Selective recognition of acetylated histones by bromodomains in transcriptional co-activators

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INTRODUCTION

The structure of the chromatin places an effective barrier on the ability of transcription machinery to access its binding sites. Many studies have described conserved protein complexes whose function it is to modify the chromatin structure and relieve its repressive effect (reviewed in [1–3]). They include ATP-dependent chromatin remodelling enzymes, as well as enzymes that post-translationally modify the N-terminal histone tails by acetylation, methylation, phosphorylation, sumoylation, ubiquitination and ADP-ribosylation [6–11]. Covalent modification of histone proteins has a well-known role in gene expression. Some of these histone modifications are short-lived, but others may persist through cell divisions and are thought to serve as stable epigenetic memory known as the ‘histone code’. These stable markers on histone tails act as binding sites for transcriptional co-activator proteins that may further modify the chromatin structure.

One evolutionarily conserved motif found in many transcriptional co-activators, including HATs (histone acetyltransferases; such as SAGA), chromatin remodelling complexes (such as SWI/SNF and RSC), and general transcription factors (such as TAFII250) is the bromodomain. It is a small domain that has been shown to interact with acetylated lysine residues in N-terminal tails of histones H3 and H4 in vitro [12–19]. It can also recognize and bind to acetylated non-histone proteins such as MyoD, the HIV-1 Tat and the p53 transcription factor [20–24]. Using immobilized template assays, we have previously shown that bromodomains were necessary for the anchoring of the SWI/SNF and the SAGA complexes to acetylated promoter nucleosomes [25,26]. In experiments in which SWI/SNF was recruited to nucleosomal templates, the retention of SWI/SNF required both acetylated histones and the Swi2/Snf2 bromodomain. Moreover, recognition by a few bromodomains of different acetylation patterns has been demonstrated in vitro [16,27–30]. Consistent with these in vitro experiments, mutant phenotypes were observed for an Swi2/Snf2 bromodomain deletion when combined with mutations in the SAGA HAT complex, and the Swi2/Snf2 bromodomain was required for the presence of SWI/SNF at the SUC2 promoter [26]. Furthermore, using fluorescence resonance energy transfer, it has been demonstrated that bromodomain proteins show selective recognition of acetylated histones in vivo [31]. More recently we have demonstrated the requirement of the Swi2/Snf2 bromodomain for the functional activity of the complex on SAGA-acetylated nucleosomes [32]. While the loss of the Swi2/Snf2 bromodomain has no effect on the remodelling and octamer transfer activity of unmodified nucleosomes, the bromodomain-deleted SWI/SNF has much reduced activity on SAGA-acetylated nucleosomes compared with the wild-type. Additionally, we have shown the displacement of SAGA by the SWI/SNF complex on acetylated nucleosomes and the requirement of the Swi2/Snf2 bromodomain for this activity. These data illustrate a novel and significant role of the Swi2/Snf2 bromodomain in remodelling of acetylated promoter nucleosomes and in displacing SAGA from promoters.

The evidence thus far suggests that bromodomains act as targeting modules to anchor chromatin-modifying proteins on promoters. Since bromodomains within various proteins have structural diversity, the present study investigated the interaction...
of some of these bromodomains with differentially acetylated histones in order to determine the preferred acetylated lysine residue as a target for their binding. We were particularly interested in determining the binding affinity of the various bromodomains within co-activator proteins to HAT-modified histone tails. Our results reveal selective recognition of acetylated histones by different bromodomains. Specifically, we show different sensitivities of binding of bromodomains to various acetylated H3 and H4 peptides as well as HAT-acetylated histones. Additionally, using hyperacetylated histones, we demonstrate that the bromodomain-deleted SWI/SNF complex cannot bind or remodel acetylated nucleosomes as efficiently as the wild-type. Together these data show that bromodomains have differing abilities to bind to and exert their modifying effects on acetylated nucleosomes.

EXPERIMENTAL

Cloning and purification of GST (glutathione S-transferase) fusion proteins

Seven DNA fragments encoding bromodomains within various yeast transcriptional co-activator proteins were PCR amplified from yeast genomic DNA and cloned into the BamHI and EcoRI sites of the GST expression vector, pGEX-2T, as described in the GST gene fusion system handbook (Amersham Biosciences). Table 1 in the supplementary data (http://www.BiochemJ.org/bj/402/bj4020125add.htm) shows the sequences of all of the primers used for cloning. The clones were screened for the insert using both PCR and restriction enzyme analysis. GST fusion proteins were then expressed in bacteria and purified using Glutathione Sepharose 4B beads as described in the manufacturer’s protocol (Amersham Biosciences), except that protease inhibitors (PMSF, leupeptin, pepstatin A and aprotinin) and DTT (dithiothreitol) were added to the resuspension buffer. Protein purity and amounts were checked and normalized by SDS/PAGE and Coomassie Blue staining.

GST pull-down assays with histone peptides and proteins

GST fusion proteins bound to Glutathione Sepharose 4B beads were separately incubated with acetylated histone H3 or H4 peptides in a pull-down buffer [50 mM Hepes (pH 7.5), 1 mM EDTA, 150 mM NaCl, 10 % glycerol, 0.1 % Tween 20, 0.5 mM DTT, 1 mM PMSF and 10 µg/ml pepstatin A] for 2 h at 4 °C while mixing on a rotation wheel. The supernatants were collected, and the beads were washed with the pull-down buffer three times and left as a 50 % slurry after the final wash. The beads were loaded on to SDS/PAGE (15 % gels), and the presence of acetylated peptide was detected by immunoblotting with acetyl-specific antibodies. The GST–bromodomain pull-downs with intact histones (control or hyperacetylated) were performed similarly, except that in these pull-down experiments, histone proteins served as substrates for the GST–bromodomains. Quantification of three independent experiments ± S.D. was performed by scanning the gels and using total histone signal intensity in the beads as the percentage pull-down. The GST–bromodomain pull-downs with intact HAT-acetylated histones were performed similarly, except that instead of incubation with acetylated peptides or control and hyperacetylated histones, histones were pre-acetylated with either SAGA or NuA4 HATs for 30 min at 30 °C with either non-radioactive or 3H-labelled acetyl-CoA (Amersham Biosciences) prior to incubation with the GST fusion proteins. After removing the supernatants, the beads were washed twice and the levels of interaction with the various GST fusion proteins were determined by performing either Western blot analysis or a HAT assay as described previously [33].

Purification of wild-type and mutant SWI/SNF complexes

Wild-type and Δbromodomain SWI/SNF (a strain lacking the Swi2/Snf2 bromodomain) complexes were purified from yeast whole cell extract using the TAP (tandem affinity purification) method using Snf6-TAP strains over two affinity columns as described previously [32,34–37]. Briefly, whole cell extracts were prepared from 6 litres of yeast cells grown in YPD medium [1 % (w/v) yeast extract/2 % (w/v) peptone/2 % (w/v) glucose] and were added to IgG resin (Amersham Biosciences). The complexes were eluted from the beads by TEV (tobacco etch virus) protease (Invitrogen) cleavage in a buffer containing 10 mM Tris/HCl (pH 8), 150 mM NaCl, 0.5 M EDTA, 0.1 % Nonidet P40, 10 % (v/v) glycerol, 1 mM PMSF, 2 µg/ml leupeptin, 1 µg/ml pepstatin A and 1 mM DTT. Following binding to calmodulin resin (Amersham Biosciences), the complexes were eluted using a buffer containing 10 mM Tris/HCl (pH 8), 150 mM NaCl, 1 mM MgAc, 1 mM imidazole, 2 mM EGTA, 0.1 % Nonidet P40, 10 % (v/v) glycerol, 1 mM PMSF, 2 µg/ml leupeptin, 1 µg/ml pepstatin A and 0.5 mM DTT. Purification was monitored by Western blot analysis using an anti-TAP antibody (Open Biosystems) as well as silver staining. The same amounts of the wild-type and the Δbromodomain SWI/SNF complexes were used in both the immobilized template binding and the restriction enzyme accessibility assays after normalization of the amounts of purified protein.

Immobilized template binding assay

pG5E4-SS containing a dinucleosome length G5E4 fragment flanked on both sides by 5 S sequences was prepared as described in [38]. The G5E4-SS fragments were produced by digesting pG5E4-SS with Asp718 and end-labelling with biotin-14-dATP using Klenow. The 2.5 kb end-labelled G5E4-SS fragments were then gel purified away from the backbone by digesting with Clal and Ecoal. This array was then reconstituted by step dilution with either control or hyperacetylated histones as described previously [26]. The nucleosomal arrays were then bound to paramagnetic beads coupled to streptavidin (Dynabeads Streptavidin, Invitrogen) as described previously [26,39]. Approx. 4 nM of either the wild-type or the mutant SWI/SNF was added to 200 ng of the above template in 20 µl of binding buffer and incubated for 1 h at 30 °C. The templates were then concentrated on a magnet, the supernatant was removed, and the beads were washed twice before performing Western blot analysis. The blots were probed with antibodies against the tag (anti-TAP antibodies). A graphical representation of the ratio of the bead to the supernatant ± S.D. is shown after quantification of three independent experiments determined by scanning the gels and using the signal intensities of the Western blots.

Restriction enzyme accessibility assay

In this assay, we used a 183 bp GUB fragment as the DNA template generated by PCR using a radiolabelled 5’ primer and the pGALUSFBEND plasmid [40–42]. This DNA fragment was reconstituted into a nucleosome using either control or hyperacetylated histones. The assay was performed as described previously [32,42]. Briefly, wild-type or Δbromodomain mutant SWI/SNF complexes were added to approximately 10 ng of the 32P-labelled GUB template in a binding buffer [10 mM Hepes (pH 7.8), 50 mM KCl, 5 mM DTT, 5 mM PMSF, 5 % glycerol, 0.25 mg/ml BSA and 2 mM MgCl2] in the presence or absence of 2 mM ATP. After incubation for 1 h at 30 °C, the binding reactions were treated with 10 units of Sall for 30 min at 30 °C. An equal volume of stop buffer [20 mM Tris/HCl (pH 7.5), 50 mM EDTA, 2 % SDS, 0.2 mg/ml protease K and 1 mg/ml glycogen] was
Interactions between bromodomains and acetylated histones

Figure 1 Purification of some of the yeast bromodomains as GST fusion proteins and their binding to hyperacetylated histones

(A) Coomassie Blue-stained SDS/PAGE of the purified yeast bromodomains. After purification, the amounts of GST and GST–bromodomain fusion proteins were normalized and used for all of the assays. The degradation of the Gcn5 bromodomain fusion proteins in GST did not significantly affect the results since the majority of the protein was the fusion protein. The relevant regions of the seven bromodomains in transcriptional co-activators in yeast were identified based on sequence homology to the Swi2/Snf2 bromodomain as well as published literature and are shown in parentheses. (B) The yeast bromodomains preferentially bind to histones with a high degree of acetylation with different affinities. A GST pull-down assay was performed with intact control (open columns) and hyperacetylated (solid columns) HeLa histones purchased from Upstate Biotechnology. After pull-down, beads were washed and bound proteins were analysed by SDS/PAGE and quantified. The quantification of the three independent experiments ± S.D. was determined by scanning the gels and using total histone signal intensity in the beads as the percentage pull-down. The relative fold increase in binding of the GST–bromodomain fusion proteins to hyperacetylated histones compared with control histones is shown at the bottom.

RESULTS

Bromodomains have a higher affinity for hyperacetylated histones

In order to determine the binding affinity of bromodomains within chromatin-modifying complexes to acetylated histones as well as acetylated lysine residues, we have cloned and purified bromodomains from several protein complexes as GST fusion proteins. Seven bromodomains from yeast co-activator proteins were cloned into a GST expression vector (pGEX-2T), including bromodomains from Swi2/Snf2 (the catalytic subunit of the SWI/SNF complex), Gcn5 and Spt7 (of the SAGA HAT), two bromodomains in the Bdf1 protein and one of the two bromodomains each in Rsc1 and Rsc2. We had difficulty in expressing and purifying to satisfaction the other bromodomains of the RSC complex. The sequences of the primers used for cloning are shown in Table 1 of the supplementary data (http://www.BiochemJ.org/bj/402/bj4020125add.htm). The GST fusion proteins were expressed in bacteria and the protein amounts and purity were monitored by SDS/PAGE (15% gels) and Coomassie Blue staining (Figure 1A). The overall purity of all of the bromodomains was good, except for the Gcn5 bromodomain, which had some degradation products. The cloned amino acid sequences of these bromodomains are shown at the top of Figure 1(A). The GST fusion bromodomain proteins bound to glutathione beads were first tested for their ability to bind to either control or hyperacetylated histones. Following incubation of the bromodomains with these histones and the separation of the supernatant from the beads, the beads were washed and the amount of total histones bound to bromodomains (the percentage pull-down) was determined by SDS/PAGE and added to the reaction mixtures, and incubated at 50°C for 1 h. Deproteinized samples were precipitated with 200 mM NaCl and 3 volumes of ethanol, and the pellet resuspended in 5 μl of the formamide dye (95% formamide, 10 mM EDTA, 0.1% xylene cyanol and 0.1% Bromophenol Blue). After heat denaturation, the samples were resolved on a 6% acrylamide (19:1 acrylamide to bis-acrylamide)/8 M urea sequencing gel at 150 V for 3 h and visualized by autoradiography. Graphical representation of the percentage cleavage ± S.D. is shown after quantification of three independent experiments determined by scanning the gels and measuring the signal intensities.
subsequent scanning and quantification of the bands. Figure 1(B) shows the percentage pull-down of the total histones for both control and hyperacetylated histones with the different GST-bromodomain fusion proteins. Although there are differences between the binding affinities of the different bromodomains, all bromodomains had a higher affinity for hyperacetylated histones compared with control unacetylated histones. This higher affinity ranged from 1.3-fold for the Bdf1-2 bromodomain to 2.7-fold for the Swi2/Snf2 bromodomain, shown at the bottom of Figure 1(B). The Swi2/Snf2, Gcn5, Bdf1-1, Rsc1-2 and the Spt7 bromodomain had the highest increases in binding to hyperacetylated histones respectively, whereas the second bromodomains of Bdf1 and Rsc2 had a weaker binding to hyperacetylated histones (1.3-fold and 1.4-fold respectively). Although these data do not differentiate between the different histones or acetylated lysine residues, a higher affinity for total histones when they are acetylated is observed in all bromodomains tested in the present study. In general, the increased affinity of the GST-bromodomains for hyperacetylated histones confirms the importance of the bromodomain as an acetylated binding domain within co-activator proteins.

Differential binding of bromodomains to acetylated H3 and H4 histone tail peptides

In order to study the binding abilities of the different bromodomains to acetylated H3 and H4 histone tails, the GST pull-down experiments were repeated using acetylated peptides. The GST-bromodomain proteins were individually incubated with various acetylated histone H3 or H4 peptides, followed by separation of the supernatant and the beads. After washing the beads, they were loaded on to SDS/PAGE (15% gels), and the presence of acetylated peptide was detected after immunoblotting with acetyl-specific antibodies. These experiments revealed differential binding affinities of the bromodomains with acetylated histone H3 and H4 peptides (Figure 2). For example, the Swi2/Snf2 bromodomain had a higher affinity for acetylated histone H3 Lys\(^9\) or Lys\(^14\) peptides compared with unmodified H3 peptide (Figure 2A, lane 3) and interacted with all acetylated H4 peptides as well (Figure 2B, lane 3). On the other hand, the Gcn5 bromodomain interacted specifically with acetylated H3 peptides (Figure 2A, lane 4) and only weakly with the tetra-acetylated H4 peptide (Figure 2B, lane 4). The other SAGA bromodomain (in its Spt7 subunits) only seemed to bind to H3 peptides acetylated at its Lys\(^9\) (Figures 2A and 2B, lane 5). This seems to be in contrast with the data in Figure 1(B) where we showed efficient enhancement of the binding of the Spt7 bromodomain to hyperacetylated histones compared with the control histones (a 1.9-fold increase in binding). The data in the present study are consistent with our previous observations where we have shown that the Spt7 bromodomain itself can bind to SAGA acetylated templates, but not in the context of the SAGA complex [29]. These results demonstrate that the bromodomain within the catalytic subunit of the SAGA complex is perhaps more important for recognition and binding to acetylated lysine residues in the histone tails, whereas the Spt7 bromodomain may have another function such as recognition of acetylated transcription factors or multiple lysine residues. The first bromodomain in the Bdf1 protein complex (Bdf1-1) seems to be important as an acetyl-lysine recognition module (Figures 2A and 2B, lane 6); however, its other bromodomain (Bdf1-2) only bound to tetra-acetylated H4 peptides (Figures 2A and 2B, lane 7), consistent with previously published data [20]. The two RSC bromodomains tested (Rsc1-2 and Rsc2-2) had weak binding properties to both acetylated H3 and H4 peptides. The Rsc1-2 bromodomain bound only to acetylated H3 Lys\(^9\)/Lys\(^14\) peptides and some of the H4 peptides, whereas the Rsc2-2 bromodomain did not seem to interact
Interactions between bromodomains and acetylated histones

Figure 3  Differential binding of the GST–bromodomains to HAT-acetylated histones

(A and B) Binding of the GST–bromodomain proteins to SAGA-acetylated histones. The GST pull-down assay with HAT-acetylated histones was performed similarly, except histones were pre-acetylated with the SAGA HATs for 30 min at 30°C with either non-radioactive or ³H-labelled acetyl-CoA (Amersham Biosciences) prior to incubation with the GST fusion proteins. After removing the supernatants, the beads were washed twice and the levels of interaction with the various GST fusion proteins were determined by performing either Western blot analysis (A) or a HAT assay (B) as described previously [33] to follow the ³H-labelled acetylated histone. The quantification of three independent pull-down experiments ± S.D. was performed by liquid-scintillation counting/filter binding of the ³H-labelled histones in the beads and is shown as the percentage pull-down with one representative fluorograph shown (bottom panel). (C and D) Binding of the GST–bromodomain proteins to NuA4-acetylated histones. The GST pull-down assay was repeated, except histones were pre-acetylated with the NuA4 HATs for 30 min at 30°C with either non-radioactive or ³H-labelled acetyl-CoA prior to incubation with the GST fusion proteins. Using either a Western blot analysis (C) or a HAT assay (D), we analysed the level of interaction of these acetylated H4 histones with the various GST fusion proteins. The quantification of three independent pull-down experiments with these templates was performed in a similar manner to above and the percentage pull-down with one representative fluorograph is shown in the bottom panel.

with any acetylated H4 peptide (Figures 2A and 2B, lanes 8 and 9). Together these results argue for the selective recognition of bromodomains within co-activator proteins and may account for the differences observed in the affinities of bromodomain-containing proteins for histones.

Differential binding of bromodomains to HAT-acetylated histones

In order to understand whether these bromodomains have similar affinities for more physiologically acetylated H3 or H4 histones, we have acetylated histones with either the H3- or H4-specific HATs SAGA or NuA4 respectively, prior to the GST pull-down assay. The level of interaction with acetylated histones (the percentage pull-down) was detected both by Western blot analysis using H3 or H4 acetyl-specific antibodies (Figures 3A and 3C) and by counting the levels of radioactive ³H-acetylated lysine residues in the pull-down assay (Figures 3B and 3D). These assays revealed a selective interaction of the GST–bromodomains with SAGA- or NuA4-acetylated histones. While interactions were observed between Swi2/Snf2, Gcn5, Bdf1-1 and Rsc1-2 and to some extent Spt7 bromodomains with SAGA-acetylated histones, no such interactions were observed between the Bdf1-2 and Rsc2-2 bromodomains with these histones (Figure 3A). This was confirmed when the GST pull-down was repeated with histones that had been acetylated with SAGA in the presence of ³H-acetylCoA, followed by fluorography (bottom panel of Figure 3B) and liquid scintillation counting on filter paper (Figure 3B). All of these results point to the differential affinities of these bromodomains for SAGA-acetylated histones. When the same experiments were repeated with histones that were pre-acetylated with NuA4, the results again showed different degrees of interaction. In this case, the Bdf1-2 bromodomain bound to these NuA4-acetylated templates in pull-down assays, whereas the Spt7 and the Rsc2-2 did not (Figure 3C). This was confirmed by the radioactive ³H-acetylation pull-down assay (Figure 3D). Together, all our data point to selective recognition and specificity of interaction of the bromodomains tested with various acetylated peptides and histones.

The Swi2/Snf2 bromodomain is important for the binding and remodelling activities of the complex on hyperacetylated nucleosomes

Previous studies have shown that the SWI/SNF complex is recruited to promoters by yeast transcriptional activators [41–47] and is also stabilized to acetylated nucleosomes through its
Swi2/Snf2 bromodomain [25,26]. Moreover, the bromodomain of Swi2/Snf2 participates in the functions of this protein complex in vivo [26]. In the present study, we show selective binding of different bromodomains to acetylated lysine residues, where the Swi2/Snf2 bromodomain is one of the most efficient in binding to various acetylated peptide tails and histones (see Figures 1–3). These results led us to investigate the role of the Swi2/Snf2 bromodomain in remodelling of acetylated nucleosomes. To test directly whether this bromodomain contributes to the binding and the functional activity of the complex on nucleosomes, we purified wild-type SWI/SNF as well as SWI/SNF from a strain lacking the Swi2/Snf2 bromodomain (termed ∆bromodomain SWI/SNF), and tested them in histone binding assays as well as chromatin remodelling assays. For all of these experiments, we used wild-type and mutant SWI/SNF complexes that had been highly purified over two affinity columns using the TAP method [32,34–37]. The loss of the Swi2/Snf2 bromodomain did not affect complex integrity as detected by silver staining (results not shown). The same amounts of the wild-type and the ∆bromodomain SWI/SNF complexes were used in these assays based on the normalization of the amounts of purified protein after Western blotting.

To investigate the binding abilities of the wild-type and the ∆bromodomain SWI/SNF complexes to nucleosomes, we utilized an immobilized nucleosomal array template, pG5E4, which contains five Gal4 binding sites and an E4 promoter flanked on both sides by 5S rDNA nucleosome positioning sequences [26,38]. Figure 4(A) shows the outline of the experiment. Briefly, the template was first biotinylated and reconstituted with either control or hyperacetylated histones, followed by the addition of either the wild-type or the ∆bromodomain SWI/SNF complexes. After pull-down, the binding of both complexes to nucleosome arrays was detected by Western blot analysis using an anti-TAP antibody (Figure 4B), therefore testing the ability of the Swi2/Snf2 bromodomain to bind to acetylated nucleosomes. Both the wild-type and the ∆bromodomain SWI/SNF complex bound equally well to the control template (Figure 4B, lanes 1 and 2). However, the wild-type SWI/SNF complex had a higher affinity for hyperacetylated templates compared with the mutant SWI/SNF complex (Figure 4B, lanes 3 and 4). Figure 4(C) shows a graphical representation of the ratio of the beads to the supernatants (B/S) for three independent immobilized binding experiments ± S.D. Again, the binding of the SWI/SNF complex to unmodified nucleosomes did not depend on the Swi2/Snf2 bromodomain of the SWI/SNF complex (Figure 4C). In contrast, when hyperacetylated nucleosome templates were used, the wild-type SWI/SNF but not the bromodomain-deleted complex could bind the templates efficiently (Figure 4C). These results show that while both an intact SWI/SNF complex and a complex that lacked the Swi2/Snf2 bromodomain could bind to unmodified promoter nucleosomes, the bromodomain-containing SWI/SNF complex binds with a greater affinity to the acetylated nucleosomes. The data clearly illustrate the importance of the Swi2/Snf2 bromodomain in binding to hyperacetylated histones and its requirement for efficient binding and anchoring of the SWI/SNF complex to these templates, leading to chromatin remodelling and subsequent gene activation.

To test directly whether the increased binding of the wild-type SWI/SNF complex to hyperacetylated nucleosomes (observed in Figure 4) leads to increased remodelling activity of the complex, a restriction enzyme digestion assay was performed. The DNA used in this assay was the 183 bp GUB fragment described above. As shown in Figure 5(A), the radiolabelled GUB DNA fragment was reconstituted into a nucleosome template by octamer transfer followed by the re-modelling reaction and digestion by the restriction enzyme SalI. We analysed the ability of the SalI restriction enzyme to digest its site in approximately the middle of the 183 bp GUB nucleosomal DNA with the wild-type and the ∆bromodomain SWI/SNF complexes in the presence of ATP. Figure 5(B) shows that both the wild-type and the mutant complexes increased the accessibility of the restriction enzyme to its site on the nucleosomal DNA. An equal amount of the control nucleosomal template was cleaved with either of these complexes, indicating that the deletion of the bromodomains in the Swi2/Snf2 subunit of the SWI/SNF complex did not significantly affect its remodelling activity on control (unmodified) histone templates. However, when hyperacetylated nucleosomes were used as the template, there was a 2-fold increase in remodelling activity of the wild-type SWI/SNF compared with the ∆bromodomain SWI/SNF complex, indicating that the Swi2/Snf2 bromodomain is important for the efficient

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have shown that when using hyperacetylated histones, there was a 2-fold increase in remodelling activity of the wild-type SWI/SNF compared with the Δbromodomain SWI/SNF complex or compared with the activity of the wild-type complex on control templates. The increase in the activity of the wild-type complex on hyperacetylated templates compared with control templates differs from when SAGA-acetylated templates were used. All these results show that the deletion of the bromodomain significantly reduced the binding as well as the functional activity of the SWI/SNF to remodel acetylated nucleosome complexes.

**DISCUSSION**

In the present paper, we have studied the binding of various bromodomains within transcriptional co-activators to differentially acetylated histone tail peptides as well as HAT-acetylated histones. Moreover, we have examined the importance of the Swi2/Snf2 bromodomain in the yeast SWI/SNF complex in the binding and remodelling activity of this complex on hyperacetylated nucleosomes. We have confirmed that the bromodomain is an acetyl binding and recognition domain as all of the bromodomains tested showed increased affinity for hyperacetylated histones compared with control histones in GST pull-down assays (Figure 1B). Moreover, we have shown that different bromodomains of yeast co-activator complexes bind to acetylated H3 and H4 peptides with different affinities (Figure 2). Our results reveal that while the Swi2/Snf2 bromodomain had the strongest binding affinity to acetylated H3 and H4 peptides of all the bromodomains tested, the other bromodomains differentially recognize the acetylated peptides. In addition, more physiological acetylation by the native yeast HATs (SAGA and NuA4) also show different degrees of binding to the bromodomains consistent with the peptide binding assay (Figure 3). Again, the Swi2/Snf2 bromodomain has the highest level of binding to both SAGA- and NuA4-acetylated histones. These data illustrate the importance of the bromodomains in binding to acetylated nucleosomes in general, and their selective binding to differentially acetylated histones, in particular. Our results reveal that the Swi2/Snf2, Gcn5, Bdf1-1, Rsc1-2 and the Spt7 bromodomain had the highest increase in binding to hyperacetylated histones, whereas the second bromodomains of Bdf1 and Rsc2 had a weaker overall binding to hyperacetylated histones. This did not seem to be the case when acetylated peptides or HAT-acetylated histones were used as the template. The two SAGA bromodomains had a higher affinity for acetylated H3 histones, whereas the Bdf1-2 bromodomain preferred binding to acetylated H4 histones consistent with previously published data [16,26]. We observed a 1.9-fold increase in binding of the Spt7 bromodomain to hyperacetylated histones compared with control, but only weak binding to acetylated H3 peptides and SAGA-acetylated histones. It is possible that the Spt7 bromodomain weakly interacts with individually acetylated lysine residues, but would have a synergistic increase in binding when multiple acetylated lysine residues are presented as the substrate, as our data shows. The differences observed between these bromodomains in binding to various acetylated histones are perhaps a result of sequence diversity that exists among them. Figure 6 compares the amino acid sequences between the different bromodomains within yeast co-activator complexes bind to acetylated H3 and H4 peptides of all of the bromodomains, we have shown that when using hyperacetylated histones, there was a 2-fold increase in remodelling activity of the wild-type SWI/SNF compared with the Δbromodomain SWI/SNF complex or compared with the activity of the wild-type complex on control templates. The increase in the activity of the wild-type complex on hyperacetylated templates compared with control templates differs from when SAGA-acetylated templates were used. All these results show that the deletion of the bromodomain significantly reduced the binding as well as the functional activity of the SWI/SNF to remodel acetylated nucleosome templates.

**Figure 5** The wild-type SWI/SNF complex has enhanced remodelling activity on hyperacetylated nucleosomes compared with the Δbromodomain SWI/SNF complex

(A) Diagram of the restriction enzyme accessibility assay as described previously [36,46]. A 32P-labelled GUB fragment was reconstituted into mononucleosomes with either control or hyperacetylated histones for use in this assay. An equal amount of wild-type or Δbromodomain SWI/SNF was added to this GUB template in the presence or absence of ATP followed by Sall digestion. After stopping the digestion reaction and ethanol precipitation, the DNA was resolved on a 6% acrylamide/8 M urea sequencing gel at 150 V for 3 h. The gels were dried and visualized by autoradiography and quantified. (B) Restriction enzyme accessibility assay showing increased activity on hyperacetylated nucleosomes when the Swi2/Snf2 bromodomain is present in the complex. The uncut and cut DNA fragments are labelled. (C) The quantification of three independent experiments is shown as the percentage cleavage of the mononucleosome templates by the restriction enzyme in the presence of the wild-type or the Δbromodomain mutant chromatin-remodelling complex SWI/SNF. The amount of digestion in the absence of ATP is subtracted from that value obtained in its presence when the percentage cleavage was calculated. Quantification of the experiments shows that a similar percentage of the unmodified nucleosome template (with control histones) is cleaved with both the wild-type and the mutant complexes. However, with hyperacetylated nucleosomes as the template, there is a 2-fold increase in remodelling activity of the wild-type SWI/SNF compared with the Δbromodomain SWI/SNF complex.

remodelling activity of the complex on acetylated nucleosomes. Quantification of three independent restriction enzyme accessibility experiments ± S.D. (Figure 5C) confirms these findings. In our recent publication [32], we have compared the activities of the wild-type SWI/SNF with that of the bromodomain-deleted complex on SAGA-acetylated templates and have shown that when the template was acetylated by SAGA in the presence of acetyl-CoA, reduced digestion by the Sall restriction enzyme was observed with the mutant SWI/SNF complex. In the present study, since we did not want the presence of the SAGA complex itself hindering the complete binding and remodelling of the templates by the wild-type or the mutant SWI/SNF complexes, we used hyperacetylated histones as the template instead. We

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The fact that the Swi2/Snf2 bromodomain binds efficiently to almost all of the acetylated histone peptides suggests low selectivity of the Swi2/Snf2 bromodomain for acetylated lysine residues. This low selectivity of the Swi2/Snf2 bromodomain in recognizing the acetylated peptides could mean that the SWI/SNF complex might play an important and a more global role in interactions with acetylated promoters and in transcription regulation than previously thought. In other words, the SWI/SNF complex could potentially interact with a variety of acetylated promoters that may have different acetylation patterns. Other more selective bromodomains may be more important in regulation of specifically acetylated promoters. It is also likely that some of the other modifying proteins that have two or more bromodomains (such as SAGA, RSC and TFIID) could bind to acetylated histones with higher affinity, in a similar manner to the Swi2/Snf2 bromodomain, as a result of co-operation between two tandem bromodomains within the same complex [28,29]. It is reasonable to assume that the multiple bromodomains within one complex enables simultaneous recognition of multiple modifications of acetylated histones, resulting in enhanced overall binding to acetylated promoter nucleosomes.

We have also shown that even though SWI/SNF purified from a Swi2/Snf2 bromodomain deletion strain bound and remodelled unacetylated nucleosome templates as well as the wild-type complex, it could not bind or remodel acetylated nucleosomes efficiently (Figures 4 and 5). The results show the requirement for the Swi2/Snf2 bromodomain in anchoring the complex on acetylated templates prior to any remodelling. The difference between the ability of the wild-type and the bromodomain-deleted SWI/SNF complexes to remodel acetylated promoter nucleosomes is due to the fact that the Swi2/Snf2 bromodomain has the ability to recognize and bind to acetylated histone [26]. We have also previously shown that the Swi2/Snf2 bromodomain deletion reduces occupancy of the SWI/SNF complex at the SUC2 promoter under derepressing conditions [26]. This is consistent with our current data showing that the Swi2/Snf2 bromodomain deletion cannot bind and remodel acetylated promoter nucleosomes. It is likely that bromodomain-containing protein complexes such as SWI/SNF, RSC and TFIID bind with greater affinity to nucleosomes that are pre-acetylated by HATs such as the SAGA or NuA4 complexes [32,48–50]. Thus acetylated nucleosomes could serve as substrates for sequential and ordered binding of these complexes to the promoter, where these complexes compete with each other for binding. SAGA itself, containing two bromodomains, could also bind to the nucleosomes (through its Gcn5 bromodomain) that it has acetylated or those that are acetylated by NuA4 and compete for binding with other complexes. This competition between the bromodomain-containing proteins depends on which lysine residue is acetylated since these would serve as docking sites for the complexes. Whereas the Swi2/Snf2 bromodomain, for example, is broader in recognition and binding to acetylated lysine residues, other bromodomains are more selective in this recognition (Figure 2) and thus it is also possible that multiple complexes can bind to promoters and exert their combined effects to regulate transcription. Therefore multiple bromodomains in proteins with different specificities and multiple complexes with bromodomains, as well as other protein motifs (such as the chromodomain or the SANT domain), add further complexity to histone recognition. Multiple post-translational modifications within the histone tails and multiple DNA-/protein-interacting domains within all the different co-activators together could provide a mechanism by which protein complexes can read the ‘histone code’ and exert control over chromatin function and gene regulation.

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REFERENCES

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