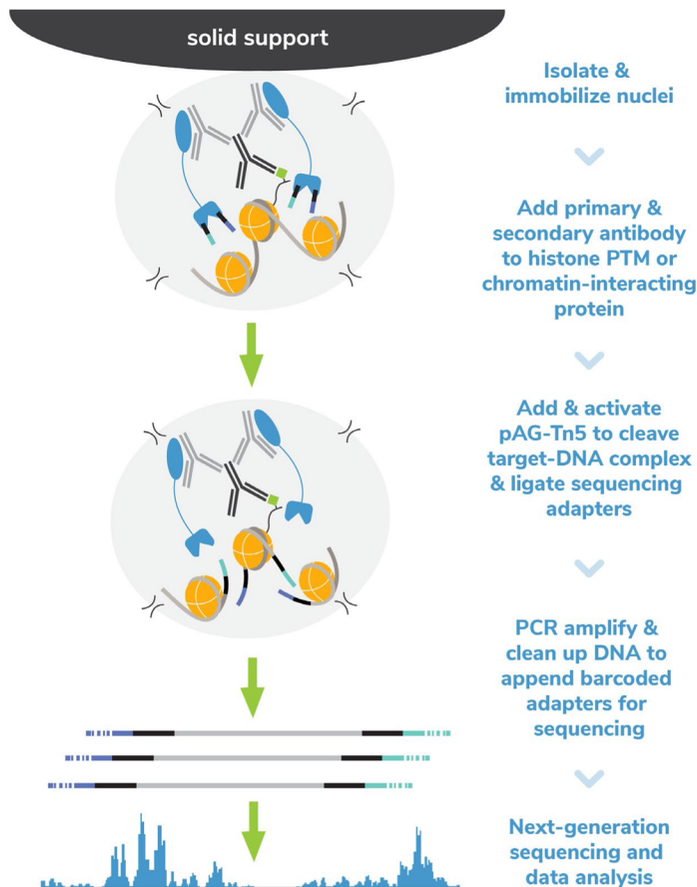


## EpiCypher® CUTANA® Direct-to-PCR CUT&Tag Protocol

Currently optimized for histone PTMs & select chromatin-associated proteins

### 1. Overview

Cleavage Under Targets & Tagmentation (CUT&Tag) is a revolutionary genomic mapping strategy developed by the group of Dr. Steven Henikoff<sup>1</sup>. It builds on Chromatin ImmunoCleavage (ChIC) from Dr. Ulrich Laemmli<sup>2</sup>, whereby a fusion of an antibody binding protein to a DNA-modifying enzyme is recruited to modify antibody-bound chromatin *in situ*<sup>3</sup>. In CUT&Tag, nuclei (recommended) or cells are immobilized to a solid support. A fusion of proteins A and G with prokaryotic transposase 5 (pAG-Tn5) is used to selectively tagment antibody-bound chromatin. The tagmentation reaction appends sequence adapters to the antibody-bound DNA, which can be directly PCR amplified *in situ* to yield sequence-ready DNA. High resolution genome-wide profiles of histone post-translational modifications (PTMs) can then be generated using as few as 3-5 million sequencing reads. The entire workflow can be conducted in a single tube using multichannel pipettes, enabling high experimental throughput (**Figure 1**).



**Figure 1.** Overview of the CUTANA® Direct-to-PCR CUT&Tag protocol.



## 2. CUTANA® Products & Services: Advantages

ChIC and CUT&Tag have revolutionized the study of chromatin regulation. Compared to ChIP-seq (the current leading approach for genome-wide mapping of histone PTMs and chromatin-associated proteins), CUT&Tag offers the following advantages:

- Requires substantially fewer cells
- More cost effective: less antibody and required sequencing depth
- Markedly improved [signal : noise]
- Rapid workflow: cells → sequence-ready libraries in < 2 days
- Empowers benchtop sequencers (e.g. Illumina® MiniSeq or MiSeq)
- Dramatically increases experimental throughput (transforming the way scientists approach experimental design for genomic mapping studies)

EpiCypher now offers CUTANA® pAG-Tn5, the essential reagent for ChIC/CUT&Tag workflows:

- Optimized fusion of Proteins A and G with hyperactive transposase 5 (pAG-Tn5) is directly compatible with a broad range of antibody isotype sources (e.g. mouse, rabbit, etc.)
- 50 and 250 reaction pack sizes, enabling greater experimental throughput
- Check [epicypher.com/resources/protocols](https://epicypher.com/resources/protocols) for regular protocol updates
- Additional CUTANA® products coming soon:
  - ChIC/CUT&RUN Kits
  - ChIC/CUT&Tag Kits
  - Spike-in controls for normalization
  - Platform validated antibodies
  - *Inquire for more information or to connect to EpiCypher scientists:*  
[info@epicypher.com](mailto:info@epicypher.com)

### 3. Protocol Notes

1. This version of the Direct-to-PCR CUT&Tag protocol is validated for histone PTMs as well as select chromatin interacting proteins (e.g. CTCF and FOXA1 transcription factors, SWI/SNF chromatin remodeling proteins). For additional information about CUT&Tag with chromatin interacting proteins, contact us at [info@epicypher.com](mailto:info@epicypher.com).
2. Always include control conditions (e.g. positive control anti-H3K4me3 or anti-H3K27me3 and negative control IgG; see Reagents & FAQ sections). This is especially critical when evaluating CUT&Tag to previously untested targets and/or antibodies.
3. Unlike CUT&RUN where cells are immobilized to ConA beads, it is recommended to use bead-immobilized nuclei for CUT&Tag. This circumvents undesirable tagmentation of mitochondrial DNA. However, cells are compatible with this protocol since digitonin remains in the buffers for cellular input flexibility and to control bead behavior. For more information on cells, see FAQ section.
4. Protocol optimized using 100,000 native (*i.e.* unfixed) human K562 nuclei per sample. However, without any further modifications, **this protocol has been validated on as few as 1,000 cells** with antibodies against H3K4me3, H3K27me3, and FOXA1.
5. PCR cycling recommendations: CUT&Tag PCR yields tend to vary by target and by cellular input amount. Rather than suggesting an arbitrary number of PCR cycles, we recommend determining the minimum number of PCR cycles that generate enough material to accurately quantify (*i.e.* by Qubit, Bioanalyzer, and/or TapeStation), which can be anywhere between 14-21 cycles. If a sample is overamplified, this can easily be corrected by bioinformatically removing PCR duplicates using [Picard](#).
6. The single best indicator of CUT&Tag success prior to sequencing is enrichment of ~300 bp fragments post-PCR, assessed by a Bioanalyzer or TapeStation.
7. Protocol optimized in 200 µL 8-strip PCR tubes to: **a)** minimize beads sticking to tubes; **b)** enable more rapid workflow with multichannel pipets; **c)** provide more consistent sample handling; and **d)** enable high-throughput sample preparation.
8. Take caution throughout to avoid ConA beads sticking to the tube sides and caps: the beads dry out easily, which can result in sample loss.
9. We recommend a 5% digitonin stock solution in DMSO (as opposed to heated H<sub>2</sub>O); this improves solubility and standardizes cell permeabilization / protocol reproducibility.
10. **IMPORTANT:** Since CUT&Tag has lower background and is compatible with low cell inputs, **it is not recommended to assess fragment size distribution by agarose gel or capillary electrophoresis prior to library preparation**. Such analysis is not indicative of the success of a CUT&Tag experiment, and further the amount of DNA recovered is often below the sensitivity of detection for each approach. Instead, assess DNA yield compared to positive (e.g. H3K4me3, H3K27me3) and negative (IgG) controls, determine fragment size distribution of sequence-ready libraries (**Figure 5**), and evaluate peak structure and expected genome-wide distribution in NGS data.

## 4. Buffers, Reagents & Materials Needed

**Buffer recipes** (note: see order information for buffer components, Section 6)

**Nuclear Extraction (NE) Buffer** (200 µL/sample)

20 mM HEPES–KOH, pH 7.9

10 mM KCl

0.1% Triton X-100

20% Glycerol

0.5 mM Spermidine

1x Roche cOmplete™, Mini, EDTA-free Protease Inhibitor (CPI-mini, 1tab/10mL)

*After spermidine and CPI are added, store @ 4°C for up to 1 week. NE buffer without spermidine and CPI is stable @ 4°C for up to 6 months*

**Bead Activation Buffer** (211 µL/sample)

20 mM HEPES, pH 7.9

10 mM KCl

1 mM CaCl<sub>2</sub>

1 mM MnCl<sub>2</sub>

*Filter sterilize*

*Store @ 4°C for up to 6 months*

**Wash150 Buffer** (use to prepare Digitonin150 Buffer)

20 mM HEPES, pH 7.5

150 mM NaCl

0.5 mM Spermidine

1x Roche cOmplete™, Mini, EDTA-free Protease Inhibitor (CPI-mini, 1tab/10mL)

*Filter sterilize*

*Store @ 4°C for up to 1 week*

**Digitonin150 Buffer** (450 µL/sample)

Wash150 Buffer + 0.01% Digitonin\*

*Prepare fresh each day and store @ 4°C*

**Antibody150 Buffer** (50 µL/sample)

Digitonin Buffer\* + 2 mM EDTA

*Prepare fresh each day and store @ 4°C*

**Wash300 Buffer** (use to prepare Digitonin300 and Tagmentation Buffers)

20 mM HEPES, pH 7.5

300 mM NaCl

0.5 mM Spermidine

1x Roche cOmplete™, Mini, EDTA-free Protease Inhibitor (CPI-mini, 1tab/10mL)

*Filter sterilize*

*Store @ 4°C for up to 1 week*

**Digitonin300 Buffer** (450  $\mu$ L/sample)  
Wash300 Buffer + 0.01% Digitonin\*  
*Prepare fresh each day and store @ 4°C*

**Tagmentation Buffer** (50  $\mu$ L/sample)  
Wash300 Buffer + 10 mM MgCl<sub>2</sub>  
*Store @ 4°C for up to 1 week*

**TAPS Buffer** (50  $\mu$ L/sample)  
10 mM TAPS, pH 8.5  
0.2 mM EDTA  
*Store @ RT for up to 6 months*

**SDS Release Buffer** (5  $\mu$ L/sample)  
10 mM TAPS, pH 8.5  
0.1% SDS  
*Store @ RT for up to 6 months*

**SDS Quench Buffer** (15  $\mu$ L/sample)  
0.67% Triton-X 100 in Molecular grade H<sub>2</sub>O  
*Store @ RT for up to 6 months*

*\*Digitonin 5% stock solution should be prepared in DMSO and aliquots stored at -20°C for six months. Note that digitonin is not necessary for nuclei permeabilization in the CUT&Tag workflow (purified nuclei are inherently permeable to antibody and pAG-Tn5). Rather, it helps to prevent the nuclei-conjugated beads from forming a thin film on tubes. Digitonin also makes the protocol compatible with cells if used, although not recommended.*

## Reagents

| Item  | Vendor                   | Catalog No.   | Notes  |
|---|--------------------------|---|--|
| Concanavalin A (ConA) Conjugated Paramagnetic Beads     | EpiCypher                | <a href="#">21-1401</a>                             | ConA is a lectin, which can cause immune activation. For technical support re. immune cell studies, contact <a href="mailto:info@epicypher.com">info@epicypher.com</a> .   |
| CUTANA® pAG-Tn5   | EpiCypher                | <a href="#">15-1017</a> and <a href="#">15-1117</a> | 50 & 250 reaction pack sizes available. Supplied as 20X stock.   |
| Rabbit IgG Negative Control Antibody                    | EpiCypher                | <a href="#">13-0042</a>                             | Use 0.5 µg in CUT&Tag  |
| SNAP-ChIP® Certified H3K4me3 Positive Control Antibody  | EpiCypher                | <a href="#">13-0041</a>                             | Use 0.5 µg in CUT&Tag  |
| SNAP-ChIP® Certified H3K27me3 Positive Control Antibody | EpiCypher                | <a href="#">13-0030</a>                             | Use 0.5 µg in CUT&Tag  |
| Antibody to histone PTM                                 | User-dependent           |   | EpiCypher continues to conduct <u>extensive</u> antibody characterization (most particularly for those against histone PTMs <sup>4</sup> ). Contact us for recommendations: <a href="mailto:info@epicypher.com">info@epicypher.com</a> |
| Secondary antibody (rabbit α-mouse immunoglobulin)      | Abcam                    | ab46540   | Required for CUT&Tag. Use with primary antibodies made in mouse.   |
| Secondary antibody (guinea pig α-rabbit immunoglobulin) | Antibodies Online        | ABIN101961  | Required for CUT&Tag. Use with primary antibodies made in rabbit.  |
| Agencourt AMPure XP magnetic beads                      | Beckman Coulter          | A63880  | For PCR cleanup and size selection ( <i>i.e.</i> adapter dimer and primer removal)   |
| Qubit™ 1x dsDNA HS Assay Kit                            | Thermo Fisher Scientific | Q33230  | For DNA quantification   |
| CUTANA® High-Fidelity 2X PCR Master Mix                 | EpiCypher                | <a href="#">15-1018</a>                             | Must be the <u>non-hot start</u> version of Q5 for CUT&Tag   |
| Universal i5 Primer                                     | IDT                      | NA  | Primer sequence provided below   |
| Uniquely Barcoded i7 Primers                            | IDT                      | NA  | Primer sequences provided below  |

**Universal i5 primer:** prepare at 10 µM in Molecular Biology grade H<sub>2</sub>O (RNase, DNase free)  
 5' AATGATACGGCGACCACCGAGATCTACACTAGATCGCTCGTCGGCAGCGTCAGATGTGTAT 3'

**Uniquely barcoded i7 primers:** prepare at 10 µM in Molecular Biology grade H<sub>2</sub>O (RNase, DNase free)

| Name  | Sequence (5' → 3')   | Barcode  | Illumina rev. compl. read |
|-------|--|----------|---------------------------|
| i7_1  | CAAGCAGAAGACGGCATAACGAGAT <b>TCGCCTTA</b> GTCTCGTGGGCTCGGAGATGTG | TCGCCTTA | TAAGGCGA                  |
| i7_2  | CAAGCAGAAGACGGCATAACGAGAT <b>CTAGTACG</b> GTCTCGTGGGCTCGGAGATGTG | CTAGTACG | CGTACTAG                  |
| i7_3  | CAAGCAGAAGACGGCATAACGAGAT <b>TTCTGCCT</b> GTCTCGTGGGCTCGGAGATGTG | TTCTGCCT | AGGCAGAA                  |
| i7_4  | CAAGCAGAAGACGGCATAACGAGAT <b>GCTCAGGA</b> GTCTCGTGGGCTCGGAGATGTG | GCTCAGGA | TCCTGAGC                  |
| i7_5  | CAAGCAGAAGACGGCATAACGAGAT <b>AGGAGTCC</b> GTCTCGTGGGCTCGGAGATGTG | AGGAGTCC | GGACTCCT                  |
| i7_6  | CAAGCAGAAGACGGCATAACGAGAT <b>CATGCCTA</b> GTCTCGTGGGCTCGGAGATGTG | CATGCCTA | TAGGCATG                  |
| i7_7  | CAAGCAGAAGACGGCATAACGAGAT <b>GTAGAGAG</b> GTCTCGTGGGCTCGGAGATGTG | GTAGAGAG | CTCTCTAC                  |
| i7_8  | CAAGCAGAAGACGGCATAACGAGAT <b>CCTCTCTG</b> GTCTCGTGGGCTCGGAGATGTG | CCTCTCTG | CAGAGAGG                  |
| i7_9  | CAAGCAGAAGACGGCATAACGAGAT <b>AGCGTAGC</b> GTCTCGTGGGCTCGGAGATGTG | AGCGTAGC | GCTACGCT                  |
| i7_10 | CAAGCAGAAGACGGCATAACGAGAT <b>CAGCCTCG</b> GTCTCGTGGGCTCGGAGATGTG | CAGCCTCG | CGAGGCTG                  |
| i7_11 | CAAGCAGAAGACGGCATAACGAGAT <b>TGCCTCTT</b> GTCTCGTGGGCTCGGAGATGTG | TGCCTCTT | AAGAGGCA                  |
| i7_12 | CAAGCAGAAGACGGCATAACGAGAT <b>TCCTCTAC</b> GTCTCGTGGGCTCGGAGATGTG | TCCTCTAC | GTAGAGGA                  |

## **Equipment**

| <b>Item</b>   | <b>Vendor</b>            | <b>Catalog No.</b>      | <b>Notes</b>   |
|---|--------------------------|-------------------------|--|
| 1.5 mL Magnetic Separation Rack                     | EpiCypher                | <a href="#">10-0012</a> | For batch processing ConA beads  |
| 8-strip PCR tube Magnetic Separation Rack           | EpiCypher                | <a href="#">10-0008</a> | For bead separation steps after batch processing ConA beads; enables multi-channel sample processing |
| 8-strip 0.2 mL PCR tubes                            | EpiCypher                | <a href="#">10-0009</a> | Compatible with 8-strip magnetic stand   |
| Qubit™ 4 Fluorometer                                | Thermo Fisher Scientific | Q33226                  | For DNA quantification   |
| Agilent 2100 Bioanalyzer                            | Agilent                  | G2939A                  | Or comparable capillary electrophoresis instrument (e.g. Agilent TapeStation)                        |
| High Performance Multi-Channel Pipettors, 8-Channel | VWR                      | 76169-252               | For aspiration and wash steps to accelerate workflow   |
| Tube Nutator  | VWR                      | 82007-202               | For bead incubation steps (overnight antibody incubation, pAG-Tn5 digest reaction)                   |
| Vortex-Genie  | Scientific Industries    | SI-0236                 | For bead mixing steps  |

## 5. EpiCypher CUTANA Direct-to-PCR CUT&Tag Protocol (~5hrs)

---Day 1---

### Section I: ConA Bead Activation (~30 min)

1. Gently resuspend the **ConA beads (Concanavalin A)** and transfer 11  $\mu\text{L}$ /sample to 1.5 mL tube for batch processing.
2. Place the tube on a 1.5 mL magnetic stand until slurry clears and pipet to remove supernatant (supe).
3. Remove from magnet and add 100  $\mu\text{L}$ /sample cold **Bead Activation Buffer** and pipet to mix. Place the tube on a magnet until slurry clears and pipet to remove supe.
4. Repeat previous step for total of two washes.
5. Resuspend beads in 11  $\mu\text{L}$ /sample cold **Bead Activation Buffer**.

Split activated ConA beads into separate tubes for different cell types and/or antibodies.

6. Aliquot 10  $\mu\text{L}$ /sample of activated bead slurry into 8-strip tube. Keep beads on ice until needed.

### Section II: Nuclei Preparation and Binding Nuclei to Activated Beads (~30 min)

7. Harvest 100,000 cells/sample by centrifugation for 3 min at 600 x g at room temperature (RT) in 1.5 mL tube, and decant supe.
8. Resuspend cells in 100  $\mu\text{L}$ /sample cold **NE Buffer**; and **incubate for 10 min on ice**.
9. Spin for 3 min at 600 x g at RT; aspirate supe.
10. Resuspend nuclei pellet in 100  $\mu\text{L}$ /sample cold **NE Buffer**.
11. Aliquot 100  $\mu\text{L}$  nuclei to each 8-strip tube containing 10  $\mu\text{L}$  of activated beads. Gently vortex (setting #7) to mix.
12. **Incubate** nuclei : bead slurry for **10 min at RT**. *Nuclei will adsorb to the activated ConA beads.*

### Section III: Binding of Primary and Secondary Antibodies (~30 min + overnight + 1 hr)

13. Place the tube on a magnet until slurry clears and pipet to remove supe.
14. Add 50  $\mu\text{L}$  cold **Antibody150 Buffer** to each sample, remove from magnet, and thoroughly pipet to resuspend.
15. Add 0.5  $\mu\text{g}$  **Primary Antibody** to each sample and gently vortex.



**Figure 2.** For ConA bead activation steps, it is recommended to batch process the full volume of beads needed for all experimental samples in a single 1.5 mL tube (11  $\mu\text{L}$  beads per sample). For these steps, beads can be washed using a 1.5 mL magnetic rack (e.g. EpiCypher 10-0012, pictured), ensuring homogeneity across samples.



**Figure 3.** For processing individual samples, it is recommended to use multi-channel pipetting using 8-strip PCR tubes and compatible magnetic rack (e.g. EpiCypher 10-0008, pictured), increasing experimental throughput and reproducibility.

16. **Incubate** 8-strip tube (**caps slightly elevated**) on nutator **overnight at 4°C** (**Figure 4**).



Bead slurry before o/n incubation @ 4°C | nutator o/n | Bead slurry after o/n incubation @ 4°C

**Figure 4.** Beads should appear homogenous and not clumpy throughout the procedure. Vortex or pipette as needed to disperse clumps. Ensure tube caps are elevated during incubation steps on tube nutator.

---Day 2---

17. Place the tube on a magnet until slurry clears and pipet to remove supe.
18. Remove from magnet, add 50  $\mu$ L cold **Digitonin150 Buffer** to each 8-strip tube, and thoroughly pipet to resuspend.
19. Add 0.5  $\mu$ g **Secondary Antibody** (matched to primary antibody animal source), and gently vortex.
20. **Incubate** 8-strip tube on nutator for **30 min at RT**.
21. Place the tube on a magnet until slurry clears and pipet to remove supe.
22. While beads are on magnet, add 200  $\mu$ L cold **Digitonin150 Buffer** directly onto beads of each sample, and then pipet to remove supe.
23. Repeat previous step for total of two washes.
24. Remove from magnet, add 50  $\mu$ L cold **Digitonin300 Buffer** to each 8-strip tube, and thoroughly pipet to resuspend. Beads are often clumpy at this point, but can easily be dispersed with gentle pipetting with a P200 pipet.
  - **Critical step:** 300 mM NaCl is critical in the Digitonin300 buffer to minimize non-specific binding of pAG-Tn5 to accessible DNA.

#### Section IV: Binding of pAG-Tn5 (~1 hr)

25. Add 2.5  $\mu$ L **CUTANA® pAG-Tn5** (20x stock) to each sample, and gently vortex.
  - **Critical step:** to evenly distribute pAG-Tn5 across nuclei, ensure beads are thoroughly resuspended by gentle vortexing and/or pipetting with a P200 pipet.
26. **Incubate** samples on nutator for **1 hr at RT**, return 8-strip tube to magnet, and pipet to remove supe.
27. Add 200  $\mu$ L cold **Digitonin300 Buffer** directly to each sample, **thoroughly resuspend by pipetting**, return to magnet, and then pipet to remove supe.
28. Repeat previous step for total of two washes.

#### Section V: Targeted Chromatin Tagmentation (~3 hrs)

29. Remove from magnet, add 50  $\mu$ L cold **Tagmentation Buffer** to each sample, and

thoroughly pipet to resuspend. Beads are often clumpy at this point, but can easily be dispersed by gentle pipetting with a P200 pipet.

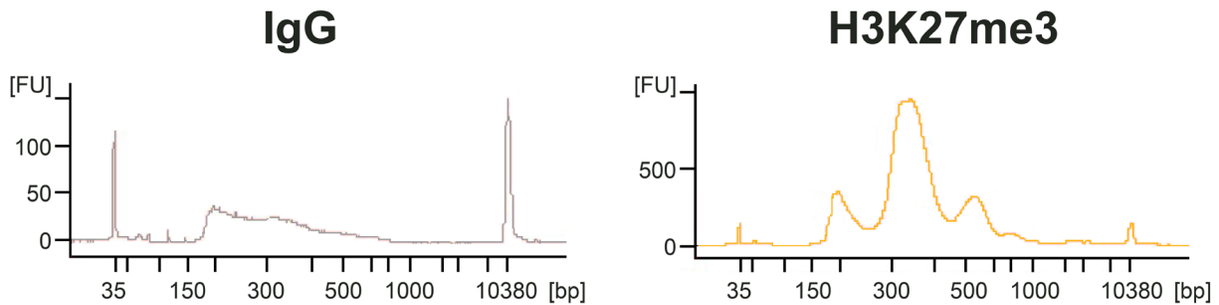
30. **Incubate** 8-strip tube for **1 hr at 37°C** in thermocycler. *Initiates tagmentation reaction.*
31. Place the tube on a magnet until slurry clears and pipet to remove supe.
32. Remove from magnet and resuspend beads in 50 µL RT **TAPS Buffer** by pipetting, return to magnet, and pipet to remove supe.
33. Remove from magnet, add 5 µL RT **SDS Release Buffer** (containing 0.1% SDS) to each sample, and vortex on max speed for 7 seconds. Quick spin to collect. *Quenches tagmentation reaction. Sample may be chunky and/or sticky, but this is normal since nuclei are being partially lysed.*
  - **Critical step:** Do not pipet to mix after adding SDS Release Buffer because it will likely lead to sample loss as the bead slurry is viscous and sticky.
34. **Incubate** 8-strip tube for **1 hr at 58°C** in thermocycler.
  - **Critical step:** Required to release tagmented chromatin fragments into solution, for both fixed and unfixed nuclei.
35. Add 15 µL RT **SDS Quench Buffer** (containing 0.67% Triton-X) to each sample, and vortex on max speed. *Neutralizes SDS, which potently inhibits PCR.*

#### Section VI: Non-hot Start PCR and Library Cleanup (~1 hr)

36. For each sample, add 2 µL each of universal i5 and barcoded i7 primers (10 µM stocks).
37. Add 25 µL **non-hot start CUTANA® High Fidelity 2x PCR Master Mix** to each sample and mix.
38. CUT&Tag-specific PCR cycling parameters:
  - a. 5 min @ 58°C → Fill-in step
  - b. 5 min @ 72°C → Extension step
  - c. 45 sec @ 98°C → DNA melting
  - d. 15 sec @ 98°C → DNA melting
  - e. 10 sec @ 60°C → hybrid primer annealing & short extension (<700 bp)
  - f. Repeat “step d-e” for a total of 14-21 cycles, depending on antibody target and cellular input amount. Use minimal number of cycles necessary to accurately quantify library.
  - g. 1 min @ 72°C → final extension
  - h. hold @ 4°C
  - **Critical step:** PCR cycling parameter is designed to amplify DNA fragments within the sequence-able range (100 bp-700 bp). Thus, contaminating higher molecular weight DNA fragments are not enriched by these PCR parameters.
39. DNA cleanup using **1.3x AMPure** beads to sample volume (e.g. 65 µL AMPure to 50 µL PCR; recovers > ~75 bp DNA fragments), as per manufacturer’s recommendations.
40. Elute DNA in 15 µL **0.1x TE buffer** and use 1 µL to quantify the purified PCR product using the **Qubit™ fluorometer** as per manufacturer's recommendations.

### Section VII: Agilent 2100 Bioanalyzer System (~1 hr)

41. For each purified PCR DNA library for Illumina sequencing, load 1  $\mu$ L of library sample on **Agilent High Sensitivity DNA Chip** (Cat# 5067-4626) as per manufacturer's recommendations.
42. Upon Bioanalyzer (or equivalent approach) confirmation that mononucleosome fragments were predominantly enriched (~300 bp with sequence adapters, **Figure 5**), proceed to Illumina sequencing as per manufacturer's recommendations.
  - **Note:** Only 3-5 million paired-end reads are needed for good coverage in CUT&Tag.



**Figure 5.** Typical Bioanalyzer traces for IgG negative control and H3K27me3 positive control CUT&Tag sequencing libraries (1  $\mu$ L PCR amplified / AMPure purified libraries loaded on Bioanalyzer). The positive control H3K27me3 library is predominantly enriched for mononucleosomes as indicated by the peak at ~300 bp (~150 bp mononucleosomes + 150 bp sequence adapters). Quality Bioanalyzer (or Tapestation / equivalent approach) traces are the best indicator of success prior to sequencing. However, this analysis should not be performed prior to NGS library preparation, since the amount of CUT&Tag DNA is likely to be below the limit of detection prior to library amplification.

## 6. Buffer components

| Components   | Source                   | Cat #            |
|--|--------------------------|------------------|
| HEPES  | Sigma-Aldrich            | H3375            |
| KCl  | Sigma-Aldrich            | P3911            |
| CaCl <sub>2</sub>  | Sigma-Aldrich            | C1016            |
| MnCl <sub>2</sub>  | Sigma-Aldrich            | 203734           |
| Molecular biology grade H <sub>2</sub> O (RNase, DNase free) | VWR                      | VWRV02-0201-0500 |
| NaCl   | Sigma-Aldrich            | S5150-1L         |
| EDTA (prepare 0.5 M stock at pH 8.0)                         | Sigma-Aldrich            | E5134            |
| Spermidine (Pure, 6.4M)                                      | Sigma-Aldrich            | S0266            |
| Digitonin  | Millipore Sigma          | 300410           |
| DMSO   | Sigma                    | D8418-100ml      |
| Trypan Blue  | Thermo Fisher Scientific | T10282           |
| cOmplete™, Mini, EDTA-free Protease Inhibitor Cocktail       | Roche                    | 11836170001      |
| Triton X-100   | Sigma-Aldrich            | X100             |
| Glycerol   | Millipore Sigma          | G5516            |
| 1 M TAPS, pH 8.5   | Boston Bioproducts       | BB-2375          |
| Sodium dodecyl sulfate (SDS)                                 | Sigma-Aldrich            | L4509            |

## 7. Frequently Asked Questions (FAQs)

### 1. What is the best way to know if a CUT&Tag experiment worked prior to sequencing?

Results from challenging cell inputs / targets may be ambiguous, so EpiCypher recommends including positive / negative controls (H3K4me3 or H3K27me3 / IgG) in every experiment. If QC checks for controls perform as expected (below), then proceeding to sequencing with all samples is recommended. If sequencing results for challenging cell inputs / targets are not satisfactory, further optimization may be necessary (e.g. cell type and / or number, antibody concentration, alternate antibodies, etc.).

First, including control antibodies (IgG and H3K4me3 or H3K27me3) in the experimental design is critical for assessing the success of a CUT&Tag experiment. In CUT&Tag, libraries are quality-controlled at two stages prior to sequencing: 1. PCR yields, and 2. Bioanalyzer DNA fragment distribution. PCR yields are typically lower than CUT&RUN, but the positive control antibody PCR yields are greater than IgG, particularly for high abundance PTMs (e.g. H3K27me3; H3K4me3 yields are often comparable or very slightly higher than IgG, since this is a lower abundance PTM). The expected PCR-amplified fragment distribution is predominantly mononucleosomes (~300 bp = 150 insert + 150 adapters). In short, prior to sequencing, enrichment of mononucleosome size fragments is the best indicator of success in CUT&Tag prior to sequencing.

### 2. Why are chromatin purification steps prior to PCR omitted from this protocol?

Direct-to-PCR CUT&Tag is a parsimonious workflow that offers many advantages over the chromatin purification option. In the direct approach, tagmented DNA is directly PCR amplified *in situ* to generate NGS libraries. The entire workflow from isolated nuclei to sequence-ready libraries can be performed in a single tube, enabling increased experimental throughput and supporting automation. By circumventing chromatin purification steps, sample loss is minimized, which increases sensitivity for low cell inputs.

### DNA spike-in controls

### 3. Can residual *E. coli* in the pAG-Tn5 prep be used for sample input normalization? What spike-in DNA control does EpiCypher recommend?

The EpiCypher pAG-Tn5 preparation is highly purified and depleted of contaminating nucleic acids, so residual *E. coli* DNA cannot be used for sample input normalization. The primary advantages of our optimized purification strategy are: 1. ensures lot-to-lot consistency; 2. maintains high specific activity of pAG-Tn5; and 3. prevents contaminating *E. coli* DNA from dominating signal in ultra-low cell input experiments. Nevertheless, EpiCypher is currently working on two spike-in solutions for CUT&Tag: exogenous DNA and fully defined semi-synthetic nucleosomes.

### 4. Does EpiCypher offer spike-in DNA controls for CUT&Tag?

EpiCypher plans to offer multiple spike-in controls for CUT&RUN and CUT&Tag, in addition to exogenous *E. coli* DNA spike-in. In particular, CUTANA® spike-in nucleosomes (*i.e.* fully-

defined semi-synthetic nucleosomes) compatible with each approach are in active development and offer multiple advantages, including enabling sample input normalization and multiple control points (1. antibody specificity; 2. antibody efficiency; 3. pAG-MNase /pAG-Tn5 activity; and 4. experimental success).

### Cell input compatibility

#### 5. Why are nuclei used in the default protocol?

Tn5 is known to efficiently tagmitochondrial DNA in the cytoplasm, which consumes sequencing bandwidth. Using nuclei in CUT&Tag circumvents this problem.

#### 6. What types of cell inputs are compatible with CUT&Tag?

While we do not recommend using whole cells (for the reasons listed in #5), these may be used in CUT&Tag with the caveat that a percentage of reads will be lost to mitochondrial DNA (degree of acceptability to be experimentally determined). We routinely use native nuclei derived from K562 cells in our CUT&Tag studies. If cells must be used, optimize the digitonin concentration to ensure cells are effectively permeabilized (e.g. perform a digitonin titration and monitor cell permeability using trypan blue staining).

#### 7. Is CUT&Tag compatible with frozen or crosslinked cells?

CUT&Tag is compatible with frozen and/or lightly crosslinked cells (*i.e.* 0.1% formaldehyde for 1 min). These protocol variations are currently under internal development and will be publicized when fully tested.

### Antibodies and targets

#### 8. Does EpiCypher's CUT&Tag protocol work on non-PTM targets?

Our internal R&D team has generated high quality CUT&Tag genomic profiles for the SWI/SNF chromatin remodeling complex and TFs (e.g. CTCF and FOXA1). In addition, Kaya-Okur et al. report several non-PTM targets in CUT&Tag (CTCF, SOX2, NPAT, and Pol II)<sup>1</sup>. However, because CUT&Tag is performed at twice the salt concentration of CUT&RUN and tagmentation is performed at 37°C, some TFs may be incompatible with CUT&Tag.

#### 9. What PTM antibodies does EpiCypher recommend for CUT&Tag?

We recommend starting with SNAP-certified PTM antibodies since they exhibit high specificity and efficiency in ChIP when tested against designer nucleosome panels (representing sets of related on- and off-target PTMs). In particular, we strongly recommend starting with positive control antibodies that have been verified to exhibit robust performance in CUT&Tag (EpiCypher H3K4me3 antibody, Catalog No. 13-0041 and EpiCypher H3K27me3 antibody, Catalog No. 13-0030). We will update our CUTANA® Compatible Antibodies product page as we continue to identify those best-in-class antibodies for CUT&Tag: <https://www.epicypher.com/antibodies/cutana-cut-run-compatible-antibodies/>.

Contact us directly for potential antibody recommendations to additional targets ([info@epicypher.com](mailto:info@epicypher.com)).

### Miscellaneous

10. Why is digitonin still used in the protocol if nuclei do not require digitonin for permeabilization? Digitonin is not necessary for nuclei permeabilization in the CUT&Tag workflow (purified nuclei are inherently permeable to antibody and pAG-Tn5). However, without digitonin, the bead-nuclei mixture tends to form a thin film on the side of the tubes, resulting in reduced yields. Digitonin also makes the protocol compatible with cells if used, although not recommended.

## 8. References

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