



The Histone H4 Tail Regulates the Conformation of the ATP-Binding Pocket in the SNF2h Chromatin Remodeling Enzyme

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Abstract

The chromatin remodeling complex ACF helps establish the appropriate nucleosome spacing for generating repressed chromatin states. ACF activity is stimulated by two defining features of the nucleosomal substrate: a basic patch on the histone H4 N-terminal tail and the specific length of flanking DNA. However, the mechanisms by which these two substrate cues function in the ACF remodeling reaction is not well understood. Using electron paramagnetic resonance spectroscopy with spin-labeled ATP analogs to probe the structure of the ATP active site under physiological solution conditions, we identify a closed state of the ATP-binding pocket that correlates with ATPase activity. We find that the H4 tail promotes pocket closure. We further show that ATPase stimulation by the H4 tail does not require a specific structure connecting the H4 tail and the globular domain. In the case of many DNA helicases, closure of the ATP-binding pocket is regulated by specific DNA substrates. Pocket closure by the H4 tail may analogously provide a mechanism to directly couple substrate recognition to activity. Surprisingly, the flanking DNA, which also stimulates ATP hydrolysis, does not promote pocket closure, suggesting that the H4 tail and flanking DNA may be recognized in different reaction steps.

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Introduction

Chromatin remodeling enzymes alter histone–DNA contacts to enable both activation and repression of transcription. The ATP-dependent chromatin assembly factor (ACF) is implicated in gene silencing [1]. The gene-silencing function of ACF is postulated to rest in its ability to generate evenly spaced nucleosomes, which are thought to enable the formation of heterochromatic structures [2,3]. At least two classes of mechanisms can explain how the activity of ACF is regulated such that it specifically acts at loci that need to be silenced [1]. One class of mechanisms entails the recruitment of ACF via sequence-specific DNA binding factors. The other class of mechanisms relies on the intrinsic ability of ACF to distinguish between different types of nucleosomes based on specific nucleosomal cues. Here we address

the latter class of mechanisms in the context of human ACF (hACF).

hACF is part of the ISWI (*imitation switch enzyme*) family of remodeling complexes, which contain one ATPase subunit with homology to the *Drosophila* ISWI ATPase and up to five accessory subunits. hACF contains the ATPase subunit SNF2h (*sucrose nonfermenting 2 homolog enzyme*) and the accessory subunit ACF1. Previous work has shown that ISWI family complexes use two distinct features of a nucleosome in their reaction cycle: the length of DNA flanking the nucleosome and a basic patch of amino acids on the N-terminal histone H4 tail [4–11]. The reliance on these two features is thought to allow down-regulation of ISWI activity in two different *in vivo* contexts. In actively transcribed genes, the H4 tail is acetylated at lysine 16, a residue in the critical epitope of the H4 tail, the basic patch [1]. This

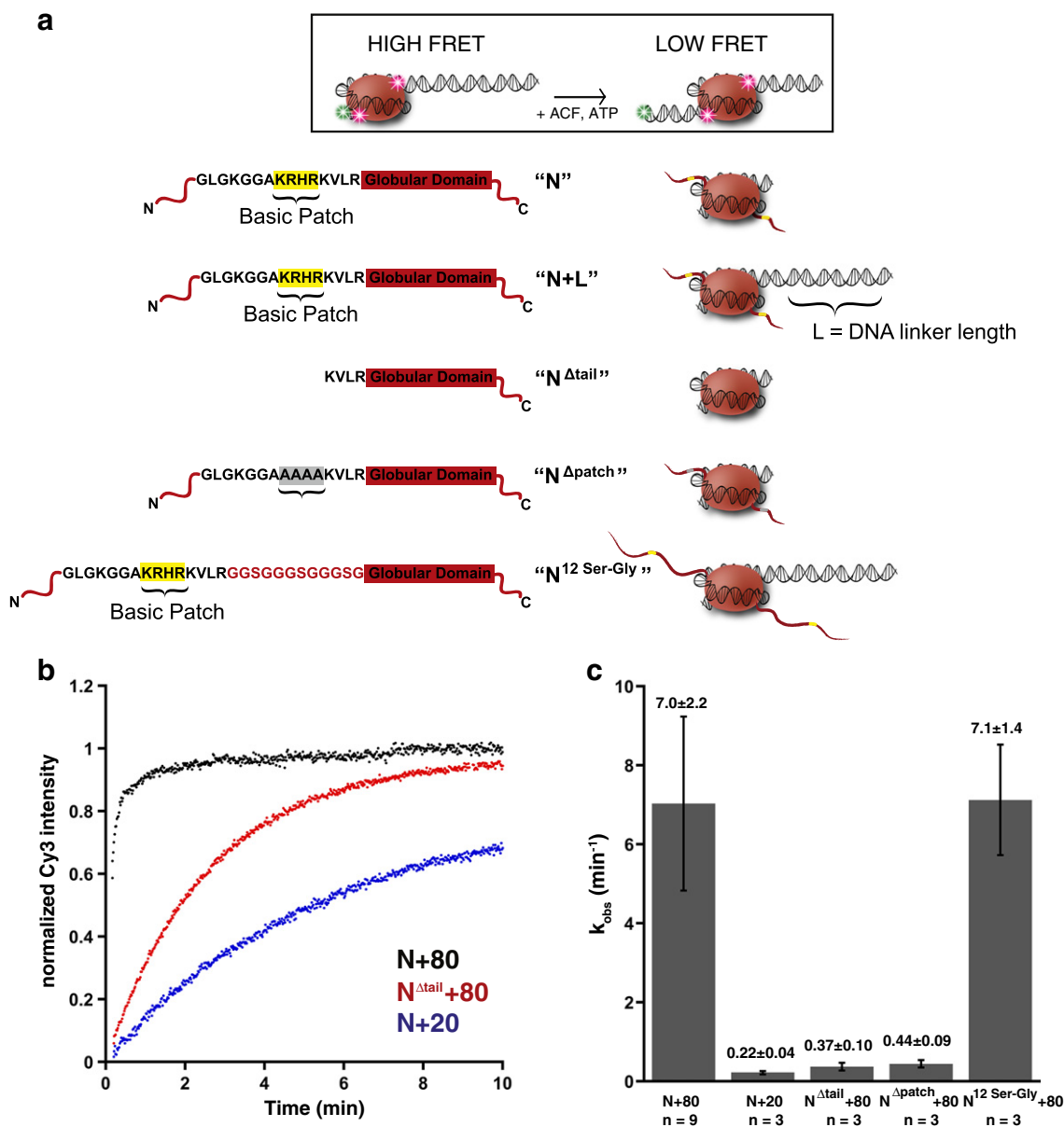


Fig. 1. H4 tail and flanking DNA length affect hACF nucleosome remodeling. (a) Nucleosome constructs used for FRET experiments. Histone octamers with a cysteine engineered at residue 120 of H2A were labeled with Cy5 and assembled onto DNA templates containing the 601 nucleosome positioning sequence and a proximal Cy3 dye and either 20 or 80 bp of DNA flanking the 601 positioning sequence on one side. The "N^{Δtail}" mutant lacks the first 19 amino acids of the H4 tail, including the basic patch of residues K₁₆R₁₇H₁₈R₁₉. In the "N^{Δpatch}" mutant, the basic-patch residues are replaced with alanines. In the "N^{12 Ser-Gly}" construct, a 12-amino-acid linker (Gly-Ser-Gly-Gly)₃ is inserted between residues 24 and 25 of histone H4. (b) Representative time courses from the FRET-based assay monitoring unquenching of the Cy3 donor dye are shown with saturating hACF and ATP-Mg. (c) Rate constants for remodeling (k_{obs}) with hACF for the nucleosome constructs under saturating hACF and ATP. The average values of the individual rate constants (min⁻¹) with standard deviation are shown on top of each bar.

post-translational mark reduces the activity of ISWI family enzymes *in vitro* and may analogously inhibit ISWI complexes at active loci *in vivo* [6,12–14]. In folded chromatin, the H4 tail is postulated to interact with an acidic patch on a neighboring nucleosome, and the flanking DNA is also thought to become

sterically less accessible [15–17]. Such physical occlusion may prevent the ISWI enzyme from mobilizing nucleosomes that are already compacted. Since the H4 tail and flanking DNA act as signals that the enzyme must recognize and integrate in order to act appropriately, we call them "substrate cues".

Interestingly, both substrate cues contribute more to the rate of remodeling by ISWI family enzymes than to their binding affinity [5,10,11,18]. Lengthening the DNA flanking the nucleosome up to 60 bp stimulates hACF remodeling rates, and beyond 60 bp, hACF remodeling rates remain largely constant [10,11]. Previous studies further imply that the HAND-SANT-SLIDE domain of ISWI ATPases interacts with flanking DNA [19–21]. Within the H4 tail, four amino acids, 16–19 (KRHR), are critical for maximal ISWI activity, with the two arginines being the most important [4,6]. Further, the specific location of the H4 tail on the nucleosome seems to be important, as swapping the locations of the H4 and H3 tails showed a significant decrease in activity [4,6]. This result may be explained by the proposed binding site of the ATPase domain of ISWI ATPases, which is two helical DNA turns from the nucleosomal dyad at superhelical location ± 2 (SHL ± 2) [8,20,22–25]. This is the region where the N-terminal H4 tail emerges from the nucleosome core particle. Complicating the mechanistic interpretation of the roles of flanking DNA and the H4 tail is the observation that hACF functions as a dimer in which the two protomers appear to bind the nucleosome on opposite sides [25]. Thus, each protomer has access to flanking DNA and one of the two H4 tails. Current models suggest that such a dimeric architecture allows hACF to achieve bidirectional movement of nucleosomes in a processive manner because the two protomers can take turns moving the nucleosome in one direction [25,26].

Given that both the flanking DNA and the H4 tail have catalytic effects, both cues must either directly or indirectly regulate the ATPase active site. However, the nature of the communication between the substrate cues and the active site is not known. More fundamentally, there is a lack of understanding about the types of conformational changes that occur in the ATP-binding pocket of chromatin remodeling enzymes and how these conformational changes correspond to changes in nucleosome structure. Because they participate in catalysis, the H4 tail and flanking DNA offer the opportunity to address (1) the

linkage between conformational changes in the ATP active site and physical changes in the nucleosome substrate and (2) the energetic contributions to catalysis of critical nucleosome epitopes that are known to lead to specificity. A better understanding of how hACF activity is regulated by variable features of a nucleosome has broader significance for understanding how chromatin remodeling enzymes as a class may be regulated by mechanisms other than simple recruitment.

In this study, we use ATP analogs containing an environmentally sensitive spin probe to detect a conformational change in the ATP-binding pocket under physiological solution conditions. Our results suggest that two structural features of the nucleosome that hACF is known to use, the H4 tail and flanking DNA, make qualitatively different contributions to catalysis raising the possibility that the two substrate cues act in different reaction steps.

Results

The effect of the H4 tail and flanking DNA on nucleosome remodeling

To better understand how these two classes of substrate cues affect activity, we first compared their catalytic effects on nucleosome remodeling. We measured remodeling rate constants for hACF with nucleosome substrates lacking each cue using a fluorescence resonance energy transfer (FRET)-based assay developed previously [11]. In this assay, we use nucleosomes containing a Cy3 label on one DNA end and a Cy5 label at amino acid 120 of H2A of the histone octamer by engineering in a cysteine substitution at this position. When an end-positioned nucleosome moves toward the center, this results in a loss of FRET (Fig. 1a). We monitor the change in FRET by following unquenching of the Cy3 dye with time (Fig. 1a and b). Removing the H4 tail in the context of nucleosomes containing 80 bp of flanking DNA reduces the

Fig. 2. Spin-labeled ATP analogs reveal conformational changes in the ATP-binding pocket of SNF2h. (a) 2'3'SLATP supports SNF2h nucleosome remodeling. Remodeling kinetics of N + 80 nucleosomes (5 nM) with 150 nM SNF2h, 1 mM MgCl₂ and 0.5 mM ATP or 2'3'SLATP (k_{obs} for $n = 3$ is $2.0 \pm 0.06 \text{ min}^{-1}$ and $1.2 \pm 0.22 \text{ min}^{-1}$, respectively, for ATP and 2'3'SLATP). Representative remodeling traces are shown. Inset: 2'3'SLATP analog (nitroxyl-radical spin probe shown in red). (b) Representative EPR spectra of 2'3'SLADP and 2'3'SLADP·BeF_x with SNF2h and N + 60 nucleosomes. The spectrum of 2'3'SLADP alone (without SNF2h and nucleosomes) is shown in black. The presence of nucleosomes and SNF2h (pink spectrum) produces a bound peak (P1) in the 2'3'SLADP spectrum with a high-field to low-field (P1–P5) splitting of 4.8 mT (116.9° cone angle). When the analog 2'3'SLADP·BeF_x is used with nucleosomes and SNF2h (blue spectrum), this produces a second, more immobilized peak (P1, blue) with a P1–P5 splitting of 6.2 mT (70.1° cone angle). Spectral peaks are truncated to enhance resolution of bound probe. (c) Representative spectra of 2'3'SLADP (pink) and 2'3'SLADP·BeF_x (blue) bound to SNF2h in the presence of saturating concentrations of a 60-bp DNA fragment. The black spectrum (2'3'SLADP alone) is from (b) for comparison. (d) Comparison of the ATPase rates of wild-type enzyme (black) and the E309A mutant (blue) in the presence of N + 60 nucleosomes. The curves level off due to depletion of 2-amino-6-mercapto-7-methylpurine riboside, the Enzchek assay detection reagent (see Materials and Methods). (e) Representative spectra of 2'3'SLADP (pink) and 2'3'SLADP·BeF_x (blue) bound to the E309A SNF2h mutant in the presence of N + 60 nucleosomes.

maximal rate constant of remodeling by 19-fold under single-turnover conditions (saturating and excess ATP and hACF over nucleosomes; Fig. 1b). Mutating the basic patch of the H4 tail, K₁₆R₁₇H₁₈R₁₉, to alanines gives a similar rate defect as removing the H4 tail (16-fold *versus* 19-fold, respectively; Fig. 1a and c). Shortening the flanking DNA from 80 to 20 bp reduces the maximal rate of remodeling by 32-fold (Fig. 1a and c). Thus, overall,

individually mutating each cue gives reductions in rate of similar magnitude. For technical reasons, we were not able to reliably measure rates of nucleosome remodeling under conditions where both cues are simultaneously removed. The larger reduction in rate demanded substantially longer time courses and made it difficult to ensure that the enzyme maintained the same specific activity for the duration of the experiment.

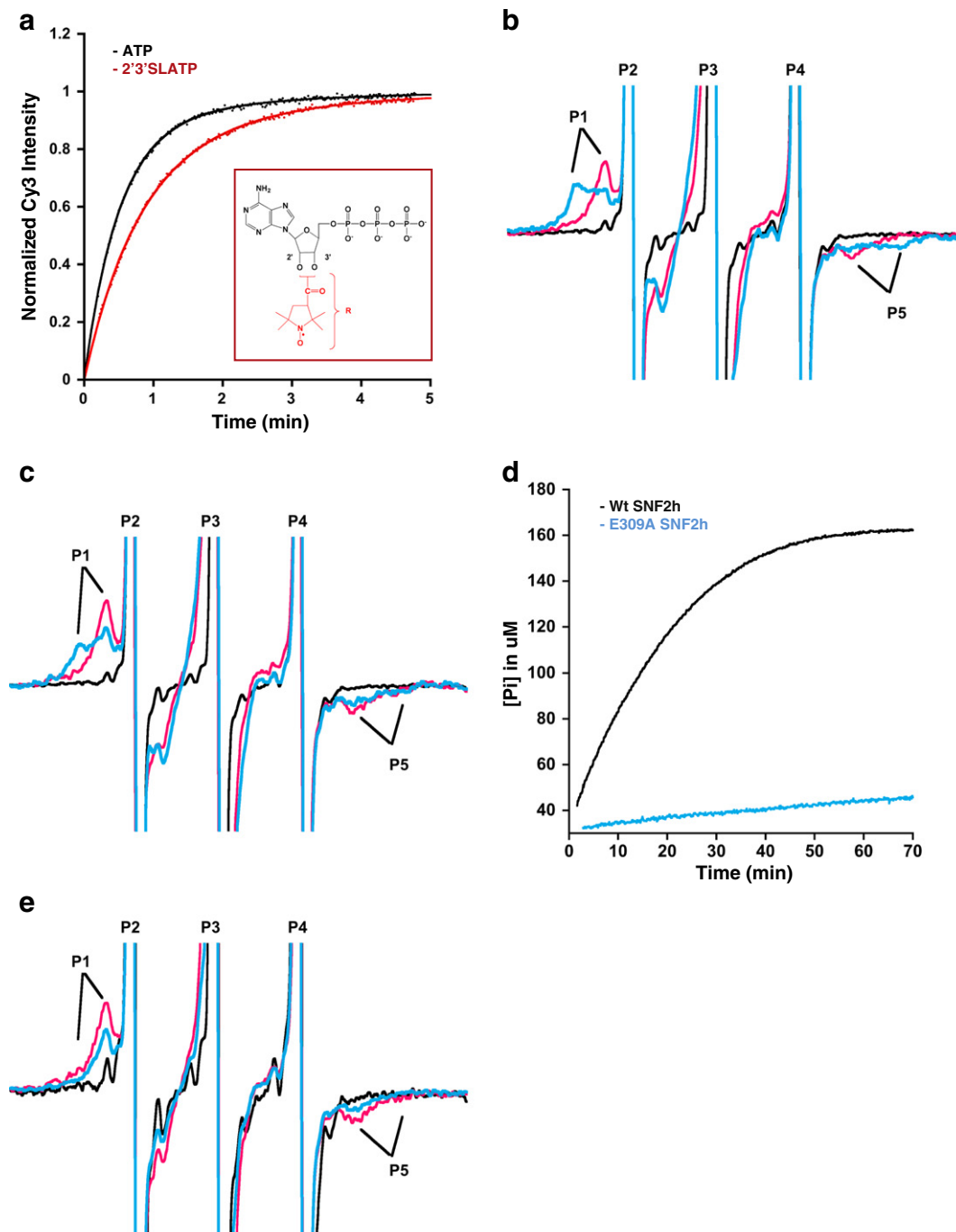


Fig. 2 (legend on previous page)

Table 1. Effects of substrate and nucleotide state on the conformation of the ATP-binding pocket of SNF2h. Fraction mobile was calculated as described in Materials and Methods. The average fraction mobile value is reported, and in parentheses, the values from independent experiments are shown. The reported K_{eq} values are averages of the K_{eq} values calculated for each independent sample spectrum (for $n = 3$ standard deviation is shown. For $n = 2$, only the average value is shown). Cone angles were calculated from splittings as described previously [39]

SNF2h	Nucleosome	Analog	Fraction mobile	K_{eq} (6.2 mT/4.8 mT)	Splitting (mT)	Cone angle (°)
Wt	N + 60	ADP	0.84 (0.83,0.85)	0.19	4.8	116.9
Wt	N + 60	ADP·BeF _x	0.25 (0.21,0.28,0.25)	3.1 ± 0.6	6.2, 4.8	70.1, 116.9
Wt	N + 60	ADP·AlF _x	0.19 (0.19,0.19)	4.3	6.4, 4.7	60.9, 120.1
Wt	None	ADP	0.87 (0.89,0.84)	0.16	4.7	120.1
Wt	None	ADP·BeF _x	0.94 (0.91,0.97)	0.065	4.7	120.1
Wt	N	ADP·BeF _x	0.12 (0.11,0.17,0.1)	7.4 ± 2.2	6.3, 4.9	65.6, 113.8
Wt	N	ADP·AlF _x	0.056 (0.04,0.07)	19	6.5, 4.8	55.8, 116.9
Wt	N ^{Δtail} + 60	ADP	0.81 (0.78,0.85)	0.24	4.8	116.9
Wt	N ^{Δtail} + 60	ADP·BeF _x	0.56 (0.50,0.56,0.63)	0.8 ± 0.2	6.3, 4.7	65.6, 120.1
Wt	N ^{Δtail} + 60	ADP·AlF _x	0.74 (0.78,0.71)	0.35	5.9, 4.7	82.1, 120.1
Wt	N ^{Δpatch} + 60	ADP·BeF _x	0.61 (0.55,0.67)	0.66	6.3, 4.8	65.6, 116.9
Wt	60mer DNA	ADP	0.83, 0.82 (0.83)	0.2	4.7	120.1
Wt	60mer DNA	ADP·BeF _x	0.37 (0.32,0.41)	1.8	6.3, 4.8	65.0, 116.9
E309A	N + 60	ADP	0.77 (0.8, 0.74)	0.3	4.8	116.9
E309A	N + 60	ADP·BeF _x	0.79 (0.76, 0.81)	0.27	4.8	116.9
E309A	N + 60	ADP·AlF _x	0.64 (0.8, 0.49)	0.65	4.8	116.9

We next investigated how the H4 tail might stimulate ATP hydrolysis. It has been suggested that interaction of the H4 tail with histone-bound DNA at SHL ± 2 may drive a region of the H4 tail (residues 16–24) containing the basic patch (16–19) into an α -helical conformation [27,28]. We therefore asked whether a particular secondary structure adopted by the H4 tail at SHL ± 2 might be required for the reaction. To test this possibility, we inserted a 12-amino-acid unstructured serine-glycine linker between the H4 tail and the globular domain (Fig. 1a and c; N^{12 Ser-Gly} + 80). Such a linker should disrupt any continuous secondary structure and also increase the total distance of the basic patch from the globular region. Previous calculations for the length of unstructured peptides suggest that the 12-amino-acid linker should increase the length between the histone octamer core and the basic patch by at least 24 Å [29–31]. Interestingly, the 12-amino-acid Ser-Gly insertion does not alter the maximal hACF remodeling rates (Fig. 1c). The comparison of the basic-patch mutant with the 12-amino-acid Ser-Gly insertion mutant (Fig. 1c, N^{Δpatch} + 80 versus N^{12 Ser-Gly} + 80) along with previous tail-swapping experiments [4,6] suggest that the location of the basic-patch epitope near the SHL ± 2 position of the nucleosome is more important for function than the precise structure of its connection to the nucleosome.

Conformational changes at the ATP-binding site measured by EPR spectroscopy

The stimulatory effects of the H4 tail basic patch and flanking DNA on hACF activity suggest that these substrate cues allosterically or directly regulate the ATP-binding site. Our goal was to determine how the two substrate cues affect the conformation of the

ATP-binding pocket. One of the challenges to understanding how substrate cues contribute to catalysis in motor proteins is that these enzymes tend to adopt multiple conformations that depend on both the nucleotide state and interactions with the second substrate [32]. Even in cases where there are multiple high-resolution crystal structures defining distinct conformations, such as with myosin and kinesin, the functional significance of these structures in a reaction cycle can be difficult to resolve [33,34]. The problem is greater for chromatin remodeling enzymes because crystallography of chromatin remodeling enzymes bound to the nucleosome substrate has been technically challenging. Electron paramagnetic resonance (EPR) spectroscopy provides a powerful means to assign functional states to static structures [33–35]. EPR spectroscopy can be used to resolve multiple protein conformations in equilibrium, under a wide variety of conditions including physiological reaction conditions. The traditional approach for EPR measurements is site-directed spin labeling where a single cysteine is incorporated into the protein at the location of interest and then is covalently modified by a reactive spin probe. We have previously used this technique to study the conformational changes of the H4 tail of the nucleosome in the presence of SNF2h, the ATPase subunit of hACF [25]. Here our goal was to monitor changes in the ATP-binding site; thus, we instead used spin-labeled ATP analogs. The analog specifically places the reporter group at the ATP-binding site without modifying the protein. The EPR spectrum of the spin probe provides information on the conformational restriction experienced by the probe, thus monitoring changes in the nucleotide-binding pocket as we have shown previously for kinesin and myosin motors [33,34]. Here we used spin-labeled ATP (2'3'

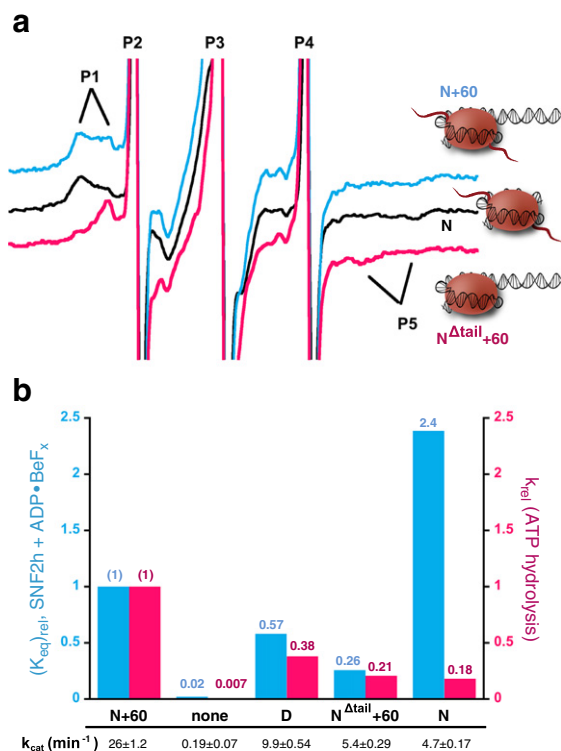


Fig. 3. Effects of substrate cues on the immobile/mobile probe equilibrium and on ATP hydrolysis. (a) Representative spectra of 2'3'SLADP·BeF_x bound to SNF2h in the presence of different types of nucleosomes. The equilibrium between the immobile and mobile populations of bound probe shifts in favor of the more immobile conformation when flanking DNA is removed (N + 60, blue spectrum *versus* N, black spectrum). In contrast, the equilibrium shifts toward the more mobile conformation when the H4 tail is removed (N + 60, blue spectrum *versus* N^{Δtail} + 60, pink spectrum). (b) Top panel: comparison of effects on ATP hydrolysis and K_{eq} . Blue bars: K_{eq} values relative to the N + 60 substrate. K_{eq} is obtained as defined in Table 1 in the presence of 2'3'SLADP·BeF_x. Pink bars: k_{cat} values for ATP hydrolysis relative to the N + 60 substrate. Bottom panel: The values of k_{cat} for ATP hydrolysis under multiple-turnover conditions.

SLATP), where the reporter spin moiety is attached to the ribose ring of ATP (Fig. 2a, inset).

In all the EPR experiments, we used SNF2h, the ATPase subunit of the hACF complex, on its own because of the technical challenge of obtaining the whole complex at the concentrations required for taking EPR spectra. SNF2h recapitulates all the core functions of the hACF complex (Fig. 1b) [11,36–38]. In addition, we confirmed that 2'3'SLATP supports remodeling by SNF2h at similar rates as ATP (Fig. 2a, time course, $k_{obs} = 2.0 \pm 0.06 \text{ min}^{-1}$ and $1.2 \pm 0.22 \text{ min}^{-1}$, for ATP and 2'3'SLATP, respectively). The spin probe in solution yields a spectrum consisting of three sharp lines (Fig. 2b, black). The spectrum of SNF2h-2'3'SLADP bound to nucleosomes with 60 bp of flanking DNA (SNF2h-N + 60

complex) shows more restricted mobility compared to when free in solution (Fig. 2b, pink). This is indicated by a broadening of the spectrum and the outward movement of the low-field peak (P1) and high-field dip (P5). The observed P1–P5 splitting is 4.8 mT (Table 1). A similar spectrum and splitting is obtained with SNF2h-2'3'SLADP alone (Table 1). The mobility of the probe can be modeled as a cone within which the probe diffuses due to thermal motion, where the vertex angle of the cone correlates with the magnitude of the P1–P5 splitting [39]. For the SNF2h-N + 60-2'3'SLADP complex, the vertex angle is 116.9° (Table 1). The P2–P4 sharp peaks corresponding to unbound nucleotide can be resolved from the broader peaks of the bound nucleotide at low and high field. Competition with unlabeled ATP eliminated the P1 and P5 components and restored the spectrum to that of freely tumbling 2'3'SLADP (data not shown), indicating that the probe is bound at the active site of SNF2h.

We next determined if we could identify changes in the conformation of the ATP-binding pocket by following changes in the mobility of the probe upon changing the ATP state. ADP·BeF_x is a stable, bipartite analog that can mimic different nucleotide states depending on the structural context. These states include ground-state ATP-like structures, transition-state-like structures and ADP-P_i product-like structures [40–42]. When we used 2'3'SLADP·BeF_x in complex with SNF2h and N + 60, we observed large changes in the EPR spectrum. The spectrum revealed a new subpopulation of spin probes with a greater P1–P5 splitting of 6.2 mT, in addition to the previously observed population with a splitting of 4.8 mT (Fig. 2b, cyan; Table 1). This new component is indicated by the additional P1 and P5 components in the spectrum. A greater splitting corresponds to a more restricted or more immobile probe, with a corresponding cone angle of mobility of 70.1°. Another bipartite nucleotide analog that mimics activated states of ATP, 2'3'SLADP·AlF_x, gives similar results as 2'3'SLADP·BeF_x (Fig. 2b, Table 1 and Supplementary Fig. 1). When BeF_x is added to the SNF2h-2'3'SLADP complex in the absence of the nucleosome, the immobile component does not increase (Table 1). The multicomponent spectra were deconvolved to determine the fraction of mobile *versus* immobile probes (Fig. 2b and Table 1, calculated $K_{eq} = \text{fraction immobile} / \text{fraction mobile}$; see Materials and Methods).

The differences in probe mobility suggest two conformations of the ATP-binding pocket: a more open (mobile probes) state and a more closed (immobile probes) state. The differences between these two states could reflect local changes in the ATP-binding pocket or large-scale changes involving domain rearrangements. By either interpretation, in the absence of the nucleosome, the closed state is largely absent. SNF2h is also largely inactive in ATP

hydrolysis in the absence of nucleosomes (Fig. 3b) [11]. The presence of the nucleosome shifts the equilibrium 48-fold toward the closed state, and ATP hydrolysis is correspondingly stimulated (Fig. 2b and Table 1, K_{eq} of 3.1 with the nucleosome, 6.5×10^{-2} without the nucleosome; Fig. 3b). This comparison raised the possibility that formation of the closed state correlates with ATPase activity.

Effect of naked DNA on conformational changes in the ATP-binding site

Naked DNA is another substrate known to stimulate ATP hydrolysis by SNF2h [10,36] (Fig. 3b). We therefore tested if this substrate can also promote formation of the closed state. We examined the spectrum of 2'3'SLADP·BeF_x in the context of a SNF2h-DNA complex, using a 60-bp-long duplex DNA (no nucleosome). We observed an increase in the immobile component for the SNF2h-DNA complex relative to SNF2h alone (Fig. 2c). The presence of DNA shifts the equilibrium 28-fold toward the more immobile component (Table 1, K_{eq} of 1.8 of SNF2h with DNA, 6.5×10^{-2} without DNA or nucleosome). Thus, naked DNA both stimulates ATP hydrolysis by SNF2h and drives the conformation of the nucleotide-binding pocket of SNF2h to the closed state.

Effects of active-site mutation on conformational changes in the ATP-binding site

To further test if the inferred conformational change in the ATP-binding pocket of SNF2h correlates with activity, we characterized the behavior of a SNF2h mutant that is deficient in ATP hydrolysis. The ATP-binding site of SNF2h contains the characteristic Walker A and Walker B motifs present in many ATPases, including the superfamily 2 (SF2) of helicases to which ATP-dependent chromatin remodeling enzymes belong [43]. The conserved aspartate and glutamate (DE) residues in the Walker B motif help coordinate the catalytic Mg²⁺ and water molecules for ATP hydrolysis [43,46]. Mutation of the D or E residue inhibits ATP hydrolysis in other SF2 ATPases [43]. We mutated glutamate 309 in the Walker B motif of SNF2h to alanine (E309A) to generate a SNF2h mutant that is defective in ATP hydrolysis (Fig. 2d). This mutant is also deficient in formation of the closed state in the presence of 2'3'SLADP·BeF_x and N + 60 nucleosomes (Fig. 2e). The observation that perturbation of a catalytic residue on the ATP-binding pocket also reduces formation of the closed state further suggests that formation of the closed state is important for catalysis.

The roles of the H4 tail and flanking DNA in nucleotide pocket closure

The apparent correlation between ATP hydrolysis and a closed conformation of the ATP-binding

pocket in the presence of ADP·BeF_x and ADP·AlF_x provided a meaningful context to test the energetic contributions of the substrate cues, the H4 tail and flanking DNA. We first looked at the effect of the H4 tail. The spectrum of 2'3'SLADP bound to SNF2h complexed with nucleosomes lacking the H4 tail (N^{Δtail} + 60) shows, as the dominant component, a P1–P5 splitting of 4.8 mT corresponding to a cone angle of mobility of 116.9° (Table 1). When BeF_x is added, an additional subpopulation of spin probes with a greater P1–P5 splitting of 6.3 mT is observed. However, the immobile/mobile equilibrium for the complex among ADP·BeF_x, SNF2h and N^{Δtail} + 60 is reduced ~4-fold relative to the corresponding complex with N + 60 (Fig. 3a, pink *versus* cyan spectra, respectively). We observed a similar reduction of probes in the immobile conformations upon mutating the basic patch of the H4 tail (Table 1, N^{Δpatch} + 60). These data suggest that the basic patch of the H4 tail promotes a closed conformation of the ATP-binding pocket of SNF2h.

The effect of the H4 tail on the equilibrium between closed and open states of the ATP-binding pocket in the presence of ADP·BeF_x (Fig. 3a, $1/K_{\text{eq}}^{\text{rel}} \sim 4$ -fold) is of similar magnitude to the effect on ATP hydrolysis (Fig. 3b, $1/k^{\text{rel}} \sim 5$ -fold). This quantitative correlation raises the possibility that the H4 tail promotes ATP hydrolysis by stabilizing a closed conformation of the ATP-binding pocket.

Like the H4 tail, the presence of DNA flanking the nucleosome promotes ATP hydrolysis by SNF2h [10,11] (Fig. 3b). We therefore next examined the effects of flanking DNA on the conformation of the nucleotide-binding pocket. The EPR spectrum of 2'3'SLADP·BeF_x bound to SNF2h with nucleosomes lacking flanking DNA but containing the H4 tail (N nucleosomes) shows primarily an immobile component, with a P1–P5 splitting of 6.3 mT corresponding to a 65.6° cone angle of mobility (Fig. 3a, black spectrum; Table 1). Deconvolution of the spectrum gives a K_{eq} value of ~7 for the closed/open equilibrium, which is slightly greater than the K_{eq} value for nucleosomes with flanking DNA (Table 1, N *versus* N + 60 with ADP·BeF_x). Thus, even though both the H4 tail and flanking DNA stimulate ATP hydrolysis, the H4 tail stabilizes the closed conformation of the ATP-binding pocket while the flanking DNA appears to slightly destabilize the closed conformation.

Discussion

A central question in understanding the mechanism of chromatin remodeling enzymes is how they are able to recognize which nucleosomes to move. Previous studies have indicated that recognition of the H4 tail and flanking DNA by ISWI enzymes is critical for catalysis but not for ground-state binding [5,10,11,18]. The mechanistic basis for how these

cues act in the reaction cycle is not understood. To understand how hACF uses the H4 tail and flanking DNA cues during catalysis, we adapted an EPR-based method to assay for conformational changes in the ATP-binding pocket of SNF2h, the ATPase subunit of hACF. Using the spin-labeled nucleotide analogs SLADP, SLADP·BeF_x and SLADP·AlF_x, we observe a conformational change in the nucleotide-binding pocket of SNF2h under physiological solution conditions. In one conformation, the spin probe on the nucleotide is more restricted (6.2 mT, more immobile) than in the other conformation (4.8 mT, more mobile). A 60-bp double-stranded DNA fragment and the nucleosome, both of which stimulate ATP hydrolysis, shift the equilibrium toward the restricted conformation of the probe (Fig. 2b and c). This shift in equilibrium requires ADP·BeF_x and is not detected with ADP alone. A Walker B mutant (E309A), which shows defects in ATP hydrolysis, shifts the equilibrium toward the more mobile conformation of the probe. While ADP·BeF_x has been shown to mimic the ATP ground state in many ATPases, it has also been shown to mimic transition-state-like structures and the ADP·P_i product in some systems [40–42]. Given the absence of high-resolution structures of chromatin remodeling ATPases bound to nucleotide analogs, we do not know which state is mimicked by ADP·BeF_x in the context of SNF2h. Nevertheless, the state mimicked by this analog stabilizes a conformation of the SNF2h active site that correlates with ATP hydrolysis. This correlation suggests that ADP·BeF_x mimics either the ground-state ATP structure or a transition-state-like structure in the context of SNF2h.

At a physical level, what could the conformational change in the active site represent? The spin probe reports on environmental changes near the 2'3' hydroxyl groups of the nucleotide. These changes could reflect local effects around the 2'3' hydroxyl groups or larger conformational changes in the active site. The ATPase domain of chromatin remodeling enzymes consists of two RecA-like lobes separated by a flexible linker. ATP binds in a cleft between these lobes. In many distantly related SF1 and SF2 family helicases, binding of DNA or RNA promotes closure of the two RecA-like lobes of the ATPase domain onto ATP, thus promoting catalysis [44–46]. Work on *Sulfolobus solfataricus* SWI2/SNF2, which is part of the chromatin remodeling superfamily, suggests that binding of the enzyme to DNA can alter the orientation of the two RecA-like lobes relative to one another [47,48].

We find that the substrate cue provided by the basic patch on the H4 tail increases ATPase rates and correspondingly shifts the equilibrium toward a more closed active site. Rate enhancement by the H4 tail is maintained upon inserting a flexible linker between the H4 tail and the globular domain. This result is consistent with a model in which the H4 tail

stabilizes a particular conformation of the enzyme through specific binding interactions rather than acting as a mechanical element for the transduction of force. In the case of the related chromatin remodeling enzyme, Chd1, the two RecA-like lobes of the enzyme are physically propped open by a third domain and thus are in an auto-inhibited state where they cannot close and hydrolyze ATP until the nucleosome is bound [49]. It is possible that binding the H4 tail helps relieve an analogous auto-inhibited conformation of SNF2h and allows for closure of the ATP-binding pocket. Indeed, a recent study of ISWI, the *Drosophila* homolog of SNF2h, suggests that the H4 tail plays a role in relieving auto-inhibition by displacing a basic-patch mimic sequence on an auto-inhibitory domain of ISWI [50]. Further, work on *Drosophila* ISWI has shown that DNA binding induces a conformational change in the ATPase domain [51]. Given these previous observations and the results presented here, we speculate that the H4 tail stabilizes a closed conformation of the two RecA-like lobes of SNF2h in which the catalytic residues are properly aligned for ATP hydrolysis, thereby directly coupling substrate recognition to activity.

In contrast to the H4 tail, the substrate cue provided by the flanking DNA increases ATP hydrolysis but does not shift the equilibrium toward a closed active site. The simplest interpretation of the difference between the H4 tail and the flanking DNA is that these substrate cues act in different reaction steps. It is possible that the flanking DNA acts in a different rate-limiting step prior to ATP hydrolysis. For example, such a step could entail a conformational change in SNF2h that allows the proper engagement of the HAND-SANT-SLIDE domain with flanking DNA. However, substantial future work will be needed to determine the exact nature of the reaction step that is stimulated by flanking DNA.

Materials and Methods

Protein purification

Both human SNF2h and human hACF complex were expressed in SF9 cells and purified as described previously [11]. For EPR experiments, SNF2h was concentrated using the centrifugal filter unit Microcon YM-100 (Millipore).

Nucleosome assembly

Nucleosomes were assembled using the 601 positioning sequence, containing a PstI site 18 bp in from one end, as well as 0, 20, 60 and 80 bp of DNA flanking the nucleosome on the other end as described previously [11]. To measure kinetics of nucleosome remodeling by FRET, we labeled H2A containing a unique engineered cysteine at residue 120 under denaturing conditions with

Cy5-maleimide and then assembled it by salt dialysis into octamers. H2A-120C-Cy5 octamers were then assembled into nucleosomes using Cy3-end-labeled DNA as described previously [11].

FRET-based nucleosome remodeling assay

Kinetics of nucleosome remodeling were measured as previously described on an ISS K2 fluorometer and were fit as previously described to two-exponential phases using Kaleidagraph [11]. Samples were excited at 520 nm, using a 400-nm-long pass filter, and emission of the donor Cy3 dye was measured at 565 nm (Fig. 1b), using a 495-nm-long pass filter. All reactions were performed at 30 °C, with 5 nM nucleosomes and varying concentrations of hACF enzyme in excess of nucleosomes, in the following reaction buffer: 60 mM KCl, 2 mM MgCl₂, 0.32 mM ethylenediaminetetraacetic acid, 12% (v/v) glycerol, 12 mM Hepes (pH 7.9), 4 mM Tris (pH 7.9), 0.02% (v/v) Nonidet P-40, 0.5 mg/ml Flag peptide, 0.4 mM DTT and 0.4 mM Benzamidine hydrochloride. After a 10-min incubation at 30 °C, reactions were initiated by the addition of 2 mM ATP-Mg. Saturation of nucleosomes by hACF was confirmed by varying the concentration of hACF.

ATPase assay

ATP hydrolysis rates were measured using the Enzchek assay kit (Invitrogen) under multiple-turnover conditions with saturating ATP and nucleic acid ligand over SNF2h (3 mM ATP-Mg, either 400 nM nucleosomes or 8 μM gel-purified, 60-bp duplex DNA and 200 nM SNF2h). Absorbance measurements were taken on a Flexstation 3 platereader (Molecular Devices). Reactions were performed at 30 °C in the following reaction buffer: 60 mM KCl, 3 mM MgCl₂, 20 mM Tris (pH 7.5), 0.02% Nonidet P-40, 200 mM 2-amino-6-mercapto-7-methyl-purine riboside and 1 U/ml purine nucleoside phosphorylase. After a 10-min incubation at 30 °C, reactions were initiated by the addition of 3 mM ATP-Mg. Using a serial dilution of KH₂PO₄ as substrate for the Enzchek reaction, we used a standard curve of A₃₆₀/P_i (μM) to convert absorbance values into the concentration of P_i in micromolar. The value of k_{cat} was obtained by taking the initial slope of each trace (μM/min) and dividing it by the concentration of SNF2h (0.2 μM).

EPR spectroscopy

EPR measurements were performed with a Bruker Instruments EMX EPR spectrometer (Billerica, MA) as described previously [25]. First-derivative, X-band spectra were recorded in a high-sensitivity microwave cavity using 50-s, 10.0-mT-wide magnetic field sweeps. For clarity, only 8.0-mT-wide fields are shown in figures. The instrument settings were as follows: microwave power, 25 mW; time constant, 164 ms; frequency, 9.83 GHz; modulation, 0.1 mT at a frequency of 100 kHz. Each spectrum used in the data analysis is an average of 50–150 sweeps from an individual experimental preparation. 2'3'SLADP was prepared as described previously [52]. 2'3'SLADP was prepared by the same method but replacing ATP with ADP as a starting material [52]. Spectra of 10 μM of the

nucleotide analog 2'3'SLADP were measured with 16 μM SNF2h (or 32 μM SNF2h for the Walker B mutant, E309A) with or without 8 μM nucleosomes or 16 μM 60-bp DNA in the following buffer conditions: 12 mM Hepes (pH 7.9), 4 mM Tris (pH 7.5), 60 mM KCl, 1–1.5 mM MgCl₂, 0.32 mM ethylenediaminetetraacetic acid, 12% glycerol, 0.4 mM DTT, 0.4 mM BenzHCl and 0.4 mg/ml Flag peptide. For experiments with beryllium fluoride, 0.5 mM BeF_x (made immediately before use by combining BeCl₂ and NaF to final concentrations of 0.5 mM and 2.5 mM, respectively) was added, and for experiments with AlF_x, 0.5 mM AlF_x (made immediately before use by combining AlCl₃ and NaF to final concentrations of 0.5 mM and 2.5 mM, respectively) was added. Samples were incubated at room temperature for 10 min before taking the first scan in the EPR machine. Nucleosome and DNA concentrations were varied to ensure that the SNF2h was saturated with either ligand.

Deconvolution of spectra

Experimental spectra were deconvolved into mobile, immobile and unbound probe components using a least-squares minimization FORTRAN program employing the Levenberg–Marquardt algorithm. Basis spectra for the immobile and mobile components in the least-squares minimization were determined by a subtraction procedure using experimentally determined spectra [53]. The unbound basis was determined from solution measurement of the spectrum of spin probe. Double integration of the first-derivative spectra was used to determine the fractions of probes involved in the individual components. As we are interested in the equilibrium between immobile and mobile components, they were weighted 100× relative to the free probe basis in the minimization.

Supplementary data to this article can be found online at <http://dx.doi.org/10.1016/j.jmb.2014.02.021>.

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Abbreviations used:

hACF, human ACF; SHL ± 2 , superhelical location ± 2 ;
FRET, fluorescence resonance energy transfer; EPR,
electron paramagnetic resonance; SF2, superfamily 2.

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