SimpleChIP® Plus Enzymatic Chromatin IP Kit (Magnetic Beads)

1 Kit
(30 Immunoprecipitations)

Storage: All components in this kit are stable for at least 12 months past the reference date indicated on the component label when stored at the recommended temperature and left unused.

Note: Buffer A (4X) #7006, Buffer B (4X) #7007, and ChIP-Grade Protein G Magnetic Beads #9006 contain 0.05% sodium azide.

Reagents not supplied:
1. Magnetic Separation Rack #7017/#14654
2. 1X PBS #9872
3. Nuclease Free Water #12931
4. Ethanol (96-100%)
5. Formaldehyde (37% Stock)
6. SimpleChIP® Universal qPCR Master Mix #88989

Please visit www.cellsignal.com/technologies/chip.html for a complete listing of recommended companion products.

Description: The SimpleChIP® Plus Enzymatic Chromatin IP Kit (Magnetic Beads) #9005 is designed to conveniently provide reagents needed to perform up to 30 chromatin immunoprecipitations from cells or tissue samples, and is optimized for 4 X 10^6 cells or 25 mg of tissue per immunoprecipitation. This kit is compatible with ChIP-Seq.

Figure 1. Chromatin immunoprecipitations were performed with cross-linked chromatin from HCT 116 cells and either TCF4/TCF7L2 (C48H11) Rabbit mAb #2569, Non-phospho (Active) β-catenin (D13A1) Rabbit mAb #8814, or Tri-Methyl-Histone H3 (Lys4) (C42D8) Rabbit mAb #9751, using SimpleChIP® Plus Enzymatic Chromatin IP Kit (Magnetic Beads) #9005. DNA Libraries were prepared using DNA Library Prep Kit for Illumina® (ChIP-seq, CUT&RUN) #56795. The figure shows binding across ACSL5, a known target gene of TCF4/TCF7L2, β-catenin, and H3K4me3.

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FIGURE 2. Chromatin immunoprecipitations were performed with cross-linked chromatin from mouse liver and either Tri-Methyl-Histone H3 (Lys4) (C42D8) Rabbit mAb #9751 or FoxA1/HNF3 (D7P9B) Rabbit mAb #58613, using SimpleChIP® Plus Enzymatic Chromatin IP Kit (Magnetic Beads) #9005. DNA Libraries were prepared using DNA Library Prep Kit for Illumina® (ChIP-seq, CUT&RUN) #56795. The figure shows binding across chromosome 16 (upper), including Snai2 (lower), a known target gene of H3K4me3 and FoxA1.

FIGURE 3. Chromatin immunoprecipitations were performed with cross-linked chromatin from mouse liver and either RING1B (D22F2) XP® Rabbit mAb #5694 or SUZ12 (D39F6) XP® Rabbit mAb #3737, using SimpleChIP® Plus Enzymatic Chromatin IP Kit (Magnetic Beads) #9005. DNA Libraries were prepared using DNA Library Prep Kit for Illumina® (ChIP-seq, CUT&RUN) #56795. The figure shows binding across HOXA (upper) and HOXD (lower), known target genes of RING1B and SUZ12.
FIGURE 4. Mouse brain was cross-linked and disaggregated into a single-cell suspension using a Dounce homogenizer. The chromatin was prepared and digested, and chromatin immunoprecipitations were performed using the indicated ChIP-validated antibodies. Purified DNA was analyzed by quantitative real-time PCR using SimpleChIP® Mouse GAPDH Intron 2 Primers #8986, SimpleChIP® Mouse RPL30 Intron 2 Primers #7015, SimpleChIP® Mouse HoxA1 Promoter Primers #7341, and SimpleChIP® Mouse HoxD10 Exon 1 Primers #7429. The amount of immunoprecipitated DNA in each sample is represented as signal relative to the total amount of input chromatin (equivalent to 1).

FIGURE 5. Mouse liver was cross-linked and disaggregated into a single-cell suspension using a tissue disaggregator. The chromatin was prepared and digested, and chromatin immunoprecipitations were performed using the indicated ChIP-validated antibodies. Purified DNA was analyzed by quantitative real-time PCR using SimpleChIP® Mouse GAPDH Intron 2 Primers #8986, SimpleChIP® Mouse AFM Intron 2 Primers #7269, SimpleChIP® Mouse HoxA1 Promoter Primers #7341, and SimpleChIP® Mouse HoxD10 Exon 1 Primers #7429. The amount of immunoprecipitated DNA in each sample is represented as signal relative to the total amount of input chromatin (equivalent to 1).

FIGURE 6. Mouse brain or mouse liver were cross-linked and disaggregated into a single-cell suspension using a Dounce homogenizer or tissue disaggregator, respectively. The chromatin was prepared and digested, and chromatin immunoprecipitations were performed using the indicated ChIP-validated antibodies. DNA was purified and 10 μl was separated by electrophoresis on a 1% agarose gel and stained with ethidium bromide. The majority of chromatin from both brain (lane 1) and liver (lane 2) was digested to 1 to 5 nucleosomes in length (150 to 900 bp).
Introduction:
The chromatin immunoprecipitation (ChIP) assay is a powerful and versatile technique used for probing protein-DNA interactions within the natural chromatin context of the cell (1,2). This assay can be used to identify multiple proteins associated with a specific region of the genome, or the opposite, to identify the many regions of the genome associated with a particular protein (3-6). In addition, the ChIP assay can be used to define the spatial and temporal relationship of a particular protein-DNA interaction. For example, the ChIP assay can be used to determine the specific order of recruitment of various protein factors to a gene promoter or to “measure” the relative amount of a particular histone modification across an entire gene locus during gene activation (3,4). In addition to histone proteins, the ChIP assay can also be used to analyze binding of transcription factors, transcription co-factors, DNA replication factors, and DNA repair proteins.

When performing the ChIP assay, cells or tissues are first fixed with formaldehyde, a reversible protein-DNA cross-linking agent that serves to fix or “preserve” the protein-DNA interactions occurring in the cell (see method overview) (1,2). Cells are lysed and chromatin is harvested and fragmented using either sonication or enzymatic digestion. The chromatin is then subjected to immunoprecipitation using antibodies specific to a particular protein or histone modification. Any DNA sequences that are associated with the protein or histone modification of interest will co-precipitate as part of the cross-linked chromatin complex and the relative amount of that DNA sequence will be enriched by the immunoprecipitation process. After immunoprecipitation, the protein-DNA cross-links are reversed and the DNA is purified. The enrichment of a particular DNA sequence or sequences can then be detected by a number of different methods.

Standard PCR methods are often employed to identify the DNA sequences or regions of the genome associated with a particular protein or histone modification (1,2). PCR is used to measure the relative abundance of a particular DNA sequence enriched by a protein-specific immunoprecipitation versus an immunoprecipitation with a non-specific antibody control. PCR products are run on an agarose or acrylamide gel to facilitate quantification, and the level of enrichment of the DNA sequence is determined relative to the total amount of input DNA (Percent Input Method). The level of enrichment can also be expressed as fold enrichment above background (enrichment relative to that of the non-specific antibody control). Real-time PCR provides a more accurate, gel-free system for the quantification of DNA enrichment. Alternatively, the ChIP assay can be combined with genomic tiling micro-array (ChIP on chip) techniques, sequencing, or cloning strategies, which allow for genome-wide analysis of protein-DNA interactions and histone modifications (5-8).

The SimpleChIP® Plus Kit contains buffers and reagents needed to perform the ChIP assay with mammalian cells and tissue samples, and works for both histone modifications and non-histone DNA-binding proteins. After cross-linking, tissues are disaggregated into a single-cell suspension. Cells are then lysed and the chromatin is fragmented by partial digestion with Micrococcal Nuclease to obtain chromatin fragments of 1 to 5 nucleosomes in size. Enzymatic digestion of chromatin is milder than sonication and eliminates problems due to variability in sonication power and emulsification of chromatin during sonication, which can result in incomplete fragmentation of chromatin or loss of antibody epitopes due to protein denaturation and degradation. The chromatin immunoprecipitations are performed using antibodies and either ChIP-Grade Protein G Agarose Beads or ChIP-Grade Protein G Magnetic Beads. After reversal of protein-DNA cross-links, the DNA is purified using DNA purification spin columns provided in the kit. The DNA purification spin columns combine the convenience of spin-column technology with the selective binding properties of a uniquely designed silica membrane that allows for efficient recovery of DNA and removal of protein contaminants without the need for phenol/chloroform extractions and ethanol precipitations. After DNA purification, the enrichment of particular DNA sequences can be analyzed by a variety of methods.

In addition to providing buffers and reagents required to perform the ChIP assay, the SimpleChIP® Plus Kit provides important controls that allow for user determination of a successful ChIP experiment. The kit contains a positive control histone H3 rabbit monoclonal antibody, a negative control normal rabbit IgG antibody, and primer sets for PCR detection of the ribosomal protein L30 (RPL30) gene locus (human and mouse primer sets are included in the kit). Histone H3 is a core component of chromatin in the cell and is bound to most DNA sequences throughout the genome, including the RPL30 locus. Thus, immunoprecipitation of chromatin with the histone H3 rabbit monoclonal antibody will enrich for the RPL30 gene, while immunoprecipitation with the normal rabbit IgG antibody will not result in RPL30 gene enrichment. This enrichment can be quantified using either standard PCR or quantitative real-time PCR methods with the RPL30 primer sets provided in the kit. Importantly, since histone H3 is bound to most DNA sequences throughout the genome, the Histone H3 Rabbit mAb serves as a positive control IP for almost any locus studied, giving the user even more confidence that their ChIP experiment was performed successfully.

The SimpleChIP® Plus Enzymatic Chromatin IP Kit (Magnetic Beads) #9005 provides enough reagents to perform up to 30 immunoprecipitations and is optimized for 25 mg of tissue or 4 X 10^6 cultured cells per immunoprecipitation. A ChIP assay can be performed in as little as two days and can easily be scaled up or down for use with more or less tissue or cells. This kit is compatible with ChIP-Seq.

Background References:
I. Tissue Cross-linking and Sample Preparation

When harvesting tissue, remove unwanted material such as fat and necrotic material from the sample. Tissue can then be processed and cross-linked immediately, or frozen on dry ice and stored at -80°C for processing later. For optimal chromatin yield and ChIP results, use 25 mg of tissue for each immunoprecipitation to be performed. The chromatin yield does vary between tissue types and some tissues may require more than 25 mg for each immunoprecipitation. Please see Appendix A for more information regarding the expected chromatin yield for different types of tissue. One additional chromatin sample should be processed for Analysis of Chromatin Digestion and Concentration (Section IV). If desired, five additional chromatin samples should be processed for Optimization of Chromatin Digestion (Appendix B).

Before starting:

1. Prepare 45 µl of 37% formaldehyde per 25 mg tissue to be processed and keep at room temperature. Use fresh formaldehyde that is not past the manufacturer’s expiration date.

2. Prepare 3 ml of Phosphate Buffered Saline (PBS) + 15 µl 200X PIC per 25 mg of tissue to be processed and place on ice.

3. Prepare 45 µl of 37% formaldehyde per 25 mg of tissue to be processed and keep at room temperature. Use fresh formaldehyde that is not past the manufacturer’s expiration date.

4. Add 2 ml of 10X glycine to each 15 cm dish containing 20 ml medium, swirl briefly and keep dish on ice. It is important to keep the tissue cold to avoid protein degradation.

5. To crosslink proteins to DNA, add 45 µl of 37% formaldehyde per 1 ml of PBS + PIC and rock at room temp for 20 min. Final formaldehyde concentration is 1.5%.

6. Stop cross-linking by adding 100 µl of 10X Glycine per 1 ml of PBS + PIC and mix for 5 min at room temperature.

7. Centrifuge tissue at 500 x g in a benchtop centrifuge for 5 min at 4°C.

8. Remove supernatant and wash one time with 1 ml PBS + PIC per 25 mg tissue.

9. Repeat centrifugation at 500 x g in a benchtop centrifuge for 5 min at 4°C.

10. Remove supernatant and resuspend tissue in 1 ml PBS + PIC per 25 mg tissue and store on ice. Disaggregate tissue into single-cell suspension using a Medimachine (Part B) or Dounce homogenizer (Part C). (SAFE STOP) Alternatively, samples may be stored at -80°C before disaggregation for up to 3 months.

B. Tissue Disaggregation Using Medimachine from BD Biosciences (part #340587)

1. Cut off the end of a 1000 µl pipette tip to enlarge the opening for transfer of tissue chunks.

2. Transfer 1 ml of tissue resuspended in PBS + PIC into the top chamber of a 50 mm tissue culture dish (part #340593).

3. Grind tissue for 2 min according to manufacturer’s instructions.

4. Collect cell suspension from the bottom chamber of the medimachine using a 1 ml syringe and 18 gauge blunt needle. Transfer cell suspension to a 15 ml conical tube and place on ice.

5. Repeat steps 2 to 4 until all the tissue is processed into a homogenous suspension.

6. If more grinding is necessary, add more PBS + PIC to tissue. Repeat steps 2 to 5 until all tissue is ground into a homogenous suspension.

7. Check for single-cell suspension by microscope (optional).

8. Centrifuge cells at 2,000 x g in a benchtop centrifuge for 5 min at 4°C.

9. Remove supernatant from cells and continue with Nuclei Preparation and Chromatin Digestion (Section III).

C. Tissue Disaggregation Using a Dounce Homogenizer

1. Transfer tissue resuspended in PBS + PIC to a Dounce homogenizer.
Chromatin Immunoprecipitation Protocol (cont.)

- Prepare 1.1 ml 1X Buffer B (275 µl 4X Buffer B #7007 + 825 µl water) + 0.55 µl 1M DTT per IP prep and place on ice.
- Prepare 100 µl 1X ChIP Buffer (10 µl 10X ChIP Buffer #7008 + 90 µl water) + 0.5 µl 200X PIC per IP prep and place on ice.

1. Resuspend cells in 1 ml ice-cold 1X Buffer A + DTT + PIC per IP prep. Incubate on ice for 10 min. Mix by inverting tube every 3 min.

2. Pellet nuclei by centrifugation at 2,000 x g in a benchtop centrifuge for 5 min at 4°C. Remove supernatant and resuspend pelleted nuclei in 1 ml ice-cold 1X Buffer B + DTT per IP prep. Repeat centrifugation, remove supernatant, and resuspend pelleted nuclei in 100 µl 1X Buffer B + DTT per IP prep. Transfer sample to a 1.5 ml microcentrifuge tube, up to 1 ml total per tube.

3. Add 0.5 µl of Micrococcal Nuclease #10011 per IP prep, mix by inverting tube several times, and incubate for 20 min at 37°C with frequent mixing to digest DNA to a length of approximately 150-900 bp. Mix by inversion every 3 to 5 min. The amount of Micrococcal Nuclease required to digest DNA to the optimal length may need to be determined empirically for individual tissues and cell lines (see Appendix B). HeLa nuclei digested with 0.5 µl Micrococcal Nuclease per 4 x 10^6 cells and mouse liver nuclei digested with 0.5 µl Micrococcal Nuclease per 25 mg of tissue gave the appropriate length DNA fragments.

4. Stop digest by adding 10 µl of 0.5 M EDTA #7011 per IP prep and placing tube on ice for 1-2 min.

5. Pellet nuclei by centrifugation at 16,000 x g in a microcentrifuge for 1 min at 4°C and remove supernatant.

6. Resuspend pellet in 100 µl of 1X ChIP Buffer + PIC per IP prep and incubate on ice for 10 min.

7. Sonicate up to 500 µl of lysate per 1.5 ml microcentrifuge tube with several pulses to break nuclear membrane. Incubate samples for 30 sec on wet ice between pulses. Optimal conditions required for complete lysis of nuclei can be determined by observing the nuclei under light microscope before and after sonication. HeLa nuclei were completely lysed after 3 sets of 20-s pulses using a VirTis Visonic 100 Ultrasonic Homogenizer/Sonicator at setting 6 with a 1/8-inch probe. Alternatively, nuclei can be lysed by homogenizing the lysate 20 times in a Dounce homogenizer; however, lysis may not be as complete.

8. Clarify lysates by centrifugation at 9,400 x g in a microcentrifuge for 10 min at 4°C.

9. Transfer supernatant to a new tube. (SAFE STOP This is the cross-linked chromatin preparation, which should be stored at -80°C until further use. Remove 50 µl of the chromatin preparation for Analysis of Chromatin Digestion and Concentration (Section IV). This 50 µl sample may be stored at -20°C overnight."

IV. Analysis of Chromatin Digestion and Concentration (Recommended Step)

1. To the 50 µl chromatin sample (from Step 9 in Section III), add 100 µl nuclease-free water, 6 µl 5 M NaCl #7010, and 2 µl RNase A #7013. Vortex to mix and incubate samples at 37°C for 30 min.

2. To each RNAse A-digested sample, add 2 µl Protamine K #10012. Vortex to mix and incubate samples at 65°C for 2 h.

3. Purify DNA from samples using DNA purification spin columns as described in Section VII. (SAFE STOP) DNA may be stored at -20°C for up to 6 months.

4. After purification of DNA, remove a 10 µl sample and determine DNA fragment size by electrophoresis on a 1% agarose gel with a 100 bp DNA marker. DNA should be digested to a length of approximately 150-900 bp (1 to 5 nucleosomes).

5. To determine DNA concentration, transfer 2 µl of purified DNA to 98 µl nuclease-free water to give a 50-fold dilution and read the OD260. The concentration of DNA in μg/ml is OD260 x 2,500. DNA concentration should ideally be between 50 and 200 μg/ml.

NOTE: For optimal ChIP results, it is highly critical that the chromatin is of appropriate size and concentration. Over-digestion of chromatin may diminish signal in the PCR quantification. Under-digestion of chromatin may lead to increased background signal and lower resolution. Adding too little chromatin to the IP may result in diminished signal in the PCR quantification. A protocol for optimization of chromatin digestion can be found in Appendix B.

V. Chromatin Immunoprecipitation

For optimal ChIP results, use approximately 5 to 10 µg of digested, cross-linked chromatin (as determined in Section IV) per immunoprecipitation. This should be roughly equivalent to a single 100 µl IP prep from 25 mg of disaggregated tissue or 4 x 10^6 tissue culture cells. Typically, 100 µl of digested chromatin is diluted into 400 µl 1X ChIP Buffer prior to the addition of antibodies. However, if more than 100 µl of chromatin is required per IP, antibodies can be added directly to the undiluted chromatin preparation for immunoprecipitation of chromatin complexes.

Before starting:

1. In one tube, prepare enough 1X ChIP Buffer for the dilution of digested chromatin into the desired number of immunoprecipitations: 400 µl of 1X ChIP Buffer (40 µl of 10X ChIP Buffer + 360 µl water) + 2 µl 200X PIC per immunoprecipitation. When determining the number of immunoprecipitations, remember to include the positive control Histone H3 (D2B12) XP® Rabbit mAb #4620 and negative control Normal Rabbit IgG #2729 samples. Place mix on ice.

2. To the prepared 1X ChIP Buffer, add the equivalent of 100 µl (5 to 10 µg of chromatin) of the digested, cross-linked chromatin preparation (from Step 9 in Section III) per immunoprecipitation. For example, for 10 immunoprecipitations, prepare a tube containing 4 ml 1X ChIP Buffer (400 µl 10X ChIP Buffer + 3.6 ml water) + 200X PIC + 1 ml digested chromatin preparation.

3. Remove a 10 µl sample of the diluted chromatin and transfer to a microtube. This is your 2% Input Sample, which can be stored at -20°C until further use (Step 1 in Section VI).

4. For each immunoprecipitation, transfer 500 µl of the diluted chromatin to a 1.5 ml microcentrifuge tube and add the immunoprecipitating antibody. The amount of antibody required per IP varies and should be determined by the user. For the positive control Histone H3 (D2B12) XP® Rabbit mAb #4620, add 10 µl to the IP sample. For the negative control Normal Rabbit IgG #2729, add 1 µl (1 µg) to 2 µl (2 µg) to the IP sample. If using antibodies from Cell Signaling Technology, please see recommended dilution listed on the datasheet or product webpage and calculate the amount (µg) of IgG antibody for negative control based on the Cell Signaling Antibody concentration for a fair comparison. Incubate IP samples 4 h to overnight at 4°C with rotation.

NOTE: Most antibodies from Cell Signaling Technology work optimally between 1 and 2 µg per IP sample. In the case where there are multiple samples with varying concentrations, it is best to match the negative control Normal Rabbit IgG #2729 to the highest antibody concentration.

5. Resuspend ChIP-Grade Protein G Magnetic Beads #9006 by gently vortexing. Immediately add 30 µl of Protein G Magnetic Beads to each IP reaction and incubate at 2 h at 4°C with rotation.

6. Pellet protein G magnetic beads in each immunoprecipitation by placing the tubes in a magnetic separation rack #7017. Wait 1 to 2 min for solution to clear and then carefully remove supernatant.

7. Wash protein G magnetic beads by adding 1 ml of cold salt wash to the beads and incubate at 4°C for 5 min with rotation. Repeat steps 6 and 7 two additional times for a total of 3 low salt washes.

8. Add 1 ml of high salt wash to the beads and incubate at 4°C for 5 min with rotation.

9. Pellet protein G magnetic beads in each immunoprecipitation by placing the tubes in a Magnetic Separation Rack. Wait 1 to 2 min for solution to clear and then carefully remove supernatant. Immediately proceed to Section VI.
VI. Elution of Chromatin from Antibody/Protein G Magnetic Beads and Reversal of Cross-links

Before starting:
(1) All buffer volumes should be increased proportionally based on the number of immunoprecipitations in the experiment.
- Remove and warm 2X ChIP Elution Buffer #7009 in a 37°C water bath and ensure SDS is in solution.
- Set a water bath or thermomixer to 65°C.
- Prepare 150 µl 1X ChIP Elution Buffer (75 µl 2X ChIP Elution Buffer #7009 + 75 µl water) for each immunoprecipitation and the 2% input sample.

1. Add 150 µl of the 1X ChIP Elution Buffer to the 2% input sample tube and set aside at room temperature until Step 6.
2. Add 150 µl 1X ChIP Elution Buffer to each IP sample.
3. Elute chromatin from the antibody/protein G magnetic beads for 30 min at 65°C with gentle vortexing (1,200 rpm). A thermomixer works best for this step. Alternatively, elutions can be performed at room temperature with rotation, but may not be as complete.
4. Pellet protein G magnetic beads by placing the tubes in a magnetic separation rack and wait 1 to 2 min for solution to clear.
5. Carefully transfer eluted chromatin supernatant to a new tube.
6. To all tubes, including the 2% input sample from Step 1, reverse cross-links by incubation can be extended overnight.
7. Immediately proceed to Section VII. (SAFE STOP) Alternatively, samples can be stored at -20°C for up to 4 days. However, to avoid formation of a precipitate, be sure to warm samples to room temperature before adding DNA Binding Buffer #10007 (Section VII, Step 1).

VII. DNA Purification Using Spin Columns

Before starting:
- (1!) Add 24 ml of ethanol (96-100%) to DNA Wash Buffer #10008 before use. This step only has to be performed once prior to the first set of DNA purifications.
- Remove one DNA Purification collection tube #10010 for each DNA sample from Section V.

1. Add 750 µl DNA Binding Buffer #10007 to each DNA sample and vortex briefly.
2. Transfer 450 µl of each sample from Step 1 to a DNA spin column in collection tube.
3. Centrifuge at 18,500 x g in a microcentrifuge for 30 sec.
4. Remove the spin column from the collection tube and discard the liquid. Replace spin column in the collection tube.
5. Transfer the remaining 450 µl of each sample from Step 1 to the spin column in collection tube. Repeat Steps 3 and 4.
6. Add 750 µl of DNA Wash Buffer #10008 to the spin column in collection tube.
7. Centrifuge at 18,500 x g in a microcentrifuge for 30 sec.
8. Remove the spin column from the collection tube and discard the liquid. Replace spin column in the collection tube.
9. Centrifuge at 18,500 x g in a microcentrifuge for 30 sec.
11. Add 50 µl of DNA Elution Buffer #10009 to each spin column and place into a clean 1.5 ml microcentrifuge tube.
12. Centrifuge at 18,500 x g in a microcentrifuge for 30 sec to elute DNA.
13. Remove and discard DNA spin column. Eluate is now purified DNA. (SAFE STOP) Samples can be stored at -20°C for up to 6 months.

VIII. Quantification of DNA by PCR:

Recommendations:
- Use Filter-tip pipette tips to minimize risk of contamination.
- The control primers included in the kit are specific for the human or mouse RPL30 gene (#7014 + #7015) and can be used for either standard PCR or quantitative real-time PCR. If the user is performing ChIPs from another species, it is recommended that the user design the appropriate specific primers to DNA and determine the optimal PCR conditions.

- A Hot-Start Taq polymerase is recommended to minimize the risk of nonspecific PCR products.
- PCR primer selection is critical. Primers should be designed with close adherence to the following criteria:

<table>
<thead>
<tr>
<th>Primer length:</th>
<th>24 nucleotides</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optimum Tm:</td>
<td>60°C</td>
</tr>
<tr>
<td>Optimum GC:</td>
<td>50%</td>
</tr>
<tr>
<td>Amplicon size:</td>
<td>150 to 200 bp (for standard PCR) 80 to 160 bp (for real-time quantitative PCR)</td>
</tr>
</tbody>
</table>

Standard PCR Method:
1. Label the appropriate number of 0.2 ml PCR tubes for the number of samples to be analyzed. These should include the 2% input sample, the positive control histone H3 sample, the negative control normal rabbit IgG sample, and a tube with no DNA to control for DNA contamination.
2. Add 2 µl of the appropriate DNA sample to each tube.
3. Prepare a master reaction mix as described below, making sure to add enough reagent for two extra tubes to account for loss of volume. Add 18 µl of master mix to each reaction tube.

<table>
<thead>
<tr>
<th>Reagent</th>
<th>Volume for 1 PCR Reaction (18 µl)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nuclease-free H2O</td>
<td>12.5 µl</td>
</tr>
<tr>
<td>10X PCR Buffer</td>
<td>2.0 µl</td>
</tr>
<tr>
<td>4 mM dNTP Mix</td>
<td>1.0 µl</td>
</tr>
<tr>
<td>5 µM RPL30 Primers</td>
<td>2.0 µl</td>
</tr>
<tr>
<td>Tag DNA Polymerase</td>
<td>0.5 µl</td>
</tr>
</tbody>
</table>

4. Start the following PCR reaction program:

- a. Initial Denaturation 95°C 5 min
- b. Denature 95°C 30 sec
- c. Anneal 62°C 30 sec
- d. Extension 72°C 30 sec
- e. Repeat steps b-d for a total of 34 cycles.
- f. Final Extension 72°C 5 min

5. Remove 10 µl of each PCR product for analysis by 2% agarose gel or 10% polyacrylamide gel electrophoresis with a 100 bp DNA marker. The expected size of the PCR product is 161 bp for human RPL30 #7014 and 159 bp for mouse RPL30 #7015.

Real-Time Quantitative PCR Method:
1. Label the appropriate number of PCR tubes or PCR plates compatible with the model of PCR machine to be used. PCR reactions should include the positive control histone H3 sample, the negative control normal rabbit IgG sample, a tube with no DNA to control for contamination, and a serial dilution of the 2% input chromatin DNA (undiluted, 1:5, 1:25, 1:125) to create a standard curve and determine the efficiency of amplification.
2. Add 2 µl of the appropriate DNA sample to each tube or well of the PCR plate.
3. Prepare a master reaction mix as described below. Add enough reagents for two extra reactions to account for loss of volume. Add 18 µl of reaction mix to each PCR reaction tube or well. (SAFE STOP) If necessary cover plate with aluminum foil to avoid light and store at 4°C up to 4 hours or -20°C overnight until machine is ready for use.

<table>
<thead>
<tr>
<th>Reagent</th>
<th>Volume for 1 PCR Reaction (18 µl)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nuclease-free H2O</td>
<td>6.0 µl</td>
</tr>
<tr>
<td>5 µM RPL30 Primers</td>
<td>2.0 µl</td>
</tr>
<tr>
<td>SimpleChIP® Universal qPCR</td>
<td>10.0 µl</td>
</tr>
<tr>
<td>Master Mix #88989</td>
<td></td>
</tr>
</tbody>
</table>

4. Start the following PCR reaction program:

- a. Initial Denaturation 95°C 3 min
- b. Denature 95°C 15 sec
- c. Anneal and Extension: 60°C 60 sec
- d. Repeat steps b and c for a total of 40 cycles.
Chromatin Immunoprecipitation Protocol (cont.)

5. Analyze quantitative PCR results using the software provided with the real-time PCR machine. Alternatively, one can calculate the IP efficiency manually using the Percent Input Method and the equation shown below. With this method, signals obtained from each immunoprecipitation are expressed as a percent of the total input chromatin.

\[
\text{Percent Input} = 2^\Delta C_T \times 100
\]

\(\Delta C_T = C_{T\text{input sample}} - C_{T\text{IP sample}}\)

IX. NG-Sequencing Library Construction

The immuno-enriched DNA samples prepared with this kit are directly compatible with ChIP-seq. For downstream NG-sequencing DNA library construction, use a DNA library preparation protocol or kit compatible with your downstream sequencing platform. For sequencing on Illumina® platforms, we recommend DNA Library Prep Kit for Illumina® (ChIP-seq, CUT&RUN) #56795 and its associated index primers Multiplex Oligos for Illumina® (Single Index Primers) (ChIP-seq, CUT&RUN) #29580 or Multiplex Oligos for Illumina® (Dual Index Primers) (ChIP-seq, CUT&RUN) #47538.

Recommendations:

- For transcription factor or co-factor ChIP-seq, use 5 ng of ChIP-enriched DNA and amplification of the adaptor-ligated DNA with 10 cycles of PCR.
- For total histone and histone modifications, or input samples, start with 50 ng of ChIP-enriched DNA and amplification of the adaptor-ligated DNA with 6 cycles of PCR.
- For library construction of ChIP-enriched DNA for all target types, perform cleanup of adaptor-ligated DNA without size selection.
- After DNA library construction, check the DNA library for presence of adaptor dimers (~140 bp) using an Agilent High Sensitivity DNA Kit (Agilent Technologies, Cat# G2938-90322), or by agarose gel electrophoresis with 50-100 ng DNA on a 2% agarose TAE gel. If adaptor dimers are present in the DNA library, repeat cleanup of PCR amplified material.
- The quality of the library can also be confirmed using qPCR and primer sets to known positive and negative target loci. Positive primer pairs should still give the same high signal compared to negative primers as seen in the original qPCR analysis of ChIP-enriched DNA.
- After final cleanup and quality checks, prepare final purified library samples at 2-10 nM for high throughput sequencing.
Chromatin Immunoprecipitation Protocol (cont.)

APPENDIX A: Expected Chromatin Yield

When harvesting cross-linked chromatin from tissue samples, the yield of chromatin can vary significantly between tissue types. The table to the right provides a range for the expected yield of chromatin from 25 mg of tissue compared to 4 x 10^6 HeLa cells, and the expected DNA concentration, as determined in Section IV of the protocol. For each tissue type, disaggregation using a Medimachine (BD Biosciences) or a Dounce homogenizer yielded similar amounts of chromatin. However, chromatin processed from tissues disaggregated using the Medimachine typically gave higher IP efficiencies than chromatin processed from tissues disaggregated using a Dounce homogenizer. A Dounce homogenizer is strongly recommended for disaggregation of brain tissue, as the Medimachine does not adequately disaggregate brain tissue into a single-cell suspension. For optimal ChiP results, we recommend using 5 to 10 µg of digested, cross-linked chromatin per immunoprecipitation; therefore, some tissues may require harvesting more than 25 mg per each immunoprecipitation.

<table>
<thead>
<tr>
<th>Tissue/Cell</th>
<th>Total Chromatin Yield</th>
<th>Expected DNA Concentration</th>
</tr>
</thead>
<tbody>
<tr>
<td>Spleen</td>
<td>20-30 µg per 25 mg tissue</td>
<td>200-300 µg/ml</td>
</tr>
<tr>
<td>Liver</td>
<td>10-15 µg per 25 mg tissue</td>
<td>100-150 µg/ml</td>
</tr>
<tr>
<td>Kidney</td>
<td>8-10 µg per 25 mg tissue</td>
<td>80-100 µg/ml</td>
</tr>
<tr>
<td>Brain</td>
<td>2-5 µg per 25 mg tissue</td>
<td>20-50 µg/ml</td>
</tr>
<tr>
<td>Heart</td>
<td>2-5 µg per 25 mg tissue</td>
<td>20-50 µg/ml</td>
</tr>
<tr>
<td>HeLa</td>
<td>10-15 µg per 4 x 10^6 cells</td>
<td>100-150 µg/ml</td>
</tr>
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APPENDIX B: Optimization of Chromatin Digestion

Optimal conditions for the digestion of cross-linked chromatin DNA to 150-900 base pairs in length is highly dependent on the ratio of Micrococcal Nuclease to the amount of tissue or number of cells used in the digest. Below is a protocol for determination of the optimal digestion conditions for a specific tissue or cell type.

1. Prepare cross-linked nuclei from 125 mg of tissue or 2 x 10^7 cells (equivalent of 5 IP preps), as described in Sections I, II, and III. Stop after Step 2 of Section III and proceed as described below.

2. Transfer 100 µl of the nuclei preparation into 5 individual 1.5 ml microcentrifuge tubes and place on ice.

3. Add 3 µl Micrococcal Nuclease stock to 27 µl of 1X Buffer B + DTT (1:10 dilution of enzyme).

4. To each of the 5 tubes in Step 2, add 0 µl, 2.5 µl, 5 µl, 7.5 µl, or 10 µl of the diluted Micrococcal Nuclease, mix by inverting tube several times and incubate for 20 min at 37°C with frequent mixing.

5. Stop each digest by adding 10 µl of 0.5 M EDTA and placing tubes on ice.

6. Pellet nuclei by centrifugation at 13,000 rpm in a microcentrifuge for 1 min at 4°C and remove supernatant.

7. Resuspend nuclear pellet in 200 µl of 1X ChiP Buffer + PIC. Incubate on ice for 10 min.

8. Sonicate lysate with several pulses to break nuclear membrane. Incubate samples 30 sec on wet ice between pulses. Optimal conditions required for complete lysis of nuclei can be determined by observing nuclei under light microscope before and after sonication. HeLa nuclei were completely lysed after 3 sets of 20-sec pulses using a VirTis Virsonic 100 Ultrasonic Homogenizer/Sonicator set at setting 6 with a 1/8-inch probe. Alternatively, nuclei can be lysed by homogenizing the lysate 20 times in a Dounce homogenizer; however, lysis may not be as complete.

9. Clarify lysates by centrifugation at 10,000 rpm in a microcentrifuge for 10 min at 4°C.

10. Transfer 50 µl of each of the sonicated lysates to new microfuge tubes.

11. To each 50 µl sample, add 100 µl nuclease-free water, 6 µl 5 M NaCl and 2 µl RNAse A. Vortex to mix and incubate samples at 37°C for 30 min.

12. To each RNAse A-digested sample, add 2 µl Proteinase K. Vortex to mix and incubate sample at 65°C for 2 h.

13. Remove 20 µl of each sample and determine DNA fragment size by electrophoresis on a 1% agarose gel with a 100 bp DNA marker.

14. Observe which of the digestion conditions produces DNA in the desired range of 150-900 base pairs (1 to 5 nucleosomes, see Figure 6). The volume of diluted Micrococcal Nuclease that produces the desired size of DNA fragments using this optimization protocol is equivalent to 10 times the volume of Micrococcal Nuclease stock that should be added to one immunoprecipitation preparation (25 mg of disaggregated tissue cells or 4 x 10^6 tissue culture cells) to produce the desired size of DNA fragments. For example, if 5 µl of diluted Micrococcal Nuclease produces DNA fragments of 150-900 base pairs in this protocol, then 0.5 µl of stock Micrococcal Nuclease should be added to one IP prep during the digestion of chromatin in Section III.

15. If results indicate that DNA is not in the desired size range, then repeat optimization protocol, adjusting the amount of Micrococcal Nuclease in each digest accordingly. Alternatively, the digestion time can be changed to increase or decrease the extent of DNA fragmentation.
### APPENDIX C: Troubleshooting Guide

<table>
<thead>
<tr>
<th>Problem</th>
<th>Possible Causes</th>
<th>Recommendation</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Concentration of the digested chromatin is too low.</td>
<td>Not enough cells added to the chromatin digestion or nuclei were not completely lysed after digestion.</td>
<td>If DNA concentration of the chromatin preparation is close to 50 µg/ml, add additional chromatin to each IP to give at least 5 µg/IP and continue with protocol. Count a separate plate of cells before cross-linking to determine an accurate cell number and/or visualize nuclei under microscope before and after sonication to confirm complete lysis of nuclei.</td>
</tr>
<tr>
<td>2. Chromatin is under-digested and fragments are too large (greater than 900 bp).</td>
<td>Cells may have been over cross-linked. Cross-linking for longer than 10 min may inhibit digestion of chromatin. Too many cells or not enough Micrococcal Nuclease was added to the chromatin digestion.</td>
<td>Perform a time course at a fixed formaldehyde concentration. Shorten the time of cross-linking to 10 min or less. Count a separate plate of cells before cross-linking to determine accurate cell number and see Appendix B for optimization of chromatin digestion.</td>
</tr>
<tr>
<td>3. Chromatin is over-digested and fragments are too small (exclusively 150 bp mono-nucleosome length). Complete digestion of chromatin to mono-nucleosome length DNA may diminish signal during PCR quantification, especially for amplicons greater than 150 bp in length.</td>
<td>Not enough cells or too much Micrococcal Nuclease added to the chromatin digestion.</td>
<td>Count a separate plate of cells before cross-linking to determine accurate cell number and see Appendix B for optimization of chromatin digestion.</td>
</tr>
<tr>
<td>4. No product or very little product in the input PCR reactions.</td>
<td>Not enough DNA added to the PCR reaction or conditions are not optimal. PCR amplified region may span nucleosome-free region. Not enough chromatin added to the IP or chromatin is over-digested.</td>
<td>Add more DNA to the PCR reaction or increase the number of amplification cycles. Optimize the PCR conditions for experimental primer set using purified DNA from cross-linked and digested chromatin. Design a different primer set and decrease length of amplicon to less than 150 bp (see primer design recommendations in Section VIII). For optimal ChIP results add 5-10 µg chromatin per IP. See recommendations for problems 1 and 3 above.</td>
</tr>
<tr>
<td>5. No product in the positive control Histone H3-IP RPL30 PCR reaction.</td>
<td>Not enough chromatin or antibody added to the IP reaction or IP incubation time is too short. Incomplete elution of chromatin from Protein G beads.</td>
<td>Be sure to add 5-10 µg of chromatin and 10 µl of antibody to each IP reaction and incubate with antibody over-night and an additional 2 h after adding Protein G beads. Elution of chromatin from Protein G beads is optimal at 65°C with frequent mixing to keep beads suspended in solution.</td>
</tr>
<tr>
<td>6. Quantity of product in the negative control Rabbit IgG-IP and positive control Histone H3-IP PCR reactions is equivalent.</td>
<td>Too much or not enough chromatin added to the IP reaction. Alternatively, too much antibody added to the IP reaction. Too much DNA added to the PCR reaction or too many cycles of amplification.</td>
<td>Add no more than 15 µg of chromatin and 10 µl of histone H3 antibody to each IP reaction. Reduce the amount of normal rabbit IgG to 1 µl per IP. Add less DNA to the PCR reaction or decrease the number of PCR cycles. It is very important that the PCR products are analyzed within the linear amplification phase of PCR. Otherwise, the differences in quantities of starting DNA cannot be accurately measured.</td>
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<tr>
<td>7. No product in the Experimental Antibody-IP PCR reaction.</td>
<td>Not enough DNA added to the PCR reaction. Not enough antibody added to the IP reaction. Antibody does not work for IP.</td>
<td>Add more DNA to the PCR reaction or increase the number of amplification cycles. Typically a range of 1 to 5 µg of antibody are added to the IP reaction; however, the exact amount depends greatly on the individual antibody. Increase the amount of antibody added to the IP. Find an alternate antibody source.</td>
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