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O AMPure Purification Protocol

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Abstract

A plethora of biological processes like gene transcription, DNA replication, DNA recombination, and chromosome segregation are mediated through protein–DNA interactions. A powerful method for investigating proteins within a native chromatin environment in the cell is chromatin immunoprecipitation (ChIP). Combined with the recent technological advancement in next generation sequencing, the ChIP assay can map the exact binding sites of a protein of interest across the entire genome. Here we describe a-step-by step protocol for ChIP followed by library preparation for ChIP-seq from yeast cells.

Guidelines

Chromatin immunoprecipitation (ChIP) is broadly used to study chromatin dynamics. Changes in occupancy of chromosomal proteins at specific loci within the genome can be measured by using ChIP-qPCR. However, this technique is costly and time consuming with high variability per experiment. Alternatively, ChIP-seq can be used to measure differences in a protein's occupancy genome wide. Finally, calibrated ChIP-seq is essential when measuring changes in occupancy between different experimental samples.

Here we describe an optimized ChIP protocol for yeast SMC proteins that can be completed within 3 days for samples analyzed by qPCR and 4 days for samples to be further processed by calibrated deep sequencing. The protocol encompasses five distinct steps: cross-linking and cell harvesting; cell lysis and sonication; immunoprecipitation, decross-linking and DNA extraction and finally determination of the size and DNA concentration of sonicated samples. These five steps are outlined here.

Materials

Yeast Strains and Growth Material:

- Haploid S. cerevisiae strains of w303 background we have used include: (a) no tag control (AM1176), (b) SCC1-6HA (AM1145), (c) BRN1-6HA(AM5708), (d) SCC2-6HIS-3FLAG (AM6006), and (e) SCC1-6HA pMET3-CDC20 (AM1105) as previously described [9,10,11,12].
- For studies of protein occupancy during meiosis we have used diploid *S. cerevisiae* strains of SK1 background including (a) *REC8-3HA ndt80*Δ (AM4015), as previously described [<u>13</u>] and (b) *REC8-6HIS-3FLAG* (AM11000).
- 3. Haploid *S. pombe* strains used for calibration are: (a) *RAD21-3HA* (spAM76), (b)*RAD21-6HA* (spAM635), (c) *RAD21-6HIS-3FLAG* (spAM1863), or (d) *CND2-6HA* (spAM1862).
- 4. YPDA media: 1% yeast extract, 2% peptone, 2% glucose.
- 5. YPG agar plates: 1% yeast extract, 2% peptone, 2.5% glycerol, 2% agar.
- 6. YPDA4% agar plates: 1% yeast extract, 2% peptone, 4% glucose, 2% agar.
- 7. BYTA media: 1% yeast extract, 2% Bacto tryptone, 1% potassium acetate, 50 mM potassium phthalate.
- 8. SPO media: 0.3% potassium acetate, pH 7.0.
- 9. YES media: 0.5% yeast extract, 3% glucose, 225 mg/L supplements.

Equipment and Reagents:

- 1. 37% formaldehyde solution for molecular biology.
- 2. 2.5 M glycine: Dissolve 93.8 g glycine in ddH_2O (may require gentle heating) and bring up to 500 ml with ddH_2O .
- 3. Diluent buffer: 0.143 M NaCl, 1.43 mM EDTA, 71.43 mM Hepes-KOH pH 7.5.
- 4. TBS buffer: 20 mM Tris-HCl pH 7.5, 150 mM NaCl.
- 5. 2× FA lysis buffer: 100 mM Hepes–KOH pH 7.5, 300 mM NaCl, 2 mM EDTA, 2% Triton X-100, 0.2% Nadeoxycholate.
- 6. FastPrep screw-cap tubes.
- 7.100 mM PMSF.
- 8. Protease inhibitor tablets Complete EDTA free.
- 9. Zirconia/Silica beads 0.5 mm diameter.
- 10. FastPrep-24 5G Homogenizer.
- 11. Bioruptor Twin.
- 12. Dynabeads Protein G.
- 13. Magnetic rack.
- 14. ChIP Wash buffer 1—low salt: 1× FA lysis buffer, 0.1%SDS, 275 mM NaCl.
- 15. ChIP Wash buffer 2—high salt: 1× FA lysis buffer, 0.1%SDS, 500 mM NaCl.
- 16. ChIP Wash buffer 3: 10 mM Tris-HCI pH 8.0, 0.25 M LiCI, 1 mM EDTA, 0.5% NP-40. 0.5% Na-deoxycholate.
- 17. ChIP Wash buffer 4 (TE): 10 mM Tris-HCl pH 8.0, 1 mM EDTA.
- 18. Chelex 100 Resin.
- 19.10 mg/ml Proteinase K
- 20. TES buffer: 50 mM Tris-HCl pH 7.5, 10 mM EDTA, 1% SDS.
- 21. Nuclease-free molecular biology grade water.
- 22. Filter tips.
- 23. Luna Universal Probe qPCR Master Mix.

- 24. LightCycler 480 Multiwell Plate 96.
- 25. LightCycler real-time PCR.
- 26. Qiagen purification kit.
- 27. LoBind DNA microcentrifuge tubes.
- 28. Quick blunting kit.
- 29. AMPure XP beads.
- 30. Klenow 3' to 5' exo minus.
- 31. Quick ligation kit (T4 DNA ligase).
- 32. NEXTflex DNA Barcodes—12 (Bioo Scientific; #NOVA-514102).
- 33. Phusion High-Fidelity DNA polymerase.
- 34. DynaMag-PCR magnet.
- 35. WizardSV Gel and PCR cleanup system.
- 36. Qubit dsDNA-HS Assay kit (Invitrogen).
- 37. Qubit Fluorometric Quantitation machine.
- 38. Agilent 2100 Bioanalyzer system.
- 39. High Sensitivity DNA Reagents kit (Agilent Technologies).
- 40. High Sensitivity DNA Chips (Agilent Technologies).
- 41. MiniSeq High throughput Reagent Kit (150-cycle) (Illumina).
- 42. Illumina Mini-seq.

Safety warnings

• For hazard information and safety warnings, please refer to the SDS (Safety Data Sheet).

Formaldehyde and PMSF are toxic if inhaled, ingested or absorbed through the skin. Always wear a lab coat and gloves, and work in a chemical hood.

AMPure Purification Protocol

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Note

AMPure purification relies on the principle of solid-phase reversible immobilization (SPRI) as previously described [16]. SPRI beads are paramagnetic and coated with carboxyl molecules, which reversibly bind DNA in the presence of polyethylene glycol (PEG) and salt (20% PEG, 2.5 M NaCl mix). PEG causes the negatively charged DNA to bind on the bead surface. This DNA immobilization is dependent on the concentration of PEG and salt in the reaction, and the volumetric ratio of SPRI beads to DNA is critical. Equal volume of beads to DNA will give an SPRI –DNA ratio of one. As this ratio is changed the length of fragments binding and/or left in solution also changes. A lower SPRI–DNA ratio results in larger fragments at elution.

2	Equilibrate an aliquot for all purifications needed of AMPure XP beads at	30m
	Room temperature for O0:30:00 before use. Vortex to resuspend.	X
3	Pipet carefully the indicated amounts so that no extra beads adhere to the outside of the tip.	Ø
4	Add the AMPure XP beads to DNA in solution and immediately mix thoroughly by repeated pipetting.	84
5	Incubate at Room temperature for 300:10:00 to allow binding of DNA to beads.	10m

6 Place on a magnetic rack for 😒 00:05:00 .

- 7 Remove and discard the supernatant taking great care not to take any beads.
- 8 Keep sample on magnetic rack and add $\angle 250 \mu L$ freshly prepared 80% ethanol without disturbing the beads.
- 9 Incubate for 🚫 00:00:30 . Remove and discard all supernatant.

30s

5m



10	Repeat the previous 2 steps once more: Keep sample on magnetic rack and add	Ø
	\triangleq 250 µL freshly prepared 80% ethanol without disturbing the beads.	~
11	Incubate for 60 00:00:30 . Remove and discard all supernatant.	30s
12	Let the beads air-dry for 📀 00:02:00 - 😒 00:03:00 at 📱 Room temperature .	5m
	Note	
	Overdrying the AMPure beads after the washing step will negatively impact on the DNA recovery. Beads are dry enough as soon as they lose their sheen.	
	recovery. Deads are dry enough as soon as they lose their sheen.	
13	Add the recommended amount of elution buffer (EB from Qiagen kit or ultrapure ddH2O)	A
	and resuspend the beads by pipetting.	<i>y</i>
14	Incubate at 📲 Room temperature for 🚯 00:03:00 .	3m
15	Place in magnetic rack for 🚫 00:02:00 .	2m
16	Transfer the supernatant to a new DNA LoBind Eppendorf tube.	d.
	Note	
	If, for example, eluting in 30 μ l, remove 28 μ l very slowly, being careful to prevent bead	
	carryover. If beads are accidentally removed, pipet the sample back into the tube and allow the beads to bind.	