MethylCollector[™] Ultra

(version D3)

Catalog No. 55005

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Overview

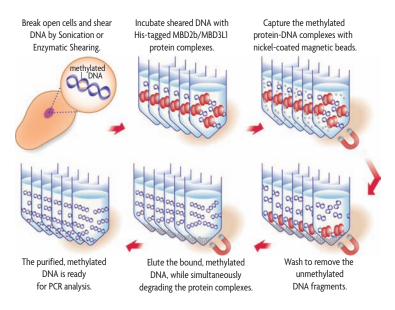
Active Motif's MethylCollector[™] Ultra Kit is an improved alternative for the enrichment of CpGmethylated DNA¹ from limited amounts of cell or tissue samples. The method is based on the Methylated CpG Island Recovery Assay (MIRA), which utilizes the high affinity of the MBD2b/MB-D3L1 complex for methylated DNA². MethylCollector Ultra is suitable for use in many downstream applications, such as real time or endpoint PCR analysis of the methylation status of particular loci in normal and diseased samples, rapid screening of the methylation status of multiple loci, bisulfite conversion followed by cloning and sequencing, or amplification and labeling for microarray analysis³. It can also be used to detect changes in DNA methylation during normal cellular differentiation and aging.

In the MethylCollector Ultra method*, His-tagged recombinant MBD2b and its binding partner MBD3L1 are combined together in order to increase the affinity of MBD2b for CpG-methylated DNA. The kit works using DNA fragments that have been prepared by enzymatic digestion or sonication from cell or tissue samples. The MBD2b/MBD3L1 protein complex is added to the DNA fragments, and it specifically binds to CpG-methylated DNA. These protein-DNA complexes are then captured with nickel-coated magnetic beads and subsequent wash steps are performed to remove fragments with little or no methylation. The methylated DNA is then eluted from the beads in the presence of Proteinase K. Due to the high efficiency of MethylCollector Ultra and the enormous amplification capability and specificity of PCR, analysis of the methylation status of a specific genomic DNA locus can be performed on DNA isolated from less than 170 cells (-1 ng DNA).

product	format	catalog no.
MethylCollector [™] Ultra	30 rxns**	55005

*Technology covered under U.S. Patent No. 7,425,415.

**MethylCollector[™] Ultra provides sufficient reagents to perform 30 reactions with excess reagents for 5 control reactions.



Flow chart of the MethylCollector Ultra process.

In MethylCollector Ultra, genomic DNA of interest is sheared by either enzymatic digestion or sonication. The sheared DNA is then incubated with a His-tagged recombinant MBD2b/MBD3L1 protein complex. The interaction of MBD2b with its binding partner MBD3L1 increases the affinity of MBD2b for CpG-methylated DNA. These protein-DNA complexes are captured with nickel-coated magnetic beads and stringent washes are then performed to remove fragments with little or no methylation. The methylated DNA is then eluted from the beads and real time or endpoint PCR is performed on the resulting supernatant using specific primers to amplify the locus of interest.

Introduction

Over the last decade, the study of DNA methylation and its role in epigenetic cell signaling has grown rapidly⁴⁷. Methylation of CpG dinucleotides, which occurs at the fifth position of the cytosine pyrimidine ring, is of particular interest.

Although CpG dinucleotides are generally methylated throughout the genome of normal somatic cells, CpG islands (clusters of CpG dinucleotides in gene regulatory regions) are usually unmethylated⁸. Aberrant hypermethylation of CpG islands and subsequent transcriptional repression is one of the earliest and most common somatic genome alterations in multiple human cancers^{9,10}. Somewhat paradoxically, a decrease in the total amount of cytosine methylation is observed in many neoplastic tissues, but the genome context of this hypomethylation has not been identified¹¹. Aberrant methylation of CpG islands thus seems to be a tumor type-specific event^{10, 12} and current efforts have concentrated on finding ways to exploit the diagnostic and therapeutic implications of these abnormalities^{13,14}.

Methyl-CpG binding proteins appear to be central players in the process of DNA methylationdependent gene silencing¹⁵. This family of proteins takes its definition from the methyl-CpG binding domain (MBD), the minimum portion with specific affinity for a single, symmetrically methylated CpG pair. The MBD was characterized by deletion studies of MeCP2¹⁶. After the recognition of the MBD, four additional genes were found to contain this domain, namely MBD1, MBD2, MBD3 and MBD4¹⁷. In general, all MBD proteins, except MBD4, have been reported to be associated with histone deacetylase subunits as part of large multi-subunit complexes^{18, 19}. A few studies support the notion of selectivity in the association of a particular MBD with particular promoters^{20, 21}, but other results indicate that the CpG distribution along the sequence may influence the interaction of each MBD protein with DNA²².

The MBD2b protein has been found to possess one of the highest affinities for methylated DNA among MBD proteins and has the greatest capacity to differentiate between methylated and unmethylated DNA²². The combination of MBD2b with its binding partner MBD3L1, methyl-CpG-binding protein 3-like-1, generates a higher affinity for methylated DNA than MBD2b protein alone^{2, 23}. The specificity of the MBD2b/MBD3L1 complex is able to enrich for methylated DNA fragments containing as few as 5 methylated CpGs, and is more efficient at methylated DNA enrichment than antibody-based immunoprecipitation methods. In addition, MethylCollector Ultra can be used in parallel with the UnMethylCollector[™] Kit (Catalog No. 55004) to provide a more thorough analysis of the methylation state at each locus of interest²⁴.

Traditional Methods to Study DNA Methylation

To date, there are several methods used for methylation analysis:

- Methylation-sensitive restriction enzyme analysis: Isoschizomers of bacterial restriction endonucleases with different sensitivities for 5-methylcytosine can be used to determine the methylation status of specific CpG dinucleotides²⁵. Methylation-sensitive restriction enzymes have several limitations, such as the fact that the methylation-sensitive restriction merely informs on the methylation status of the cytosine residues which are recognized by the restriction enzymes used, but do not provide information about other methylation sites.
- 2. Bisulfite conversion: Bisulfite conversion²⁶ consists of the treatment of double-stranded genomic DNA with sodium bisulfite, leading to deamination of unmethylated cytosines into uracil. PCR is then performed with primers that differentiate between methylated and unmethylated sequences. Bisulfite-based techniques can be cumbersome, involving time-and labor-intensive chemical treatments that damage DNA and limit throughput.
- 3. Methylated DNA Immunoprecipitation (MeDIP): In this assay, an antibody specific for methylated cytosines (anti-5-methylcytosine antibody) is used to immunoprecipitate methylated DNA from genomic DNA fragmented by enzymatic digestion or sonication²⁷. The resulting enrichment is usually analyzed by PCR based methods; thus MeDIP can be combined with DNA microarrays for genome-wide analysis of CpG methylation. However, this technique is relatively time-consuming, requires a large amount of fragmented DNA starting material and only works with denatured DNA.

The MethylCollector Ultra Kit is for research use only. Not for use in diagnostic procedures.

Range of detection: MethylCollector Ultra can be performed on 1 ng - 1 µg of fragmented genomic DNA.

Sensitivity: Enriches methylated DNA fragments with as few as 5 methylated CpGs.

Nature of the MethylCollector Ultra Assay: MethylCollector Ultra is an improved technique for the enrichment of methylated CpG islands. The methylation status of specific promoters contained within CpG islands can be analyzed using either endpoint or real time PCR analysis of the locus of interest with customer designed PCR primers. Control human, male genomic DNA that was digested with *Mse* I is included in the kit along with PCR primers specific for both unmethylated and methylated promoters.

- GAPDHGlyceraldehyde-3-phosphate dehydrogenase is important for metabolism. Because
this gene is often constitutively expressed, it is considered to be an actively tran-
scribed housekeeping gene containing an unmethylated promoter in healthy tissues.
The region amplified by this primer pair is 69 base pairs and contains 7 CpGs.
- *Xist* X inactive specific transcript is a methylated promoter in human male genomic DNA. The region amplified by this primer pair is 178 base pairs and contains 8 CpGs.
- NBR2 Neighbor of BRCA1 gene 2 is located near the breast cancer gene BRCA1. Evidence indicates that NBR2 and BRCA1 share a bi-directional promoter. This region of the NBR2 gene is methylated in healthy tissues. The region amplified by this primer pair is 103 base pairs and contains 7 CpGs.

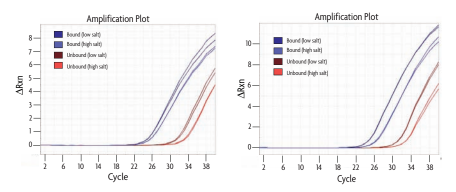
With high salt binding conditions, the *Mse* I digested control human, male genomic DNA provided in the kit should have at least a 10-fold enrichment of methylated DNA bound and eluted from the protein complex as detected with the NBR2 PCR Primer Mix. This means that of the total DNA recovered, there is ten times as much methylated DNA recovered in the eluted fraction as compared to the unbound fraction for the same locus.

With low salt binding conditions, the *Mse* I digested control human, male genomic DNA provided in the kit should have at least a 10-fold enrichment of methylated DNA bound and eluted from the protein complex as detected with the Xist PCR Primer Mix. This means that of the total DNA recovered, there is ten times as much methylated DNA recovered in the eluted fraction as compared to the unbound fraction for the same locus.

There should be less than 5% of methylated DNA detected in the eluted fraction using the GAPDH PCR Primer Mix for either low or high salt binding conditions.

A. NBR2 Amplification

B. Xist Amplification



C. GAPDH Amplification

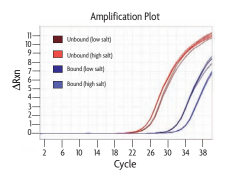


Figure 1: Real time PCR analysis of control PCR primer sets.

MethylCollector Ultra was performed using 100 ng of *Mse* I digested human, male genomic DNA under both low and high salt conditions. Eluted DNA was purified and analyzed using real time PCR for both methylated and unmethylated promoters. **A)** Amplification plot using the provided NBR2 PCR primer mix with the unbound and eluted fractions. NBR2 is methylated in the control DNA and shows early amplification of the eluted fractions. **B)** Amplification plot using the provided Xist PCR primer mix with the unbound and shows early amplification in the eluted fractions. **C)** Amplification plot using the provided GAPDH PCR primer mix with the unbound and eluted fractions. **C)** Amplification plot using the provided GAPDH PCR primer mix with the unbound and eluted fractions. GAPDH is unmethylated in the control DNA and shows late amplification of the eluted fractions.

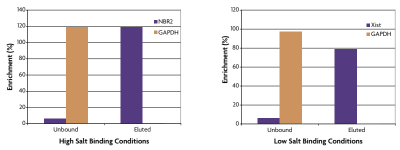
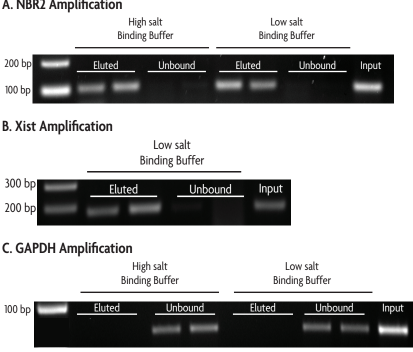


Figure 2: Fold enrichment of MethylCollector Ultra binding reactions.

MethylCollector Ultra was performed with 100 ng of Mse I digested human, male genomic DNA using both high and low salt binding conditions. Unbound and eluted DNA was cleaned and analyzed in real time PCR. The amount of DNA recovered in the eluted fraction was divided by the amount of input DNA used in the binding reaction to produce a percent enrichment. The methylated promoters NBR2 and Xist had greater than 10-fold enrichment of eluted DNA as compared to unbound DNA for the same locus, while the unmethylated GAPDH promoter was found almost exclusively in the unbound fraction.



A. NBR2 Amplification

Figure 3: Specificity of the MBD2b/MBD3L1 protein complex for CpG methylated DNA.

MethylCollector Ultra was performed with 100 ng of Mse I digested human, male genomic DNA using both high and low salt conditions. Eluted DNA was cleaned and analyzed in endpoint PCR for 36 cycles. The methylated NBR2 (A) and Xist (B) promoters are highly enriched in the eluted fraction while the unmethylated GAPDH (C) promoter is only detected in the unbound fraction, indicating the high specificity of MBD2b/MBD3L1. The Xist PCR Primer Mix has been optimized for use with low salt binding conditions.

Kit Components and Storage

Kit components arrive on dry ice. Upon receipt, we recommend storing each component at the temperatures listed in the table below. The magnetic beads can be stored frozen upon arrival, however, we recommend long-term storage at 4°C. **Once the magnetic beads have been thawed, do not re-freeze the beads.**

Reagents	Quantity	Storage / Stability
His-MBD2b/MBD3L1 protein complex	350 µl	-20°C for 6 months
High Salt Binding Buffer	35 ml	-20°C for 6 months
Low Salt Binding Buffer	35 ml	-20°C for 6 months
Elution Buffer AM1	3.5 ml	-20°C for 6 months
Protease Inhibitor Cocktail	100 µl	-20°C for 6 months
Proteinase K	70 µl	-20°C for 6 months
Proteinase K Stop Solution	70 µl	-20°C for 6 months
Human, male genomic DNA (<i>Mse</i> I digested) (20 ng/µl)	250 µl	-20°C for 6 months
GAPDH PCR Primer Mix (2.5 pmol/µl)	400 µl	-20°C for 6 months
Xist PCR Primer Mix (2.5 pmol/µl)	400 µl	-20°C for 6 months
NBR2 PCR Primer Mix (2.5 pmol/µl)	400 µl	-20°C for 6 months
10X PCR Buffer	1.5 ml	-20°C for 6 months
10X PCR Loading Dye	1.5 ml	-20°C for 6 months
Magnetic Nickel Beads	350 µl	4°C for 6 months
Glycogen (20 mg/ml)	35 µl	-20°C for 6 months
Bar Magnet	1	Room temperature
Mini Glue Dots	2 Dots	Room temperature
8-strip PCR tubes and caps	12 strips	Room temperature

Additional Materials Required

- Fragmented DNA sample
- Sample PCR primer sets
- Sterile DNase-free water
- Filter pipette tips
- Microcentrifuge tubes and microcentrifuge
- Magnetic stand. You can assemble a magnetic stand using the provided bar magnet and glue

dots (see Appendix - Section C) or use commercially available stands

- Rotisserie shaker
- Phenol/chloroform
- 5 M Ammonium acetate (see Troubleshooting Guide, Appendix Section D, for details regarding the use of 3 M sodium acetate, pH 5.2)
- 100% ethanol
- 70% ethanol
- dNTP mixture (5 mM each)
- Taq polymerase (5 U/µl) (Example: New England Biolabs M0267L or GeneSpin STS-T1000)
- SYBR Green mix for real time PCR analysis
- PCR cycler
- (Optional) 5M Betaine stock solution for use with endpoint PCR

NOTES BEFORE STARTING

Fragmentation of Genomic DNA

The provided His-MBD2b/MBD3L1 protein complex has an enhanced affinity to bind methylated cytosines, particularly DNA fragments that contain five or more methylated cytosines. To enable clear interpretation of results, genomic DNA should be prepared such that DNA fragments containing a CpG region of interest do not contain methylated cytosines outside of this region (see "Appendix D. Troubleshooting" for further discussion). DNA can be fragmented by restriction digest or by mechanical means (*e.g.*, sonication).

Restriction digestion is especially useful for analysis of individual CpG islands. The genomic DNA is cut with a methylation-insensitive restriction enzyme(s) so that only CpGs of interest are contained within a particular restriction fragment. This fragment should be long enough (75 bp or longer) to allow for PCR analysis. Some useful methylation-insensitive restriction enzymes are shown in the below table. As might be expected, the enzymes whose recognition sites contain G and C bases cut more frequently in CpG islands than enzymes whose sites are composed only of A and T bases.

	Recognition Sequence	Number of fragments (per kb) in CpG islands	Number of fragments (per kb) in non-CpG islands
Mse I	TTAA	0.80	2.88
Bfa I	CTAG	1.56	1.55
Tas I	AATT	0.80	2.88
Csp6 I	GTAC	2.23	1.41

Mechanical fragmentation is ideal when a single DNA sample will be used for simultaneous analysis of many CpG islands (*e.g.*, when the isolated DNA will be analyzed by microarray methods) or when a CpG region of interest is not flanked by suitable restriction sites. In general, the DNA should be sheared to an average fragment size of less than 500 bp to minimize the number of CpG islands on each fragment.

Example fragmentation protocols are provided for both restriction digest and mechanical fragmentation on page 10. We suggest using 4 μ g of purified genomic DNA when performing restriction digestion and 20 μ g of purified genomic DNA for sonication. MethylCollector Ultra reactions can be performed on Ing - 1 μ g of fragmented genomic DNA.

Example Fragmentation Protocols

Restriction digest

This protocol can be modified depending on the amount of isolated genomic DNA or the restriction enzyme being used. We recommend preparing high-quality genomic DNA using a commercially available kit or a standard established protocol. The quality of the genomic DNA can be assessed by agarose gel electrophoresis and DNA concentration can be determined by UV spectrophotometry.

a) Set up the following restriction digest (with *Mse* I as an example, New England Biolabs (NEB)):

Genomic DNA (400 ng/µl)	10 µl
10X NEB Buffer 4	10 µl
100X BSA	1µl
Mse I (10 U/µl)	1µl
dH,O	78 µl
Total volume	100 µl

Note 1: The DNA volume may vary depending on its initial concentration.

- Note 2: MethylCollector Ultra has been used with as little as 1 ng of restriction-digested genomic DNA. As a reference, a human cell contains about 6 picograms DNA; 1 ng of genomic DNA corresponds to 170 cells.
- b) Mix well by pipetting and incubate at 37°C for 2 hours to overnight.
- c) Heat-inactivate *Mse* I by incubating the reaction mixture at 65°C for 20 minutes. If using an alternative restriction enzyme that cannot be heat-inactivated, the DNA can be purified by phenol/chloroform extraction and precipitation, or through use of a DNA purification column. See "Appendix D. Troubleshooting" on page 25 for comments about heat-inactivation.

Note 1: For greater accuracy, the digested DNA should be quantified.

Note 2: This digested DNA should be stored at -20°C until use.

Mechanical fragmentation (sonication)

Because *Mse* I or other restriction enzymes cannot always be used to fragment and isolate the DNA sequences of interest, sonication of the genomic DNA is an alternative method. Sonication is recommended for DNA fragmentation in preparation for Next-Gen sequencing analysis.

- a) Pipette 20 µg genomic DNA into a 1.5 ml microcentrifuge tube and adjust final volume to 300 µl by addition of 10 mM Tris-HCl pH 8.5.
- b) Using a tip probe sonicator, sonicate on ice with 15 pulses of 20 seconds (30% amplitude if using Active Motif's EpiShear[™] Sonicator, Catalog Nos. 53051 & 53052), with a 20-second pause on ice between each pulse. The sheared DNA can be visualized by ethidium staining after electrophoresis on a 3% agarose gel. The majority of the DNA fragments should be between 100 and 350 bp in length.

MethylCollector Ultra Protocol

PLEASE READ THE ENTIRE PROTOCOL BEFORE STARTING!

Step 1: Planning the experiment

- 1. Determine the number of reactions needed. For PCR or sequencing analysis, one reaction per sample is usually sufficient. For recommendations on downstream applications, such as whole genome amplification or Next-Gen sequencing, please refer to Appendix A.
- 2. Two different binding buffers are included in the kit. For high salt conditions (high stringency reactions) we recommend using High Salt Binding Buffer. For low salt conditions (lower stringency reactions) we recommend using Low Salt Binding Buffer. It may be necessary to perform experiments with both binding conditions to determine the optimal conditions for the target of interest.

Step 2: Binding reaction

- 1. Thaw components from storage as needed for preparation. Keep all components on ice when not in use.
- 2. Prepare Complete Binding Buffer for the desired binding conditions.

For high salt binding conditions (high stringency): Prepare the appropriate amount of Complete Binding Buffer according to the table below. Store on ice.

Reagent	One rxn	8 rxns
High Salt Binding Buffer	100 µl	800 µl
Protease Inhibitor Cocktail	0.5 µl	4 µl
Total Volume	100.5 μl	804 μl

For low salt binding conditions (lower stringency): Prepare the appropriate amount of Complete Binding Buffer according to the table below. Store on ice.

Reagent	One rxn	8 rxns
Low Salt Binding Buffer	100 µl	800 µl
Protease Inhibitor Cocktail	0.5 µl	4 µl
Total Volume	100.5 µl	804 μl

Note: The provided High and Low Binding Buffers are optimal for efficient capture of DNA fragments that contain five or more methylated CpGs.

3. Dilute the fragmented DNA in water if necessary. The MethylCollector Ultra protocol can be performed on a large range of sample DNA amounts (1 ng to 1 µg). We recommend 100 ng because it gives robust results without requiring a large amount of DNA.

- 4. In this step, the Input DNAs that will be used in the final PCR step are prepared.
 - a. If performing real time PCR: For the control genomic DNA provided in the kit, it is recommended that several Input DNA concentrations be run in triplicate. Input DNA should be tested at 0.01, 0.1, 1 and 10 ng/µl.
 - b. If performing endpoint PCR: For the control genomic DNA provided in the kit, PCR analysis is performed for 36 cycles on 25 ng of control DNA. The control DNA (provided at 20 ng/ μ l) should be diluted to 5 ng/ μ l for use in Input PCR. This can be done by diluting the DNA 1/4 in dH₂O (*e.g.*, 5 μ l of 20 ng/ μ l DNA + 15 μ l dH₂O to make 5 ng/ μ l DNA). 5 μ l of the 5 ng/ μ l DNA is used for Input PCR (see page 21).
 - **Note:** Customer sample Input DNA can be treated similarly. If your locus-specific PCR primers are efficient and PCR will be performed for 36 cycles, 25 ng of sample DNA can be used for the Input PCRs. However, PCR primer efficiency varies and you may want to try several amounts of Input DNA to be sure to obtain PCR products from reactions still in the linear phase of amplification.
- Using the PCR tubes provided, fully resuspend magnetic beads by inverting, then aliquot a 10 μl slurry into each tube. If preparing more than 4 reactions, cap and re-invert the beads after every 4 aliquots. (Note: When working with magnetic beads, pipette gently.)
- 6. Binding Reaction: Add the remaining components in the order shown below to each PCR tube. Pipet the His-MBD2b/MBD3L1 protein complex up and down several times to ensure homogeneity before use. Prepare a positive control reaction using the provided *Mse* I digested human, male genomic DNA.

Reagent	One rxn	Positive Control
Magnetic beads	10 µl	10 µl
Complete Binding Buffer	70 µl	75 µl
Fragmented genomic DNA	from 1 ng-1 µg (in a final volume of 10 µl)	-
Control Human, male genomic (<i>Mse</i> I digested)	DNA -	5 µl
His-MBD2/MBD3L1 protein com	ıplex 10 μl	10 µl
Total Volume	100 µl	100 µl

Note: It is recommended to aliquot the provided His-MBD2b/MBD3L1 protein complex into several small fractions for future use. Pipet the His-MBD2b/MBD3L1 protein complex up and down several times to ensure homogeneity before making aliquots. Store at -20°C.

8. Cap tubes and shake to mix thoroughly. Incubate on a rotisserie shaker for 1 hour at 4°C.

Step 3: Wash beads

- After the capture step is complete, spin the PCR tubes briefly and place tubes on a magnetic stand to pellet beads to the side of the tube. If further analysis of the unbound fraction will be performed, such as a comparison of the level of enrichment of methylated DNA, place supernatant in a microcentrifuge tube and set aside at 4°C for DNA clean up in Step 5 of the protocol. Otherwise, remove and discard the supernatant. To use the magnet provided in the kit, please see Appendix C.
- 2. Wash beads four times with 200 µl Binding Buffer. Pipette 2-3 times gently to resuspend. Use the same stringency Binding Buffer for the wash steps as was used for the binding reactions.
 - a. Place tubes on magnetic stand and allow beads to pellet on the side of the tube.
 - b. Carefully remove the supernatant and any residual bubbles.
 - c. Add 200 µl Binding Buffer and resuspend the pellet completely by pipetting several times. Ensure that the beads do not stick to the pipette tips. Depending on the strength of the magnet being used, it may be necessary to remove the tubes from the magnet and place in a separate rack to fully resuspend the beads.
 - d. Repeat steps a-c.
- In a microcentrifuge tube, prepare Complete Elution Buffer by adding 2 µl of Proteinase K to 98 µl of Elution Buffer AM1 for each reaction.
- 4. After the final wash, place tubes on magnetic stand and allow beads to settle to the side. Remove and discard supernatant without disturbing the beads.

Step 4: Recovery of methylated DNA fragments

- 1. Resuspend washed beads with 100 µl Complete Elution Buffer by pipetting 2-3 times.
- 2. Incubate samples at 50°C for 30 minutes. Every 10 minutes, invert the tubes to ensure the beads are resuspended.
- 3. During this incubation warm the Proteinase K Stop Solution at 37°C for 10 minutes.
- 4. Quick spin the tubes. Place tubes in magnetic stand and allow beads to pellet onto tube sides.
- 5. Carefully transfer the supernatant to a new microcentrifuge tube.
- 6. Add 2 µl of Proteinase K Stop Solution to each supernatant. Pipet 2-3 times to mix.
- 7. Proceed to Step 5, DNA clean up, or else DNA can be stored at -20°C. If the DNA is stored at -20°C, it recommended to be reheated at 37°C for 10 minutes prior to use.

Step 5: DNA clean up

Prior to use in downstream applications it is necessary to clean up the DNA. DNA clean up can be performed using one of the methods listed below:

Purification columns – use PCR clean up kits such as Active Motif's Chromatin IP DNA Purification Kit (Catalog No. 58002). Samples should be eluted in 50 μl volume, or other appropriate volume of water or buffer as needed for specific downstream applications.

For microarray analysis, use MinElute PCR Purification Kit (Qiagen part no. 28004) and elute in 10 μ l volume to obtain more concentrated DNA.

For MIRA-Seq, use MinElute PCR Purification Kit (Qiagen part no. 28004), elute into 24 μ l volume of EB and use 20 μ l of the purified DNA for the library preparation. Refer to Section A of the Appendix for more information on processing samples for MIRA-Seq.

• Phenol/chloroform extraction followed by ethanol precipitation – follow the protocol listed below

Phenol/Chloroform Extraction & Ethanol Precipitation

- 1. Add an equal volume of Phenol:Chloroform:Isoamyl Alcohol (25:24:1, v/v/v) to the eluted sample and, if desired, the unbound fraction from Step 2 No. 1.
- 2. Vortex the tube at maximum speed for 15 seconds.
- 3. Centrifuge the tube for 5 minutes at 12,000 x g at room temperature.
- 4. Carefully transfer the top, aqueous phase to clean microcentrifuge tube without collecting any of the lower organic phase or precipitate that may occur between the phases.
- 5. To each sample add:

1 µl Glycogen (20 mg/ml) (included in the kit)

1 sample volume of 5 M ammonium acetate

2.5 sample volumes of 100% ethanol

- 6. Mix well and incubate at -80°C for at least 2 hours.
- 7. Centrifuge the tube for 20 minutes at 12,000 x g, 4°C.
- 8. Carefully discard the supernatant without disturbing the pellet.
- 9. Add 500 µl of cold 70% ethanol. Do not disturb the pellet.
- 10. Centrifuge the tube for 10 minutes at 12,000 x g, 4°C.
- 11. Carefully discard the supernatant without disturbing the pellet.
- 12. Air-dry the pellet for 5 minutes (do not completely dry the pellet).
- Resuspend the DNA pellet in 50 µl sterile DNase-free water., or use other appropriate volumes of buffer or water as needed for specific downstream applications.
- This eluted DNA can be used immediately in PCR or stored at -20°C. If the DNA is stored at -20°C, it will need to be reheated at 37°C for 10 minutes prior to use in downstream applications.

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 - **Notes:** The polymerase chain reaction (PCR) process for amplifying nucleic acid is covered by U.S. Patent Nos. 4,683,195 and 4,683,202 assigned to Hoffmann-La Roche. Patents pending in other countries.

Use of methylation-specific PCR (MSP) is protected by U.S. Patent Nos. 5,786,146, 6,017,704, 6,200,756 & 6,265,171 and International patent WO97/46705. No license under these patents to use the MSP process is conveyed to the purchaser by purchasing this product.

MethylCollector[™] Ultra is covered under U.S. Patent No. 7,425,415.

Appendix

Section A. Downstream Applications

The heterodimeric MBD2b/MBD3L1 protein used in the MethylCollector Ultra Kit can be used to selectively bind and enrich for methyl CpG dinucleotides that are often found in the promoter/ regulatory regions of genes. The binding affinity of the MBD2b/MBD3L1 protein increases with the density of methyl CpG dinucleotides. The amount of DNA recovered from the MethylCollector Ultra Kit will depend on the global methylation status of the sample DNA. Usually only a small percentage of the starting material will be recovered, yet the recovered sample is specific for CpG-methylated DNA.

PCR Analysis

By amplifying a specific target site using PCR, it is possible to determine if there is an increase in methylation at that single site. Alternatively, the same site can be compared across different sample types, but it will be necessary to run MethylCollector Ultra on the same amount of each DNA sample and generate a separate standard curve for each sample.

Next-Gen Sequencing Analysis (MIRA-Seq)

To analyze whole-genome methylation profiles, researcher may chose to perform Next-Gen sequencing following the enrichment of methylated DNA with MethylCollector Ultra. This process has been validated when DNA enriched from MethylCollector Ultra is end adapted for use in sequencing. For instructions on how to prepare eluted DNA for Next-Gen sequencing, please refer to the Illumina ChIP-Seq DNA Sample Prep Kit protocol that can be viewed and downloaded at http://support.illumina.com/sequencing/sequencing_kits/chip-seq_dna_sample_prep_kit/best_practices.ilmn. Please refer to page 14 for recommended DNA clean-up and volume specifications for MIRA-Seq.

Microarray Analysis

Researchers who are interested in the global changes to methylation pattern due to an experimental treatment rather than specific target analysis may prefer to analyze the enriched methylated DNA by microarray, also known as MIRA-chip. For an overview of this technology, please visit http://www.ncbi.nlm.nih.gov/About/primer/microarrays.html.

MethylCollector Ultra can be used prior to microarray analysis. The final elution of the MBD2b/ MBD3L1 pulled-down DNA should be passed through a DNA clean-up column, such as Qiagen MinElute PCR Purification Kit (part no. 28004), to clean the DNA and minimize the final recovery volume for a more concentrated DNA sample. To perform the microarray analysis, microgram quantities of DNA will be needed. Since the amount of DNA recovered from the MethylCollector Ultra kit depends on the global methylation status of the sample, it may be necessary to pool multiple samples together or to perform whole genome amplification on the recovered material in order to obtain microgram quantities. Once microgram quantities of DNA have been obtained, the MethylCollector Ultra DNA can be labeled with Cy5 and the input DNA can be labeled with Cy3 for hybridization to the microarray.

Whole genome amplification

Active Motif's GenoMatrix[™] Whole Genome Amplification Kit (Catalog No. 58001) has been successfully validated for use with the GC-rich DNA from the MethylCollector Ultra Kit and shown to maintain the sequence representation of the starting material. It is recommended to start with 500 ng - 1 µg of fragmented DNA per binding reaction, alternatively, multiple small reactions can be pooled together. 10 ng is the minimum starting material required for the GenoMatrix[™] Whole Genome Amplification Kit.

To amplify DNA without using the GenoMatrix[™] Whole Genome Amplification Kit, a Ligated Mediated PCR will need to be performed. This will involve blunt ending the enriched, purified DNA, ligating blunt adaptors and PCR amplifying for 14-20 cycles. Amplicons produced from this method are generally between 300-500 bp due to a bias for smaller PCR products. The ligation reaction also has a tendency to be less efficient. However, after amplification you should obtain the microgram quantities of DNA needed for microarray analysis.

Non-amplification

If whole genome amplification will not be performed, multiple MethylCollector Ultra reactions will need to be combined in order to achieve microgram yields. It may be necessary to combine 10 or more separate binding reactions from the same sample material.

Bisulfite Conversion

Active Motif's MethylDetector Bisulfite Modification Kit (Catalog No. 55001) can be used to perform DNA conversion with bisulfite. The MethylDetector Kit works in the range of 200 pg to 2 µg of DNA, although special considerations need to be taken when working with samples containing less than 100 ng DNA. Following bisulfite conversion, DNA is purified and cloned into a DNA vector of choice. Cloned DNA is then transformed into competent bacterial cells, such as Active Motif's RapidTrans TAM 1 (Catalog No. 11096) and colonies are screened for positive clones. Plasmid DNA is isolated from the positive clones and submitted for sequencing. A comparison of bisulfite converted sequence data against the unconverted DNA sequence will provide the number of methylation sites for the region of interest.

Section B. PCR Analysis

PCR Primer Design

MethylCollector Ultra includes PCR primers for use with the provided *Mse* I digested human, male genomic DNA. If possible, real time PCR is recommended for analysis of DNA isolated with MethylCollector Ultra. To design primers specific to the CpG region of interest in your sample, please follow the recommendations below.

Primer design considerations

- Primers should flank the CpG region of interest and produce an amplicon between 100-350 bp in length for endpoint PCR or an amplicon of 100-150 bp for real time PCR. The CpG region needs to contain at least five methylated CpGs.
- ii. Each primer should be approximately 18-22 nucleotides long, contain 50% GC content and have a Tm between 55°C-60°C.
- iii. Restriction-digested DNA: PCR primer pairs should amplify a restriction fragment (or portion of a restriction fragment) that contains a CpG-rich region of interest. Each amplicon must also be free of internal sites for the restriction enzyme.
- iv. Sonicated DNA: PCR primers should flank the CpG-containing region of interest and the amplicon should not contain any CpG-dinucleotides that are outside of this region. This will minimize amplification of fragments that are isolated as a result of methylated CpGs that are near, but not within, the CpG-rich region of interest.
- v. PCR primers should be designed with the aid of a reliable primer design computer program (*e.g.*, http://frodo.wi.mit.edu/cgi-bin/primer3/primer3_www.cgi). Due to the technical limitations of PCR, it is sometimes necessary to design more than one primer pair for a given fragment of interest.
- vi. Potential primer pairs can be evaluated via computer simulation using a program such as UCSC Genome Browser (http://genome.ucsc.edu/) to ensure the primers selected will produce a single amplicon in a CpG region of the species being amplified.

Determining Primer Efficiency

It is advised to determine the efficiency of the PCR primers being used. Primers with an efficiency less than 90% will have poor reproducibility. To calculate primer efficiency:

Primer efficiency (%) = $[10^{(-1/slope)}-1] \times 100\%$

To obtain the slope value, follow the instructions for generating and graphing a standard curve in the Data Analysis and Use of Input DNAs section on page 21. Use the slope of the plotted standard curve in the primer efficiency equation above.

Endpoint PCR Analysis

A typical endpoint PCR protocol example follows below. This protocol was optimized for the control samples. For each new set of primers amplifying the promoter region of interest, the PCR conditions have to be optimized carefully (optimal T_m , number of cycles, *etc.*).

1. For one PCR reaction:

Reagent	One rxn
Sterile water	11.8 µl
10X PCR Buffer	2.5 µl
10X PCR compatible loading dye	2.5 µl
dNTP mixture (5 mM each dNTP)	1 µl
Forward Primer* (5 pmol/µl)	1 µl
Reverse Primer* (5 pmol/µl)	1µl
Taq (5 U/μl)	0.2 µl
DNA sample (eluted or Input)	5 µl
(Optional) 1 M Betaine can be added to PCR reactions as an enhancer to improve PCR results	
Total Volume	25 µl

* The provided PCR Primer Mixes contain both Forward and Reverse primers for use with the provided control DNA. Use 2 μ l of the PCR primer Mix in the PCR protocol described above. The Xist PCR primer mix has been optimized for use with low salt binding conditions, while NBR2 and GAPDH primer mixes can be used with either low or high salt binding conditions.

2. Place tubes in a PCR thermocycler and program as below:

94°C for 3 minutes (94°C for 20 seconds, 55°C for 30 seconds, 72°C for 30 seconds) for 36 cycles Hold at 4°C

- 3. Endpoint PCR can be analyzed by agarose gel electrophoresis. Run reactions by loading 10 µl from each of the PCRs on a thin 3% agarose gel at 100 V for 40 minutes in parallel with an appropriate DNA ladder. Post-stain the gel with 1 µg/ml ethidium bromide in 1X TAE buffer for 20 minutes. Observe gel under UV.
 - *GAPDH* Glyceraldehyde-3-phosphate dehydrogenase should be unmethylated in the control human, male genomic DNA and is not expected to produce a 69 base pair PCR product in the eluted DNA fraction.
 - *Xist* X inactive specific transcript is methylated in the control human, male genomic DNA and will produce a 178 base pair PCR product with the eluted fraction.
 - *NBR2* Neighbor of BRCA1 gene 2 is methylated in the control human, male genomic DNA and is expected to produce a 103 base pair PCR product with the eluted fraction.

Real Time PCR Analysis

This is an example PCR reaction. Please follow the specific instructions for your real time PCR instrument.

1. For one PCR Reaction:

Reagent	10 µl PCR reactions	20 µl PCR reactions
Fast SYBR Green master mix	5 µl	10 µl
Forward primer* (5 pmol/µl)	0.5 µl	1μl
Reverse primer* (5 pmol/µl)	0.5 µl	1μl
Sterile water	1μl	3 µl
DNA sample (eluted or Input)	3 µl	5 µl
Total volume	10 µl	20 µl

* The provided PCR Primer Mixes contain both Forward and Reverse primers for use with the provided control DNA. Use 1 µl of the PCR Primer Mix in the 10 µl reaction or 2 µl of the PCR primer mix in the 20 µl reaction for the PCR protocol described above. The Xist PCR primer mix has been optimized for use with low salt binding conditions, while NBR2 and GAPDH primer mixes can be used with either low or high salt binding conditions.

- Note: It is recommended to prepare triplicates of each sample and Input reaction. Input DNA should be tested at 0.01, 0.1, 1 and 10 ng/μl to obtain a standard curve.
- Place tubes in a Real Time PCR instrument and program as below. The amplification conditions should be optimized for each target locus and PCR instrument. A suggested starting point is:

95 °C for 2 minutes (95 °C for 10 seconds, 60 °C for 30 seconds) for 40 cycles

- Analyze the results. Data analysis varies depending on the instrument used. Obtain the standard curve from the Input samples. Use the standard curve to quantify the DNA in each sample.
 - *GAPDH* Glyceraldehyde-3-phosphate dehydrogenase should be unmethylated in the control human, male genomic DNA and will not amplify in the eluted fraction.
 - *Xist* X inactive specific transcript is a methylated promoter in the control human, male genomic DNA and will amplify early in the eluted fraction.
 - NBR2 Neighbor of BRCA1 gene 2 is a methylated promoter in the control human, male genomic DNA and will amplify early in the eluted fraction.

Data Analysis and Use of Input DNAs

Methylated DNA isolated using MethylCollector Ultra is usually analyzed by PCR amplification of the loci of interest. However, if the goal is to compare the methylation status of particular loci in different DNA samples, it is essential that MethylCollector Ultra be performed on the same amount of each DNA sample. Thus, DNA samples should be carefully quantified before use. In addition, Input DNA should be prepared for each of the different DNA samples (see Step 2. No. 4 in the Protocol) to clearly indicate the relative concentrations of the DNA samples.

For real time PCR, generating a standard curve using the input DNA enables accurate determination of the enriched DNA concentration.

- Produce a standard curve at 10, 1, 0.1 and 0.01 ng/µl in triplicate using the Input DNA from Step2, No. 4. We recommend running a standard curve every time MethylCollector Ultra is performed. However, if the primer efficiency has been determined to be greater than 90% according to the calculations on page 18, the standard curve can be generated once and stored for future use with the same DNA sample.
- 2. Run each sample with the appropriate DNA standard (*i.e.* prepare a separate standard curve for each cell line or species tested).
- Every gene will generate a different amplification profile. CT = Threshold Cycle or cycle number where the signal exceeds the background threshold level. CT values should be plotted for each gene to create a linear regression plot.
- 4. Plot CT versus log DNA concentration. See Figure 4 below.

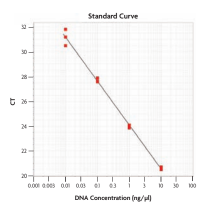


Figure 4: Example standard curve linear regression plot.

A standard curve for the human, male genomic DNA is provided as a reference only. Input DNA was tested in triplicate and plotted against the CT value. A new standard curve should be generated each time the assay is performed.

- 5. Using the CT value of the sample, extrapolate the DNA concentration of the sample DNA using the standard curve plot. To determine the amount of enriched DNA in the sample simply multiple the DNA concentration by the volume of enriched DNA.
- 6. Calculate the percent enrichment. Use the sample DNA quantity calculated above and compare it with the amount of DNA used in the initial binding reaction (Step 2, No. 6).

Enrichment (%) = Amount of enriched Sample DNA material Amount of fragmented DNA used in the binding reaction x 100%

7. *Optional:* Calculate the fold enrichment of methylated DNA. If the unbound material was collected in Step 3, No.1 and purified for analysis in PCR, the eluted samples can be compared to the unbound samples for the same locus in order to determine the fold enrichment.

Fold enrichment = Sample DNA concentration in the eluted fraction
Sample DNA concentration in the unbound fraction

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Section C. Use of Magnetic Beads and Included Bar Magnet

Caution: The included neodymium bar magnet is extremely powerful and is easily broken if handled incorrectly.

- 1. The magnet should be stored in the provided tube.
- 2. Be careful when working near metal objects or surfaces. A free magnet will jump great distances onto nearby metal surfaces with surprising speed. This can break the magnet.
- 3. If the magnet becomes attached to a flat metal surface, it should be removed by sliding it off the edge of surface. The magnet may be broken if you attempt to pull one end away from the metal.

Assembly of Magnetic Stands

The provided Mini Glue Dots can be used to attach the bar magnet to an empty tip box to create an effective magnet stand.

Creating a magnetic stand for 8-well PCR strips:

Note: 8-well strip tubes for use with standard 96-well PCR cyclers are appropriate.

- 1. Remove the covering tape from one side of two glue dots.
- Place a strip of PCR tubes in the wells of an empty tip box (200 µl tips) and place the magnet directly against the tubes. This is the way the magnet will be positioned when the glue dots are used to affix it to the box.
- 3. Attach the glue dots on the bar magnet (the uncovered face of the dot is placed on the magnet) as shown below.

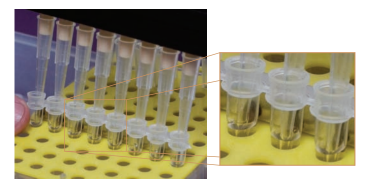


- 4. Remove the covering tape from the exposed side of the glue dots. Fix the magnet to the tip box so that it is against the PCR tubes. The magnetic stand is now ready for use.
 - **Note:** Familiarize yourself with using the magnetic stand before performing with PCR tubes for the first time. Add 5 µl of magnetic beads to 100 µl High Salt Binding Buffer in one tube of an 8-well strip of PCR tubes. Use this tube with the assembled bar magnet stand to become familiar with use of the beads and

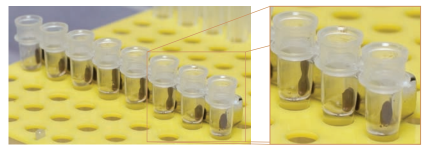
magnet. It is difficult to re-suspend the beads if the tubes are directly adjacent to the magnet, so it is usually best to move the tubes away from the magnet for resuspension.

Washing should be performed as follows:

a. Place the tubes in the rack against the magnet and allow the beads to be pinned to the side of the tube, as shown below.



b. Remove supernatant with a 200 µl pipetteman or a 200 µl eight-channel pipetteman.



- c. Move the tube strip into a row that is not adjacent to the magnet.
- d. Add wash buffer and pipette up and down to fully resuspend the beads. Ensure that a minimal amount of beads cling to the tips when the resuspension is complete.
- e. Repeat steps a-d until desired washing steps are complete.

Centrifugation of 8-well PCR strip tubes:

When working with 8-well PCR strip tubes, it may be desirable to centrifuge the tubes to collect the liquid and beads from the inside of the caps. This is easily accomplished in a centrifuge fitted with adaptors for spinning microtiter plates. In this case, a standard 96-well plate can be placed in the adaptor to hold the tubes in place. Take care to ensure the rotor is balanced (*e.g.*, place a microtiter plate and tubes of appropriate mass in the rotor's opposing 96-well plate adaptor). Spin the plates briefly to let the rotor reach a speed of 1000 x g before allowing the rotor to stop.

Section D. Troubleshooting Guide

Problem/question	Recommendation
The target DNA fragment has less than 5 methylated CpGs.	The provided Binding Buffers are optimal for efficient capture of DNA fragments that have five or more methylated CpGs. MethylCollector [™] Ultra is not recommended for isolation of DNA fragments containing less than 5 methylated CpGs.
Little or no enrichment of methylated DNA in the bound fraction	The binding conditions may be too stringent. Try using the Low Salt Binding Buffer to set up binding reactions for that particular target.
PCR amplification	It has been determined that using a hot-start polymerase (<i>i.e.</i> Phusion [™] from NEB) instead of a classic <i>Taq</i> polymerase may increase the sensitivity of the assay.
	Selection of an appropriate qPCR master mix is important to achieve good amplification. Since all commercially available mixes have different compositions regarding chemical enhancers or inhibitors for non-specific amplification, results may vary depending on the master mix used.
	The addition of 1 M Betaine to endpoint PCR reactions can be used to enhance the PCR and improve the results. Betaine lowers the melting temperature of GC-rich regions making them comparable to AT regions.
Storage of DNA	Once DNA is prepared using MethylCollector Ultra, samples may be stored at -20°C prior to PCR analysis. However, we recommend heating the frozen material to 37°C for 10 minutes before use in PCR, as heat-treatment releases any DNA bound to the tube during storage.
Can I use 3 M sodium acetate, pH 5.2 instead of 5 M ammonium acetate in the ethanol precipitation?	Yes, 3 M sodium acetate, pH 5.2 can be used at 1/10th sample volume along with 2 sample volumes of 100% ethanol during the precipitation step. However, we have noticed that the ammonium acetate precipitation had better yield of recovery than the sodium acetate in a direct comparison of several samples.
Should I use Restriction Digest or Sonication to fragment my DNA?	Restriction Digest is very precise and reproducible, however, the DNA must be well puri- fied and analysis of several loci may also require use of different enzymes. In addition, the region of interest may not be flanked by suitable restriction sites and single-nucleotide polymorphisms (SNPs) between different cell types may confound results. In contrast, Sonication is random, which enables analysis of many loci simultaneously (microarray), but it may not be possible to shear DNA small enough to isolate CpG islands of interest. In addition, results may vary from shearing to shearing depending on sonicator used. Also it is difficult to prepare DNA from a small number of cells.
Heat inactivation or removal of restriction enzyme used to fragment DNA	After restriction digest, we recommend that samples be treated for 20 minutes at 65°C. Some enzymes (such as <i>Mse</i> I) will be inactivated by this treatment, while those that are not will be forced off the DNA. In most cases (even when using enzymes that are not heat-inactivated), DNA treated in this fashion should be suitable for use in the MethylCollector Ultra protocol. In some situations (<i>e.g.</i> , when the DNA used in a digest is contaminated with cellular proteins or when a large amount of restriction enzyme is required for the digest) it may be desirable to purify the digested DNA by purification columns or through phenol extraction/ethanol precipitation.
10X PCR Loading Dye	If PCR is performed using the 10X PCR Loading Dye provided, it is not necessary to add additional loading dye to the samples before running samples on agarose gel.

Section E. Related Products

DNA Methylation	Format	Catalog No.
MethylDetector™	50 rxns	55001
MethylCollector™ Ultra	30 rxns	55005
UnMethylCollector™	30 rxns	55004
Hydroxymethyl Collector	25 rxns	55013
MeDIP	10 rxns	55009
hMeDIP	10 rxns	55010
DNMT Activity / Inhibition Assay	1 x 96 rxns	55006
Methylated DNA Standard Kit	3 x 2.5 μg	55008
5-Carboxylcytosine DNA Standard Kit	0.5 µg	55014
Fully Methylated Jurkat DNA	10 µg	55003
Recombinant Tet1 protein	25 µg	31363
β-Glucosyltransferase enzyme	500 Units	55012
PvuRts1l restriction enzyme	50 Units	55011

Whole Genome Amplification	Format	Catalog No.
GenoMatrix [™] Whole Genome Amplification Kit	50 rxns	58001

Antibodies	Application	Format	Catalog No.
3-Methylcytosine (3-mC) pAb	DB	100 µg	61111
5-Carboxylcytosine (5-caC) pAb	DB, IF	100 µl	61225
5-Formylcytosine (5-fC) pAb	DB, IF	100 µl	61223
5-Hydroxymethylcytosine pAb	DB, IF, IHC, MeDIP	100 µl	39769
5-Hydroxymethylcytosine mAb	DB, MeDIP	100 µg	39999
5-Methylcytosine mAb	FACS, IHC, IP, MeDIP	50 µg	39649
5-Methylcytosine pAb	DB, MeDIP	100 µg	61255
DNMT1 mouse mAb	ChIP, IHC, IP, WB	100 µg	39204
DNMT2 rabbit pAb	WB	100 µg	39205
DNMT3A mouse mAb	ChIP, IF, IHC, WB	100 µg	39206
DNMT3B mouse mAb	ChIP, IF, IP, WB	100 µg	39207
DNMT3L rabbit pAb	WB	100 µl	39907
MBD1 mouse mAb	WB	100 µg	39215
MBD2 mouse mAb	WB	100 µg	40965
MBD3 mouse mAb	WB	100 µg	39216
MBD4 mouse mAb	WB	100 µg	39217
MeCP2 rabbit pAb	WB	100 µg	39218

Active Motif also offers a growing list of application validated antibodies, including antibodies for histones and histone modifications, transcription factor antibodies, DNA methylation-related antibodies and ChIP validated antibodies. For a complete list go to www.activemotif.com/abs

Histone Purification	Format	Catalog No.
Histone Purification Kit	10 rxns	40025
Histone Purification Mini Kit	20 rxns	40026
Chromatin Assembly	Format	Catalog No.
Chromatin Assembly Kit	10 rxns	53500
Hel a Core Histones	36 µg	53501

Histone Acetyltransferase and Deacetylase Activity	Format	Catalog No.
HAT Assay Kit (Fluorescent)	1 x 96 rxns	56100
Recombinant p300 protein, catalytic domain	5 µg	31205
HDAC Assay Kit (Fluorescent)	1 x 96 rxns	56200
HDAC Assay Kit (Colorimetric)	1 x 96 rxns	56210

ChIP-IT [™] Kits	Format	Catalog No.
ChIP-IT [™] Express	25 rxns	53008
ChIP-IT [™] Express Enzymatic	25 rxns	53009
ChIP-IT [™] Express HT	96 rxns	53018
ChIP-IT [™] Protein G Magnetic Beads	25 rxns	53014
Re-ChIP-IT [™]	25 rxns	53016
ChIP-IT [™]	25 rxns	53001
ChIP-IT [™] w/o controls	25 rxns	53004
ChIP-IT [™] Shearing Kit	10 rxns	53002
ChIP-IT [™] Enzymatic	25 rxns	53006
ChIP-IT [™] Enzymatic w/o controls	25 rxns	53007
Enzymatic Shearing Kit	10 rxns	53005
Salmon Sperm DNA/Protein G agarose	25 rxns	53003
ChIP-IT [™] Control Kit – Human	5 rxns	53010
ChIP-IT [™] Control Kit – Mouse	5 rxns	53011
ChIP-IT™ Control Kit – Rat	5 rxns	53012
Ready-to-ChIP HeLa Chromatin	10 rxns	53015
Ready-to-ChIP Hep G2 Chromatin	10 rxns	53019
Ready-to-ChIP K-562 Chromatin	10 rxns	53020
Ready-to-ChIP NIH/3T3 Chromatin	10 rxns	53021

Recombinant Methylated Histones	Format	Catalog No.
Recombinant Histone H2A	50 µg	31251
Recombinant Histone H3 (C110A)	50 µg	31207
Recombinant Histone H3 monomethyl Lys4	50 µg	31208
Recombinant Histone H3 dimethyl Lys4	50 µg	31209
Recombinant Histone H3 trimethyl Lys4	50 µg	31210
Recombinant Histone H3 monomethyl Lys9	50 µg	31211
Recombinant Histone H3 dimethyl Lys9	50 µg	31212
Recombinant Histone H3 trimethyl Lys9	50 µg	31213
Recombinant Histone H3 monomethyl Lys27	50 µg	31214
Recombinant Histone H3 dimethyl Lys27	50 µg	31215
Recombinant Histone H3 trimethyl Lys27	50 µg	31216
Recombinant Histone H3 monomethyl Lys36	50 µg	31217
Recombinant Histone H3 dimethyl Lys36	50 µg	31218
Recombinant Histone H3 trimethyl Lys36	50 µg	31219
Recombinant Histone H3 monomethyl Lys79	50 µg	31220
Recombinant Histone H3 dimethyl Lys79	50 µg	31221
Recombinant Histone H3 trimethyl Lys79	50 µg	31222
Recombinant Histone H4	50 µg	31223
Recombinant Histone H4 monomethyl Lys20	50 µg	31224
Recombinant Histone H4 dimethyl Lys20	50 µg	31225
Recombinant Histone H4 trimethyl Lys20	50 µg	31226

Technical Services

If you need assistance at any time, please call Active Motif Technical Service at one of the numbers listed below.

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