

ACCEL-NGS® METHYL-SEQ DNA LIBRARY KIT

The Accel-NGS Methyl-Seq DNA Library Kit maximizes DNA recovery of bisulfite-converted samples and constructs libraries that accurately represent sample composition. The Accel-NGS Methyl-Seq workflow maximizes DNA recovery through a post-bisulfite library preparation, utilizing a highly efficient adapter attachment that is compatible with single-stranded, bisulfite-converted DNA. Library yields from this kit are up to 100x greater than those from methods that bisulfite convert after library construction. Additionally, the template-independent adapter attachment chemistry of the Accel-NGS Methyl-Seq Kit provides a more complete, less biased library as observed from comprehensive methylome coverage by Whole Genome Bisulfite Sequencing (WGBS).



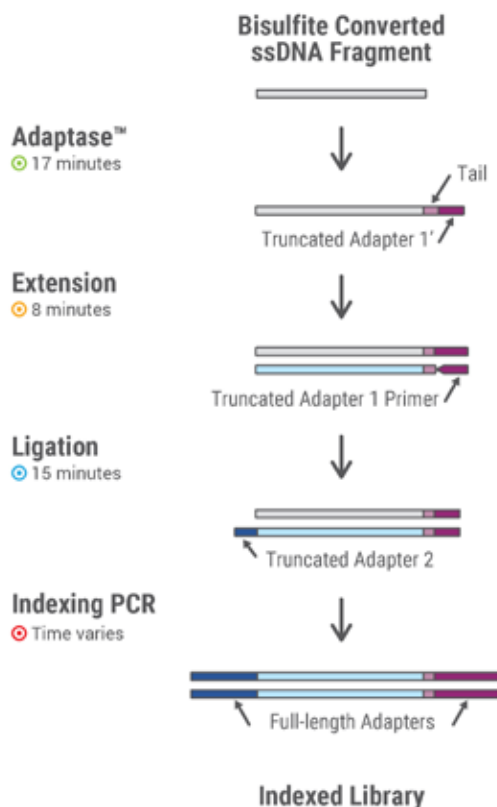
Features

- High recovery of input DNA
- Low bias library preparation
- Simple, 2-hour library prep
- Compatible with Illumina® platforms
- Minimal PCR cycles required
 - 4 cycles for 100 ng
 - 7 cycles for 10 ng
 - 11 cycles for 1 ng
 - 14 cycles for 100 pg

Applications

- WGBS
- Reduced Representation Bisulfite Sequencing (RRBS)
- Detecting genome-wide methylation in 5 ng of cfDNA
- Hybridization capture using NimbleGen™ SeqCap™ Epi Enrichment System
- Bisulfite-converted DNA enriched by MeDIP, ChIP or other methods

Workflow Superior to the Leading Kits



The Accel-NGS Methyl-Seq workflow constructs libraries from single-stranded DNA fragments. The Adaptase step simultaneously performs end repair, tailing of 3' ends, and ligation of the first truncated adapter. The Extension and Ligation steps add the second truncated adapter to the bottom strand only. The Indexing PCR step increases yield and incorporates full length adapters.

Superior Library Complexity and Coverage from 1 ng

To demonstrate the efficient and unbiased performance of the Accel-NGS Methyl-Seq Kit, a titration experiment using 100 ng, 10 ng, or 1 ng of *Arabidopsis* genomic DNA was performed and compared to two alternative methods (random primer and traditional) using these same inputs.

Input	Sample	% Reads Aligned	Genome Coverage	% Duplicate Reads	Estimated Library Size (Millions)	% CpX Missing	% CpX Covered $\geq 10X$
100 ng <i>Arabidopsis</i>	Accel-NGS Methyl-Seq	89.6	22.0X	1.9	714	0.56	92.2
	Traditional	80.2	21.0X	2.7	604	0.57	88.1
	Random Primer	71.4	16.0X	22.1	48	7.70	39.4
10 ng <i>Arabidopsis</i>	Accel-NGS Methyl-Seq	87.8	22.0X	2.7	406	0.58	90.4
	Traditional	76.7	19.0X	11.9	70	0.57	83.9
	Random Primer	71.9	16.0X	22.2	45	5.20	45.2
1 ng <i>Arabidopsis</i>	Accel-NGS Methyl-Seq	83.3	18.0X	18.2	38	0.59	77.1
	Traditional	80.7	10.0X	62.3	6	2.00	17.0
	Random Primer	73.4	12.0X	46.1	12	6.60	31.3
10 ng Human	Accel-NGS Methyl-Seq	86.4	8.9X	7.9	1,393	N/A	N/A

Each *Arabidopsis thaliana* file was normalized to 30.2 million reads and data reported as an average of duplicate bisulfite-converted samples. To assess coverage for the human genome, an Accel-NGS Methyl-Seq library was constructed using HapMap NA12878 DNA and the sequencing data was normalized to 183.5 million reads.

Nimblegen's SeqCap Epi: CpGiant

The Accel-NGS Methyl-Seq Library Kit was compared to Kapa Biosystems' library prep at 80M reads per sample. The coverage metrics used were evaluated at 1 μ g and 100 ng (within input specifications) and at low inputs. An average of two duplicate libraries is shown.

Coverage Metrics

Input	Method	% Aligned	% on Target	% Duplication	Mean Coverage	% Covered $\geq 2X$	% Covered $\geq 20X$	None Covered
100 ng	SWIFT	90	73	6.5	49x	98.6	78.6	0.8
1 μ g	Kapa	90	80	9.4	51x	98.6	81.1	0.8
10 ng	SWIFT	91	77	26.0	35x	98.5	71.0	0.8
10 ng	Kapa	87	78	71.0	1x	24.7	0.2	47.7
1 ng	SWIFT	90	73	62.0	8x	93.6	2.3	1.0

Référence	Désignation	Conditionnement
Kits de préparation de banques Méthyl-Seq		
SW30024	Accel-NGS MethyI-Seq DNA Library Kit for Illumina	24 rxns
SW30096	Accel-NGS MethyI-Seq DNA Library Kit for Illumina	96 rxns
Index pour les kits de préparation de banques Méthyl-Seq		
SW36024	MethyI-Seq Set A Indexing Kit (12 indices, 2 reactions each)	24 rxns
SW38096	MethyI-Seq Dual Indexing Kit for Illumina (96 combinations)	96 rxns
SW390384	Accel-NGS MethyI-Seq Unique Dual Indexing Kit (96 indices, 384 rxns)	384 rxns
SW39096	Accel-NGS MethyI-Seq Unique Dual Indexing Kit (24 indices, 96 rxns)	96 rxns
Produits complémentaires : kits de traitement au bisulfite		
ZD5030	EZ DNA Methylation-Lightning Kit	50 rxns
ZD5030T	EZ DNA Methylation-Lightning Kit	10 Preps
ZD5031	EZ DNA Methylation-Lightning Kit	200 rxns
ZD5032	EZ-96 DNA Methylation-Lightning Kit (Shallow-Well)	2 x 96 rxns
ZD5033	EZ-96 DNA Methylation-Lightning Kit (Deep-Well)	2 x 96 rxns
ZD5046	EZ-96 DNA Methylation-Lightning MagPrep	4 x 96 rxns
ZD5047	EZ-96 DNA Methylation-Lightning MagPrep	8 x 96 rxns

Commande en ligne



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