ChIP-IT® Express
Magnetic Chromatin
Immunoprecipitation Kit
&
Sonication Shearing Kit

(version F5)

Catalog Nos. 53008 & 53032

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Overview

Chromatin Immunoprecipitation (ChIP) is a powerful tool for studying protein/DNA interactions\(^1\,^2\). Intact cells are fixed with formaldehyde, which cross-links and preserves protein/DNA interactions. The DNA is then sheared into small, relatively uniform fragments using either sonication or enzymatic digestion, and specific protein/DNA complexes are immunoprecipitated with an antibody directed against the DNA-binding protein of interest. Following immunoprecipitation, cross-linking is reversed, the proteins are removed by treatment with Proteinase K, and the DNA is recovered. The DNA is then analyzed to determine which DNA fragments were in complex with the protein of interest (see Figure 1 on page 2).

ChIP is extremely useful for the study of chromatin biology and transcriptional regulation because it enables the localization of chromatin proteins, modified histones and transcription factors to specific DNA loci. Furthermore, because protein/DNA interactions are fixed while in an endogenous, chromosomal context, ChIP results reflect the influence of chromosomal topology and the effects of cellular regulatory proteins\(^3\,^4\,^5\).

ChIP can be technically demanding. The method requires high-quality antibodies to recognize the fixed, target-bound proteins of interest, and an efficient reagent (usually protein A or G beads) to precipitate the antibody/chromatin complex. In addition, specialized buffers, inhibitor cocktails and blocking reagents are required to minimize non-specific enrichment and reduce protein degradation.

Active Motif’s ChIP-IT\(^\text{®}\) Express Kits contain proven ChIP reagents that provide a complete solution for convenient, accurate monitoring of protein/DNA interactions. The ChIP-IT Express Kits utilize protein G-coupled magnetic beads, which have greatly simplified and streamlined the ChIP protocol. A number of steps have been reduced in length or completely eliminated (see page 3 for details), making ChIP-IT Express an extremely rapid and efficient way to perform ChIP.

ChIP-IT Express Kits contain reagents sufficient to make 10 chromatin preparations, as well as 2 shearing optimizations, and to perform 25 ChIP reactions. If you need to shear more chromatin, the stand-alone ChIP-IT Express Shearing Kit provides just the reagents to perform and validate the shearing portion of the protocol. For additional protocols and to learn about products related to ChIP-IT Express, please visit [www.activemotif.com/chipitdocs](http://www.activemotif.com/chipitdocs).

<table>
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<tr>
<th>product</th>
<th>format</th>
<th>catalog no.</th>
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<tr>
<td>ChIP-IT(^\text{®}) Express</td>
<td>25 rxns</td>
<td>53008</td>
</tr>
<tr>
<td>ChIP-IT(^\text{®}) Express Shearing Kit</td>
<td>10 rxns</td>
<td>53032</td>
</tr>
<tr>
<td>ChIP-IT(^\text{®}) Protein G Magnetic Beads</td>
<td>25 rxns</td>
<td>53014</td>
</tr>
</tbody>
</table>
Flow Chart of Process

Cross-link protein to DNA in living cells with formaldehyde.

Break open cells and shear DNA by Sonication or Enzymatic Shearing.

Add primary antibody of interest.

Reverse cross-links, digest with Proteinase K, then stop reaction.
DNA is ready for analysis.

Elute chromatin, pellet beads with magnet and transfer supernatant to a fresh tube.

Capture antibody-bound protein/DNA complexes and wash using Magnetic Beads.

Figure 1: Improved chromatin immunoprecipitation using ChIP-IT Express.

In ChIP-IT Express, cells are treated with formaldehyde to fix protein/DNA interactions and then the fixed chromatin is sheared by either sonication or enzymatic digestion. The sheared chromatin is incubated with an antibody directed against a protein of interest, and antibody-bound protein/DNA complexes are precipitated through use of magnetic Protein G-coupled beads. The captured chromatin is then eluted, the cross-links are reversed, and the recovered DNA is analyzed by PCR to identify DNA loci associated with the protein of interest.

References

ChIP-IT Express Advantages

**Complete kits for rapid and efficient ChIP**
- Shorter protocol and dramatically reduced hands-on time
- The possibility to perform numerous ChIP experiments simultaneously
- Compatible with multi-channel pipetting

The ChIP-IT Express and ChIP-IT Express Enzymatic Kits provide reagents and protocols to simplify all aspects of the chromatin immunoprecipitation procedure. The kits can be used to prepare chromatin, determine optimal conditions for shearing chromatin and perform ChIP reactions.

ChIP-IT Express Kits contain components to make 10 chromatin preparations as well as to perform 2 shearing optimizations, and quantities of all other components to perform 25 ChIP reactions. The included protein-G coated magnetic beads are provided ready to use. These beads have a high binding capacity for IgG and low non-specific binding. As a result, these magnetic beads require fewer washing steps than agarose beads, and it is not necessary to pre-clear the chromatin prior to ChIP. An added advantage is that the magnetic beads pellet much more quickly than standard agarose beads, which must be pelleted by centrifugation steps. In addition, magnetic stands (and the included bar magnet) are designed to pellet the beads onto the side of the tube. This makes it easier to remove buffers without disturbing the beads, so washing can be performed using multi-channel pipettors. This dramatically reduces hands-on time and ensures sample-to-sample consistency. The provided siliconized microcentrifuge tubes (1.7 ml) simplify wash steps and ensure a minimal loss of Protein G beads and DNA.

Other steps in the ChIP protocol have also been optimized. The specialized Elution Buffer in combination with the Reverse Cross-linking Buffer saves time and eliminates the DNA loss that can occur during manipulations.

These improvements greatly reduce hands-on time and 8-, 16- or 24-ChIP experiments can easily be performed at the same time. This is not possible with traditional ChIP methods, which are time- and labor-intensive. For even higher throughput ChIP, Active Motif offers ChIP-IT Express HT (Catalog No. 53018), which enables you to perform 96 ChIPs simultaneously.

**Shearing options – ChIP-IT Express (Sonication) and ChIP-IT Express Enzymatic (Enzymatic)**

The ChIP-IT Express Kit provides reagents sufficient to prepare 10 sonication-sheared chromatin preparations as well as to perform 2 shearing optimizations. Using this protocol, each preparation of sheared chromatin requires one 15 cm tissue culture plate of cells and yields chromatin sufficient for up to 6 ChIP-IT Express reactions (one ChIP reaction is considered to be the incubation of one sample of chromatin with one antibody). However, the chromatin preparation protocols can be scaled up or down depending on how many cells you would like to work with (see Appendix – Section D of this manual). If you want to prepare additional samples using sonication, the ChIP-IT Express Shearing Kit is sold separately (Catalog No. 53032).
The ChIP-IT Express Enzymatic Kit is similar to the ChIP-IT Express Kit, but it utilizes a proprietary Enzymatic Shearing Cocktail and Digestion Buffer to shear chromatin using enzymatic digestion instead of sonication. Because enzymatic shearing is solely time and temperature dependent, the problems associated with sonication (overheating and emulsification) are eliminated. Thus, enzymatic shearing is not only simpler, it is also easier to optimize shearing conditions and to get more reproducible shearing from prep to prep, which improves your ChIP results.

Companion Products

Please visit our website for complete information on the items below; Catalog Nos. of these and other useful products are listed in the Related Products section in the Appendix of this manual:

Because appropriate controls make ChIP interpretation and troubleshooting easier, Active Motif offers its ChIP-IT Control Kits for human, mouse and rat samples. These useful kits contain positive & negative control antibodies and species-specific positive control PCR primers. The kits are invaluable when validating antibodies for use in ChIP.

Validated ChIP Control qPCR primer sets are also available separately for use as positive and negative controls with many of the more common ChIP targets to confirm the success of the ChIP reactions. Use of our primer sets will save you the time and effort required to synthesize and test your own gene/species-specific control primers.

Active Motif’s Ready-to-ChIP Chromatin is another useful reagent that will save you time. These high-quality chromatin samples are available from a variety of different cell types (HeLa, Hep G2, K-562 & NIH/3T3 cells); they have been sheared by sonication and are ready for use in ChIP.

For users who need to make more sheared chromatin samples than is possible with the reagents provided in the ChIP-IT Express Kits, ChIP-IT Express Shearing Kits are also available separately.

If you are using an isotype of mouse monoclonal antibody that does not bind well to protein G, consider our Bridging Antibody for Mouse IgG; it binds with a strong affinity to both the protein G beads and the mouse primary antibody. This maximizes capture of mouse antibody-immune complexes, which improves results of ChIP and IP experiments that use mouse primary antibody.

Finally, one difficult aspect of ChIP is finding an antibody that recognizes the target protein when it is bound to DNA and fixed by formaldehyde. Antibodies that perform well in Western blotting or other applications may not work well in ChIP. For this reason, Active Motif offers an ever-increasing number of ChIP-validated antibodies that have been verified to work in ChIP. See Appendix – Section G of this manual or go to www.activemotif.com/chipabs to generate an up-to-date list of antibodies that will help make your ChIP successful.

Active Motif’s ChIP-IT High Sensitivity Kit is ideal for use when studying low abundance transcription factor targets, working with antibodies with sub-optimal binding affinities, or when starting ChIP with a limited number of cells. This assay provides higher sensitivity to overcome these
obstacles. The ChIP-IT High Sensitivity Kit is compatible with the ChIP-IT qPCR Analysis Kit which can be used to simplify qPCR analysis and to enable normalization of data across multiple sample types and experiments.

The ChIP-IT Express HT was designed for users who have many ChIPs to perform. It enables true high-throughput ChIP by providing you with the reagents and protocols needed to adapt the magnetic bead-based ChIP-IT Express Kit method to a format that makes possible 96-well ChIP.

A common experimental objective is to establish that two epigenetic marks or chromatin-associated proteins are present at the same locus in a single chromatin sample. This is done by sequential ChIP, where the sample is subjected to a second ChIP with a different antibody. Because sequential ChIP can be a complicated process, Active Motif offers its Re-ChIP-IT Kit, which uses the methodology developed for ChIP-IT-Express to simplify the technique of sequential ChIP.

For researchers looking to combine ChIP with genome-wide analysis using Next-Generation sequencing (ChIP-Seq), Active Motif offers our ChIP-IT ChIP-Seq assay. This kit includes proven reagents, streamlined protocols and validation controls to perform the ChIP enrichment, validate the ChIP reaction and prepare sequencing libraries for use in next-generation sequencing.

ChIP is traditionally performed using formaldehyde-fixed tissues or cultured cell lines. In order to utilize the valuable retrospective data that is available from clinical samples, Active Motif has developed the ChIP-IT FFPE Kit for use with formalin-fixed paraffin-embedded (FFPE) tissue blocks and histology slides.
Kit Components and Storage  
ChIP-IT Express Kit (Cat. No. 53008)

Please store each component at the temperature indicated in the table below. **Do not re-freeze the Protein G Magnetic Beads after you have received this kit.**

<table>
<thead>
<tr>
<th>Reagents</th>
<th>Quantity</th>
<th>Storage / Stability</th>
</tr>
</thead>
<tbody>
<tr>
<td>RNase A (10 µg/µl)</td>
<td>40 µl</td>
<td>-20°C for 6 months</td>
</tr>
<tr>
<td>5 M NaCl</td>
<td>200 µl</td>
<td>-20°C for 6 months</td>
</tr>
<tr>
<td>100 mM PMSF</td>
<td>475 µl</td>
<td>-20°C for 6 months</td>
</tr>
<tr>
<td>Proteinase K (0.5 µg/µl)</td>
<td>250 µl</td>
<td>-20°C for 6 months</td>
</tr>
<tr>
<td>Proteinase K Stop Solution</td>
<td>150 µl</td>
<td>-20°C for 6 months</td>
</tr>
<tr>
<td>Protease Inhibitor Cocktail (PIC)</td>
<td>2 x 100 µl</td>
<td>-20°C for 6 months</td>
</tr>
<tr>
<td>1X Lysis Buffer</td>
<td>16 ml</td>
<td>-20°C for 6 months</td>
</tr>
<tr>
<td>10X Glycine</td>
<td>33 ml</td>
<td>-20°C for 6 months</td>
</tr>
<tr>
<td>10X PBS</td>
<td>120 ml</td>
<td>-20°C for 6 months</td>
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<tr>
<td>Shearing Buffer</td>
<td>10 ml</td>
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</tr>
<tr>
<td>Elution Buffer AM2</td>
<td>1.6 ml</td>
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</tr>
<tr>
<td>Reverse Cross-linking Buffer</td>
<td>1.6 ml</td>
<td>-20°C for 6 months</td>
</tr>
<tr>
<td>ChIP Buffer 1</td>
<td>70 ml</td>
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<tr>
<td>ChIP Buffer 2</td>
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<tr>
<td>Protein G Magnetic Beads*</td>
<td>650 µl</td>
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<tr>
<td>Siliconized 1.7 ml microcentrifuge tubes</td>
<td>25</td>
<td>Room temperature</td>
</tr>
<tr>
<td>Bar Magnet</td>
<td>1</td>
<td>Room temperature</td>
</tr>
<tr>
<td>Mini Glue Dots</td>
<td>1 sheet</td>
<td>Room temperature</td>
</tr>
</tbody>
</table>

* The Protein G Magnetic Beads are shipped on dry ice, but **should not be re-frozen** by the customer. Upon receipt of this kit, the beads should be stored at 4°C.
Kit Components and Storage
ChIP-IT Express Shearing Kit (Catalog No. 53032)

This component list is for the stand-alone ChIP-IT Express Shearing Kit, which contains only shearing reagents. The component list for the complete ChIP-IT Express Kit is on the preceding page. Please store each component at the temperature indicated in the table below.

<table>
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<tr>
<th>Reagents</th>
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<th>Storage / Stability</th>
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<td>100 mM PMSF</td>
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<tr>
<td>Proteinase K (0.5 µg/µl)</td>
<td>250 µl</td>
<td>-20°C for 6 months</td>
</tr>
<tr>
<td>Proteinase K Stop Solution</td>
<td>150 µl</td>
<td>-20°C for 6 months</td>
</tr>
<tr>
<td>Protease Inhibitor Cocktail (PIC)</td>
<td>2 x 100 µl</td>
<td>-20°C for 6 months</td>
</tr>
<tr>
<td>1X Lysis Buffer</td>
<td>16 ml</td>
<td>-20°C for 6 months</td>
</tr>
<tr>
<td>10X Glycine</td>
<td>33 ml</td>
<td>-20°C for 6 months</td>
</tr>
<tr>
<td>10X PBS</td>
<td>120 ml</td>
<td>-20°C for 6 months</td>
</tr>
<tr>
<td>Shearing Buffer</td>
<td>10 ml</td>
<td>-20°C for 6 months</td>
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Additional materials required

- A ChIP-validated antibody directed against the protein of interest
- Dounce homogenizer with a small clearance pestle (e.g. Active Motif Catalog Nos. 40401 & 40415 with the tight-fitting pestle). Use of a homogenizer is strongly recommended for shearing chromatin by sonication and required for enzymatic shearing. Dounce homogenization greatly improves your chances for successful ChIP.
- Magnetic stand. You can assemble a magnetic stand using the provided bar magnet (see Appendix – Section E) or use commercially available stands (e.g. the Promega MagneSphere® Technology twelve-position Magnetic Separation Stand).
- 37% formaldehyde solution (formalin) with 10-15% methyl alcohol to prevent polymerization. We do not recommend paraformaldehyde.
- Phase contrast/tissue culture microscope and hemocytometer
- Phenol/chloroform (1:1) TE saturated pH 8 (DNA Purification, Molecular Biology Grade)
- 3 M Sodium Acetate pH 5.2 (purification of Input DNA and purification of sheared DNA prior to checking concentration by spectrophotometry or gel electrophoresis)
- 100% ethanol
- 70% ethanol
- DNase-free H₂O (purification of Input DNA)
- Rocking platform for culture plates
- Apparatus to rotate tubes end-to-end at 4°C (e.g. a Labquake from Barnstead/Thermolyne with a tube holder for 1.7 ml microcentrifuge tubes)
- Microcentrifuge (table top centrifuge 4°C) and microcentrifuge tubes
- Spectrophotometer for DNA quantitation
- Pipettors and tips (filter tips are recommended)
- Sonicator (e.g. Active Motif’s EpiShear™ Sonicator with a 1/8” probe (Catalog No. 53051) with the EpiShear™ Cooled Sonication Platform (Catalog No. 53080))
- Agarose gel electrophoresis apparatus
- Minimal cell culture media (growth media without serum)
- Cell scraper (rubber policeman)

Optional materials

- 8-well PCR strips (e.g. Thermo Fisher Part No. AB-0451)
- 1.7 ml siliconized Eppendorf tubes (e.g. Active Motif Catalog No. 53036)
- Chromatin IP DNA Purification Kit (e.g. Active Motif Catalog No. 58002)
ChIP-IT Express Experimental Design

PLEASE READ THE ENTIRE PROTOCOL BEFORE STARTING!

Points to consider:

• **Cell growth and chromatin preparation.** When planning an experiment, calculate the number of chromatin preparations you will require and determine the number of ChIP reactions you plan to perform on each chromatin preparation. Be sure to include the appropriate control ChIP reactions in your calculations. Also, note that if you wish to analyze the effect of particular compounds or culturing conditions on transcription factor/DNA interactions, you should prepare chromatin from control (untreated) cells as a reference sample.

• **Protein G-coated magnetic beads.** The supplied magnetic beads are ready to use once fully resuspended to form a homogeneous slurry. There is no need to pre-block the beads or pre-clear the sample. For best results, gently shake and roll the tube. The beads settle quickly, and therefore should be resuspended just before pipetting. **Protein G Magnetic Beads are shipped on dry ice, but should not be re-frozen by the customer. Upon receipt, the beads should be stored at 4°C.** The ChIP-IT Protein G Magnetic Beads are also sold separately (Catalog No. 53014).

• **Antibodies must be suitable for ChIP.** ChIP antibodies must recognize fixed, native protein that is bound to DNA and/or complexed with other proteins. Many antibodies that perform well in other applications do not perform in ChIP. Thus, ChIP performed with an antibody that has not been ChIP-validated must include appropriate controls (such as Active Motif’s RNA pol II antibody, Catalog No. 39097) and a negative control IgG (mouse IgG for mouse monoclonals and rabbit IgG for rabbit polyclonals) to validate the chromatin preparation and the ChIP methodology. For your convenience, Active Motif sells ChIP-IT Control Kits for human, mouse and rat samples; these kits contain positive and negative control antibodies, appropriate positive PCR primers, PCR buffer and loading dye (see Related Products in Appendix).

• **Siliconized tubes.** Perform the ChIP reactions in the provided siliconized 1.7 ml microcentrifuge tubes or in 8-well PCR strips. (Do not use these tubes for preparing chromatin or isolating Input DNA, as you will then not have enough of them for the ChIP reactions.)
• **Bar magnet.** The provided bar magnet can be used with the provided siliconized 1.7 ml microcentrifuge tubes or 8-well PCR strips (see Appendix – Section E for detailed instructions). Commercially available side-pulling magnetic stands (e.g. Promega MagneSphere® Technology twelve-position Magnetic Separation Stand – 1.7 ml microcentrifuge tube format) can also be used.

• **Resuspend solutions completely.** Thaw the PMSF and the Proteinase K Stop Solution at room temperature until fully dissolved, which takes about 30 minutes. Vortex gently and spin down briefly before use.

• **Maximum volume of chromatin.** Chromatin shearing (sonication) buffers usually contain detergents (e.g. 0.1% SDS and 0.5% sodium deoxycholate is typical). If you plan to use more than 60 µl sonicated chromatin in a ChIP reaction, use the 200 µl reaction volume in Table 1 (page 12). This will ensure that the detergent in the shearing buffer does not interfere with antibody binding. Do not use more than 60 µl of sheared chromatin in a 100 µl reaction.

• **Quantity of antibody.** Optimal results are typically achieved with 1-3 µg of antibody. However, this will vary according to the affinity of the antibody and the quality of the chromatin; you may need to use more of a particular antibody.

• **Safety precautions.** Formaldehyde and PMSF are highly toxic chemicals. Appropriate safety precautions (i.e. safety glasses, gloves and lab coat) should be used. Also, formaldehyde is highly toxic by inhalation and should be used only in a ventilated hood. Finally, chromatin sonication should be performed in a biosafety hood if the chromatin is extracted from biohazardous or infectious materials.
Protocols – Preparation of Sheared Chromatin

A. Cell Fixation & Shearing

This protocol describes fixation and sonication shearing of cells from one 15 cm plate (approximately 1.5 x 10^7 cells). (Appendix – Section D includes information on scaling the protocol for use with other amounts of cells.) This shearing protocol assumes that you have already optimized shearing conditions for your specific cell line and treatment. If you have not, do not use this protocol now as it does not prepare enough chromatin to test for optimal conditions. Instead, use the protocols in Appendix – Sections A-C, which use 3 plates of cells. This will generate sufficient chromatin to test multiple shearing conditions for your cell type (and treatment) to determine the optimal conditions. The optimized conditions can then be used with the protocols below.

Shearing tips: ChIP experiments usually require chromatin that has been sheared to a size of 200-1500 bp. In general, shearing efficiency is improved through the use of a small shearing volume and a V-bottom tube rather than a round-bottom tube. Also, note that shearing is inefficient if the chromatin sample becomes emulsified with air bubbles. This can be avoided by using lower shearing power and by turning the power up gradually. If a chromatin preparation becomes emulsified inadvertently, discontinue shearing and centrifuge the sample for 4 minutes at 8,000 rpm at 4°C in a microcentrifuge to remove trapped air. Finally, to prevent overheating and denaturation of chromatin, samples should be kept on ice as much as possible during shearing, and shearing should be performed discontinuously (i.e. sonicate for 20 seconds, then place on ice/water for 30 seconds, sonicate again for 20 seconds, etc.). If possible, shear while on ice.

Note: Several of the buffers used below require addition of PMSF and protease inhibitors (PIC). Thaw these reagents before starting the chromatin preparation (i.e. 30 minutes at room temperature), then add to the buffers immediately before use.

1. Grow cells to 70-80% confluency in one 15 cm plate. Stimulate cells as desired to activate the pathway of interest.

2. When cells are ready to harvest, freshly prepare the following solutions. The volumes below are calibrated to one 15 cm plate:
   a. **Fixation Solution:** Add 0.54 ml 37% formaldehyde to 20 ml minimal cell culture medium and mix thoroughly. Leave at room temperature.
   b. **1X PBS Solution:** Add 2.33 ml 10X PBS to 21 ml dH_2O, mix and place on ice.
   c. **Glycine Stop-Fix Solution:** Combine 1 ml 10X Glycine Buffer, 1 ml 10X PBS and 8 ml dH_2O. Mix well and leave at room temperature.
   d. **Cell Scraping Solution:** Add 0.6 ml 10X PBS to 5.4 ml dH_2O, mix and place on ice.

3. Pour medium off the cells and add 20 ml Fixation Solution to each plate. Incubate on a shaking platform for 10 minutes at room temperature.

   Note: In standard protocols, chromatin is fixed for 10 minutes prior to shearing. While these are typical fixation conditions, some antibody/chromatin combinations may work better with shorter fixation times.
4. Pour Fixation Solution off and wash by adding 10 ml ice-cold 1X PBS to each plate. Rock the plate for 5 seconds, then pour off the PBS.

5. Stop the fixation reaction by adding 10 ml Glycine Stop-Fix Solution to each of the plates. Swirl to cover, and then rock at room temperature for 5 minutes.

6. Wash each plate by pouring off the Glycine Stop-Fix Solution, then adding 10 ml ice-cold 1X PBS. Rock the plate for 5 seconds, then pour off the PBS.

7. Just before use, add 30 µl 100 mM PMSF to Cell Scraping Solution. Add 5 ml of this ice-cold Cell Scraping Solution to each plate and scrape cells with a rubber policeman. Hold the plate at an angle and scrape cells down to collect them at the bottom edge of the plate. Use a 1 ml pipette to transfer the cells to a 15 ml conical tube on ice.

8. Pellet the cells from step 7 by centrifugation for 10 minutes at 2,500 rpm (720 RCF) at 4°C.

9. Remove the supernatant and discard. At this point the protocol can be continued or the pellet can be frozen. If freezing the pellet, add 1 µl 100 mM PMSF and 1 µl PIC and freeze at -80°C.

Shearing by Sonication

The section below describes the isolation and preparation of chromatin using sonication shearing.

1. Thaw pellet (if necessary) on ice and resuspend cells in 1 ml ice-cold Lysis Buffer supplemented with 5 µl PIC + 5 µl PMSF. Incubate on ice for 30 minutes.

2. Transfer the cells to an ice-cold dounce homogenizer. Dounce on ice with 10 strokes to aid in nuclei release.

   **Monitor Cell Lysis:** To ensure cell lysis, take 10 µl of the cell lysate from the dounce and look at it under a phase contrast microscope using a hemocytometer to verify that the nuclei have been released. It is often helpful to look at the cells before and after the lysis step as this makes it easier to identify the nuclei versus whole cells. Intact cells should have a dark central region (nucleus) surrounded by a halo of less dense cytoplasm. In lysed cells, the nuclei will appear as dots surrounded by asymmetric debris. If the cells are not lysed, then dounce on ice with an additional 10 strokes, or until the cells are lysed.

3. Transfer cells to a 1.7 ml microcentrifuge tube and centrifuge for 10 minutes at 5,000 rpm (2,400 RCF) in a 4°C microcentrifuge to pellet the nuclei.

4. Carefully remove the supernatant and discard. Resuspend the nuclei pellet in 350 µl Shearing Buffer (supplemented with 1.75 µl PIC and 1.75 µl PMSF) and place the samples on ice. It may be necessary to briefly dounce the nuclei to resuspend them.

5. Shear the DNA with your sonicator using the conditions that were previously determined to provide optimally sheared chromatin for your cell line (Appendix – Sections A-C).

6. Centrifuge the sheared chromatin samples for 10 minutes at 15,000 rpm (18,000 RCF) in a 4°C microcentrifuge. Carefully transfer supernatant to a fresh 1.7 ml microcentrifuge tube. This is the sheared chromatin. It can be used right away or stored at -80°C. Before freezing, remove 50 µl for use in assessing the efficiency of your DNA shearing and determining the DNA concentration. The remaining chromatin (approximately 350 µl) is sufficient for up to 6 ChIP reactions and should be aliquoted before freezing to minimize freeze-thaw cycles.
Note: Use the 50 µl sample removed above in the DNA Clean Up protocol in Appendix – Section C to check the DNA concentration and confirm that the chromatin has been sheared. You will need the concentration to set up you ChIP reactions.

Protocols – Chromatin Immunoprecipitation

B. Immunoprecipitation

1. Thaw chromatin (if necessary) on ice. Transfer 10 µl to a microcentrifuge tube; this tube is the “Input DNA” that will be processed in Step D6 and then be used as a control in PCR analysis. Store this sample at 4°C if it will be used within 6 hours; otherwise, store it at -20°C.

2. Set up the ChIP reactions by adding the components shown in Table 1 below to the provided siliconized 1.7 ml microcentrifuge tubes, or to PCR tubes. Be sure to use the DNA concentration that was determined for your sheared chromatin sample to calculate the volume to use. Before pipetting the magnetic beads, they should be fully resuspended by inverting and/or vortexing the bottle. The antibody should be the final component added to the reaction.

<table>
<thead>
<tr>
<th>Reagent</th>
<th>One reaction (if using less than 60 µl of chromatin)</th>
<th>One reaction (if using more than 60 µl of chromatin)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Protein G Magnetic Beads</td>
<td>25 µl</td>
<td>25 µl</td>
</tr>
<tr>
<td>ChIP Buffer 1</td>
<td>10 µl</td>
<td>20 µl</td>
</tr>
<tr>
<td>Sheared Chromatin (7-25 µg)*</td>
<td>20-60 µl</td>
<td>61-100 µl</td>
</tr>
<tr>
<td>Protease Inhibitor Cocktail (PIC)</td>
<td>1 µl</td>
<td>2 µl</td>
</tr>
<tr>
<td>dH₂O</td>
<td>Add enough so that the final reaction volume will be 100 µl</td>
<td>Add enough so that the final reaction volume will be 200 µl</td>
</tr>
<tr>
<td>Antibody (added last)</td>
<td>1-3 µg</td>
<td>1-3 µg</td>
</tr>
<tr>
<td>Total Volume</td>
<td>100 µl</td>
<td>200 µl</td>
</tr>
</tbody>
</table>

*Note: Depending on the application, ChIP can be performed using from 1-50 µg of chromatin. An important factor is the volume of the chromatin, especially if the chromatin was prepared by sonication, as the detergents used during sonication will decrease antibody binding. Use the 200 µl ChIP reaction shown in the right column above if the volume of chromatin will be > 60 µl. See the Appendix for discussions on the amount of chromatin to use, and for methods to quantify DNA in chromatin.

3. Cap tube and incubate on an end-to-end rotator for 4 hours at 4°C. In some cases, sensitivity may be improved if the incubation is performed overnight.

4. Spin tube briefly to collect liquid from the inside of the cap.
5. Place tube on magnetic stand to pellet beads on the tube side.
6. Carefully remove and discard supernatant.

C. Wash Magnetic Beads

Note: Do not allow the beads to “dry out”. Allow no more than 1 minute to elapse between removing buffer and then adding the next wash or the elution buffer. For suggestions regarding bead washing methods, see Appendix – Section E.

For 1.7 ml microcentrifuge tubes:
1. Wash beads one time with 800 µl ChIP Buffer 1.
2. Wash beads two times with 800 µl ChIP Buffer 2.
3. After the final wash, remove as much supernatant as possible without disturbing the beads. Use a 200 µl Pipetteman if necessary.

For 8-well PCR strips:
1. Wash beads three times with 200 µl ChIP Buffer 1.
2. Wash beads two times with 200 µl ChIP Buffer 2. After the final wash, remove as much supernatant as possible without disturbing the beads.

D. Elute Chromatin, Reverse Cross-links and Treat with Proteinase K

1. Resuspend the washed beads with 50 µl Elution Buffer AM2.
2. Incubate 15 minutes at room temperature on an end-to-end rotator.
3. Briefly spin tubes to collect liquid from caps.
4. Add 50 µl of the Reverse Cross-linking Buffer to the eluted chromatin and mix by pipetting up and down. Place tubes in magnetic stand; allow beads to pellet to the sides of the tubes.
5. Transfer the supernatant, which contains the chromatin, to a fresh tube.
6. It is now time to process the “Input DNA” sample: take the 10 µl Input DNA aliquot (that was set aside in Step B1 above) and thaw on ice, if needed. Add 88 µl ChIP Buffer 2 and 2 µl 5M NaCl to the Input DNA sample only, so that its final volume is 100 µl.
7. Incubate the ChIP and Input DNA samples at 95°C for 15 minutes in a thermocycler.

Note: If you are using larger, thicker-walled microcentrifuge tubes, perform a 2.5 hour incubation at 65°C. More proteinaceous samples may need a longer incubation time. The sample may be stored at -20°C at this point.

8. Return tubes to room temperature, spin the tubes briefly if liquid has collected on the inside of the caps, then add 2 µl Proteinase K.
9. Cap the tubes, mix well and incubate at 37°C for 1 hour. During this incubation, place the Proteinase K Stop Solution at room temperature for 30 minutes to 1 hour.
10. Return the tubes to room temperature and add 2 µl Proteinase K Stop Solution. Briefly centrifuge the tubes to collect liquid from the caps. The DNA can be used immediately in PCR or stored at -20°C.

**Protocols – PCR Analysis**

**E. End Point PCR**

The protocol below is a guideline for optimizing PCR analysis of DNA collected by ChIP. In order to obtain reliable comparisons of the DNA amounts collected in different ChIP reactions, end point PCR must be stopped during the linear phase of amplification. As this window of amplification differs between samples and PCR primer sets, the correct number of PCR cycles must be determined empirically. Real-time PCR analysis simplifies these considerations, as it is possible to determine the Ct value of each sample, representing the cycle number at which linear amplification begins. Due to these advantages, we advise that you use real-time PCR whenever possible.

PCR should be performed on four DNA templates: DNA from ChIP performed using positive and negative control antibodies (such as the positive control RNA pol II antibody and Negative Control IgG supplied in Active Motif’s ChIP-IT Control Kits), the Input DNA sample and DNA from ChIP with the test antibody. A water-only control should also be performed to ensure the PCR reagents are not contaminated.

**Design of the primers**

Analyze your potential primer pairs using an *in silico* PCR program (*i.e.* the UCSC Genome Browser at http://genome.cse.ucsc.edu/cgi-bin/hgPcr) to ensure that the primers selected will produce a single amplicon from the genomic DNA of the species from which the DNA is being amplified. Ideally, the amplicons should be 150-400 bp in length for standard PCR. Use of PCR design programs can be helpful in selecting good primer pairs.

**Note:** PCR is extremely sensitive and all precautions should be taken to guard against contamination. Gloves should be worn and filter-tip pipettes should be used.

**Setting up and performing PCR**

In the example below, PCR reactions are set up using 2 different PCR cocktails, which contain positive & negative control PCR primer sets. If you are using a positive control antibody from one of Active Motif’s ChIP-IT Control Kits, use only positive control PCR primers with that antibody; our positive control antibodies were chosen because they bind to many regions in the genome, making it impractical to design “negative control” primers for our positive control antibodies. For analysis of ChIP performed with other antibodies, it may be possible, depending on the antibody, to include both positive and negative PCR primer sets that are appropriate for that antibody.

1. Program the thermocycler. The program should start with an initial melt step at 94°C for 3 minutes, then 30-36 cycles of [94°C for 20 seconds, 59°C for 30 seconds and 72°C for 30 seconds], then a hold cycle at 10°C. The total volume of each PCR will be 25 µl. 36 cycles is a good starting point; you may need to optimize the number of cycles for your system.
2. **IMPORTANT:** Dilute the Input DNA sample 1:10 by adding 20 µl Input DNA to 180 µl dH₂O.

3. Use the table below to label PCR tubes and add the PCR templates and water-only control, keeping the tubes on ice. Add the PCR cocktails you will make in Step 4:

<table>
<thead>
<tr>
<th>Reaction No.</th>
<th>PCR Template (5 µl each)</th>
<th>PCR cocktail (20 µl each)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>ChIP DNA – Positive control antibody</td>
<td>Positive PCR cocktail</td>
</tr>
<tr>
<td>2</td>
<td>ChIP DNA – Negative control IgG</td>
<td>Positive PCR cocktail</td>
</tr>
<tr>
<td>3</td>
<td>Input DNA (diluted 1:10)</td>
<td>Positive PCR cocktail</td>
</tr>
<tr>
<td>4</td>
<td>ChIP DNA – Test antibody</td>
<td>Positive PCR cocktail</td>
</tr>
<tr>
<td>5</td>
<td>H₂O (no DNA control)</td>
<td>Positive PCR cocktail</td>
</tr>
<tr>
<td>6</td>
<td>ChIP DNA – Positive control antibody</td>
<td>Negative PCR cocktail</td>
</tr>
<tr>
<td>7</td>
<td>ChIP DNA – Negative control IgG</td>
<td>Negative PCR cocktail</td>
</tr>
<tr>
<td>8</td>
<td>Input DNA (diluted 1:10)</td>
<td>Negative PCR cocktail</td>
</tr>
<tr>
<td>9</td>
<td>ChIP DNA – Test antibody</td>
<td>Negative PCR cocktail</td>
</tr>
<tr>
<td>10</td>
<td>H₂O (control)</td>
<td>Negative PCR cocktail</td>
</tr>
</tbody>
</table>

4. Set up the Positive PCR cocktail and the Negative PCR cocktail on ice according to the tables below. Add the dH₂O first and the Taq polymerase last. Mix thoroughly and keep on ice. This ensures that the reaction mixture is inactive until the cycling is started. As discussed above, in Active Motif’s ChIP-IT Control Kits only positive control PCR primers are provided with the positive control antibody and Negative control IgG. (These are provided as a mixture of forward and reverse primers, so use 4 µl). However, for your test antibody we recommend to design and test both positive and negative PCR primer sets, if possible.

**Positive PCR cocktail:**

<table>
<thead>
<tr>
<th>Reagent</th>
<th>1 reaction</th>
<th>5 reactions</th>
</tr>
</thead>
<tbody>
<tr>
<td>DEPC H₂O</td>
<td>12.3 µl</td>
<td>61.5 µl</td>
</tr>
<tr>
<td>Positive Forward primer (5 pmol/µl)</td>
<td>2.0 µl</td>
<td>10 µl</td>
</tr>
<tr>
<td>Positive Reverse primer (5 pmol/µl)</td>
<td>2.0 µl</td>
<td>10 µl</td>
</tr>
<tr>
<td>dNTP mixture (5 mM each dNTP)</td>
<td>1.0 µl</td>
<td>5.0 µl</td>
</tr>
<tr>
<td>10X PCR Buffer</td>
<td>2.5 µl</td>
<td>12.5 µl</td>
</tr>
<tr>
<td>Taq (5 U/µl)</td>
<td>0.2 µl</td>
<td>1.0 µl</td>
</tr>
<tr>
<td><strong>Total Volume</strong> (Not including DNA template)</td>
<td>20 µl</td>
<td>100 µl</td>
</tr>
</tbody>
</table>
Negative PCR cocktail:

<table>
<thead>
<tr>
<th>Reagent</th>
<th>1 reaction</th>
<th>5 reactions</th>
</tr>
</thead>
<tbody>
<tr>
<td>DEPC H₂O</td>
<td>12.3 µl</td>
<td>61.5 µl</td>
</tr>
<tr>
<td>Negative Forward primer (5 pmol/µl)</td>
<td>2.0 µl</td>
<td>10 µl</td>
</tr>
<tr>
<td>Negative Reverse primer (5 pmol/µl)</td>
<td>2.0 µl</td>
<td>10 µl</td>
</tr>
<tr>
<td>dNTP mixture (5 mM each dNTP)</td>
<td>1.0 µl</td>
<td>5.0 µl</td>
</tr>
<tr>
<td>10X PCR Buffer</td>
<td>2.5 µl</td>
<td>12.5 µl</td>
</tr>
<tr>
<td>Taq (5 U/µl)</td>
<td>0.2 µl</td>
<td>1.0 µl</td>
</tr>
<tr>
<td><strong>Total Volume (Not including DNA template)</strong></td>
<td>20 µl</td>
<td>100 µl</td>
</tr>
</tbody>
</table>

5. Add 20 µl of the appropriate PCR cocktail to each of the 5 µl PCR templates (on ice) prepared in Step 3, for a total volume of 25 µl. Mix each reaction by pipetting up and down. Cap PCR tubes carefully and ensure that each reaction mixture is in the bottom of the tube.

6. Place PCR tubes in thermocycler and start the PCR program described in Step 1. After the cycles are complete, remove the tubes and place on ice.

7. These PCR reactions can be immediately analyzed as described below, or stored at -20°C.

**Analysis of PCR products**

1. Run ~8 µl of each PCR product on a 3% agarose gel. Save remaining PCR product in case additional gels must be run. Use gel combs with 2.5 mm-wide wells.

2. PCR products obtained with the human GAPDH positive control primers are 166 bp; those from the mouse EF1α primers are 233 bp, while the rat actin primers produce 223 bp PCR products. Use either a 50 or 100 bp ladder as the migration standard. Run the gel until the PCR products are well separated from the primers and primer dimers. Stain gel and analyze.

Figure 2: PCR of chromatin immunoprecipitation performed on 100,000 cells using ChIP-IT Express.

HeLa cells were fixed for 10 minutes with 1% formaldehyde and then chromatin was prepared by sonication shearing (5 pulses). ChIP was performed in duplicate on chromatin isolated from 100,000 and 750,000 cells using a Negative Control IgG and an RNA pol II antibody. The DNA isolated through these ChIP reactions was then analyzed by 36 cycles of PCR using GAPDH positive control primers. (These antibodies and primers are available as the ChIP-IT Control Kit – Human. Kits for mouse and rat are also available; see Related Products.) Ten µl of each PCR was separated on a 1% agarose gel and visualized by UV-illumination following ethidium bromide staining. PCR using the GAPDH primers on DNA isolated with the RNA pol II antibody reproducibly generated more product than similar reactions performed on DNA isolated using the Negative Control IgG. These results demonstrate that ChIP performed with RNA pol II antibody greatly enriched for GAPDH promoter DNA, while ChIP performed with negative IgG did not.
F. Real-time PCR

Following the final elution, cross-link reversal and proteinase K digestion of the immunoprecipitated chromatin, the samples should be subjected to a DNA clean-up step prior to certain downstream applications. For real-time PCR, we suggest using the Active Motif Chromatin IP DNA Purification Kit (Catalog No. 58002) prior to amplification. These columns yield 50 µl; 2 µl will be used for each PCR, giving you enough DNA for 25 PCR reactions. We suggest that you use a commercially available SYBR Green PCR Kit and a real-time thermocycler.

A. Design of the primers

• Analyze your potential primer pairs using an in silico PCR program (i.e. the UCSC Genome Browser at http://genome.cse.ucsc.edu/cgi-bin/hgPcr). Please note that primer sets that have worked in end point PCR may not work in real-time PCR.

• Primers that dimerize should be avoided, as they will be bound by SYBR Green, which will compromise accurate quantitation. You can test your primers for self-complementarity and secondary structure at http://frodo.wi.mit.edu/cgi-bin/primer3/primer3_www.cgi.

• Ideally, the amplicons should be 50-150 bp in length for real-time PCR.

• G/C stretches at the 3´ ends of primers should be avoided.

• The difference in melting temperature between the forward and reverse primers should not exceed 3°C.

B. Generation of a standard curve to accurately determine fold enrichment

Note: This step is not essential, but many downstream calculations will rely on a standard curve.

1. To test the efficiency of your primers, produce a standard curve by performing qPCR with your primer set on known DNA quantities of Input DNA in triplicate. Run three to five samples that are 10-fold dilutions, e.g. 0.005 ng, 0.05 ng, 0.5 ng, 5 ng and 50 ng.

2. Run the ChIP and IgG samples along with the dilution series of the Input DNA standards. Every primer set will have a different amplification profile, so the Ct values can then be plotted to create a linear regression plot. This may not be necessary as most real-time thermocyclers will create a standard curve automatically.

3. Ct = Threshold Cycle (cycle number where the signal exceeds the background threshold level).

4. Plot Ct vs. DNA quantity (log scale) of the dilutions to produce the standard curve (Figure 3).

Two methods for calculating the fold enrichment using the slope of the standard curve. The first method is sufficient for most circumstances, but an optional second method is presented as well.
Method 1:
1. This first method entails (a) solving for the DNA quantity of the ChIP and IgG samples, then (b) calculating the Fold Enrichment of the ChIP sample relative to the IgG sample:
   
   \[ X = \text{DNA quantity} \]
   \[ Y = \text{Ct} \]
   \[ M = \text{slope of the standard curve line.} \]
   \[ B = \text{the Ct value where } X = 1, (\text{e.g. 27.46 cycles in Figure 3}) \]
   
   ChIP Ct = 22.77 cycles (from Figure 3)
   IgG Ct = 30.22 cycles (from Figure 3)

   (a) \[ Y = M (\log X) + B \]
   \[ \log(X) = (Y - B) ÷ M \]

   (b) Fold Enrichment = ChIP DNA quantity ÷ IgG DNA quantity

   Using the data from Figure 3, calculations are performed as follows:
   Solve for X using the known values of the slope, y-intercept, ChIP Ct and IgG Ct:
   
   (a) \[ \log(X) = (Y - B) ÷ M \]
   \[ i.e.: \log(\text{DNA quantity}) = (\text{Ct} - \text{y-int}) ÷ \text{slope} \]
   
   For the ChIP sample, \[ \log(X) = (22.77 - 27.463) ÷ -3.508 \], so \[ X = 21.767 \text{ ng} \]
   
   For the IgG sample, \[ \log(X) = (30.22 - 27.463) ÷ -3.508 \], so \[ X = 0.1637 \text{ ng} \]

   (b) Solve the Fold Enrichment using the calculated DNA quantities:
   \[ 21.767 \text{ ng} ÷ 0.1637 \text{ ng} = 133 \text{ fold enrichment} \]

Method 2:
2. This second method calculates enrichment as a ratio of the amplification efficiency of the ChIP sample over that of the IgG.

   The efficiency of the primers can be calculated using the slope of the standard curve and the following formula:
   
   \[ \% \text{Efficiency} = [10(-1/slope) - 1] x 100\% \]

   In Figure 3, the slope = -3.508
   
   \[ \% \text{Efficiency} = [10(-1/-3.508) - 1] x 100\% = 92.8\% \]

   The ideal efficiency is 100% ±10% (i.e. 90-110%, where -3.32 = 100% efficiency). In this example, the primer set falls within the acceptable range. Therefore, the amplification efficiency of the primer can be used to determine the amplification efficiency of the ChIP sample and the IgG sample.

   \[ \text{AE} = \text{amplification efficiency (is a factor of the primer efficiency)} = 10^{(-1/\text{slope})} \]
   \[ \text{Fd} = \text{dilution factor (i.e. 1)} \]

   (a) \[ \% \text{ChIP} = \text{AE}^{(\text{Input Ct} - \text{ChIP Ct})} \times \text{(Fd)(100)} \]
   (b) \[ \% \text{IgG} = \text{AE}^{(\text{Input Ct} - \text{IgG Ct})} \times \text{(Fd)(100)} \]
   (c) Fold Enrichment = \% ChIP ÷ \% IgG
Using the data from Figure 3, calculations are performed as follows:

Slope = -3.508 (from Figure 3)

\[ AE = 10^{\frac{-1}{\text{slope}}} = 10^{\frac{-1}{-3.508}} = 1.928 \]

Input Ct = 21.36 cycles (use the value of the Input DNA standard closest to the ChIP sample, which is the 50 ng Input DNA sample in Figure 3)

ChIP Ct = 22.77 cycles (from Figure 3)

IgG Ct = 30.22 cycles (from Figure 3)

(a) \%

\[ \% \text{ ChIP} = (1.928(21.36 - 22.77))(1)(100) \]

\[ \% \text{ ChIP} = (1.928(-1.41))(1)(100) \]

\[ \% \text{ ChIP} = 39.62\% \]

(b) \%

\[ \% \text{ IgG} = (1.928(21.36 - 30.22))(1)(100) = 0.3\% \]

\[ \% \text{ IgG} = (1.928(-8.86))(1)(100) \]

\[ \% \text{ IgG} = 0.3\% \]

(c) Solve the Fold Enrichment using the calculated amplification efficiencies:

39.62\% + 0.3\% = 132 fold enrichment
Figure 3: Standard curve produced from ten-fold dilutions of Input DNA.

Five 10-fold dilutions of Input DNA were qPCR amplified in triplicate along with the ChIP and IgG samples using GADPH primers. The values for each amount of Input DNA were plotted and used to produce the standard curve. Its slope and y-intercept values were used with the Ct values of the samples to calculate the fold enrichment.

Figure 4: ChIP enrichment of GAPDH promoter DNA using RNA pol II antibody and Negative Control IgG.

HeLa cells were fixed for 10 minutes with 1% formaldehyde and then chromatin was prepared by sonication shearing (5 pulses). ChIP was performed using ChIP-IT Express on chromatin isolated from 750,000 cells using a Negative Control IgG and an RNA pol II antibody (Catalog No. 39097). Real-time PCR was performed on DNA purified from each of the ChIP reactions using a primer pair specific for the GAPDH gene. (These antibodies and primer set are available together in the ChIP-IT Control Kit – Human (Catalog No. 53010)). These results demonstrate that ChIP performed with RNA pol II antibody greatly enriched for GAPDH promoter DNA, while ChIP performed with negative IgG did not.
Appendix

Optimizing the Shearing Conditions

Chromatin shearing conditions can vary significantly depending on the cell type and, occasionally the cell culture & cell stimulation conditions. However, after shearing has been optimized for a given cell type, those conditions usually give consistent results with that cell type. For this reason, we recommend use of the following protocols to determine the optimal shearing conditions the first time you make chromatin from a cell line. This requires you to grow and work with three times the number of cells for optimization than you will need for subsequent ChIP experiments, where you will use your optimized conditions on a single plate of cells.

Section A. Cell Fixation to Optimize Shearing Conditions

In the protocols below, chromatin is prepared from cells grown in three 15 cm plates (approximately 4.5x 10^7 cells) and the chromatin is sheared using 3 different conditions. Generally, at least one of these conditions yields chromatin suitable for use in ChIP. Because only 50 µl of each sample is used for shearing analysis, one or more of them (usually ~300 µl for each shearing condition) may be used for approximately 6 ChIP experiments, provided that protease inhibitor cocktail (PIC) and PMSF are included in the buffers. Use of PIC and PMSF during optimization will, however, reduce the number of shearing and ChIP reactions that can be performed once you have established optimal shearing conditions and are ready to perform ChIP.

Note: If you wish to use the unused chromatin in ChIP, thaw the PIC and PMSF before starting the protocol and add to the buffers immediately before use. Freeze the unused chromatin during shearing analysis. If you are performing optimization simply to identify shearing parameters, do not add PIC and PMSF to the buffers. This will conserve these reagents so that you can prepare 10 samples of sheared chromatin using your optimized conditions, then perform 25 ChIP reactions.

1. Grow cells to 70-80% confluency on three 15 cm plates. If applicable, treat all three plates equally to influence the pathway of interest, as they will be combined into a single sample.

2. When cells are ready to harvest, prepare fresh Fixation Solution, ice-cold 1X PBS, Glycine Stop-Fix Solution and Cell Scraping Solution as follows:
   a. **Fixation Solution:** Add 1.62 ml of 37% formaldehyde to 60 ml minimal cell culture medium and mix thoroughly. Leave at room temperature.
   b. **1X PBS:** Add 7 ml 10X PBS to 63 ml dH₂O, mix and place on ice.
   c. **Glycine Stop-Fix Solution:** Combine 3 ml 10X Glycine Buffer, 3 ml 10X PBS and 24 ml dH₂O. Mix well and leave at room temperature.
   d. **Cell Scraping Solution:** Add 1.8 ml 10X PBS to 16.2 ml dH₂O, mix and place on ice.

3. Pour medium off the three plates and add 20 ml Fixation Solution to each plate. Incubate on a shaking platform for 10 minutes at room temperature.
Note: In standard protocols, chromatin is fixed for 10 minutes prior to shearing. While these are typical fixation conditions, some antibody/chromatin combinations may work better with shorter fixation times.

4. Pour Fixation Solution off the plates and wash by adding 10 ml ice-cold 1X PBS to each plate. Rock the plate for 5 seconds, then pour off the PBS.

5. Stop the fixation reaction by adding 10 ml Glycine Stop-Fix Solution to each plate. Swirl to cover the cells and then rock at room temperature for 5 minutes.

6. Wash each plate by pouring off the Glycine Stop-Fix Solution, then adding 10 ml ice-cold 1X PBS. Rock the plate for 5 seconds, then pour off the PBS.

7. Just before use, add 90 µl 100 mM PMSF to Cell Scraping Solution. Add 5 ml of this ice-cold Cell Scraping Solution to each plate and scrape cells with a rubber policeman. Hold the plate at an angle and scrape cells down to collect them at the bottom edge of the plate. Use a 1 ml pipette to transfer the cells to a 15 ml conical tube on ice. Do the same for the other two plates, pooling the cells from all three plates in one tube.

8. Pellet the pooled cells by centrifugation for 10 minutes at 2,500 rpm (720 RCF) at 4°C.

9. Remove the supernatant and discard. At this point the protocol can be continued or the pellet can be frozen. If freezing the pellet, add 1 µl 100 mM PMSF and 1 µl PIC and freeze at -80°C. When you are ready, continue with Section B below.

Section B. Optimization of Chromatin Shearing by Sonication

Please read Shearing tips on page 10 before beginning. In the protocol below, chromatin is sheared for 5, 10 and 15 pulses. Generally, at least one of these conditions yields chromatin suitable for use in ChIP. Because only a fraction of each preparation is used for to analyze the shearing efficiency, all remaining optimal preparation(s) can be used for ChIP experiments, provided that PIC and PMSF are used in the buffers below, and those above for cell fixation.

Our sonication optimization protocol was developed using Active Motif’s EpiShear™ Probe Sonicator with a 1/8” probe in combination with an EpiShear™ Cooled Sonication Platform (Catalog No. 53080) to maintain probe height and temperature consistency between samples. Samples were processed at 25% power in a volume of approximately 300 µl.

1. Thaw pellet (if necessary) on ice and resuspend cells in 3 ml ice-cold Lysis Buffer (supplemented with 15 µl PIC + 15 µl PMSF). Incubate on ice for 30 minutes.

2. Transfer the cells to an ice-cold dounce homogenizer. Dounce on ice with 10 strokes to aid in nuclei release.

Monitor Cell Lysis: To ensure cell lysis, take 10 µl of the cell lysate from the dounce and look at it under a phase contrast microscope using a hemocytometer to verify that the nuclei have been released. It is often helpful to look at the cells before and after the lysis step as this makes it easier to identify the nuclei versus whole cells. Intact cells should have a dark central region (nucleus) surrounded by a halo of less dense cytoplasm. In lysed cells, the
nuclei will appear as dots surrounded by asymmetric debris. If the cells are not lysed then
dounce on ice with an additional 10 strokes, or until the cells are lysed.

3. Transfer cells to a 1.7 ml microcentrifuge tube and centrifuge for 10 minutes at 5,000 rpm
(2,400 RCF) in a 4°C microcentrifuge to pellet the nuclei.

4. Carefully remove the supernatant and discard. Resuspend the nuclei pellet in 1.0 ml Shearing
Buffer (supplemented with 5 µl PIC and 5 µl PMSF), aliquot into equal volumes into three
1.7 ml microcentrifuge tubes, then place on ice. Each aliquot should be approximately 300 µl.

5. Shear the three aliquots of fixed chromatin at 25% power using three different conditions:
   a. Five pulses of 20 seconds each, with a 30-second rest on ice between each pulse.
   b. Ten pulses of 20 seconds each, with a 30-second rest on ice between each pulse.
   c. Twenty pulses of 20 seconds each, with a 30-second rest on ice between each pulse.

6. Centrifuge the sheared chromatin samples for 10 minutes at 15,000 rpm (18,000 RCF) in a 4°C
microcentrifuge. Transfer the supernatants to fresh 1.7 ml microcentrifuge tubes and save
50 µl aliquots from each, which will be used to determine shearing efficiency. The sheared
chromatin and the aliquots can be stored at -80°C. Or, use the 50 µl aliquots immediately
in Section C below to reverse cross-links and purify the chromatin prior to gel analysis and
spectrophotometry.

Section C. DNA Clean Up to Assess Shearing Efficiency and
DNA Concentration

1. If necessary, thaw the 50 µl aliquots of each sheared chromatin sample.
2. Add 150 µl dH₂O, then 10 µl 5 M NaCl to each tube.
3. Heat all samples at 65°C in a water bath or a thermocycler for 4 hours to overnight to reverse
the cross-links, taking care to prevent the lids from popping open if you use a water bath.
4. Add 1 µl RNase A to each sample and incubate at 37°C for 15 minutes.
5. Add 10 µl Proteinase K to each sample and incubate at 42°C for 1.5 hours.

Note: If you intend to use a spectrophotometer to determine the DNA concentration,
the DNA must first be cleaned up. Column purification is not recommended as
the high protein content may clog the column. Therefore, the DNA should be
phenol/chloroform extracted and precipitated, which is performed as follows:

   a. Add 200 µl 1:1 phenol/chloroform TE saturated pH 8 to the sample, vortex to mix
      completely and centrifuge for 5 minutes at maximum speed in a microcentrifuge.
   b. Transfer the aqueous phase to a fresh microcentrifuge tube, then add 20 µl 3 M Sodium
      Acetate pH 5.2 and 500 µl 100% ethanol. Vortex to mix completely and place at -80°C
      for at least 1 hour. Alternatively, the sample can be left at -20°C overnight.
   c. Centrifuge at maximum speed for 10 minutes in a microcentrifuge at 4°C.
   d. Carefully remove and discard supernatant. Do not disturb the pellet.
e. Add 500 µl 70% ice cold ethanol without disturbing the pellet and spin for 5 minutes at maximum speed in a 4°C microcentrifuge.

f. Carefully remove and discard supernatant. Do not disturb pellet. Allow pellet to air-dry.

g. Resuspend pellet in 30 µl dH₂O and use a spectrophotometer to measure the absorbance at 260 nm to determine the DNA concentration (1.0 A₂₆₀ unit = 50 µg/ml). The DNA concentration of the aliquot can be used to back calculate the concentration of the sheared chromatin sample. **Optional, but recommended:** If you are performing ChIP on multiple chromatin samples, such as non-treated and treated samples, use the DNA concentrations so that the initial quantity of DNA is equal for all ChIP reactions (7-25 µg is recommended). This ensures that equal amounts of chromatin are used per IP, and that the relative differences between the treatment groups are comparable.

6. We recommended loading two different quantities of each sheared sample on the gel to ensure one falls within an acceptable range. Add 4 µl of a 6X Loading Buffer to 16 µl of sample, then load 5 µl & 10 µl of each sample on a 1% TAE agarose gel. Run the gel at 100V for 45 minutes to 1 hour, until the loading dye reaches 3/4 of the way to the end of the gel.

7. Optimal sonication shearing should result in a 200-1500 bp smear similar to that shown in lanes 2 or 3 of Figure 5 below.

8. If PIC and PMSF were used during the optimization experiment, the remaining chromatin of the preparation(s) that produced chromatin sheared to the appropriate length can be used in ChIP experiments.

**Figure 5: Gel analysis of sonication shearing (ChIP-IT Express).**

HeLa cells were fixed for 10 minutes with 1% formaldehyde and then chromatin was prepared using the ChIP-IT Express Kit. Three samples of approximately 300 µl each were sheared with 5, 10 and 20 pulses at 25% amplitude using the Active Motif EpiShear™ Probe Sonicator with a 1/8” probe. Each pulse consisted of a 20-second sonication followed by a 30-second rest on ice to prevent heat build up. The sheared and un-sheared chromatin samples were subjected to cross-link reversal, treated with Proteinase K and RNase A, then phenol/chloroform extracted and precipitated as described. Samples were separated by electrophoresis through a 1% agarose gel. Optimally sheared chromatin will yield a smear between 200-1500 bp.

- **Lane 1:** 100 to 1000 bp ladder.
- **Lane 2:** HeLa DNA sheared for 5 pulses (optimal).
- **Lane 3:** HeLa DNA sheared for 10 pulses (optimal).
- **Lane 4:** HeLa DNA sheared for 20 pulses (over-sheared).

**Note:** From this experiment, the DNA sonicated for both 5 and 10 pulses are suitable for use in ChIP.
Section D. Scale Up/Down of Chromatin Preparation

Our standard chromatin preparation protocols use cells grown in one 15 cm tissue culture plate (approximately $1.5 \times 10^7$ cells) and yield enough material to perform up to 6 ChIP reactions. Depending on your experiments, you may wish to work with different volumes of cells. The comments and table below are designed to help adapt our protocols for using different cell numbers.

- It is not recommended to use a sample with less than 500 µl or more than 2 ml of the Lysis Buffer for the cell lysis and dounce homogenization steps.
- If you intend to compare ChIP results from various samples, treat the samples equally. For example, grow induced and uninduced cells in the same size plate and to the same density, then use equal volumes and shearing conditions. This will help ensure that the chromatin preparations are equivalent in terms of cell numbers (genome equivalents), DNA shearing efficiency, etc. Quantify the DNA in the chromatin preparations (by following the DNA Clean Up protocol above) and then use equal mass quantities of chromatin in each ChIP assay.
- Each human diploid cell contains 6.6 picograms of DNA. This can be used to estimate DNA in a chromatin preparation if the number of cells in the starting material is known. We estimate DNA recovery of chromatin shearing to be about 60-70%, depending on the cell/tissue type.
- When using a sonicator with a 3 mm (or similar) probe, shearing should be performed in a 1.5 or 1.7 ml Eppendorf in 300-400 µl. Expect to lose 50 µl of chromatin (due to aerosolized sample). Smaller volumes (50-100 µl) will require a 1 mm probe for effective shearing.
- Perform sonication shearing so the chromatin produced is at an appropriate concentration for ChIP. ChIP reactions should be performed in a small volume (ideally 100 µl, and not more than 200 µl total volume). Because a 200 µl ChIP reaction contains 25 µl beads, 20 µl ChIP Buffer 1, and several µl of antibody and PIC, the maximum volume of chromatin is ~150 µl. The detergent in the chromatin preparation will negatively impact the ChIP, which limits how much chromatin can be added. Keep the volume of chromatin used for ChIP to no more than 100 µl per 200 µl ChIP reaction.

<table>
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<th>1 well of a 24-well plate</th>
<th>10 cm plate</th>
<th>15 cm plate</th>
<th>3 x 15 cm plates</th>
</tr>
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<tr>
<td>Number of Cells (tissue weight)</td>
<td>130,000 (-)</td>
<td>0.66 x $10^7$ (-)</td>
<td>1.5 x $10^7$ (-75 mg)</td>
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<td>Fixation Solution</td>
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<td>500 µl + 2.5 µl PMSF</td>
<td>1 ml + 5 µl PMSF</td>
<td>5 ml + 30 µl PMSF</td>
<td>15 ml + 90 µl PMSF (5 ml + 30 µl/plate)</td>
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<td>Pool the 3 plates</td>
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<tr>
<td>Lysis Buffer + PIC + PMSF</td>
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<td>500 µl + 2.5 µl PIC + 2.5 µl PMSF</td>
<td>1 ml + 5 µl PIC + 5 µl PMSF</td>
<td>3 ml + 15 µl PIC + 15 µl PMSF</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Pool the 3 plates</td>
</tr>
<tr>
<td>Shearing Buffer + PIC + PMSF</td>
<td>50-100 µl + 0.25-0.5 µl PIC + 0.25-0.5 µl PMSF</td>
<td>300 µl + 1.5 µl PIC 1.5 µl PMSF</td>
<td>350 µl + 1.75 µl PIC 1.75 µl PMSF</td>
<td>1000 µl + 5 µl PIC 5 µl PMSF</td>
</tr>
</tbody>
</table>
Section E. Use of Magnetic Beads and Included Bar Magnet

- The magnet should be stored in the provided tube.
- Be careful when working near metal objects or surfaces. A free magnet will jump great distances onto nearby metal surfaces with surprising speed. This can break the magnet.
- Use the provided Mini Glue Dots to attach the bar magnet to an empty pipette tip box to create an effective magnetic stand for use with either PCR strips or microcentrifuge tubes.
- If the magnet becomes attached to a flat metal surface, it should be removed by sliding it off the edge of the surface. The magnet may break if you attempt to pull one end or pry it away from the metal.

Caution: The included neodymium bar magnet is extremely powerful and is easily broken if handled incorrectly.

Creating a magnetic stand for 8-well PCR strips:

Note: 8-well strip tubes for use with standard 96-well PCR cyclers are recommended (e.g. Thermo Fisher AB-0451).

1. Place a strip of PCR tubes in the wells of an empty tip box (200 µl tips) and place the magnet directly against the tubes. This is the way the magnet will be positioned when the glue dots are used to affix it to the box.
2. Remove the covering tape from one side of two glue dots and attach the glue dots on the bar magnet (the uncovered face of the dot is placed on the magnet) as shown below.
3. Remove the covering tape from the exposed side of the glue dots. Fix the magnet to the tip box so that it is against the PCR tubes. The magnetic stand is now ready for use.

Note: Familiarize yourself with using the magnetic stand before performing with PCR tubes for the first time. Add 5 µl of magnetic beads to 100 µl ChIP Buffer 1 in one tube of an 8-well strip of PCR tubes. Use this tube with the assembled bar magnet stand to become familiar with use of the beads and magnet. It is difficult to re-suspend the beads if the tubes are directly adjacent to the magnet, so it is best to move the tubes away from the magnet for resuspension steps.
Washing should be performed as follows:

a. Place the tubes in the rack against the magnet and allow the beads to be pinned to the side of the tube, as shown below.

b. Remove supernatant with a 200 µl pipette or a 200 µl eight-channel pipette.

c. Move the tube strip into a row that is not adjacent to the magnet.

d. Add wash buffer and pipet up and down to fully re-suspend the beads. Ensure that a minimal amount of beads cling to the tips when the re-suspension is complete.

e. Repeat steps a-d until desired washing steps are complete.

Centrifugation of 8-well PCR strip tubes:
When working with 8-well PCR strip tubes, it may be desirable to centrifuge the tubes to collect the liquid and beads from the insides of the caps. This is easily accomplished in a centrifuge fitted with adaptors for spinning microtiter plates. Place a standard 96-well plate in the adaptor to hold the tubes in place. Be sure to balance the rotor (i.e. place a microtiter plate and tubes of appropriate mass in the rotor’s opposing 96-well plate adaptor). Spin the plates briefly to let the rotor reach a speed of 1000 x g before allowing the rotor to stop.

Creating a magnetic stand for 1.7 ml microcentrifuge tubes:
1. Remove the covering tape from one side of two glue dots.
2. Place two 1.7 ml microcentrifuge tubes in the wells of an empty tip box (1000 µl) and place the magnet directly against the tubes. This is the way the magnet will be positioned when the glue dots are used to affix it to the box.
3. Attach the glue dots on the bar magnet (the uncovered face of the dot is placed on the magnet) as shown above.

4. Remove the covering tape from the exposed side of the glue dot. Fix the magnet to the tip box so that it is against the tubes. The magnetic stand is now ready for use.

**Note:** 1.7 ml microcentrifuge tubes are held less securely in this assembled tube stand than in a typical commercial magnetic stand. This is not a problem if the below washing protocol is followed. Work with 1 tube at a time, and keep the tubes in the standard tube rack unless you are holding the tube next to the magnet.

Washing is best performed one tube at a time, as follows:

1. Place the tube in a standard 1.7 ml microcentrifuge tube rack and open the cap.
2. Place the opened tube in the assembled magnetic stand. The beads will pellet more rapidly if the bottom of the tube is held against the magnet, as shown below, and then slowly lowered into the well. This will pellet the beads up onto the side of the tube.

3. Allow the beads to pellet completely and remove supernatant with a 1000 µl pipette man. You can either leave the tube in the rack or pull it out when you remove the buffer. The beads will remain on the side of the tube, even when not next to the magnet.
4. Return the tube to the standard microcentrifuge tube rack, add 800 µl wash buffer and fully resuspend the beads by pipetting up and down.
5. Repeat steps 2-4 until desired washing steps are complete. After the final wash has been removed, the last traces of wash buffer should be removed with a 200 µl pipette man.

**Section F. Troubleshooting Guide**

<table>
<thead>
<tr>
<th>Problem/question</th>
<th>Recommendation</th>
</tr>
</thead>
<tbody>
<tr>
<td>At what points in the protocol can I stop?</td>
<td>The protocol may be stopped and samples stored at the times and temperatures below: 1. After formaldehyde fixation and centrifugation (intact cell pellet), -80°C. 2. After chromatin shearing, -80°C. 3. After the cross-link reversal, -20°C. 4. After DNA clean up, -20°C.</td>
</tr>
<tr>
<td>Problem/question</td>
<td>Recommendation</td>
</tr>
<tr>
<td>------------------</td>
<td>----------------</td>
</tr>
<tr>
<td>After sonication shearing and centrifugation, a viscous or cloudy layer is visible in the chromatin.</td>
<td>Depending upon the cell type, lipid or glycogen layers may be seen after centrifugation. For example, liver tissue may have a glycogen layer and a milky appearance, while fatty tissues may have a lipid layer. Avoid such layers when you remove the supernatant. However, if the whole supernatant is cloudy, it should not interfere with the IP reaction.</td>
</tr>
<tr>
<td>Poor yield of sheared chromatin.</td>
<td>Nuclei not released. It is highly recommended to perform dounce homogenization, even when using sonication. Use a dounce homogenizer with a small clearance pestle (see reference in Optional materials on page 7). Monitor cell lysis under a microscope. Generally, the more cells that are lysed, the higher the sheared chromatin yield. Decrease the fixation time. Over-fixed cells are often very resistant to lysis and shearing. Cross-linking for longer periods of time tends to cause cells to form into a giant cross-linked aggregate that is not sheared efficiently. Decrease the incubation time of the formaldehyde fixation step to 5 minutes. Sonication samples were emulsified. Avoid emulsification by turning up the power of the sonicator gradually. If a chromatin preparation becomes emulsified inadvertently, discontinue shearing and centrifuge the sample for 4 minutes at 8,000 rpm in a 4°C microcentrifuge to remove trapped air. Use fresh formaldehyde when preparing Fixation Solution. Buffers were not scaled proportionally to the size of the sample. Use the chart in Appendix – Section D to scale up or down chromatin preparation.</td>
</tr>
<tr>
<td>Shearing efficiency is not clear from gel analysis.</td>
<td>Material is stuck in the wells, and smears or streaks are seen from the top to bottom of the lane. The sheared chromatin needs to have the cross-links reversed, protein removed (Proteinase K) and RNA removed (RNase), followed by DNA purification using phenol/chloroform. Follow the DNA Clean Up protocol in Appendix – Section C. DNA purification columns are not recommended, as the high protein content may clog the columns. For samples with a lot of chromatin, you may need to reverse cross-links for a longer period (4 hours to overnight at 65°C). Lost DNA during the purification step. Phenol should be saturated with TE pH 8. Lower pH solutions will degrade the DNA. Column purification is not recommended due to the high protein content of the sample, which may clog the column. High molecular weight products. Decrease the size of the fragments by re-sonicating the sample.</td>
</tr>
<tr>
<td>Performing ChIP with a large volume of chromatin.</td>
<td>This is not recommended. It is better to set up several small ChIP reactions (200 µl each) and pool the samples at the end, rather than trying to ChIP a single large sample. Do not perform a single scaled-up reaction, as the capture efficiency is lower.</td>
</tr>
<tr>
<td>Poor or no enrichment with target antibody.</td>
<td>Too little chromatin. Generally, we recommend using 7-10 µg of chromatin for a “regular” ChIP (with highly abundant, DNA-associated targets such as histones). For moderately abundant transcription factors, 7-25 µg of chromatin is recommended; for very low abundance transcription factors, a maximum of 50 µg chromatin is recommended for each IP reaction. For small-scale ChIP, chromatin quantities have been reported as low as 1 µg, but this is largely dependant upon transcription factor abundance and the affinity strength of the antibody, and may need to be adjusted accordingly. Be sure to quantitate the concentration of the sheared chromatin sample(s) being ChIP’d to ensure both that adequate chromatin is used per sample, and that equal mass quantities of chromatin are used in each ChIP. Antibody is not ChIP validated. The antibody does not efficiently recognize fixed proteins, either because the epitope is destroyed by fixation or because the epitope is masked by other proteins in a larger complex. To assist in ChIP validating an antibody, it is very useful to use a positive control antibody such as RNA Pol II and a negative IgG from the same species, and primers that have been proven to work in the type of PCR being used.</td>
</tr>
<tr>
<td>Problem/question</td>
<td>Recommendation</td>
</tr>
<tr>
<td>------------------</td>
<td>----------------</td>
</tr>
<tr>
<td>Poor or no enrichment with target antibody (continued).</td>
<td>Low-affinity antibody. Increase the incubation time of the ChIP reaction to overnight at 4°C on an end-to-end rotator. Antibody affinity to protein G is weak. Individual monoclonals have variable binding affinities to protein G, which are pH dependent; the optimal pH may vary for each Ig. For those with low to medium affinity, capture efficiency by protein G can be dramatically improved through use of our Bridging Antibody (Catalog No. 53017). This antibody is a rabbit anti-mouse pAb that recognizes all subclasses of mouse immunoglobulins. If your IgG has a weak/medium affinity to protein A or G, the Bridging Antibody will increase antibody capture by the beads without increasing background. Problems with PCR. DNA to be used in real-time PCR must be purified prior to amplification. Primer issues. Confirm the species specificity of your primers. You may need to redesign your primers. Primers that work in end point PCR do not always work in real-time PCR.</td>
</tr>
<tr>
<td>The PCR products are the correct size, but are very faint.</td>
<td>Load more PCR product, and/or use smaller wells for the agarose gel. It should be noted that because the PCR reactions are stopped in log phase of amplification, the yield of PCR product may be lower than in typical PCR amplifications, which are performed for maximum product yield. You can also perform more PCR cycles. <strong>Note:</strong> a primer dimer band may be visible underneath the correct size PCR product.</td>
</tr>
<tr>
<td>No PCR bands with Input DNA samples (but the ChIP’d samples have the correct PCR product).</td>
<td>Perform the 1:10 dilution on the Input DNA prior to PCR, as indicated in the manual. Proteinase K may not have been completely inhibited. Warm the Proteinase K Stop Solution to room temperature for 30 minutes and then vortex briefly to get the material in solution, then add the recommended 2 µl and perform the recommended incubation. Purification of the Input DNA with a DNA purification column prior to PCR will effectively address either of the above issues.</td>
</tr>
<tr>
<td>No PCR products for the ChIP’d samples (but the Input DNA yields the correct PCR product)</td>
<td>Increase the amount of chromatin used in the ChIP reaction, the amount of antibody used, or both. Use a different antibody.</td>
</tr>
<tr>
<td>No PCR products with real-time PCR</td>
<td>The DNA should be purified before performing real-time PCR. We recommend Active Motif’s Chromatin IP DNA Purification Kit (Catalog No. 58002) prior to amplification. Its columns yield 50 µl; 2 µl is used for each PCR, providing enough DNA for 25 PCR reactions.</td>
</tr>
<tr>
<td>High background.</td>
<td>Chromatin not sheared enough. Shearing should produce DNA fragments that are small enough to exclude background from neighboring chromosomal sequences, but still large enough that there is a good possibility your amplicon remains intact. We recommend 200-1500 bp fragments. If the DNA fragments are too large, the background is increased. Consider increasing the number of pulses for sonication. Check the fragment size on a gel to assess your shearing efficiency. Antibody issue. Too much antibody relative to the amount of chromatin in the ChIP reaction. Excess antibody will result in more non-specific binding, which will be detected as increased background.</td>
</tr>
<tr>
<td>Too much template DNA. Reduce the amount of DNA in the PCR reaction.</td>
<td></td>
</tr>
</tbody>
</table>
Problem/question Recommendation

High background (continued).

Increase washes. In most cases, the washing protocol in this manual is appropriate. However, if the background is high you can increase washing stringency in several ways:

1. After adding ChIP Buffer 1 and/or ChIP Buffer 2 during the wash steps, gently agitate the samples for several minutes before removing the buffer.
2. Perform additional washes. Sufficient ChIP Buffer 1 is provided for two “extra” washes per sample. Sufficient ChIP Buffer 2 is provided for one additional wash.
3. Add two washes using a high-salt buffer (20 mM Tris-Cl, 1 mM EDTA, 0.1% SDS, 1% Triton X-100, 500 mM NaCl pH 7.4), which is not provided. These additional washes should be performed after the washes with ChIP Buffer 1. Then, proceed with the ChIP Buffer 2 washes, as outlined in the protocol.

Block the magnetic beads. The magnetic beads provided are ready to use for most ChIPs. However, for applications highly sensitive to non-specific binding (such as when cloning ChIP DNA or using antibodies that require extra blocking in other applications), you may add blocking reagents to the ChIP reaction.

Blocking reagents: 2.5 µg/µl BSA (e.g. Sigma Cat. No. 4503) and either 1.25 µg/µl tRNA (e.g. Sigma Cat. No. R3629) or 2.5 µg/µl Salmon sperm DNA (e.g. Sigma Cat. No. A-7888) (final concentrations shown for each) can be added directly to the ChIP reaction. These amounts are a good starting point, and can be increased as needed.

---

**Section G. Related Products**

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<td>RNA ChIP-IT®</td>
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<td>Chromatin IP DNA Purification Kit</td>
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ChIP-validated Antibodies
For an up-to-date list of over 125 ChIP-validated antibodies, please visit www.activemotif.com/chipabs.

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<td>Universal Magnetic Co-IP Kit</td>
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<table>
<thead>
<tr>
<th>Modified Histones Array</th>
<th>Format</th>
<th>Catalog No.</th>
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<tbody>
<tr>
<td>MODified™ Histone Peptide Array</td>
<td>1 array</td>
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<thead>
<tr>
<th>Histone Modification FP Binding Assay</th>
<th>Format</th>
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<tbody>
<tr>
<td>HiLite™ Histone H3 Methyl-Lys9 / Lys27 FP Binding Assay</td>
<td>1 kit</td>
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<tr>
<th>Histone ELISAs</th>
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<tr>
<td>Histone H3 monomethyl Lys4 ELISA</td>
<td>1 x 96 rxns</td>
<td>53101</td>
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<tr>
<td>Histone H3 dimethyl Lys4 ELISA</td>
<td>1 x 96 rxns</td>
<td>53112</td>
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<tr>
<td>Histone H3 trimethyl Lys4 ELISA</td>
<td>1 x 96 rxns</td>
<td>53113</td>
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<tr>
<td>Histone H3 acetyl Lys9 ELISA</td>
<td>1 x 96 rxns</td>
<td>53114</td>
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<tr>
<td>Histone H3 dimethyl Lys9 ELISA</td>
<td>1 x 96 rxns</td>
<td>53108</td>
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<tr>
<td>Histone H3 trimethyl Lys9 ELISA</td>
<td>1 x 96 rxns</td>
<td>53109</td>
</tr>
<tr>
<td>Histone H3 phospho Ser10 ELISA</td>
<td>1 x 96 rxns</td>
<td>53111</td>
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<tr>
<td>Histone H3 acetyl Lys14 ELISA</td>
<td>1 x 96 rxns</td>
<td>53115</td>
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<tr>
<td>Histone H3 monomethyl Lys27 ELISA</td>
<td>1 x 96 rxns</td>
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<tr>
<td>Histone H3 trimethyl Lys27 ELISA</td>
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<td>Histone H3 phospho Ser28 ELISA</td>
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<table>
<thead>
<tr>
<th>Histone Purification &amp; Chromatin Assembly</th>
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<tr>
<td>Histone Purification Kit</td>
<td>10 rxns</td>
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<tr>
<td>Histone Purification Mini Kit</td>
<td>10 rxns</td>
<td>40026</td>
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<tr>
<td>Chromatin Assembly Kit</td>
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<td>HeLa Core Histones</td>
<td>36 µg</td>
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Recombinant Methylated, Acetylated and Phosphorylated Histone Proteins
For an up-to-date list of Recombinant Histone Proteins, please visit www.activemotif.com/recombhis.

<table>
<thead>
<tr>
<th>Histone Acetyltransferase and Deacetylase Activity</th>
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<tbody>
<tr>
<td>HAT Assay Kit (Fluorescent)</td>
<td>1 x 96 rxns</td>
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<tr>
<td>Recombinant p300 protein, catalytic domain</td>
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<td>31205</td>
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<tr>
<td>Recombinant GCN5 protein, active</td>
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<td>31204</td>
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<tr>
<td>HDAC Assay Kit (Fluorescent)</td>
<td>1 x 96 rxns</td>
<td>56200</td>
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<tr>
<td>HDAC Assay Kit (Colorimetric)</td>
<td>1 x 96 rxns</td>
<td>56210</td>
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<tr>
<th>Histone Demethylase Activity</th>
<th>Format</th>
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</thead>
<tbody>
<tr>
<td>Histone Demethylase Assay (Fluorescent)</td>
<td>48 rxns</td>
<td>53200</td>
</tr>
</tbody>
</table>
Technical Services

If you need assistance at any time, please call Active Motif Technical Service at one of the numbers listed below.

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