Recruitment of the SWI/SNF chromatin remodeling complex by transcriptional activators

Natalya Yudkovsky,1,4 Colin Logie,2,4 Steven Hahn,3 and Craig L. Peterson2,5

1Molecular and Cellular Biology Program, University of Washington, The Fred Hutchinson Cancer Research Center; 2Program in Molecular Medicine and Department of Biochemistry and Molecular Biology, University of Massachusetts Medical School, Worcester, Massachusetts 01605 USA; 3Fred Hutchinson Cancer Research Center and the Howard Hughes Medical Institute, Seattle, Washington 98109 USA; 4Program in Molecular Medicine and Department of Biochemistry and Molecular Biology, University of Massachusetts Medical School, Worcester, Massachusetts 01605 USA

SWI/SNF is a chromatin remodeling complex that facilitates expression of a number of yeast genes. Here we demonstrate that SWI/SNF can be recruited from yeast nuclear extracts by a transcriptional activator. Recruitment is dependent on an activation domain but not on promoter sequences, TBP, or RNA polymerase II holoenzyme. We also show that acidic activation domains can target SWI/SNF remodeling activity. These results demonstrate that SWI/SNF activity can be targeted by gene-specific activators and that this recruitment can occur independently of Pol II holoenzyme.

Received June 29, 1999; revised version accepted August 6, 1999.

The regulation of transcription initiation requires that transcription factors function in the context of eukaryotic chromatin. Two classes of chromatin remodeling enzymes have been identified that facilitate transcription from chromatin templates in vivo, the histone acetyltransferases and the ATP-dependent remodeling enzymes. The founding member of the transcription-associated histone acetyltransferases is yeast Gcn5p, which was initially identified as a transcriptional coactivator (Georgakopoulos and Thireos 1992; Marcus et al. 1994) and then subsequently as the catalytic subunit of several histone acetyltransferase complexes (Grant et al. 1997; Pollard and Peterson 1997; Saleh et al. 1997). The yeast SWI/SNF complex is a paradigm of the ATP-dependent chromatin remodeling enzymes and is required for expression of a number of genes and the full functioning of a variety of transcriptional activators (Burns and Peterson 1997). Genetic studies initially suggested that the transcriptional requirement for SWI/SNF reflected the ability of the complex to antagonize chromatin-mediated transcriptional repression (Winston and Carlson 1992; Kruger et al. 1995). Purified yeast SWI/SNF is a 2-MD enzyme that can use the energy of ATP hydrolysis to increase the accessibility of nucleosomal DNA to DNase I (Côté et al. 1994), restriction enzymes [Logie and Peterson 1997, 1999], or DNA-binding transcription factors in vitro (Côté et al. 1994; Utley et al. 1997).

The inactivation of SWI/SNF results in a decrease in transcription of only a small subset of yeast genes (Burns and Peterson 1997; Holstege et al. 1998), indicating that SWI/SNF might need to be targeted to particular genes and to specific nucleosomes in vivo. Three different models have been proposed to explain SWI/SNF. The nontargeting model proposes that SWI/SNF introduces transient changes in chromatin structure by a catalytic and random fashion throughout the genome, and that persistent, targeted changes in chromatin only occur in the presence of a DNA-binding transcription factor (Owen-Hughes et al. 1996). Alternatively, SWI/SNF might be targeted to specific genes by virtue of its association with an RNA polymerase II [Pol II] holoenzyme that was itself recruited to the promoter region via transcriptional activator proteins. Indeed, both yeast SWI/SNF and its mammalian homolog, BRG1, have been reported to copurify with Pol II holoenzyme [Wilson et al. 1996; Cho et al. 1998; Neish et al. 1998]. However, other Pol II holoenzyme preparations do not contain SWI/SNF, suggesting that only a fraction of holoenzyme contains SWI/SNF [Myers et al. 1998]. Finally, SWI/SNF might be targeted through direct interactions with gene-specific activators. Consistent with this model, yeast SWI/SNF can interact with the mammalian glucocorticoid receptor in yeast whole-cell extracts (Yoshinaga et al. 1992), and recruitment of SWI/SNF to the HO promoter in vivo requires the SWI5 activator (Cosma et al. 1999).

Here we have tested directly whether transcriptional activators can recruit the SWI/SNF complex and if recruitment requires an obligatory association with a Pol II holoenzyme. We find that a DNA-bound activator can recruit SWI/SNF from a yeast nuclear extract (NE) under conditions in which RNA Pol II holoenzyme and other general transcription factors (GTFs) are not sequestered. Moreover, the acidic activators, GAL4–AH and GAL4–VP16, can target SWI/SNF remodeling activity in a purified system containing only DNA, histones, SWI/SNF, and activator proteins. By contrast, the GAL4 DNA-binding domain or a nonacidic activator, GAL4–proline, are unable to recruit SWI/SNF activity. These results indicate that SWI/SNF may be targeted to a subset of yeast genes by direct interactions with gene-specific transcriptional activators.

Results and Discussion

Recruitment of the SWI/SNF complex can occur independently of RNA Pol II holoenzyme and TBP

To investigate the targeting of SWI/SNF to a promoter,
we used yeast nuclear transcription extracts [NEs] in combination with modified His4 promoter templates immobilized on magnetic beads [Fig. 1A; Ranish et al. 1999]. Preinitiation complexes [PICs] were formed by incubating NE with the immobilized template, followed by washing of the complexes, and liberation of the PIC with PsI restriction enzyme digestion. In this system, PIC assembly depends on the presence of a promoter, TATA box, TBP, TFIIA, and holoenzyme subunits, and is modestly stimulated by activators [Ranish et al. 1999].

Figure 1B shows a Western blot of a PIC assembly experiment with Gal4–AH. Two subunits of the SWI/SNF complex, Swi3p and Snf5p, were recruited to the template DNA with extracts from wild-type cells [lane 1]. To determine whether SWI/SNF is recruited to DNA as part of Pol II holoenzyme, we used a NE made from a srb2 deletion (srb2Δ) strain in our immobilized template assay. This extract is defective for in vitro transcription in this assay, and its activity can be partially restored by the addition of recombinant Srb2 [rSrb2] [Ranish et al. 1999]. In addition, deletion or mutation of genes that encode holoenzyme components, such as Srb2p, prevents the recruitment of the entire holoenzyme complex to the promoter, but allows recruitment of TFIIID and TFIIA in vitro [Ranish et al. 1999]. Figure 1B shows the levels of Rpb3p, a Pol II subunit, and Gal11p, a holoenzyme subunit, are severely decreased in the case of the srb2Δ NE, as compared with wild type [lane 1 vs. lane 2]. Importantly, the levels of both of these components increased significantly on the addition of rSrb2 [lane 3]. Although the levels of Swi3p and Snf5p were lower in the case of the srb2Δ extract, as compared with wild type, their binding was not stimulated by the addition of rSrb2 [Fig. 1B, lanes 1–3]. Thus, the lower binding of SWI/SNF in this extract is probably due to a lower specific activity of this extract and not a consequence of the srb2 mutation. To test whether SWI/SNF was recruited to the template by a GTF not in the holoenzyme, such as TFIIID, we used an extract made from a TBP temperature-sensitive mutant strain in the immobilized template assay. The I143N mutation in TBP abolishes TBP–DNA binding and disrupts all PIC formation [Fig. 1B, lane 4; Reddy and Hahn 1991]. NE made from this strain is defective for in vitro transcription assays, but its activity can be restored by the addition of rTBP [Ranish et al. 1999]. This TBP mutation did not affect the recruitment of either Swi3p or Snf5p to the promoter, and addition of rTBP did not stimulate the levels of these factors as it did Rpb3p and Gal11p [Fig. 1B, lanes 4,5]. We therefore conclude that in this transcription system, the recruitment of SWI/SNF to the promoter occurs independently of the recruitment of holoenzyme and of other GTFs.

**SWI/SNF is recruited to DNA by activators**

Next we tested directly whether SWI/SNF was recruited to the templates by activator in the yeast NEs. Assays were performed either without activator, with the Gal4[1–94] DNA-binding domain, or with Gal4–AH or Gal4–VP16, with both wild-type and TBP1I143N NEs. In the wild-type extract, Gal4–AH stimulated recruitment of Rpb3p, Srb4p, TFIIA, and Toa2p 4- to 10-fold, as compared with recruitment in the absence of activator [Fig. 2, lane 2 vs. lane 6]. Gal4–AH did not significantly recruit TBP [Ranish et al. 1999]. Gal4–VP16, however, stimulated recruitment of all these components 2- to 20-fold [Fig. 2, lane 2 vs. lane 8]. Recruitment of Swi3p and Snf5p by these activators was stronger than the recruitment of holoenzyme components. Gal4–AH increased Swi3p and Snf5p levels >13-fold, whereas Gal4–VP16 increased Swi3p and Snf5p levels >21-fold; both Swi3p and Snf5p were barely detectable in the absence of activator [Fig. 2, lanes 2,6,8]. In contrast, the binding of subunits of two other SWI/SNF-like complexes, Isw1p, a component of ISW1 [Tsukiyama et al. 1999], and Sth1p, a component of RSC [Cairns et al. 1996], was constitutive and was not influenced by the presence or absence of activators, promoter sequences, or Pol II holoenzyme [Fig. 2, data not shown]. The decrease in Sth1p levels seen on the ΔPromoter template is most likely due to the fact that this template is ~150 bp shorter than the wild-type template. To show that the recruitment of SWI/SNF was dependent on the activation domains, we used Gal4[1–94] as a control [Fig. 2, lane 4]. Although it weakly stimulated binding of Rpb3p, Srb4p, TFIIA, and Toa2p levels, we saw no stimulation of either Swi3p or Snf5p recruitment. Both Gal4–AH and Gal4–VP16 stimulated Swi3p and Snf5p recruitment to the same extent in TBP1I143N NE as in wild type [Fig. 2, lanes 10,12,14]. Importantly, Swi3p and Snf5p recruitment was unaffected by the addition of rTBP to restore PIC assembly [Fig. 2, lanes 16,18,20]. In the case of Gal4–AH, SWI/SNF recruitment clearly occurred in the absence of recruitment of other PIC components. However, Gal4–VP16 was also able to recruit nearly wild-type levels of Rpb3p, Srb4p, and TFIIA in the absence of rTBP, as noted previously [Fig. 2, lane 14; Ranish et al. 1999]. This is because Gal4–VP16, unlike Gal4–AH, can recruit holo-
We also performed an immobilized template assay using a template that terminated 14-bp downstream of the Gal4 site. Digestion of this template with Pst I after incubation with NE produced a 53-bp fragment. Although approximately twofold less Gal4–AH bound to this template, we saw no recruitment of SWI/SNF [data not shown]. These data, together with evidence that SWI/SNF can only bind nonspecifically to pieces of DNA >80 bp (Quinn et al. 1996), suggest that the formation of a stable complex between SWI/SNF and activator requires DNA, but not specific promoter sequences.

Transcriptional activators can target SWI/SNF remodeling activity

Next, we wished to determine whether a transcriptional activator could target the chromatin remodeling activity of purified SWI/SNF complex. To investigate this possibility, we exploited a sensitive nucleosomal array remodeling assay in which the activity of a nucleosome remodeling complex is coupled to restriction enzyme activity [Logie and Peterson 1997, 1999]. The DNA template that we used is composed of 11 head-to-tail repeats of a L. variegatus 5S rRNA gene [208-11S; Fig. 3A]. Each repeat can position a nucleosome after in vitro reconstitution with purified histone octamers, yielding a homogeneous array of positioned nucleosomes. The sixth repeat of this DNA template contains a 5S sequence that bears a unique SalI–HincII restriction site close to the dyad axis of symmetry of the reconstituted nucleosome [Polach and Widom 1995], thereby allowing for a quantitative kinetic assay for nucleosome remodeling within nucleosomal arrays [Logie and Peterson 1999]. To investigate whether a Gal4 activator can target SWI/SNF remodeling activity, we modified the DNA template so that it contains five high affinity GAL4-binding sites adjacent to the 5S repeat that harbors the SalI–HincII restriction site [208-11S–Gal4; Fig. 3A]. Reconstitution of the 208-11S–Gal4 DNA template into a nucleosomal array positions the Gal4-binding sites in the linker region between two positioned nucleosomes [Fig. 3A].

We assembled four sets of remodeling reactions in which targeting of remodeling activity would occur on radiolabeled arrays competing against a 15-fold excess of unlabeled competitor arrays. [Fig. 3B]. HincII digestion of these four array mixtures was essentially equivalent in the absence of SWI/SNF ([Fig. 3B] first order rate = 1 × 10^{-3}/min; similar results are seen in the presence of SWI/SNF but in the absence of ATP [Logie and Peterson 1997; Logie et al. 1999]]. Then, we assessed the capacity of SWI/SNF to remodel the labeled nucleosomal arrays in the absence of a Gal4 derivative. Addition of a limiting concentration of SWI/SNF complex (0.8 nM) stimulated HincII kinetics three- to fivelfold (first order rate = 3 × 10^3 to 5 × 10^{-3}/min). Thus, SWI/SNF can remodel both types of nucleosomal arrays, and the presence or absence of Gal4-binding sites on the labeled or competitor array does not influence the kinetics [Fig. 3B; data not shown]. Next, we tested whether Gal4–VP16 could target SWI/SNF activity to an array.
bearing Gal4 sites. This was achieved by including 10 nM Gal4–VP16 in the four array mixtures. The presence of Gal4–VP16 did not affect remodeling kinetics when both the labeled and unlabeled arrays lacked Gal4 sites (first order cleavage rate = 3 × 10⁻³/min; Fig. 3B, top). When remodeling of the labeled array lacking Gal4 sites (208-11S) was measured in the presence of a 15-fold excess of unlabeled array bearing Gal4 sites (208-11S–GAL4), we observed a ~3-fold reduction in the first order rate of restriction enzyme cleavage induced by Gal4–VP16 (1 × 10⁻³/min; Fig. 3B, second panel), indicating that the Gal4–VP16 that was bound to the competitor array sequestered SWI/SNF activity. When both the labeled and competitor array bore GAL4 sites, we observed an approximately two- to threefold increase in rate compared with the rate in the absence of Gal4–VP16 (5 × 10⁻³ vs. 14 × 10⁻³/min; Fig. 3B, third panel). Finally, when 10 nM Gal4–VP16 was added to reactions containing a labeled array bearing GAL4 sites (208-11S–GAL4) immersed in a 15-fold excess of unlabeled array lacking GAL4 sites (208-11S), we observed a dramatic enhancement of the cleavage rate when compared with SWI/SNF alone [23 × 10⁻³/min; Fig. 3B, bottom]. This stimulation of restriction enzyme cleavage rate by activators was absolutely dependent on the continuous presence of ATP (data not shown; see also Logie and Peterson 1997) and on the presence of SWI/SNF complex (Fig. 4A). Together, these experiments demonstrate that the Gal4–VP16 activator can recruit SWI/SNF remodeling activity in vitro to nucleosomal arrays bearing GAL4-binding sites.

Recruitment of SWI/SNF remodeling activity requires an acidic activation domain

Next we tested the effects of four different GAL4 derivatives on remodeling kinetics—GAL4 (1–94), GAL4–AH, GAL4–VP16, and the nonacidic activator, GAL4–proline. Each reaction contained 0.2 nM labeled 208-11S–GAL4 array and 3 nM unlabeled 208-11S competitor array. Under these conditions, SWI/SNF remodeling is barely detectable [HincII cleavage rate 1–2 × 10⁻³/min]. Addition of 10 nM GAL4–AH or GAL4–VP16 increased the apparent first order rate of SWI/SNF remodeling in this experiment more than sixfold (14–17 × 10⁻³/min). However, the presence of GAL4(1–94) or GAL4–proline—even when present at 60 nM (data not shown)—had no effect on the first order rate of cleavage (1–2 × 10⁻³/min; Fig. 4A). Thus, targeting of SWI/SNF nucleosome remodeling activity requires an acidic activation domain.

We have shown that SWI/SNF can be recruited from a NE by a DNA-bound activator in the absence of an obligatory association with a Pol II holoenzyme or TBP. Furthermore, an activator can target SWI/SNF activity in reactions that contain only DNA, histones, SWI/SNF, and the activator protein. These results are consistent with the hypothesis that SWI/SNF is recruited in vivo by direct interactions with gene-specific activators. SWI/SNF may only be recruited by activators that control expression of the small number of SWI/SNF-dependent genes, or, alternatively, SWI/SNF may be recruited to many genes even though SWI/SNF activity might not be rate determining for expression. Our data does not rule out the possibility that SWI/SNF can be recruited to target genes via other mechanisms. In fact, recently it has been shown that recruitment of SWI/SNF to the HTA1–HTB1 locus requires HIR1 and HIR2, which encode transcriptional repressors (Dimova et al. 1999).

We also found that the activator-dependent recruitment of SWI/SNF to an immobilized template required >53 bp of DNA adjacent to the GAL4-binding site. These results indicate that SWI/SNF may not interact stably
with activators in solution, but that SWI/SNF is targeted by a DNA-bound activator. This possibility is consistent with the observation that recruitment of SWI/SNF to the HO locus requires SWI5, which binds without the assistance of SWI/SNF to two sites located between positioned nucleosomes (Cosma et al. 1999; J. Krebs and C.L. Peterson, unpubl.). SWI/SNF might also be targeted by transcription factors bound to nucleosomes because some transcription factors are able to occupy nucleosomal sites (Workman and Kingston 1992; Li and Wrangle 1993). Alternatively, other remodeling activities may facilitate the binding of an activator to a nucleosomal site, which would then allow subsequent recruitment of SWI/SNF.

How do activators coordinate interactions with the myriad of transcription factors that can serve as regulatory targets? In the case of chromatin remodeling factors, an acidic activation domain can target either the ATP-dependent SWI/SNF complex (this work) or the SAGA histone acetyltransferase (Utley et al. 1998). The ability of activators to recruit either SWI/SNF or SAGA may explain the functional redundancy of these two complexes in vivo (Pollard and Peterson 1997; Roberts and Winston 1997; Biggar and Crabtree 1999). However, in the case of the HO gene, whose expression is absolutely dependent on both SWI/SNF and SAGA (Pollard and Peterson 1997), it appears that the SWI5 activator can recruit SWI/SNF, but not SAGA complex. Thus, whereas generic activators like GAL4–AH or GAL4–VP16 may be able to interact with all possible targets, different gene-specific activators may have a restricted spectrum of binding partners to better coordinate the sequence of events that lead to gene activation.

Materials and methods

**Immobilized templates and PIC assay**

Wild-type template was prepared and biotinylated by PCR from pSH515 as described previously (Ranish et al. 1999). The ΔPromoter template was made by digesting pSH515 with XhoI and BamHI to remove a ~150-bp HIS4 promoter fragment. The cut plasmid was then purified, filled in with Klenow fragment (BIBCO-BRL), and religated to create pSH515ΔP. The ΔPromoter template was prepared and biotinylated by PCR from pSH515ΔP with the same primers and reaction conditions as for pSH515. The templates were immobilized on M-280 Streptavidin Dynabeads (Dynal) as described previously (Ranish et al. 1999).

The PIC assay was performed in 100-µl reactions as described in Ranish et al. (1999), except that no BSA was included in the transcription buffer used to wash the templates. The wild-type and mutant yeast strains used are described in Ranish et al. (1999). Each reaction contained 480 µg of wild-type, 480 µg of Srb2, or 360 µg of TBP143N nuclear extract, as indicated.

**Reconstitution of nucleosomal arrays**

DNA templates were prepared by restriction enzyme digestion of pCL7c [208-11S] or pCL6b [208-11S-Gal4], purified by gel filtration, and labeled by Klenow fill-in reaction with [α-32P]dATP (Logie and Peterson 1999). Reconstitution of histone octamers onto 208-11 array templates were performed by salt gradient dialysis with a ratio of histone octamer to 5S rRNA repeat of 1.0 (Logie and Peterson 1999). Coupled remodeling-restriction reactions were performed in 25 µl, with a final concentration of 5 nM Hist1, 125 mM NaCl, 10 mM Tris-HCL (pH 8.0), 1 mM DTT, 0.1 mg/ml BSA, and 1 mM ATP. HinII (New England Biolabs) was added to reactions at a final concentration of 500 units/ml. HinII cleavage was quantified by PhosphorImager analysis, and first-order rates were determined by curve fitting. In multiple independent experiments, the first-order rates of restriction enzyme cleavage for each particular combination of array, remodeler, and activator varied by ±30%.

**Protein purification**

SWI/SNF complex was purified from yeast strain CY396 (Logie and Peterson 1999) and lacked detectable levels of Pol II holoenzyme as assayed by Western blot analysis with antibodies to RPB1, SRR2, SRB4, and SRB5 (data not shown). In contrast, high levels of these components were detectable early in the fractionation scheme (i.e., the Ni-NTA eluate. GAL4[1–94], GAL4[1–147]–AH, and GAL4[1–147]–VP16 were purified from bacteria as described previously (Lin et al. 1988; Chasman et al. 1989). The GAL4-proline was purified from bacteria with the protocol for GAL4[1–94]. The binding activities of GAL4 derivatives were determined by gel shift assays with a 154-bp DNA fragment containing a single, high affinity GAL4-binding site (Côté et al. 1994) in the presence of 5 nM unlabeled 208-11S–Gal4 array DNA template. In these assays ~10 nM GAL4–AH was required for half-maximal binding. The concentration of each GAL4 derivative was adjusted to yield an equivalent level of DNA-binding activity.

**Acknowledgments**

We thank members of the Hahn and Reeder laboratories for help and suggestions, and J. Ranish and T. Tsukiyama for thoughtful discussions. We also thank H. Sakurai for Gal11 antibodies, Nancy Thompson for Rpb3 antibodies, J. Movius for Srb4 antibodies, B. Laurent for Sth1 antibodies, and R. Young for Srb5 antibodies. We also thank J. Ranish for the TRP temperature-sensitive extract, J. Geiger for rTBP, and J. Flanagan for GST–SWI3 protein. We thank T. Imbalzano for the kind gift of GAL4[1–94] protein and N. Tanese for the GAL4–proline expression plasmid. This work was supported by grants from the National Institutes of Health to C.L.P. (GM496054) and to S.H. (GM42551), a grant from the National Cancer Institute to N.Y. (T32CA09657), and by a fellowship to C.L. from the Human Frontiers Science Organization. C.L.P. is a Scholar of the Leukemia Society of America, and S.H. is an Associate Investigator of the Howard Hughes Medical Institute.

The publication costs of this article were defrayed in part by payment of page charges. This article must therefore be hereby marked ‘advertisement’ in accordance with 18 USC section 1734 solely to indicate this fact.

**References**

Biggar, S.R. and G.R. Crabtree. 1999. Continuous and widespread roles for the Swi-Snf complex in transcription. **EMBO J.** 18: 2254–2264.

Burns, L. and C.L. Peterson. 1997. Protein complexes for remodeling chromatin. **Biochem. Biophys. Acta** 1350: 159–168.
