

INSTRUCTION MANUAL

EZ DNA Methylation-Lightning[™] Kit Catalog Nos. D5030 & D5031

Highlights

- Fastest method for complete bisulfite conversion of DNA for methylation analysis.
- Ready-to-use conversion reagent is added directly to DNA.
- High-yield, converted DNA is ideal for PCR, MSP, array, bisulfite and Next-Gen sequencing.

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For Research Use Only

Product Contents:

EZ DNA Methylation-Lightning™ Kit	D5030 50 rxns.	D5031 200 rxns.	Storage Temperature
Lightning Conversion Reagent*	5 tubes	20 tubes	Room Temp.
M-Binding Buffer	30 ml	125 ml	Room Temp.
M-Wash Buffer**	6 ml	24 ml	Room Temp.
L-Desulphonation Buffer	10 ml	40 ml	Room Temp.
M-Elution Buffer	1 ml	4 ml	Room Temp.
Zymo-Spin™ IC Columns	50 columns	200 columns	Room Temp.
Collection Tubes	50 tubes	200 tubes	Room Temp.
Instruction Manual	1	1	_

Note - Integrity of kit components is guaranteed for one year from date of purchase. Reagents are routinely tested on a lot-to-lot basis to ensure they provide maximal performance and reliability.

* The **Lightning Conversion Reagent** is in a ready-to-use liquid format. The reagent should be stored tightly capped at room temperature with minimum exposure to light.

** Add 24 ml of 100% ethanol to the 6 ml **M-Wash Buffer** concentrate (D5030) or 96 ml of 100% ethanol to the 24 ml **M-Wash Buffer** concentrate (D5031) before use.

EZ DNA Methylation-Lightning[™] Kit technologies are patent pending.

Use of Methylation Specific PCR (MSP) is protected by US Patents 5,786,146 & 6,017,704 & 6,200,756 & 6,265,171 and International Patent WO 97/46705. No license under these patents to use the MSP process is conveyed expressly or by implication to the purchaser by the purchase of this product.

Note - [™] Trademarks of Zymo Research Corporation. This product is for research use only and should only be used by trained professionals. Some reagents included with this kit are irritants. Wear protective gloves and eye protection. Follow the safety guidelines and rules enacted by your research institution or facility. Freedom EVO[®] is a registered trademark of Tecan Group Ltd. Pyrosequencing[®] is a registered trademark of Biotage.

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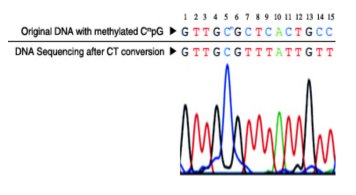
Introduction to DNA Methylation:

Cytosine methylation is a naturally occurring base modification, in both prokaryotic and eukaryotic organisms, consisting of the addition of a methyl group to the fifth carbon position of the cytosine pyrimidine ring via a methyltransferase enzyme (1). In prokaryotes DNA methylation provides a way to protect host DNA from digestion by restriction endonucleases that are designed to eliminate foreign DNA. DNA methylation in higher eukaryotes functions in the regulation/control of gene expression (2).

The majority of DNA methylation in mammals occurs in 5'-CpG-3' dinucleotides, although other patterns do exist. About 80 percent of all 5'-CpG-3' dinucleotides in mammalian genomes are found to be methylated, and the majority of the twenty percent that remain unmethylated are within promoters or in the first exons of genes. It has been demonstrated that aberrant DNA methylation is a widespread phenomenon in cancer and may be among the earliest changes to occur during oncogenesis (3). DNA methylation has also been shown to play a central role in gene imprinting, embryonic development, X-chromosome gene silencing, and cell cycle regulation.

The ability to detect and quantify DNA methylation efficiently and accurately has become essential for the study of cancer, gene expression, genetic diseases, and many other important aspects of biology. To date, a number of methods have been developed to detect/quantify DNA methylation including: high-performance capillary electrophoresis (4) and methylation-sensitive arbitrarily primed PCR (5). However, the most common techniques used today still rely on bisulfite conversion (6).

Treating DNA with bisulfite chemically modifies non-methylated cytosines into uracil, methylated cytosines remain unchanged. Once converted, the methylation profile of the DNA can be determined using the desired downstream application. For single locus analysis, the region of interest is generally amplified following bisulfite conversion (i.e., bisulfite PCR) and then sequenced or processed for Pyrosequencing[®]. Recent advances in methylation detection also allow the investigation of genome-wide methylation patterns using technologies including array-based methods, reduced representation bisulfite sequencing (RRBS), and whole genome bisulfite sequencing (7).



DNA sequencing results following bisulfite treatment. DNA with methylated C at nucleotide position #5 was processed using the **EZ DNA Methylation™ Kit**. The recovered DNA was amplified by PCR and then sequenced directly. The methylated cytosine at position #5 remains intact while the unmethylated cytosines at positions #7, 9, 11, 14 and 15 are completely converted into uracil following bisulfite treatment (detected as thymine following PCR).

References:

1. Adams RL. Bioessays. 1995; 17(2): 139-145.

2. Costello JF, Plass CJ. Med. Genet. 2001; 38(5): 285-303.

3. Stirzaker C. Cancer Res. 1997; 57(11): 2229-2237.

4. Fraga MF, *et al.* Electrophoresis. 2000; 21(14): 2990-2994.

5. Gonzalgo ML. Cancer Res. 1997; 57(4): 594-599.

6. Frommer M. Proc. Natl. Acad. Sci. USA. 1992; 89(5): 1827-1831.

7. Rakyan VK, *et al.* Nat. Rev. 2011, 12(8): 529-541. Note: 96-Well spin-plate formats are available for processing larger numbers of samples. Also, MapPrep kits are available (p. 8) for adaptation to liquid handling robots (e.g., Tecan – Freedom EVO[®]) and automated sample prep.

Select Citations:

1. Ehrich M, *et al.* Nuc. Acids Res. 2007; 35 (5): e29

2. Kaneda M, *et al.* Nature. 2004; 429: 900-903

3. Zhang F, *et al.* Proc. Natl. Acad. Sci. USA. 2007; 104 (11): 4395-4400.

4. Oda M, *et al.* Genes & Dev. 2006; 20: 3382-3394.

5. England RPM, *et al.* Nature Meth. 2005; 2: 1-2.

6. Berman BP, et al. Nature Gen. 2012; 44: 40-46.

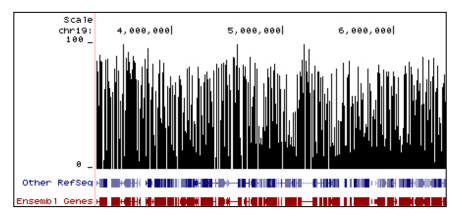
Leung DC, et al. Proc.
 Natl. Acad. Sci. USA. 2011;
 108 (14): 5718-5723.

8. Hesselink AT, et al. Clin. Cancer Res. 2011; 17: 2459-2465.

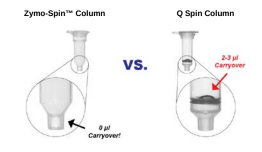
9. Campan M, et al. PLoS ONE. 2011, 6 (12): e28141.

Product Description:

The **EZ DNA Methylation-Lightning™ Kit** features rapid and reliable bisulfite treatment and conversion of DNA for methylation analysis. Key to the fast workflow is the ready-to-use **Lightning Conversion Reagent**. No preparation is necessary, simply add this unique reagent to a DNA sample, wait about an hour, and let the reaction proceed to completion. DNA denaturation and bisulfite conversion processes are combined with added heat to facilitate rapid denaturation. Desulphonation and clean-up of the converted DNA is performed using a unique low-elution spin column. High yield, converted DNA is ideal for PCR, array, bisulfite and next generation sequencing, etc.



Methylation Plot From Reduced Representation Bisulfite Sequencing (RRBS). Data shows the relative percentage of methylation at individual CpG sites in mouse DNA. Methylation percentage is shown across a ~3 Mb region of mouse chromosome 19. Bisulfite sequencing libraries were prepared using mouse genomic DNA prepped with the Genomic Clean & Concentrator™ (D4010, D4011 – Zymo Research) and bisulfite converted using EZ DNA Methylation™ technology prior to Next-Gen sequencing.



Zymo-Spin[™] Columns Ensure No Buffer Retention

Zymo-Spin[™] IC Design Characteristics. The image above shows the unique design of the Zymo column that facilitates extremely small elution volumes (≥10 µl) without buffer carryover. This is unlike other columns that can retain liquid (binding/wash buffer residue) leading to carryover into the eluate.

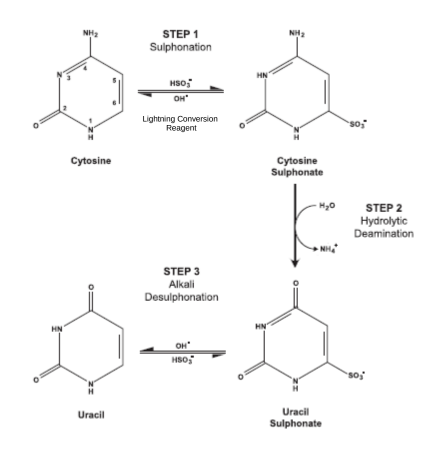
Specifications:

- **DNA Input:** Samples containing between 100 pg to 2 µg of DNA. For optimal results, the amount of input DNA should be from 200 to 500 ng.
- **Conversion Efficiency:** > 99.5% of non-methylated C residues are converted to U; > 99.5% protection of methylated cytosines.
- DNA Recovery: >80%

Reagent Preparation:

• Preparation of M-Wash Buffer

Add 24 ml of 100% ethanol to the 6 ml **M-Wash Buffer** concentrate (D5030) or 96 ml of 100% ethanol to the 24 ml **M-Wash Buffer** concentrate (D5031) before use.



Overview of Bisulfite Conversion. Steps 1 and 2 occur during bisulfite conversion, while Step 3 is performed as the DNA is bound to the column matrix. For the reaction to proceed to completion, it is essential the DNA be fully denatured.

Samples >20 µl must be processed using multiple conversion reactions. Replicate reactions can be cleaned using the same column for each by repeating steps 3-5.

The capacity of the collection tube with the column inserted is $800 \ \mu$ l. Empty the collection tube whenever necessary to prevent contamination of the column contents by the flow-through.

Alternatively, water or TE $(pH \ge 6.0)$ can be used for elution if required for your experiments.

Protocol:

 Add 130 μl of Lightning Conversion Reagent to 20 μl of a DNA sample in a PCR tube. Mix, then centrifuge briefly to ensure there are <u>no</u> droplets in the cap or sides of the tube.

Note: If the volume of DNA is less than 20 $\mu I,$ compensate with water.

- 2. Place the PCR tube in a thermal cycler and perform the following steps:
 - 1. 98°C for 8 minutes
 - 2. 54°C for 60 minutes
 - 3. 4°C storage for up to 20 hours

Note: The 4°C storage step is optional.

- 3. Add 600 µl of **M-Binding Buffer** to a **Zymo-Spin™ IC Column** and place the column into a provided **Collection Tube**.
- 4. Load the sample (from Step 2) into the **Zymo-Spin™ IC Column** containing the **M**-**Binding Buffer**. Close the cap and mix by inverting the column several times.
- 5. Centrifuge at full speed (\geq 10,000 x *g*) for 30 seconds. Discard the flow-through.
- 6. Add 100 µl of **M-Wash Buffer** to the column. Centrifuge at full speed for 30 seconds.
- Add 200 µl of L-Desulphonation Buffer to the column and let stand at room temperature (20-30°C) for 15-20 minutes. After the incubation, centrifuge at full speed for 30 seconds.
- 8. Add 200 µl of **M-Wash Buffer** to the column. Centrifuge at full speed for 30 seconds. Repeat this wash step.
- Place the column into a 1.5 ml microcentrifuge tube and add 10 μl of M-Elution Buffer directly to the column matrix. Centrifuge for 30 seconds at full speed to elute the DNA.

The DNA is ready for immediate analysis or can be stored at or below -20°C for later use. For long term storage, store at or below -70°C. We recommend using 1-4 μ l of eluted DNA for each PCR, however, up to 10 μ l can be used if necessary. The elution volume can be > 10 μ l depending on the requirements of your experiments, but small elution volumes will yield higher DNA concentrations.

details)

Appendix: Bisulfite Conversion and PCR Optimization

1.		of Double Stranded DNA Templates. The a DNA template during bisulfite conversion.	he following	
	Template:	A: 5'-GACCGTTCCAGGTCCAGCAGTGCGCT-3' B: 3'-CTGGCAAGGTCCAGGTCGTCACGCGA-5'		Note: Methylated "C" is underlined in the examples.
	Bisulfite Converted:	<pre>A: 5'-GATCGTTTTAGGTTTAGTAGTGCGTT-3' B: 3'-TTGGCAAGGTTTAGGTTGTTATGCGA-5'</pre>		Note: Following bisulfite conversion, the strands are
2.	•		reated as Ts	no longer complementary.
	Bisulfite Converted: Primers: Reverse: Forward:	A: 5'-GATCGTTTTAGGTTTAGTAGTGCGTT-3' 3'-ATCATCACRCAA-5' 5'-GATYGTTTTAGGT-3'	R= G/A Y= C/T	Note: Only one strand (A) is amplified by a given primer set. Only the reverse primer binds to the converted DNA, the forward primer will bind
		es primer design assistance with its <u>Bisulfite Pri</u> ww.zymoresearch.com/tools/bisulfite-primer-seel		the strand generated by the reverse primer. If the primer contains CpG dinucleotides with uncertain
3.	human or mouse genom amplification is 100 pg. 500 ng. Although, up to	uired for Bisulfite Conversion. The minima nic DNA required for bisulfite treatment and subs The optimal amount of DNA per bisulfite treatme 2 μg of DNA can be processed, it should be not y result in incomplete bisulfite conversion for so	sequent PCR ent is 200 to ted that high	methylation status, then mixed bases with C and T (or G and A) can be used. Usually, there should be no more than one mixed position per primer and it should be located toward the 5' end of the primer. It is not recommended to have mixed bases located at the
4.	amplification of bisulfite 150-300 bp; however la	cually, 35 to 40 cycles are required for succ converted DNA. Optimal amplicon size should rger amplicons (up to 1 kb) can be generated b nealing temperatures between 55-60°C typically	be between by optimizing	3' end of the primer.
	treated DNA usually is amplification is relatively	d cytosine residues are converted into uracil, t AT-rich and has low GC composition. Non-s common with bisulfite treated DNA due to its AT plymerases is strongly recommended for the am	specific PCR	Zymo Taq [™] is a "hot start" DNA polymerase <u>specifically</u> <u>designed</u> for the amplification of bisulfite treated DNA. (see page 9 for

5. Quantifying Bisulfite Treated DNA. Following bisulfite treatment of genomic DNA, the original base-pairing no longer exists since non-methylated cytosine residues are converted into uracil. Recovered DNA is typically A, U, and T-rich and is single stranded with limited non-specific base-pairing at room temperature. The absorption coefficient at 260 nm resembles that of RNA. Use a value of 40 μ g/ml for Ab₂₆₀ = 1.0 when determining the concentration of the recovered bisulfite-treated DNA.

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Frequently Asked Questions:

- Q: Should the input DNA be dissolved in TE, water, or some other buffer prior to its conversion?
- **A:** Water, TE or modified TE buffers can be used to dissolve the DNA and do not interfere with the conversion process.
- Q: Which *Taq* polymerase(s) do you recommend for PCR amplification of converted DNA?
- **A:** We recommend a "hot start" DNA polymerase (e.g., ZymoTaq[™], page 9).

Q: Why are there two different catalog numbers for the EZ-96 DNA Methylation-Lightning[™] Kit?

A: The two different catalog numbers are used to differentiate between the binding plates that are included in the kit. Deep and shallow-well binding plates are available to accommodate most rotors and microplate carriers. Below is a comparison of the two binding plates.



Binding Plate	Silicon-A™ Plate	Zymo-Spin™ I-96 Plate
Style	Shallow-Well	Deep-Well
Height of Binding Plate	19 mm (0.75 inches)	35 mm (1.38 inches)
Binding Plate/Collection Plate Assembly	43 mm (1.69 inches)	60 mm (2.36 inches)
Binding Cap./Minimum Elution Volume	5 µg/30 µl	5 µg/15 µl
Catalog Numbers	D5032	D5033

Ordering Information:

Product Description	Catalog No.	Kit Size
EZ DNA Methylation-Lightning™ Kit	D5030 D5031	50 rxns. 200 rxns.
EZ-96 DNA Methylation-Lightning™ Kit (Shallow-Well)	D5032	2 x 96 rxns.
EZ-96 DNA Methylation-Lightning™ Kit (Deep-Well)	D5033	2 x 96 rxns.
EZ-96 DNA Methylation-Lightning™ MagPrep*	D5046 D5047	4 x 96 rxns. 8 x 96 rxns.

For Individual Sale	Catalog No.	Amount(s)
Lightning Conversion Reagent	D5030-1 D5032-1	1 tube 1 bottle
M-Binding Buffer	D5005-3 D5006-3 D5040-3	30 ml 125 ml 250 ml
M-Wash Buffer	D5001-4 D5002-4 D5007-4 D5040-4	6 ml 24 ml 36 ml 72 ml
L-Desulphonation Buffer	D5030-5 D5031-5 D5046-5	10 ml 40 ml 80 ml
M-Elution Buffer	D5001-6 D5002-6 D5007-6 D5041-6	1 ml 4 ml 8 ml 40 ml
Zymo-Spin™ IC Columns (capped)	C1004-50 C1004-250	50 columns 250 columns
Collection Tubes	C1001-50 C1001-500 C1001-1000	50 tubes 500 tubes 1,000 tubes
MagBinding Beads	D4100-2-6 D4100-2-8 D4100-2-12 D4100-2-16 D4100-2-24	6ml 8 ml 12 ml 16 ml 24 ml
Zymo-Spin™ I-96 Binding Plates	C2004	2 plates
Silicon-A™ Binding Plates	C2001	2 plates
Conversion Plates w/ Pierceable Cover Film	C2005	2 plates/films
Collection Plates	C2002	2 plates
Elution Plates	C2003	2 plates

* **MagPrep** kits are adaptable to liquid handling robots (e.g., Tecan – Freedom EVO[®]) making them ideal for automated sample prep.

Epigenetics Products From Zymo Research

THE Epigenetics COMPANY™

Product	Description	Kit Size	Cat No. (Format)
	Bisulfite Kits for DNA Methylation Detection	on	
EZ DNA Methylation™ Kit	For the conversion of unmethylated cytosines in DNA to uracil via the <u>chemical-denaturation</u> of DNA and a specially designed CT Conversion Reagent. <i>Fast-Spin</i> technology ensures ultra-pure, converted DNA for subsequent DNA methylation analysis. Magnetic bead format for adaptation to automated liquid handling platforms.	50 Rxns. 200 Rxns. 2x96 Rxns. 2x96 Rxns. 4x96 Rxns. 8x96 Rxns.	D5001 (spin column) D5002 (spin column) D5003 (shallow-well plate) D5004 (deep-well plate) D5040 (magnetic bead) D5041 (magnetic bead)
EZ DNA Methylation- Gold™ Kit	For the fast (3 hr.) conversion of unmethylated cytosines in DNA to uracil via <u>heat/chemical-denaturation</u> of DNA and a specially designed CT Conversion Reagent. <i>Fast-Spin</i> technology ensures ultra-pure, converted DNA for subsequent DNA methylation analysis. Magnetic bead format for adaptation to automated liquid handling platforms.	50 Rxns. 200 Rxns. 2x96 Rxns. 2x96 Rxns. 4x96 Rxns. 8x96 Rxns.	D5005 (spin column)D5006 (spin column)D5007 (shallow-well plate)D5008 (deep-well plate)D5042 (magnetic bead)D5043 (magnetic bead)
EZ DNA Methylation- Direct™ Kit	Features simple and reliable DNA bisulfite conversion directly from blood, tissue (FFPE/LCM), and cells without the prerequisite for DNA purification in as little as 4-6 hrs. The increased sensitivity of this kit makes it possible to amplify bisulfite converted DNA from as few as 10 cells or 50 pg DNA. Magnetic bead format for adaptation to automated liquid handling platforms.	50 Rxns. 200 Rxns. 2x96 Rxns. 2x96 Rxns. 4x96 Rxns. 8x96 Rxns.	D5020 (spin column) D5021 (spin column) D5022 (shallow-well plate) D5023 (deep-well plate) D5044 (magnetic bead) D5045 (magnetic bead)
EZ DNA Methylation- Lightning™ Kit	Complete bisulfite conversion in about an hour using a unique liquid format conversion reagent that requires no preparation. <i>Fast-Spin</i> technology ensures ultra-pure, converted DNA for subsequent DNA methylation analysis. Magnetic bead format for adaptation to automated liquid handling platforms.	50 Rxns. 200 Rxns. 2x96 Rxns. 2x96 Rxns. 4x96 Rxns. 8x96 Rxns.	D5030 (spin column) D5031 (spin column) D5032 (shallow-well plate) D5033 (deep-well plate) D5046 (magnetic bead) D5047 (magnetic bead)
EZ DNA Methylation- Startup™ Kit	Designed for the first time user requiring a consolidated product to perform DNA methylation analysis. Includes technologies for sample processing, bisulfite treatment of DNA, and PCR amplification of "converted" DNA for methylation analysis.	1 Kit	D5024
	Methylated DNA Standards		
Universal Methylated Human DNA Standard	Human (male) genomic DNA having all CpG sites methylated. To be used for the evaluation of bisulfite-mediated conversion of DNA. Supplied with a control primer set.	1 set	D5011
Universal Methylated Mouse DNA Standard	Mouse (male) DNA having all CpG sites methylated. To be used for the evaluation of bisulfite-mediated conversion of DNA. Supplied with a control primer set.	1 set	D5012
	Other		
ChIP DNA Clean & Concentrator™	Clean and concentrate DNA from any reaction or "crude" preparation in 2 min. A 6 μ l minimum elution volume allows for highly concentrated DNA. Designed for samples containing up to 5 μ g of DNA.	50 Preps. 50 Preps.	D5201 (uncapped column) D5205 (capped column)
Genomic DNA Clean & Concentrator™	Genomic DNA clean-up in minutes. Unique spin column technology for recovery of ultra-pure large-sized DNA (100 bp to ≥200 kb) DNA from any impure preparation (e.g., Proteinase K digestion).	25 Preps. 100 Preps.	D4010 D4011
Zymo <i>Taq</i> ™ DNA Polymerase	Zymo <i>Taq</i> [™] "hot start" DNA Polymerase is specifically designed for the amplification of "difficult" DNA templates including: bisulfite-treated DNA for methylation detection. The product generates specific amplicons with little or no by-product formation. Available either as a single buffer premix or as a polymerase system with components provided separately.	50 Rxns. 200 Rxns. 50 Rxns. 200 Rxns.	E2001 (system) E2002 (system) E2003 (premix) E2004 (premix)
Methylated-DNA IP Kit	IP with a highly specific anti-5-methylcytosine monoclonal antibody. Designed for the enrichment of 5-methylcytosine-containing DNA from any pool of fragmented genomic DNA for use in genome-wide methylation analysis.	10 Rxns.	D5101
	Services		
Available	for DNA Methylation and Hydroxymethylation at http://www.zymoresearch.com powered by the latest Next-Gen Sequence		