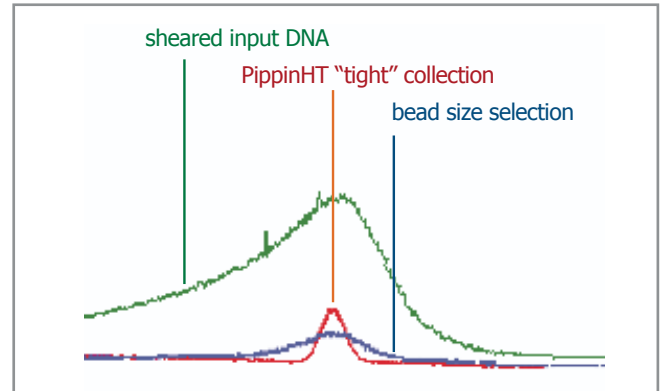
Smaller DNA fragments will preferentially copy when amplified. This can create bias during clustering or emulsion preps. The presence of larger fragments can also lead to sequencing inefficiencies.

### Remove Dimers and Unwanted Artifacts



Using a PippinHT, unwanted artifacts can be removed from microRNA libraries using a range setting. 24 samples can be processed with a 30 minute run time.

### Improve Library Quality



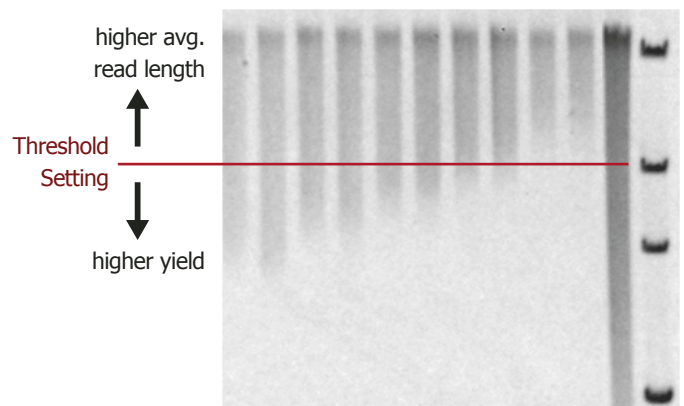
Bead-based size selection produces a significantly wider size distribution than Pippin. The product yield with beads is lower than Pippin, and the fragment range is not easily optimized.

## ...And Long-Range Genomics

Pulsed-field enabled BluePippin and PippinHT systems are standard equipment for working with high molecular weight DNA.

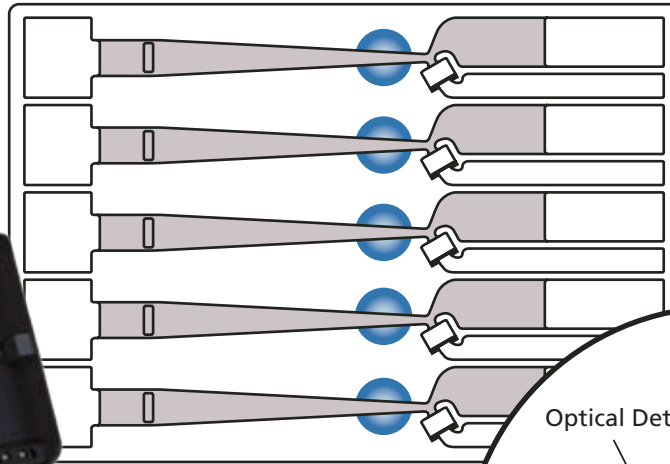
- **Single Molecule Sequencing**
- **Nanopore Sequencing**
- **Droplet Barcode Sequencing**

### High-Pass Size Selections

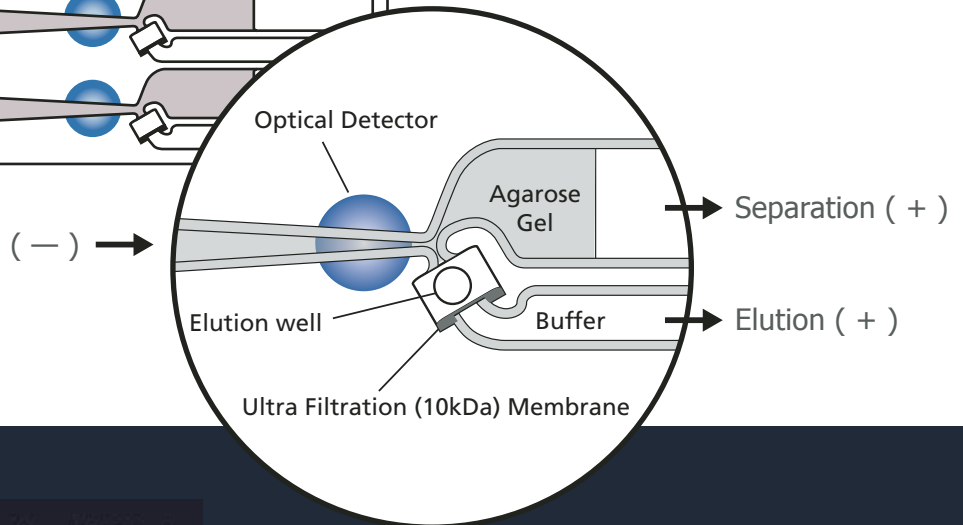


High-pass removal of smaller DNA lets 3rd-gen sequencers focus on the longest fragments.

Pre-cast agarose gel cassette



## The Principle Behind Automated Preparative Gel Electrophoresis



### Set Your Collection Range with Easy-to-Use Software

Tight	Range	BP Target	BP Start	BP End
<input type="checkbox"/>	<input type="checkbox"/>	100	92	108
<input type="checkbox"/>	<input type="checkbox"/>	350	100	600

Selection Chart	Pippin Prep	BluePippin	PippinHT
Max. DNA input	5 µg	5 µg	1.5 µg
Max. Sample Capacity	5	5	24
Pulsed-Field	No	Yes	Yes
Run Times (100 bp - 1.5 kb)	50-90 min	50-90 min*	25-45 min*
Size Selection Range	100 bp - 1.5 kb	100 bp - 50 kb	100 bp - 1.5 kb
High-Pass Size Filtering	No	Yes	Yes
Low-Pass Size Filtering	No	No	Yes

#### Specifications\*

Accuracy	> 93%
Reproducibility	> 92%
Min. Size Distribution as CV	8%
Recovery	50-80%

\* Pulsed-field protocols for HMW size selection may require up to several hours. Inquire at [info@sagescience.com](mailto:info@sagescience.com) for run times and performance specifications.



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