

OCT 16, 2023

Protocol SAM-Seq Zea Mays

basile.leduque¹, Quadrana Leandro¹

¹CNRS



basile.leduque

ABSTRACT

OPEN ACCESS



DOI:

dx.doi.org/10.17504/protocol s.io.kqdq3x25zq25/v1

Protocol Citation: basile.led uque. Ouadrana Leandro 2023. Protocol SAM-Seg Zea Mays. protocols.io https://dx.doi.org/10.17504/p rotocols.io.kqdg3x25zg25/v1

License: This is an open access protocol distributed under the terms of the Creative Commons Attribution License. which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited

Protocol status: Working We use this protocol and it's working

Created: Aug 28, 2023

Last Modified: Oct 16, 2023

https://dx.doi.org/10.17504/protocols.io.kqdg3x25zg25/v1

Background: Epigenetic modifications, including chromatin accessibility, nucleosome positioning, and DNA methylation (5mC), are pivotal in shaping genome function. However, current short read sequencing approaches present challenges in characterising epigenetic patterns along repetitive genomic regions.

Results: We developed Simultaneous Accessibility and DNA Methylation Sequencing (SAM-seq), a robust method utilising bacterial adenine methyltransferases (m6A-MTases) to mark accessible regions in purified plant nuclei. Coupled with Oxford Nanopore Technology sequencing, SAM-seq enables high-resolution profiling of cytosine methylation and m6A-tagged chromatin accessibility along individual chromatin fibres in A. thaliana and maize. Importantly, using naked genomic DNA we uncovered significant sequence preferences of m6A-MTases, which we show must be taken into account in order to obtain reliable accessibility profiles, particularly for the analysis of highly repetitive sequences such as centromeric repeats. Using this method we found intriguing antagonism between accessibility and DNA methylation within single molecules, somatic epigenetic variation at bivalent chromatin domains, periodicity of nucleosomal strings over centromeric repeats, and high-resolution methylation profiles over nucleosomes.

Conclusions: Our study highlights the importance of considering intrinsic substrate preferences of DNA modifying enzymes for their use in chromatin profiling. Hence, SAM-seq is a robust and cross-species method to chart high-resolution accessibility and DNA methylation genome-wide. This method and findings will enable the investigation of chromatin-based regulation across plant species, with implications for the study of non-model plant species with limited genomic and epigenomic information.

Oct 16 2023

PROTOCOL integer ID:

87044

Funders Acknowledgement:

European Research Council

Grant ID: 948674

Reagent Preparation

1 Extraction Buffer (EB) 1

To prepare 50mlm of fresh EB1 buffer

Component	Amount	Final Concentration
2M Sucrose	10 ml	0.4M
1M Tris-HCl pH 8	0.5 ml	10mM
1M MgCl2	0.5ml	10mM
14.3M Ethidium bromide	17.5 μΙ	5mM
+ PIs (1 tablet per 50mls)		

2 Extraction Buffer (EB) 2

To prepare 4ml of fresh EB2:

Component	Amount	Final Concentration
Ethidium bromide	1,4 µL	5mM
20% Triton X- 100	200 μL	1%
H20	2818,6 μL	
1M MgCl2	40 μL	10mM
1M Tris-HCl pH 8	40 μL	10mM
Pls (1 mini- tablet diluted in 1 ml)	400 μL	
2M Sucrose	500 µl	0.25M

3 Extraction Buffer (EB) 3

To prepare 3ml of fresh EB 3:

Component	Amount	Final Concentration
1M Tris-HCl pH 8	30 µL	10mM
1M MgCl2	6 μL	2mM
20% Triton X- 100	25 μL	0.15%
+ Pis (1 mini- tablet diluted in 1 ml)	300 μL	
2M sucrose	2550 μL	1.7M
Ethidium bromide	1,05 µL	5mM
H20 to volume	90,3µL	

4

Nuclear Preparation Buffer (NPB)

To prepare 10ml of fresh NPB:

Component	Amount	Final Concentration
0.5M MOPS pH7	400µl	20 mM
5M NaCl	80 μΙ	40 mM
1M KCI	900µI	90 mM
0.5 M EDTA	40µl	2 mM
250mM EGTA	20 μΙ	0.5 mM
3.5M spermidine	1.2µl	0.5 mM
70 mM spermine	28.5µl	0.2 mM
1 mini-tablet diluted in 1ml	1ml	1X

5 Plant-Tween-Wash-Buffer (PTWB)

To prepare 3ml of fresh PTWB:

Component	Amount	Final Concentration	

Component	Amount	Final Concentration
20% Tween- 20	30 µl	0.2%
1M pH7,5 HEPES-KOH	60 µl	20 mM
5M NaCl	90 µl	150 mM
PI Mini	300 μΙ	
Spermidine 3.5M	0,4 μΙ	0,5 mM
H20	2530 µl	

6

Activation Buffer (AB)

To prepare 3ml of freash AB:

Component	Amount	Final Concentration
1M Tris-HCl pH 8	750 µl	15 mM
5M NaCl	150 µl	15 mM
1M KCI	3 ml	60 mM
0.5 M EDTA	100µl	1 mM
250mM EGTA	100 µl	0.5 mM
3.5M spermidine	0.715 μΙ	0.05 mM
BSA	50 mg	0.1%
H2O	fill to 50 ml	
SAM 32 mM	(add before reaction)	800μΜ

7 Reagents:

for m6A reaction

EcoGII/ rCutSmart/ SAM (M0603S New England Biolabs)

for library preparation:

NEBNext Companion Module for Oxford Nanopore Ligation Seq (E7180S New England Biolabs) SQK-LSK110 (Oxford Nanopore Technologies)

for nanopore sequencing:

Plant nuclei purification and permeabilization

8	Starting material: 0.75 g of powder (7 days old leaves)
9	Add the powder to 12.5 ml of Extraction Buffer (EB) 1 in a 50 ml falcon tube. Let sit on ice for 5 min.
10	Add 1% Formaldehyde for crosslinking (i.e. 338 µl Formaldehyde 37% in 12.5ml of EB1). Incubate 5 minutes
11	Add 1.25 M glycine to stop the crosslinking (i.e. 850ml of Glycine 2M per 338 μl Formaldehyde 37%)
12	Filter the solution through Miracloth into a new 50 ml falcon tube. Repeat once.
13	Filter the solution through a 0.4µm filter into a new 50 ml falcon tube.
14	Centrifuge the filtered solution for 20 minutes at 4,000 rpm at 4°C.

15 Gently remove supernatant and resuspend the pellet in 2 ml of Extraction Buffer (EB) 2. Transfer the solution into one 1.5ml Eppendorf tube. 16 Centrifuge at 12,000 g for 10 minutes at 4°C. Repeat step 7 & 8 once 17 Remove supernatant and resuspend pellet in 300µl of Extraction Buffer (EB) 3. 18 In a clean Eppendorf tube, add 300ul of EB3. Take the 300µl solution (resuspended pellet) from step 7 and carefully layer it on top of the clean 300µl of EB3. 19 Centrifuge for 45 min at 12,000 at 4 degrees. 20 Resuspend chromatin pellet in 300µl Nuclear Preparation Buffer (NPB) and incubate on ice for 5 min m6A-MTases reaction 21 Spin down nuclei at 1000g for 5 min at 4°C and resuspend in Plant-Tween-Wash-Buffer (PTWB). 22 Resuspend pellet in 38 µl 1X rCutSmart, containing 10µl SAM and 6µl EcoGII (i.e. 7.5U for 100µl). Incubate 30 minutes at 30°C

- 23 Add 20 µl NaCl (5M) and incubate overnight at 65 °C (or 1h30min if not crosslinked).
- 24 Add 5µl of RNAse A (20mg/ml) and 2µl Proteinase K. Incubate 30 minutes at 42°C.

Phenol/Chloroform extraction of genomic DNA

- 25 Add 1 volume of 1:1 phenol:chloroform (i.e. 400µl)
- 26 Centrifuge at 12,000 g for 10 minutes at 4°C
- 27 Recover the aqueous phase (upper) in a new LoBind Eppendorf tube
- 28 Add 1 volume of chloroform (i.e. 400µl)
- 29 Centrifuge at 12,000 g for 10 minutes at 4°C.
- Recover aqueous phase (upper) in a new LoBind Eppendorf tube

31 Precipitate DNA by adding 1/10 volume of NaAcétate 3M pH5,2 + 2,5 volume of EtOH 100% 32 Incubate at −20°C from 45 minutes to overnight. 33 Centrifuge at 12000g for 30 minutes at 4°C to precipitate DNA. 34 Wash the pellet with 300µ EtOH 70%. 35 Remove EtOH and dry pellets. 36 Elute in 20µl TE pH 8.0 **Library preparation**

37

Follow the protocol provided by Oxford Nanopore technology (Genomic_DNA_by_Ligation_SQK-LSK110) with the following modification:

- -Use 0.3 to 1 μg of DNA per library
- -For a better library preparation increase the End-Prep step up to 20 minutes at 20°C and Ligation step incubation to at least 1h
- Elute in Tris-HCl 10 mM.
- LFB was used for final size selection step
- Quantify DNA using the Qubit dsDNA HS Assay Kit (Q33230) to check yield.

Sequencing

38 Sequence during 72h on v9.4.1 flow cell FLO-MIN106.1 or FLO-PRO002 in MinION or PromethION sequencer, respectively.