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P Bench top CUT&RUN with antibodies-online™ CUT&RUN Sets v.5 ⊕

Forked from Bench top CUT&Tag

Antibodies Online Gmbh1

¹antibodies-online





ABSTRACT

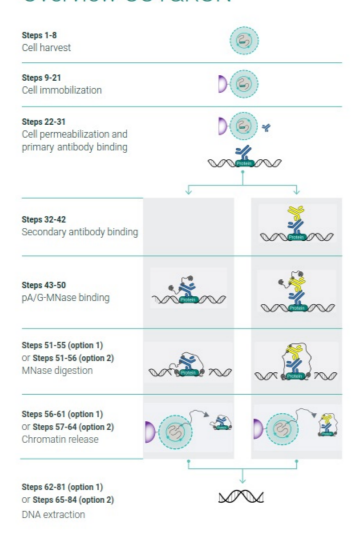
CUT&RUN (Cleavage Under Targets and Release Using Nuclease) offers a novel approach to pursue epigenetics. The method is designed to map genome wide transcription factor binding sites, chromatin-associated complexes, and histone variants and post-translational modifications.

Here we provide an updated CUT&RUN protocol that incorporates two variants, one is optimized to further reduce background noise which especially helps when working with low cell numbers and abundant antigens.



03/19/2020

Overview CUT&RUN



In CUT&RUN is performed in situ on immobilized, intact cells without crosslinking. DNA fragmentation is achieved using micrococcal nuclease that is fused to Protein A and/or Protein G (pA/G-MNase). The fusion protein is directed to the desired target through binding of the Protein A/G moiety to the Fc region of an antibody bound to the target. DNA under the target is subsequently cleaved and released and the pA/G-MNase-antibody-chromatin complex is free to diffuse out of the cell. DNA cleavage products are extracted and then processed by next generation sequencing (NGS).

All steps from live cells to sequencing-ready libraries can be performed in a single tube on the benchtop or a microwell in a high-throughput pipeline, and the entire procedure can be performed in one day.

EXTERNAL LINK

https://campaigns.antibodies-online.com/cutandrun-sets/?utm_source=protocolsio&utm_medium=protocol&utm_content=cutrun

THIS PROTOCOL ACCOMPANIES THE FOLLOWING PUBLICATION

 $\textbf{Citation:} \ \, \textbf{Antibodies Online Gmbh (03/19/2020)}. \ \, \textbf{Bench top CUT\&RUN with antibodies-online} \\ \tilde{\textbf{A}} \\ \hat{\textbf{A}} \\ \hat{\textbf{C}} \\ \textbf{CUT\&RUN Sets.} \\ \underline{\textbf{https://dx.doi.org/10.17504/protocols.io.bdwni7de}}$

The protocol is based on Skene; Henikoff JG; Henikoff S (2018): "Targeted in situ genome-wide profiling with high efficiency for low cell numbers"

GUIDELINES

Frequently Asked Questions

- Why is the DNA yield so low?
- How can I validate that my primary antibody is working for CUT&RUN?
- Why do I need a negative control antibody? Why not just use a no-antibody control?
- Can I replace the antibody negative control using a knock-out (or knock-down) of my protein?
- Do I need to use a secondary antibody? Other CUT&RUN protocols do not use a secondary.
- Should I include heterologous spike-in DNA for quantitation?
- Are there protocols to make this work for tissue, where I cannot necessarily isolate single cells?
- Is it possible to use the CUT&RUN product sets with plant tissue samples?
- Can CUT&RUN be adapted for RIP-seq?
- Is it possible to use a cell free extract of a bacterial cell instead of immobilizing and permeabilizing the cell?
- Instead of the proteinase K digestion can I denature the proteins in the CUT&RUN product complexes by heat?
- What is preferable for DNA extractions prior to library preparation: phenol-chloroform extraction or affinity purification using a column?

Why is the DNA yield so low?

CUT&RUN is performed using low cell numbers and the background signal is considerable lower than e.g. for ChIP. Due to these two factors the amount of recovered DNA is often times too low to be reliably measured based on a fluorometric assay or by capillary electrophoresis. PCR amplification of small CUT&RUN products, i.e. less than 50 bp, can be problematic and is therefore not any option. In order to assess the success of the CUT&RUN method each of the antibodies-online CUT&RUN product sets includes the CUT&RUN Positive Control antibody ABIN6923144•against the abundant H3K27me3 histone modification. DNA fragments prepared using this antibody can be measured by sensitive electrophoresis on a Bioanalyzer or Tapestation or fluorometrically on a Qubit or Nanodrop fluorometer. When using the CUT&RUN Positive Control antibody ABIN6923144•(or any other antibody specific for nucleosomal markers) a ladder corresponding to multiples of the 147 bp long nucleosomes should be visible by capillary electrophoresis.

How can I validate that my primary antibody is working for CUT&RUN?

For a CUT&RUN experiment the validation data could include e.g. a Tapestation or Bionalyzer plot showing the size distribution and qPCR data showing target enrichment.

As mentioned above, the DNA yield of an CUT&RUN experiment appears typically very low compared e.g. to ChIP-seq because of the lower initial sample size and the substantially lower DNA background. In particular for less abundant target protein the concentration is often times too low to be reliably measured using a fluorometric assay or by capillary

electrophoresis. PCR amplification of small CUT&RUN products, i.e. less than 50 bp, can be problematic and is therefore not an option. Once a sequencing library has been generated and sequenced map sequencing reads and verify the accumulation of reads at known binding sites.

Why do I need a negative control antibody? Why not just use a no-antibody control?

MNase is an endo- and exonuclease that will unspecifically bind and cleave unprotected DNA in hyper-accessible DNA, e.g. in regions surrounding regulatory elements. Free MNase will preferentially cut DNA within these hyper-accessible regions, thus potentially causing false positives and increased background signal in general.

In order to avoid this undesired effect of untethered MNase, the chromatin is randomly coated with the CUT&RUN Negative Control (ABIN6923140•) prior to the addition of pA/G-MNase is added to the samples. pA/G-MNase is then tethered via its Protein A or Protein G portion to the antbodies' Fc fragment and background DNA fragmentation is dictated by the random antibody binding as opposed to the nuclease digestion of hyper-accessible DNA regions.

Can I replace the antibody negative control using a knock-out (or knock-down) of my protein?

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Both controls are useful but address different aspects of the experiment and are therefore not interchangeable.

The CUT&RUN Negative Control (ABIN6923140) antibody is used to establish a reference background for peak calling. This is necessary because of the sparse background signal in CUT&RUN samples compared to ChIP-seq samples. The ko (or kd) control on the other hand gives an impression of unspecific binding of the antibody specific for the prote-

in of interest to other proteins. It is useful to avoid identification of false positive signals.

Do I need to use a secondary antibody? Other CUT&RUN protocols do not use a secondary.

Depending on the host species and isotype of the antibody and the Protein A and/or Protein G MNase fusion protein, a secondary antibody may be necessary for pA/G-MNase binding.

Protein A has good high affinity to all rabbit IgG antibodies but low affinity to rat, goat and sheep IgG isotype antibodies and certain mouse IgG antibody subclasses, in particular IgG1. Protein G on the other hand binds well to the Fc region of mouse, goat, sheep, and most rat IgG. Its affinity to rabbit IgG however is lower than that of Protein A. When using pAG-MNase introduced with the improved CUT&RUN protocol it is therefore generally not necessary to use a secondary antibody. Use of the pA-MNase of the original protocol however might require the use of a secondary antibody raised in rabbit to assure efficient binding of the fusion protein to the antibody.

Should I include heterologous spike-in DNA for quantitation?

Our protocol is largely based on the improved CUT&RUN protocol. Here, the authors show that accurate quantitation is possible using heterologous spike-in DNA or carry-over *E. coli* DNA from the pA/G-MNase purification.

Are there protocols to make this work for tissue, where I cannot necessarily isolate single cells?

The Henikoff lab that published the original CUT&RUN method in 2017 has since developed several variations. One of these modification is AutoCUT&RUN which outlines a workflow to automate CUT&RUN for high-throughput characterization of single cells but also patient samples and xenografts.

In short, the tissue is processed manually or enyzmatically into single, intact cell that are subsequently bound to ConA beads. The improved (and standard) CUT&RUN protocol that serve as basis for the protocol outlined in our CUT&RUN handbook can be adapted accordingly. Presently there is no work published documenting the use of intact tissue.

Is it possible to use the CUT&RUN product sets with plant tissue samples?

The CUT&RUN method can be applied without a problem to plant tissue samples (see e.g. <u>PMID30719569</u>). One essential step in addition to those lined out in the protocol is the generation of spheroblasts so that it becomes possible to permeabilize the plasma membrane for the application of the antibodies and the MNase fusion protein.

The positive control H3K27me3 antibody and the negative control guinea pig anti-rabbit IgG antibody as well as the ConA beads are suitable for use with plant samples. The antibodies that are included in some of the <u>sets</u>, such as the anti-DYKDDDDK antibodies or the secondary antibodies, can also be used with plant samples. Whether they are needed or not depends on your experiment.

Can CUT&RUN be adapted for RIP-seq?

It should be possible to adapt the CUT&RUN protocol for use with RNA as an alternative to RIP-seq. RNA in the cytoplasm will attract the degradation machinery if it is lacking the 5' cap and the 3' poly-A tail. Thus, it is advisable to use isolated nuclei as sample material. Consequently, no digitonin in the different buffers is needed since the nuclear envelope does not contain cholesterol. Isolated nuclei may be immobilized via glycoproteins on the nuclear envelope to the ConA beads like for a CUT&RUN experiment. An antibody against the protein of interest is added and subsequently the pA/G-MNase is tethered to the antibody, thus bringing the MNase into proximity of the RNA of interest. Similar to RIP-seq, isolated RNA is then translated into cDNA and can be sequenced and mapped.

Is it possible to use a cell free extract of a bacterial cell instead of immobilizing and permeabilizing the cell?

It should indeed be possible to use a bacterial extract as starting material instead of immobilized eukaryotic cells. There are a couple of things that should be considered:

One of the factors that makes CUT&RUN so much better than ChIP-seq in a eukaryotic system is that it is done in situ. The intact cells (or isolated nuclei) retain the large, undigested chromatin. Consequently, much fewer off-sites are present in the prepared DNA which is one of the reasons why CUT&RUN data sets have so much less background signal when compared to ChIP-seq data. This enrichment step will be missing when using a cell lysate as starting material. The CUT&RUN result should still be superior to the CHIP-seq result for the same starting material. However, it will be more "noisy" than CUT&RUN sequencing data from intact eukaryotic cells/nuclei. The buffers described in the CUT&RUN protocol are designed to maintain the cells in the samples intact. The binding buffer contains some divalent cations that are necessary for the binding of the cells to the Concanavalin on the beads that are used for the immobilization. Subsequently, this buffer is changed for an antibody buffer that contains EDTA to chelate the divalent cations and

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prevent DNA cleavage prior to pA/G-MNase antibody binding. Since you are working with cell lysates as opposed to intact cells there is no need to include digitonin in any of the buffers. You don't need the binding buffer used to immobilize cells on the ConA beads. In order to control premature DNA cleavage by the MNase I would include some EDTA and/or EGTA in the lysis buffer.

Probably, you will have to titrate the amount of pA/G-MNase and try different digestion times to find the best balance between efficient cleavage of the intended sites and too much off-site cleavage.

The original CUT&RUN protocol foresees the use of heterologous spike-in DNA, e.g. from *E. coli*. In its latest iteration it is argued that there is no need to add this DNA since the pA/G-MNase preparation contains sufficient levels of *E. coli* DNA to serve as a standard for quantitation. In a prokaryotic system, spike-in DNA from a eukaryotic organism (e.g. *S. cerevisiae*) is recommended. Also keep in in mind that the *E. coli* DNA carried over with the pA/G-MNase preparation may contain sequences that show homology to the bindings sites of your protein of interest.

Our product sets are optimized for use with eukaryotic cells. The positive control in all sets is a recombinant rabbit anti-H3K27me3 antibody. This control is not suitable for use with prokaryotic samples.

Instead of the proteinase K digestion can I denature the proteins in the CUT&RUN product complexes by heat?

Proteinase K digest followed by a phenol-chloroform extraction has historically been the go-to method to prepare high-molecular weight DNA, primarily from tissues. Subsequently to the diffusion of the CUT&RUN products out of the cells the DNA is present already in a relatively clean form, in particular when compared to a cell lysate like in ChIP-seq. Therefore, it is reasonable to assume that it is possible to forgo the Proteinase K treatment of the cells and instead proceed with the PCI extraction. As a reference, nucleic acid preparation using TRIzol does not include a Proteinase K digest. However, it has also been reported, that a proteinase K digest improves subsequent PCR amplification. Regarding heat inactivation instead of proteinase K digest in step, we recommend againstthis option. Your DNA of interest is at his point present in a complex consisting of the DNA, your antigen, the corresponding antibody, and the pA/G-MNase. If at this stage you boil your sample denature proteins you risk precipitation of your DNA with the protein compounds in this complex upon denaturation. This will also primarily affect the short CUT&RUN products, thus decreasing the signal to noise ratio in your library and potentially also reducing the library's complexity.

What is preferable for DNA extractions prior to library preparation: phenol-chloroform extraction or affinity purification using a column?

In the original publication describing the CUT&Tag method the authors mention the use of AMPure XP beads for the purification of the DNA subsequently to tagmentation and Proteinase K digest. A potential issue is the carry-over of active Proteinase K, which can interfere with the downstream PCR amplification. Therefore, the authors recommend now the phenol-chloroform extraction to assure complete denaturation of Proteinase K.

MATERIALS

NAME Y	CATALOG #	VENDOR ~
antibodies-online™ CUT&RUN Pro Complete Set	ABIN6923135	antibodies-online
antibodies-online™ CUT&RUN Negative Control	ABIN6923140	antibodies-online
antibodies-online™ CUT&RUN Positive Control	ABIN6923144	antibodies-online
antibodies-online™ CUT&RUN Concanavalin A Beads	ABIN6923139	antibodies-online
antibodies-online™ CUT&RUN Secondary	ABIN6923141	antibodies-online
0.5 M EDTA pH 8.0	ABIN925554	antibodies-online
Trypan Blue (0.4%)	ABIN413910	antibodies-online
Proteinase K	ABIN920948	antibodies-online
CUT&RUN Pro Sec Set	ABIN6923137	antibodies-online
Spike-in Chromatin	ABIN4889666	antibodies-online
CUTANA™ pAG-MNase for ChIC/CUT&RUN Assays	ABIN6950951	antibodies-online

STEPS MATERIALS

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Citation: Antibodies Online Gmbh (03/19/2020). Bench top CUT&RUN with antibodies-online⢠CUT&RUN Sets. https://dx.doi.org/10.17504/protocols.io.bdwni7de

NAME ~	CATALOG #	VENDOR V
antibodies-online™ CUT&RUN Concanavalin A Beads	ABIN6923139	antibodies-online
antibodies-online™ CUT&RUN Positive Control	ABIN6923144	antibodies-online
antibodies-online™ CUT&RUN Negative Control	ABIN6923140	antibodies-online
antibodies-online™ CUT&RUN Secondary	ABIN6923141	antibodies-online
CUTANA™ pAG-MNase for ChIC/CUT&RUN Assays	ABIN6950951	antibodies-online

MATERIALS TEXT

Reagents required

CUT&RUN Pro Sec Set (antibodies-online, cat no.ABIN6923137) with following components:

- Positive control Recombinant Rabbit anti-H3K27me3 Antibody (ABIN6923144)
- Negative control Polyclonal Guinea Pig anti-Rabbit IgG Antibody (ABIN6923140)
- Concanavalin A Beads (ABIN6923139)
- Secondary Rabbit anti-Mouse IgG (H&L) Antibody (ABIN6923141)
- Antibody to an epitope of interest
- Distilled, deionized or RNAse-free H2O
- 1 M Manganese Chloride (MnCl₂)
- 1 M Calcium Chloride (CaCl₂)
- 1 M Potassium Chloride
- 1 M HEPES pH 7.5 HEPES (NaOH)
- 5 M NaCl
- 0.5 M EDTA (ABIN925554)
- 0.2 M EGTA
- 2 M Spermidine
- Protease Inhibitor Cocktail
- 5% Digitonin (ABIN1304051)
- 20 mg/ml Glycogen
- Trypan Blue (ABIN413910)
- RNase A (DNase and protease free)
- 10% Sodium dodecyl sulfate (SDS)
- Proteinase K (ABIN920948)
- Phenol-chloroform-isoamyl alcohol (PCI)
- 7.5 M Ammonium Acetate (NH₄OAc)
- 70 % Ethanol
- 100% Ethanol
- 1 mM Tris-HCl pH 8.0

Materials not provided with the package

CUTANA™ pAG-MNase for ChIC/CUT&RUN Assays ABIN6950951/ABIN6950952 Spike-in DNA 200 bp mean size (e.g, D. melanogaster, ABIN4889666)

SAFETY WARNINGS

Storage

Short term storage of the antibodies for up to two weeks at 4 °C. For long term storage for up to one year at -20 °C, prepare 20 μ L aliquots of the antibodies to avoid repeated freeze-thaw-cycles.

CUT&RUN Concanavalin A Beads must be stored at 4C. Do not freeze CUT&RUN

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BEFORE STARTING

General remarks

• The original CUT&RUN protocol in Skene et al. (2018)(Skene, PJ; Henikoff JG; Henikoff, 2018) recommends sample sizes of 100 to 1000 mammalian cells for abundant antigens such as H3K27me3 or CTCF. This protocol adapted from Meers et al. (2019)(Meers, Bryson, et al., 2019) is suitable for up to 500.000 cells.

This protocol is intended to give a general outline of the CUT&RUN protocol. It has to be adjusted according to the:

- » Cell type. Your specific cell type might necessitate different treatments prior to the CUT&RUN procedure, e.g. disintegration of tissue, generation of spheroblasts,
- » MNase digestion time points during the optimization. Different samples, approaches, and digestion time points are uniformly referred to in the protocol as "samples".
- To minimize DNA breakage during sample preparation, avoid cavitation through vigorous resuspension and vigorous vortexing.
- Keep cells at room temperature during all steps prior to the addition of antibody to minimize stress on the cells and DNA breakage.
- ${}^{\bullet}$ All steps from the incubation with the primary antibodies on should be carried out at 4°C.

Antibody selection

An antibody specific for your protein of interest is crucial to direct the pA/G-MNase mediated nucleic acid cleavage to the intended site. The Protein A/G portion tethers the fusion protein to the Fc region of the antibody bound to its antigen. This allows the pA/G-Mnase nuclease portion to cleave the nucleic acid under the targeted protein and to release the nucleic acid.

Depending on the host species and isotype of the antibody and the Protein A and/or Protein G MNase fusion protein, it can be necessary to include a secondary antibody for pA/G-MNase binding (Skene & Henikoff, 2017). If the pA-MNase is used in conjunction e.g. with a primary mouse IgG1 or goat IgG antibody it is recommended to use a rabbit secondary antibody (Section V). Protein A binds well to rabbit or guinea pig IgG antibodies but only poorly to mouse IgG1 or goat IgG. No additional secondary antibody is needed when using pA/G-MNase (Meers, Bryson, et al., 2019).

The positive Control H3K27me3 antibody (ABIN6923144) and Negative Control Guinea Pig anti-Rabbit IgG antibody (ABIN6923140) are important to assess cleavage and chromatin release without the need to sequence the released DNA fragments. Do not use a no-antibody negative control: untethered pA/G-MNase will unspecifically bind and cleave any accessible DNA, thus increasing background signal.

REAGENT SETUP (for 8 samples)

1

» Wash buffer (110 mL)

Component	Volume	Final concentration
ddH2O	103 mL	-
1 M HEPES pH 7.5	2.2 mL	20 mM
5 M NaCl	3.3 mL	150 mM
2 M Spermidine	27.5 μL	0.5 mM

- Store Wash Buffer without protease inhibitors for up to one week at 4 °C.
- Add protease inhibitors fresh before use, e.g.:
- 1.1 mL Protease Inhibitor Cocktail 100x

» Binding Buffer (30 mL)

Component	Volume	Final concentration
ddH2O	29 mL	-
1 M HEPES pH 7.5	600 μL	20 mM
1 M KCl	300 µL	10 mM
1 M CaCl2	30 µL	1 mM
2.5 M MnCl2	12 μL	1 mM

[•] Store Binding Buffer for up to six months at 4 °C.

» Digitonin Wash Buffer (55 mL)

Component	Volume	Final concentration
5 % Digitonin	550 μL	0.05 %
Wash Buffer	54.5 mL	-

- Store Digitonin Wash Buffer for up to one day at 4 °C.
- Recommended Digitonin concentration ranges from 0.025% to 0.1%.
- The effectiveness of Digitonin varies between batches. Test cell permeability using Trypan Blue to determine the optimal concentration to use.

» Antibody Buffer (1 mL)

Component	Volume	Final concentration
0.5 M EDTA	4 μL	2 mM
10 % BSA	10 μL	0.1 %
Digitonin Wash Buffer	986 μL	-

[•] Store Antibody Buffer for up to one day at 4 °C until use.

» Low Salt Rinse Buffer (18 mL)

Component	Volume	Final concentration
ddH2O	17.5 mL	-
1 M HEPES pH 7.5	360 µL	20 mM
2 M Spermidine	4,5 µL	0.5 mM
5% Digitonin	180 µL	0.05 %

[•] Store Low Salt Rinse Buffer for up to one week at 4 °C until use.

» Low Salt Incubation Buffer (2 mL)

Component	Volume	Final concentration
ddH2O	1953 μL	-
1 M HEPES pH 7.5	7 μL	3.5 mM
1 M CaCl2	20 μL	10 mM
5% Digitonin	20 μL	0.05%

 $[\]bullet$ Store Low Salt Incubation Buffer for up to one week at 4 $^{\circ}\text{C}$ until use.

» Low Salt Stop Buffer (2 mL)

Component	Volume	Final concentration
ddH2O	1700 mL	-
5 M NaCl	68 µL	170 mM
0.2 M EGTA	200 μL	20 mM
Store Low Salt Stop Buffer at 4 °C unt	il use.	
Add fresh before use:		
5% Digitonin	20 μL	0.05%
RNase A (10 mg/mL)	10 μL	50 μg/mL
Glycogen (20 mg/mL)	2.5 µL	25 μg/mL
Optional:		
heterologous spike-in DNA	-	100 pg/mL

I.Cell Harvest – at room temperature

2	Harvest 10 000 to 500 000 cells for each sample at	# Room temperature	Keen cells for each sample in separate tubes

3 Centrif	age cell solution	© 00:03:00	at :: 600 x c	at	Room temperature	. Remove the lic	quid carefully	
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4	Gently resuspend cells in \Box 1 ml Wash Buffer by pipetting and transfer cell solution to a \Box 1.	5 ml
	microcentrifuge tube	

5	Centrifuge cell solution	@00:03:00	at @600 x g	at	A Room temperature	and discard the supernatant

6	Repeat steps 4-5 thrice for a total of four washes	. 🐧	go t	o ste	p #4	4
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⁷ Resuspend cell pellet for each sample in **1 ml Wash Buffer** by gently pipetting.

II. Concanavalin A beads preparation 8 Prepare one 1.5 ml microcentrifuge tube for each sample. Gently resuspend the antibodies-online™ CUT&RUN 10 Pipette 10 µl CUT&RUN Concanavalin A Beads slurry for each Concanavalin A Beads sample into the 1.5 ml microcentrifuge tubes. by antibodies-online Catalog #: ABIN6923139 Place the tubes on a magnet stand until the fluid is clear. Remove the 11 liquid carefully. Remove the microcentrifuge tube from the magnet stand. 12 13 Pipette 1 ml Binding Buffer into each tube and resuspend CUT&RUN Concanavalin A Beads by gentle pipetting. 14 Spin down the liquid from the lid with a quick pulse in a table-top centrifuge (max 100 x g). Place the tubes on a magnet stand until the fluid is clear. Remove the liquid carefully. 15 Remove the microcentrifuge tube from the magnet stand. 16 17 Repeat steps 13-16 **go to step #13** twice for a total of three washes. Gently resuspend the CUT&RUN Concanavalin A Beads in a volume of Binding Buffer corresponding to the original 18 volume of bead slurry, i.e. 10 μl per sample. III. Cell immobilization – binding to Concanavalin A beads 19 Carefully vortex the cell suspension from step ⊕ go to step #7 and add □10 µl of the CUT&RUN Concanavalin A Beads in Binding Buffer prepared in section II to each sample. 20 Close tubes tightly and rotate for © 00:05:00 to © 00:10:00 at & Room temperature. IV. Cell permeabilization and primary antibody binding Place the microcentrifuge tubes on a magnet stand until the fluid is clear. Remove the liquid carefully. Remove the microcentrifuge tubes from the magnet stand.

- Place each tube at a low angle on the vortex mixer set to a low speed (approximately 31100 rpm) and add $\blacksquare 100 \mu l$ Antibody Buffer containing digitonin.
- 24 Gently vortex the microcentrifuge tubes until the beads are resuspended.
- 25 Add **11 μl primary antibody** against your antigen of interest, the



, and the



- corresponding to a 1:100 dilution (or a volume corresponding to the manufacturer's recom- mended dilution for immunofluorescence).
- Rotate the microcentrifuge tubes for © 00:05:00 to © 00:10:00 at & Room temperature or © 02:00:00 to © Overnight at & 4 °C.
- 27 Spin down the liquid and place the tubes on a magnet stand until the fluid is clear. Remove the liquid carefully.
- 28 Remove the microcentrifuge tubes from the magnet stand.
- Resuspend with **1 ml Digitonin Wash Buffer** and mix by inversion. If clumping occurs, gently remove the clumps with a **1 ml** pipette tip.
- 30 Repeat steps 27-29 **go to step #27** once for a total of two washes.
 - If no secondary antibody is used proceed directly to section VI. pA/G-MNase binding (Step 41).

V. Secondary antibody binding (optional)

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is raised in rabbit against mouse IgG (H&L). In case a **primary antibody** from a different host species or isotype is used a different **secondary** must be selected accordingly.

Place the tubes on a magnet stand until the fluid is clear. Remove the liquid carefully.

- 32 Remove the microcentrifuge tubes from the magnet stand.
- Vortex the sample at low speed (approximately 31100 rpm) and add $\Box 100 \mu l$ Digitonin Wash Buffer per sample along the side of the tube.
- 34 Tap to remove the remaining beads from the tube side.
- 35 Add **□1 μl**



corresponding to a 1:100 dilution (or a volume corresponding to the manufacturer's recommended dilution for immunofluorescence).

- 36 Rotate the microcentrifuge tubes for \circlearrowleft 01:00:00 at ~& 4 °C .
- 37 Spin down the liquid and place the tubes on a magnet stand until the fluid is clear. Remove the liquid carefully.
- 38 Remove the microcentrifuge tubes from the magnet stand.
- Resuspend with **1 ml Digitonin Wash Buffe**r and mix by inversion. If clumping occurs, gently remove the clumps with a 1 mL pipette tip.

40 Repeat steps 37-39 **go to step #33** once for a total of two washes.

VI. Protein A-MNase or Protein AG-MNase Binding

- 41 Place the tubes on a magnet stand until the fluid is clear. Remove the liquid carefully.
- 42 Remove the microcentrifuge tubes from the magnetic stand.
- 43 Vortex the sample at low speed (approximately **§ 1100 rpm**) and add **50 μl Digitonin Wash Buffer** per sample along the side of the tube. Add **2.5 μl**



Alternatively:

Vortex the sample at low speed (approximately **(3)** 1100 rpm) and add **150 μl Digitonin Wash Buffer** containing **700 ng/mL** of your own **pA/G-MNase** preparation per sample along the side of the tube.

- Rotate the microcentrifuge tubes for © 01:00:00 at § 4 °C.
- 45 Spin down the liquid and place the tubes on a magnet stand until the fluid is clear. Remove the liquid carefully.
- 46 Remove the microcentrifuge tubes from the magnetic stand.
- 47 Resuspend with **1 ml Digitonin Wash Buffer** and mix by inversion. If clumping occurs, gently remove the clumps with a 1 ml pipette tip.
- 48 Repeat steps 45-47 **go to step #45** once for a total of two washes.

VII. MNase digestion and release of pA/G bound chromatin fragments

- Spin down the liquid from the lid with a quick pulse in a table-top centrifuge (max \bigcirc 100 x q).
- 50 Place the tubes on a magnet stand until the fluid is clear. Remove the liquid carefully.
- Resuspend with **1 ml** Low Salt Rinse Buffer and mix by inversion. If clumping occurs, gently remove the clumps with a 1 mL pipette tip.

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03/19/2020

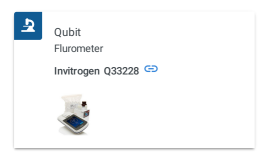
Spin down the liquid from the lid with a quick pulse in a table-top centrifuge (max @100 x g). Place the tubes on a magnet stand until the fluid is clear. Remove the liquid carefully. 53 54 Repeat steps 51-53 **go to step #51** once for a total of two washes. 55 Place each tube at a low angle on the vortex mixer set to a low speed (approx. ■1100 rpm) and add ■200 µl ice cold Low Salt Incubation Buffer per sample along the side of the tube. 56 Incubate tubes at § 0 °C for the desired time (default is © 00:05:00). Place the tubes on a cold magnet stand until the fluid is clear. Remove the liquid carefully. 57 Remove the microcentrifuge tubes from the magnet stand. 58 59 Resuspend with 200 µl Low Salt Stop Solution and mix by gentle vortexing. 60 Incubate tubes at § 37 °C for © 00:30:00. Place the tubes on a magnet stand until the fluid is clear. 62 Transfer the supernatant containing the pA/G-MNase-bound digested chromatin fragments to fresh \subseteq 1.5 ml microcentrifuge tubes. VIII. DNA extraction 63 Add 2 4 10% SDS to a final concentration of 0.1% and 5 4 Proteinase K ([M]10 mg/mL) to a final concentration of [M]2.5 mg/mL to each supernatant from step 62. 64 Gently vortex tubes at a low speed of approximately **31100 rpm**. 65 Incubate tubes at § 50 °C for © 01:00:00 or at § 37 °C © Overnight 66 Add 200 µl PCI to tube. Vortex tubes thoroughly at high speed until the liquid appears milky.

Optional: Transfer liquid to a phase-lock tube. 68 69 Centrifuge tubes in a table-top centrifuge at $\$ 16000 x g at $\$ 4 °C for $\$ 00:05:00 . 70 Carefully transfer the upper aqueous phase to a fresh ☐1.5 ml microcentrifuge tube containing ☐200 µl Chloroform:Isoamyl Alcohol 24:1. 71 Vortex tubes thoroughly at high speed until the liquid appears milky. 72 Centrifuge tubes in a table-top centrifuge at \$\infty\$16000 x g at \$4 °C for \$\infty\$00:05:00. 73 Carefully transfer the upper aqueous phase to a fresh **1.5 ml** microcentrifuge tube containing □2 µI glycogen (diluted 1:10 to □2 mg/mL from the □20 mg/mL stock solution). 74 Add 20 µl [M]3 Molarity (M) NaOAc or 2100 µl [M]5 Molarity (M) NH₄OAc 75 Add **500 μl** 100% ethanol. Place tubes for © 00:10:00 in a dry ice/Ethanol mix or © Overnight at & -20 °C. 77 Centrifuge tubes in a table-top centrifuge at @16000 x g at & 4 °C for © 00:05:00. Remove the liquid carefully with a pipette. 78 79 Add 11 ml 70% ethanol. 80 Centrifuge tubes in a table-top centrifuge at @16000 x g at & 4 °C for © 00:01:00. 81 Remove the liquid carefully with a pipette. Air-dry the pellet or dry the pellet in a SpeedVac. 82

Dissolve the pellet in 30 µl [M]1 Milimolar (mM) Tris-HCl, [M]0.1 Milimolar (mM) EDTA.

IX. Sample quality control

84 Size distribution and concentration of the CUT&RUN products can be assessed at this point, e.g. using a



or



or a Bioanalyzer or Tapestation. It is possible that the concentration of the recovered DNA is below the instrument's detection limit. It is also to be expected that the extracted DNA includes some large DNA fragments that will mask the signal of the CUT&RUN products. In this case it may be useful to PCR-amplify the DNA and check the library on a Bioanalyzer or Tapestation.

X. Sequencing library preparation

Prepare the CUT&RUN products sequencing libraries according to your established workflow. Because of the very low background with CUT&RUN, typically 5 million paired-end reads suffice for epitopes with a multitude of genomic binding sites, e.g. transcription factors or nucleosome modifications.

XI. Peak calling

The sparse background signal in CUT&RUN samples compared to ChIP-seq samples represents challenge for peak callers that employ statistical models relying on a high sequencing depth and high recall to identify true positives and avoid false positives. In contrast, peak calling for CUT&RUN data sets requires high specificity for true signal peaks. To this end, the Henikoff lab developed the Sparse Enrichments analysis for CUT&RUN (SEACR) peak caller that can be easily accessed using their web server at https://seacr.fredhutch.org/.

Alternatively, the Orkin and Yuan labs have streamlined processing of CUT&RUN data using their CUT&RUNTools pipeline https://bitbucket.org/qzhudfci/cutruntools/

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