



ELLIS BIO

User Manual

SuperMethyl™ Max Bisulfite Conversion Kit

6-Reactions, Magnetic Bead Purification

MAX-6R-BEAD

Version 2026.6.1

Questions? We're ready to help!

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1. Key Features

Ellis Bio is committed to revolutionizing DNA methylation analysis with unparalleled speed and precision.

- ✓ **Ultra-Mild Bisulfite Conversion:** Achieve complete and ultra-mild bisulfite conversion to minimize DNA damage.
- ✓ **High Efficiency:** Consistently delivers over 99.8% C-T conversion ratios of unmethylated cytosines for accurate analysis while preserving methylated cytosines.
- ✓ **High Compatibility with low-input DNA:** Minimized DNA damage enables robust application to ultra-low input, starting from 100 pg to 2 ug of gDNA or cfDNA.
- ✓ **Low Background Noise:** Reduces false positives and enhances the accuracy of 5mC signal detection.
- ✓ **Versatile Applicability:** Ideal for a wide range of applications, including methylation-specific PCR (MSP), microarrays, and NGS.

This kit offers an unmatched combination of maximized DNA integrity, accuracy, and reliability, setting a new benchmark in DNA methylation analysis.

2. Product Description

The **SuperMethyl™ Max Bisulfite Conversion Kit** from Ellis Bio offers the revolutionarily mild yet powerful bisulfite conversion for DNA methylation analysis. Featuring a newly engineered **ultra-mild bisulfite conversion formulation** and **a magnetic bead-based DNA purification method**, this kit simplifies the methyl conversion workflow and achieves efficient conversion with minimal DNA degradation. Complete bisulfite conversion and DNA purification can be accomplished in 3 hours or less, preserving DNA integrity even with ultra-low input samples, while achieving **> 99.5%** conversion efficiency. Its broad applicability makes it a valuable tool for both research and translational settings.

Designed for compatibility and consistent performance in sensitive and/or low-input DNA samples, the SuperMethyl™ Max Bisulfite Conversion Kit delivers reliable, reproducible results, making it the preferred choice for precision-focused researchers.

Minimal DNA damage: Ultra-Mild Bisulfite Technology preserves DNA integrity, ideal for cfDNA and low-input samples.

Exceptional C-to-T Conversion Efficiency: Users typically achieve 99.8% conversion rates.

High Library Complexity: Produces libraries with much greater complexity and larger insert sizes compared to conventional bisulfite conversion methods.

Enhanced CpG Detection: Detects more CpG sites with fewer sequencing reads.

Uniform GC Coverage: Delivers consistent coverage across GC-rich and GC-poor regions.

Wide Input Range: Compatible with 100 pg to 2 ug of DNA input.

Low Background Noise: Reduces false positives and enhances the accuracy of 5mC signal detection.

Versatile DNA Fragmentation Compatibility: Supports both sonication and enzymatic fragmentation workflows.

3. Kit Components

Number of tests per kit: 6 reactions

Your Kit includes:

Component	Volume/Quantity	Storage
Kit Capacity	6 Tests	Please see storage recommendations for individual components
Conversion Reagent ●	0.8 mL	4 °C (recommended)
Preparation Reagent ●	100 µL	Room temperature
Enhancer A ●	100 µL	Room temperature
Enhancer B ●	100 µL	4 °C (recommended)
Binding Buffer	7 mL	Room temperature
Wash Buffer (concentrate)*	3 mL*	Room temperature
Desulphonation Buffer	2 mL	Room temperature
Elution Buffer	0.5 mL	Room temperature
Purification Beads ●	150 µL	Room temperature
* Wash Buffer requires the addition of 12 mL 100% ethanol (EtOH) before first use.		

4. Notes for Users

User-supplied materials

- 100% ethanol
- Nuclease-free H₂O
- 1.5 mL low-adhesion microcentrifuge tubes and PCR tubes.
- Microtube rotator
- Magnetic tube racks
- Positive and negative control samples such as unmethylated DNA and fully methylated pUC19 DNA
- Lambda DNA (dam-, dcm-)

Storage

We recommend storing **Conversion Reagent** ● and **Enhancer B** ● at 4 °C and avoid exposure to air and light. All other kit components can be stored at room temperature. The kit is stable for up to 12 months. Please refer to the product label for the expiration date.

Applications

The kit is compatible with DNA from various sources including cell-free DNA (cfDNA), genomic DNA (gDNA) extracted from cells or tissues, and formalin-fixed paraffin-embedded (FFPE) samples derived DNA. Other DNA sources might be compatible; we encourage users to run quality control tests on samples to confirm.

Input DNA Requirements

The kit requires an input DNA amount of 100 pg** to 2 ug. We advise quantifying the DNA with a precise instrument such as a Qubit Fluorimeter (Thermo).

**Users may be able to use inputs lower than 100 pg. We recommend users run quality control tests on samples that are outside the recommended range.

Product Performance Indicators

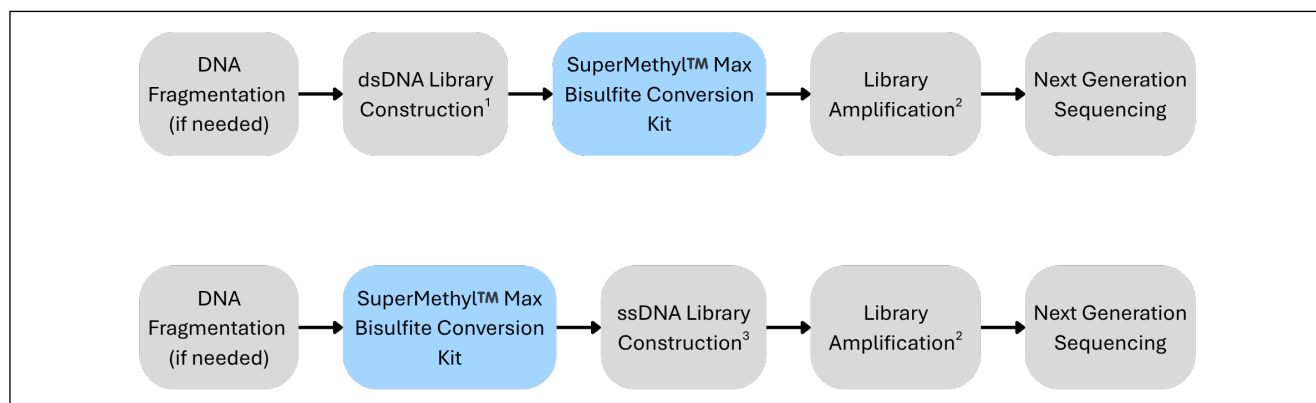
The C-to-T conversion rate at all unmethylated cytosines in both CpG and non-CpG contexts of λDNA exceeds 99.5%. The estimated methylation levels at all CpG sites in fully methylated pUC19 DNA is consistently above 95%.

Caution

This kit is for research use only and is not intended for use in diagnostic applications. **Preparation Reagent** ●, **Conversion Reagent** ●, **Enhancer A** ●, **Enhancer B** ●, **Desulphonation Buffer**, and **Wash Buffer** contain volatile ingredients. Cap the bottles tightly after use and store at recommended temperatures. Safety Data Sheets are available upon request.

5. Experimental Planning

1. If your application requires next-generation sequencing library preparation, the bisulfite conversion protocol can occur downstream or upstream of library preparation, depending on your experimental design.



¹ Available through Ellis Bio or other dsDNA library preparation suppliers. Use with methylated adapters.

² Use uracil-tolerant library amplification system, available through Ellis Bio or other suppliers.

³ Available from other suppliers.

Kit Compatibility

We recommend testing quality control samples with your preferred library construction kit to determine compatibility. The SuperMethyl™ Max Bisulfite Conversion Kit is compatible with the following library preparation kits:

- Pre-bisulfite conversion:
 - Ellis Bio SuperMethyl™ dsDNA Library Kit for Illumina
 - KAPA EvoPrep Kit (Roche)
 - KAPA HyperPrep Kit (Roche)
 - NEBNext® Ultra™ II Ligation Module (New England Biolabs)
- Post-bisulfite conversion:
 - SRSLY® PicoPlus Uracil+ Kit (Claret Bioscience)
 - SRSLY® NanoPlus Uracil+ Kit (Claret Bioscience)
 - SRSLY® MethylPlus Kit (Claret Bioscience)
 - xGen™ Methylation-Sequencing DNA Library Preparation Kit (IDT)

Methylation controls

It is recommended to include appropriate positive and negative controls in the experiment. Spike-in controls such as unmethylated lambda DNA and methylated pUC19 DNA are useful options.

6. Experimental Protocol

1. Reagent Preparation

- 1.1. **Prepare Wash Buffer:** Add 12 mL of 100% ethanol (EtOH) to the **Wash Buffer** before first use. Invert to mix thoroughly and ensure the bottle cap is tightly sealed to prevent ethanol evaporation, which could impact the effectiveness of the **Wash Buffer**.
- 1.2. **Prior to using materials in 2 mL screwcap vials:** For any kit component provided in a 2 mL screwcap vial (e.g., **Conversion Reagent** ●, **Purification Beads** ●, **Elution Buffer**, etc), briefly centrifuge the vial to remove material that may have pooled in the cap. Use a micropipette to gently resuspend the contents so that all settled material is evenly mixed before use.
- 1.3. **Conversion Reagent** ● **Notes:** Immediately prior to use, please inspect the **Conversion Reagent** ● with any signs of crystallization. If crystals are present, heat the vial at 55 °C or vortex until fully dissolved, then allow the reagent to equilibrate to room temperature before use. Failure to dissolve the crystals may significantly impair conversion efficiency. Minor crystallization on the vial cap is normal.

2. Sample Preparation

- 2.1. **For gDNA or FFPE-derived DNA:** Fragment the DNA using either sonication or enzymatic digestion. Purify the fragmented DNA using magnetic beads and perform size selection. Submit the processed DNA for quality control (QC) and determine the sample concentration.¹

For cfDNA: Fragmentation is not required.²

¹Note: Refer to Section 5 for guidance on including positive and negative methylation controls. We also recommend using a fragment analysis method, such as the TapeStation (Agilent), to assess fragment size distribution. For accurate DNA quantification, use a fluorometric method such as the Qubit assay (Thermo Fisher Scientific).

²Note: Please see our Supplementary Section 7 about gDNA contamination in cfDNA for tips for clean-up.

- 2.2. **Library preparation (see Section 5 above):** If your workflow involves a double-stranded DNA (dsDNA) library preparation, as illustrated in the figure in Section 5, proceed with the library construction steps: perform end repair, dA-tailing, ligation of methylated adapters, and carry out the post-ligation purification steps according to the manufacturer's protocol. **Do not begin library amplification at this stage.**

3. Bisulfite Conversion

- 3.1. In a 0.2 mL nuclease-free PCR tube, pipette the volume to obtain 100 pg – 2 ug of input DNA. Add nuclease-free H₂O up to a total volume of 20 µL³.

³Note: SuperMethyl Max kit is also compatible with higher DNA input volume (20-40 µL), please reach out to support@ellisbio.com for more information.

- 3.2. Add 2 µL of **Preparation Reagent** ● to the tube per **Protocol Table 1**. Mix thoroughly by vortexing or gentle pipetting. Briefly centrifuge to collect the liquid at the bottom of the PCR tube.

Protocol Table 1	
Component	Volume
Input DNA	20 µL (100 pg - 2 ug)
Preparation Reagent ●	2 µL
Total Volume	22 µL

- 3.3. Place the capped PCR tube in a thermal cycler and incubate at 42 °C (with a 55 °C heated lid) for 20 minutes using the program shown in **Protocol Table 2**. After the incubation, transfer the sample immediately to ice until used in Step 3.5.

Protocol Table 2		
Step	Temperature	Time
1. Sample Prep incubation	42 °C	20 min
2. Transfer immediately to ice for at least 2 minutes		

- 3.4. In a 1.5 mL nuclease-free microcentrifuge tube, prepare the **Conversion Mastermix**⁴ by adding components in the following order: first, add **Conversion Reagent** ●, then add **Enhancer A** ● and vortex immediately for 10 seconds. Next, add **Enhancer B** ● and vortex for an additional 10 seconds. Refer to **Protocol Table 3** for component volumes:

Protocol Table 3

Component (add in the order listed below)	Volume
Conversion Reagent ⁴ ●	77 µL
Enhancer A ●	2 µL
Enhancer B ●	8 µL
Total volume	87 µL



⁴Note: See Step 1.3. above for an important note regarding crystallization of the Conversion Reagent. We recommend preparing the Conversion Master Mix after Step 3.3. If the Conversion Reagent and Enhancer B are stored at 4 °C, allow them to equilibrate to room temperature before use. We recommend using the Conversion Master Mix immediately after its preparation. However, if the Conversion Master Mix is not used immediately after its preparation, incubate it at 55 °C to minimize the risk of precipitation prior to Step 3.5.

- 3.5. Prepare the bisulfite conversion reaction according to **Protocol Table 4**. After adding the **Conversion Mastermix**⁵ to the **Post-sample Preparation DNA** for each sample, immediately mix thoroughly by pipetting 5 times or vortexing for 5 seconds. Briefly centrifuge to collect liquid at the bottom of the PCR tubes. If the sample volume exceeds the thermocycler settings, please split the sample into additional PCR tubes as needed.

Protocol Table 4

Component	Volume
Post-sample Preparation DNA (Step 3.3)	22 µL
Conversion Mastermix (Step 3.4)	87 µL
Total volume	109 µL



⁵Note: Check the Conversion Master Mix for visible precipitation before use. If precipitation is observed, prepare a fresh Conversion Master Mix.

- 3.6. Place the capped PCR tubes with flat caps in a thermal cycler and incubate at 55 °C (with a 75 °C heated lid) using the following program in **Protocol Table 5**.

Protocol Table 5

Step	Temperature	Time
1. Bisulfite	55 °C	90 mins
2. Hold ⁶	20 °C	< 5 mins
3. Proceed to next step immediately		



⁶Note: This step is optional; we recommend proceeding directly to the next step. Please do not place the sample on ice or store it at 4 °C, as this may lead to white crystallization formation, which can reduce the DNA recovery rate. We also do not recommend additional vortexing of the conversion solution in the tube, as this may induce precipitation.

4. Purification of Bisulfite-converted DNA

- 4.1. Add 800 μ L of **Binding Buffer** and 15 μ L of **Purification Beads**⁷  to a 1.5 mL low-adhesion microcentrifuge tube.



⁷Note: Purification Beads settle quickly, pipette or vortex thoroughly to keep them well-suspended. When working with multiple samples manually, we recommend vortexing the Purification Bead and Binding Buffer solution frequently (for example, every 4 samples) to maintain consistent suspension.

- 4.2. Transfer the bisulfite converted reaction solutions from the PCR tubes (from Step 3.6) into the 1.5 mL microcentrifuge tube containing the **Binding Buffer** and **Purification Beads**  (Step 4.1). Mix thoroughly by pipetting up and down or by gently vortexing at low speed for 10 seconds. Incubate at room temperature (15 – 25 °C) for 10 minutes with continuous rotation. **A microtube rotator is strongly recommended for this incubation step.**

- 4.3. Briefly centrifuge the microcentrifuge tube to collect the liquid and beads at the bottom, then place it on a magnetic stand for 5 minutes or until beads pellet has collected on magnet and the supernatant is clear. While the microcentrifuge tube on the magnetic stand, remove and discard the supernatant⁸.

⁸Note: If some Purification Beads adhere to the non-magnetic side of the tube, remove the supernatant slowly, allowing beads to migrate to the magnetic side as the liquid volume decreases.

- 4.4. Remove the microcentrifuge tube from the magnetic stand for this step and each subsequent buffer addition.
- 4.5. Add 400 μ L of **Wash Buffer** (ensure 100% ethanol was added in Step 1.1 before first use) to the beads and resuspend by pipetting or by vortexing the microcentrifuge tube at low speed for 10 seconds.
- 4.6. Briefly centrifuge the microcentrifuge tube to collect the liquid and beads, then place it back on the magnetic stand for 3 minutes. Carefully remove and discard the supernatant.
- 4.7. Add 200 μ L of **Desulphonation Buffer** to the beads. Resuspend by pipetting or by vortexing the microcentrifuge tube at low speed for 10 seconds.
- 4.8. Incubate at room temperature for 10 minutes with continuous rotation. **A microtube rotator is strongly recommended for this incubation step.**

- 4.9. Briefly centrifuge the microcentrifuge tube to collect the liquid and beads, then place it back on the magnetic stand for 3 minutes. Carefully remove and discard the supernatant.
- 4.10. Add 400 μ L of **Wash Buffer** to the beads. Resuspend by pipetting or by vortexing the microcentrifuge tube at low speed for 10 seconds.
- 4.11. Briefly centrifuge the microcentrifuge tube to collect the liquid and beads, then place it back on the magnetic stand for 3 minutes. Carefully remove and discard the supernatant.
- 4.12. Repeat Steps 4.10 and 4.11. After the final wash and discarding the supernatant, use a 10 μ L tip to remove as much **Wash Buffer** as possible. This will help ensure efficient drying of the purification beads.
- 4.13. Place the uncapped microcentrifuge tubes on a heating element to **completely dry the Purification Beads**⁹ at 55 °C for 5 - 10 minutes.



⁹Note: **Ensure the Purification Beads are completely dry before proceeding to the elution step (Step 5).** An example photo of dried beads is shown in the Supplementary Material (Section 7). If needed, extending the heating time may improve recovery of bisulfite-converted DNA. If the heating element has a lid, please keep the heating element lid open during this step. Because heating efficiency may vary between heating blocks, we recommend optimizing the heating time to achieve optimal recovery.

5. Elution

- 5.1. Add 25-30 μ L of **Elution Buffer**¹⁰ directly to the dried beads and resuspend by pipetting or vortexing the microcentrifuge tube at low speed for 10 seconds.

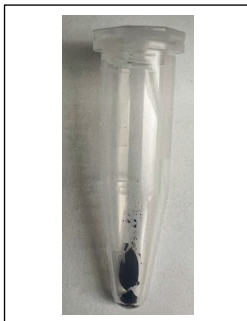
¹⁰Note: Please note that approximately 5 μ L elution buffer may remain bound to dried beads. The volume of Elution Buffer can be adjusted according to the requirements of downstream applications. Varying the elution volume may affect elution efficiency: lower volumes typically yield more concentrated DNA but may lower overall recovery.

- 5.2. Incubate the elution at 55°C for 4 minutes with mixing at 1000 RPM. Then place the microcentrifuge tube on a magnetic stand for 1 - 3 minutes, or until the solution becomes clear.
- 5.3. Carefully transfer the supernatant (eluate) to a clean microcentrifuge tube or PCR tubes for downstream amplifications.

Optional stopping point: The eluate, containing bisulfite-converted DNA, is immediately ready for downstream applications such as PCR analysis or next-generation sequencing. For storage, keep the eluate at -20 °C for short-term use or at -80 °C for long-term use.

7. Supplementary Material

Image of dried purification beads (Section 6 Experimental Protocol, Step 4)



Following incubation of beads at 55 °C for 5 - 10 minutes, sufficiently dried beads have matte and slightly cracked finish.

Library Amplification with 3rd Party Kits

In this section, you will find a supplemental library amplification protocol using KAPA HiFi Uracil+ Ready Mix (Roche). Please refer to the manufacturer's protocol for additional details.

Materials Required

- 2X KAPA HiFi Uracil+ Ready Mix
- P5 Index Primer (15 μ M)
- P7 Index Primer (15 μ M)
- PCR-grade nuclease-free water (if needed)

Reaction Setup

1. Thaw all reagents on ice and mix thoroughly by gentle vortexing or pipetting.
2. Add nuclease-free water to the bisulfite-converted DNA eluted in Section 6 (Main protocol, Step 5) to bring the total volume to 22 μ L.
3. In a PCR tube or plate, prepare the following amplification reaction mix on ice according the table below:

Component	Volume
Bisulfite Converted DNA in water (from Step 2 above)	22 μ L
2X KAPA HiFi U+ ReadyMix	25 μ L
P7 Index Primer (15 μ M)	1.5 μ L
P5 Index Primer (15 μ M)	1.5 μ L
Total Volume	50 μ L

4. Gently mix the reaction by pipetting. Briefly centrifuge the tube to collect the liquid at the bottom of PCR tubes.

5. Place the reaction in a thermocycler and run the following program:

Step	Temperature	Time	Cycles
Initial Denaturation	98 °C	45 sec	1
Denaturation	98 °C	15 sec	7 – 10 cycles ⁹
Annealing	62 °C	30 sec	
Extension	72 °C	30 sec	
Final Extension	72 °C	1 min	1
Hold	4 °C	∞	1
⁹ Note: Use 7–10 cycles depending on input amount and desired yield. Fewer cycles help preserve library complexity.			

6. Purify the amplified library using 1X AMPure XP beads (Beckman Coulter Life Sciences), or an alternative bead source per cleanup (according to their recommended library clean-up protocol).

Cell-Free DNA Sample Considerations

By running a fragment analysis (such as TapeStation or Bioanalyzer) on input cfDNA, one can determine if the majority of fragments are in the expected fragment range or if there is potential high molecular weight genomic DNA contamination. We recommend using size selection beads, such as SPRI beads to remove high molecular weight species above 700 bp. Follow the manufacturer's protocol for the size selection beads.