Nucleosome spike-in controls enable reliable next-generation epigenomic mapping



Andrea L. Johnstone¹, Danielle N. Maryanski¹, Keli L. Rodriguez¹, Ellen Weinzapfel¹, Bryan J. Venters¹, Katherine Novitzky¹, Matthew R. Marunde¹, Carolina P. Lin¹, Zachary B. Gillespie¹, Augustus Adeleke¹, Saarang Gopinath¹, Leslie Lewis¹, Marcus A. Cheek¹, Matthew J. Meiners¹, Zu-Wen Sun¹, Michael-Christopher Keogh¹

¹Epicypher Inc., Research Triangle Park, Durham NC 27709, USA

The Problem: Improved epigenomics assays are needed to advance cancer research

- > Epigenomic mapping for histone post-translational modifications (PTMs) can be used to discover gene regulatory mechanisms in cancer
- > ChIP-seq is the most widely used epigenomic mapping assay, but has major limitations:
 - X Depends on PTM antibodies which are notoriously cross-reactive ^{1,2}
 - X Requires large cell numbers unsuitable for clinical or rare cell samples
 - X Poor data quality low S:N, poor reproducibility
 - X Expensive and laborious making it difficult to scale
 - X Lacks defined controls crucial for reliable, quantitative results
- > Defined controls and improved methods are needed to fully leverage epigenomics in cancer research

Nucleosome spike-in controls show that most histone PTM antibodies fail in ChIP-seq

(A) Fully defined ChIP controls mimic physiological targets

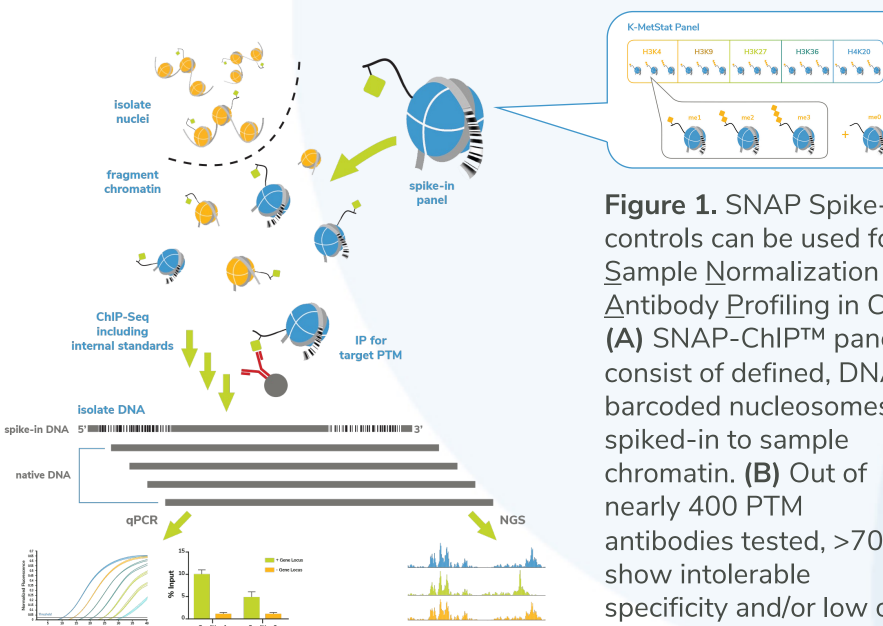


Figure 1. SNAP Spike-in controls can be used for Sample Normalization & Antibody Profiling in ChIP. (A) SNAP-ChIP™ panels consist of defined, DNAbarcoded nucleosomes spiked-in to sample chromatin. (B) Out of nearly 400 PTM antibodies tested, >70% show intolerable specificity and/or low ontarget efficiency.

(B) Most commercial antibodies are unfit for epigenomics

Antibodies Tested by SNAP-ChIP™	Lysine Methylation	Lysine Acylation	Total
Total Tested	263	129	392
Failure Rate	74.5%	64.5%	71.2%
\$ Spent	\$105,200	\$51,600	\$156,800
\$ Wasted	\$78,374	\$33,282	\$111,656
Top Cited : Citations w/"ChIP"	3,720 (N=18)	1,031 (N=6)	4,751 (N=24)
Top Cited : Failure Rate	74.5%	64.5%	79.2%

Data available at: www.ChromatinAntibodies.com³

CUTANA™ CUT&RUN has major advantages over ChIP

(A) CUT&RUN Workflow

(B) CUT&RUN vs. ChIP-seq

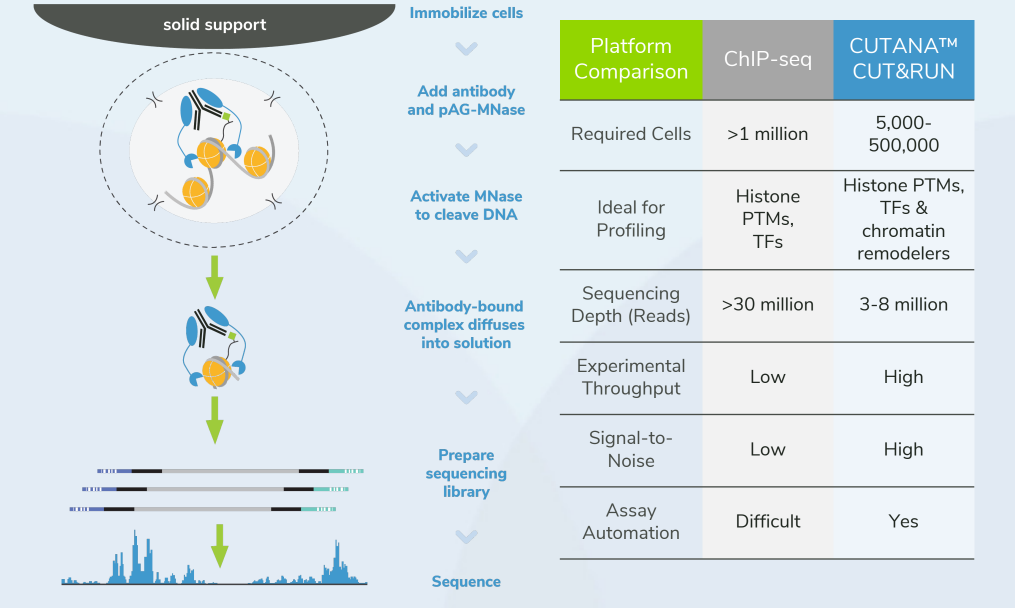
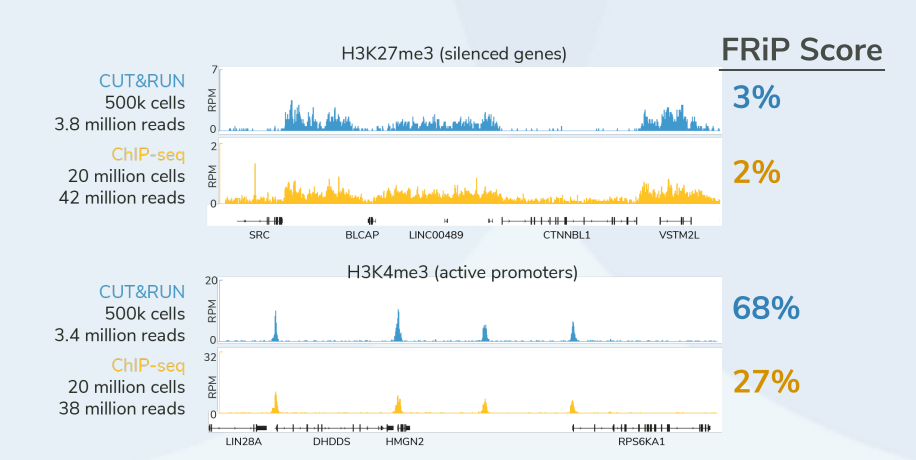


Figure 2. Overview of the CUTANA™ CUT&RUN workflow and advantages compared to ChIP-seq. Because CUT&RUN releases antibodybound fragments into solution (A), it has improved signal-to-noise even with significantly reduced cell numbers and sequencing depth (B).

CUT&RUN enables ultra-sensitive epigenomic profiling from low cell numbers

(A) CUT&RUN vs. ChIP-seq: better data, low sequencing costs



(B) CUT&RUN generates reliable profiles down to 5,000 cells

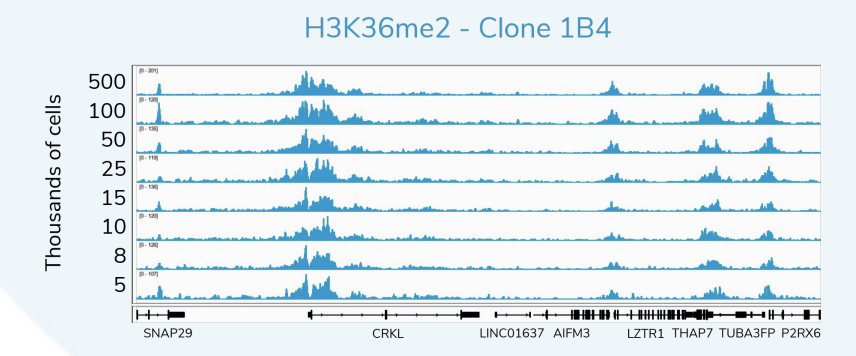
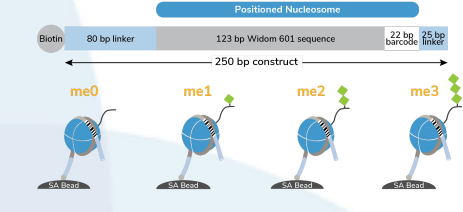


Figure 3. (A) The Fraction of Reads in Peaks (FRiP) score, a measure of signal-to-noise (S:N), is higher in CUT&RUN vs. ChIP-seq, despite using fewer cells and reduced sequencing depths. (B) Enrichment profiles are conserved when inputs are titrated from 500k to 5k cells.

SNAP-CUTANATM Spike-ins are critical controls for reliable chromatin mapping

(A) SNAP Spike-ins for CUT&RUN



(B) SNAP testing identifies specific antibodies for CUT&RUN

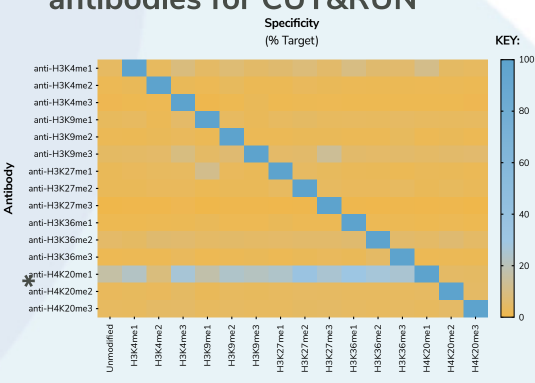
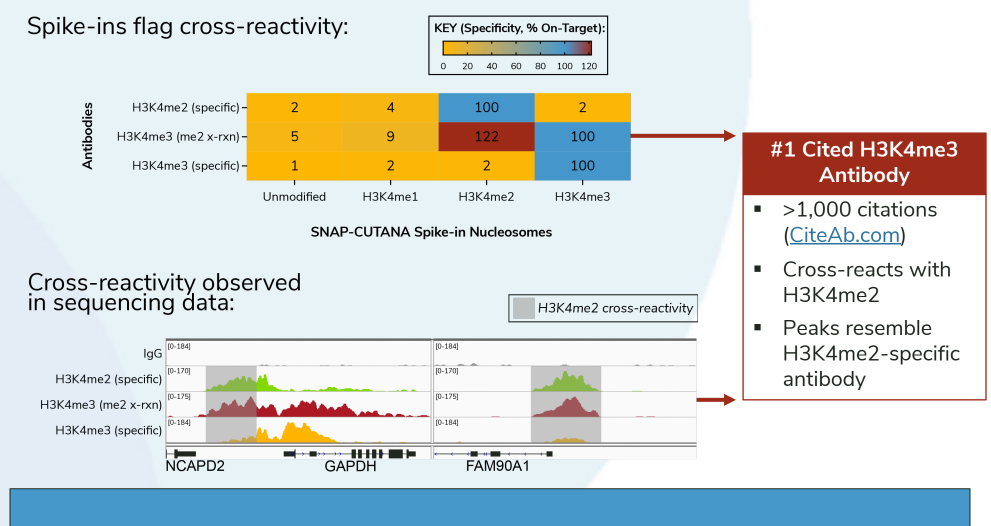


Figure 4. SNAP-CUTANA spike-in controls optimized for use in CUT&RUN (A) can be used to identify highly specific antibodies to widely studied histone PTMs (B). By spiking them into CUT&RUN workflows just prior to antibody addition, they provide a quantitative readout of onvs. off-target recovery that predicts non-specific peaks in genomic data (C). *Ongoing testing for H4K20me1. Inquire at info@epicypher.com for up-to-date information.

(C) SNAP Spike-ins predict non-specific recovery in CUT&RUN



Conclusions

- ➤ CUTANA™ CUT&RUN is poised to rapidly replace ChIP-seq
- > Our data illustrate how CUT&RUN could be applied for cancer epigenomics, particularly for low sample inputs and/or highthroughput applications
- > SNAP Spike-in controls address pervasive antibody specificity problems while enabling a direct readout of assay success and quantitative normalization (see below!)

References

- Shah et al. Examining the roles of H3K4 methylation states with systematically characterized antibodies. Mol. Cell 72, 162-177 (2018).
- 2. Small et al. Chromatin Immunoprecipitation (ChIP) to study DNA-Protein Interactions. Methods Mol. Biol. 2261, 323-343 (2021).
- 3. Maryanski et al. Novel nucleosome-based methods for rapid screening and identification of best-in-class antibodies: a community resource to improve genomic mapping. In preparation. www.ChromatinAntibodies.com

Ultra-sensitive epigenomics advances cancer research

Uncover novel biology with highly specific antibodies

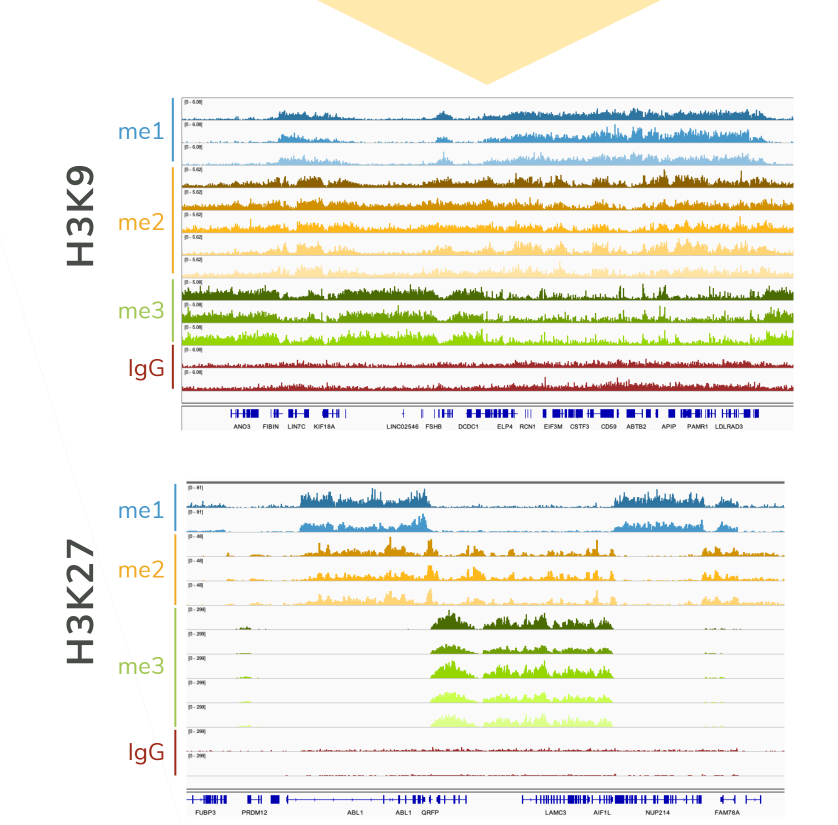


Figure 5. GENOMICALLY DISTINCT = FUNCTIONALLY DISTINCT. The use of highly specific histone PTM antibodies in CUT&RUN (Figure 4B) enables novel insights into the histone code. Distinct genomic profiles observed by mono-, di– and tri-methyl states imply distinct biological functions not previously appreciated.

Quantitative readout enables drug & clinical development applications

