The nucleosomal acidic patch relieves auto-inhibition by the ISWI remodeler SNF2h

Nathan Gamarra¹,², Stephanie L. Johnson¹, Michael J. Trnka³, Alma L. Burlingame³, Geeta Narlikar¹*

*Corresponding Author: Geeta.Narlikar@ucsf.edu

¹Department of Biochemistry and Biophysics, University of California, San Francisco, San Francisco, United States
²Tetrad Graduate Program, University of California, San Francisco, San Francisco, United States
³Department of Pharmaceutical Chemistry, University of California, San Francisco, San Francisco, United States
Abstract

ISWI family chromatin remodeling motors use sophisticated autoinhibition mechanisms to control nucleosome sliding. Yet how the different autoinhibitory domains are regulated is not well understood. Here we show that an acidic patch formed by histones H2A and H2B of the nucleosome relieves the autoinhibition imposed by the AutoN and the NegC regions of the human ISWI remodeler SNF2h. Further, by single molecule FRET we show that the acidic patch helps control the distance travelled per translocation event. We propose a model in which the acidic patch activates SNF2h by providing a landing pad for the NegC and AutoN auto-inhibitory domains. Interestingly, the acidic patch also inhibits the INO80 complex, indicating that this substrate feature can regulate remodeling enzymes with substantially different mechanisms. We therefore hypothesize that regulating access to the acidic patch of the nucleosome plays a key role in coordinating the activities of different remodelers in the cell.
Eukaryotic genomes are packaged into chromatin, enabling large amounts of DNA to fit into the spatial constraints of the nucleus. This packaging has long been appreciated as a passive barrier to DNA access by nuclear factors. The discovery that chromatin regulators play critical roles in virtually all nuclear processes has informed a more nuanced view of chromatin as a dynamic regulatory platform that coordinates access to the genetic material. The smallest unit of chromatin is the nucleosome, a DNA-protein complex composed of ~150 bp of DNA wrapped around an octamer of histone proteins (Luger et al., 1997). Nucleosomes can further interact with each other and with other factors to form higher-order structures (Luger et al., 2012). Consequently, cells have evolved several sophisticated strategies to regulate chromatin structure at the nucleosome level. These include the covalent modification of histone proteins and DNA, as well as non-covalent changes to the position or composition of nucleosomes at specific genomic loci. Many of the non-covalent transformations, ranging from sliding nucleosomes to the complete disassembly of the histone octamer, are catalyzed by ATP-dependent chromatin remodeling enzymes (Zhou et al., 2016). Underscoring their central role in chromatin regulation, remodeling enzymes play essential roles in many processes including transcription, DNA replication, and DNA repair (Falbo and Shen, 2006; Hota and Bruneau, 2016; Price and D'Andrea, 2013). How a relatively small number of remodeler types carry out such diverse regulatory functions remains an area of active research, not least because much remains unknown regarding remodeler mechanisms for substrate recognition and the coupling of that recognition to activity.

Chromatin remodelers are members of the SF2 superfamily of nucleic acid motors, which catalyze noncovalent changes to nucleic acid substrates (Zhou et al., 2016). Chromatin remodelers, however, are unique in that they specifically mobilize DNA in the context of the nucleosome, where DNA is tightly bound to histone proteins. Remodelers are further classified into families based on the domain architecture of their ATPase subunit. These families differ in their specific biochemical activities (Zhou et al., 2016). Substantial progress in our general understanding of remodeling mechanisms has been made by asking what elements of the nucleosome are important for remodeling in different families. For example, maximal remodeling by ISWI family remodelers, which primarily slide nucleosomes, requires DNA flanking the nucleosome and the N-terminal tail of histone H4 (Clapier et al., 2001; Yang et al., 2006). Conversely, maximal remodeling by SWI/SNF family remodelers, which carry out the most diverse set of changes to nucleosome structure, does not require these nucleosomal epitopes
(Guyon et al., 1999). Less is known about how these substrate cues are recognized and mechanistically coupled to remodeling. Some important insights have come from biochemical analyses and structures of remodelers in the absence of nucleosomes, which suggest that in the ground state, chromatin remodelers are held in an inactive conformation by family-specific autoinhibitory motifs (Clapier and Cairns, 2012; Hauk et al., 2010; Xia et al., 2016; Yan et al., 2016). Binding to specific nucleosomal epitopes is thought to relieve this autoinhibition via conformational changes in the remodeler, but the details of this process remain unclear.

At the same time, structures of several nucleosome-protein complexes are revealing that many of these proteins interact with a conserved acidic patch formed by histones H2A and H2B on the top surface of the nucleosome (McGinty and Tan, 2016). These proteins interact with the acidic patch using an “arginine anchor” which nestles into a pocket formed by the α1-2 helices of histone H2A (McGinty and Tan, 2016). It is therefore plausible that chromatin remodelers also recognize the acidic patch. Indeed recent work has indicated that mutating the acidic patch reduces the activity of ISWI, SWI/SNF and some CHD family remodeling enzymes (Dann et al., 2017). To investigate the mechanistic role of the acidic patch in nucleosome remodeling by the ISWI family of enzymes we used a combination of ensemble and single molecule methods in the context of the human ISWI enzyme, SNF2h. We observe that interactions with the acidic patch activate SNF2h by relieving auto-inhibition mediated by two conserved domains of ISWI enzymes. Our results further suggest that contacts with the acidic patch helps control the distance the nucleosome is moved during translocation events. Finally we find that the acidic patch also stimulates the activity of the INO80 complex. Together, these results highlight the broad and essential role the acidic patch plays in chromatin regulation.

Results

The acidic patch of the nucleosome is important for nucleosome sliding by SNF2h

The human ISWI ATPase, SNF2h, preferentially slides nucleosomes toward longer flanking DNA (Yang et al., 2006). As a consequence of this activity, SNF2h slides mononucleosomes towards the center of a DNA strand, an activity that can be detected by a native gel mobility assay (Yang et al., 2006). Recent work has shown that mutating residues in the nucleosome acidic patch reduces the ability of SNF2h to expose nucleosomal DNA to restriction enzymes (Dann et al., 2017). To investigate the mechanism by which such mutations affect the nucleosome sliding activity of SNF2h and to determine if these mutations affected SNF2h
binding or catalysis, we first used a mutant H2A in which 4 key residues in the H2A acidic patch are replaced with alanines (E61A, E64A, D90A, and D92A) (Figure 1A). We call nucleosomes in which all four of these H2A residues have been mutated to alanines “APM” (acidic patch mutant) nucleosomes. Using a native gel mobility assay, we measured remodeling rates of wild type (WT) and APM mononucleosomes containing 60 bp of DNA flanking one end of the nucleosome (0/60) (Figure 1B). APM nucleosomes are centered substantially slower than WT nucleosomes under conditions where SNF2h is in excess of nucleosomes (Figure 1B and 1C). Since the acidic patch has been shown to be critical for nucleosome binding by several chromatin proteins, we expected the apparent $K_m$, $K_m^{app}$, to be increased with APM nucleosomes. Interestingly however, $K_m^{app}$ is affected substantially less by mutation of the acidic patch than the maximal rate constant for remodeling, $k_{max}$ (Figure 1D, 61nM to 280 nM, corresponding to ~4.5-fold increased $K_m^{app}$, versus 2.5 min$^{-1}$ to 0.012 min$^{-1}$, corresponding to ~200-fold reduced $k_{max}$). These results indicate that the acidic patch plays a larger role in regulating maximal nucleosome sliding activity than nucleosome binding. We also measured ATP hydrolysis under conditions where nucleosomes are in excess of SNF2h. At saturating nucleosome concentrations, APM nucleosomes stimulate ATP hydrolysis 4-fold less than WT (Figure 1G, Figure 1-Supplement 2). Together, these results imply that the acidic patch plays a critical role in coupling ATP hydrolysis to remodeling after the binding step.

To further investigate the importance of the acidic patch, we asked whether binding by another factor to the acidic patch could compete for nucleosome sliding by SNF2h. We used a peptide derived from the latency associated nuclear antigen (LANA) from Kaposi’s sarcoma-associated herpesvirus that has previously been shown to interact directly with the acidic patch via an arginine anchor (Barbera et al., 2006). We carried out these experiments using sub-saturating concentrations of SNF2h. Under these conditions most of the nucleosomes are unbound, allowing direct competition between the LANA peptide and SNF2h for the acidic patch. The presence of the LANA peptide dramatically slows remodeling by SNF2h in a dose-dependent manner (~200-fold reduction in rate constant at 50µM LANA peptide), while a peptide with a mutant arginine anchor does not have a detectable inhibitory effect (Figures 1E and F). We measured a $K_i$ of 1.2 µM for inhibition by the LANA peptide, within 5-fold of its published affinity for the nucleosome (Figure 1F) (Fang et al., 2016). These results indicate that remodeling by SNF2h requires an interaction with the acidic patch that is mutually exclusive with the binding of acidic patch interacting factors such as the LANA peptide. The results are also consistent with
recent work showing that the LANA peptide can inhibit the restriction enzyme accessibility activity of the SNF2h containing complex, ACF (Dann et al., 2017).

We next investigated if the acidic patch residues act independently or cooperatively by measuring the effects of individual mutations. Interestingly, except for residue E64, all the single alanine mutants have comparable defects as the four point mutants combined (Figure 1-Supplement 3). This observation suggests that three of the four acidic patch residues tested here act cooperatively to promote sliding by SNF2h.

The AutoN and NegC regions of SNF2h cooperate with the acidic patch to enable maximal remodeling

We next investigated which regions of SNF2h might functionally interact with the acidic patch. In principle, these would include (i) regions that directly contact the acidic patch and (ii) regions that do not directly contact the acidic patch, but whose function is energetically coupled to the presence of the acidic patch.

To investigate regions that may directly contact the acidic patch we carried out cross-linking mass spectrometry using the zero-length, carbodiimide based reagent EDC. This method catalyzes the formation of new amide bonds between protein carboxylates, such as the side chains of aspartate and glutamate residues, and amino groups and is therefore well suited to probing electrostatic interactors of the acidic patch. Hundreds of high confidence cross-linked residue pairs were identified using this approach (Supplementary File 1). To focus on mechanistically meaningful domain-domain interactions, we employed a semi-quantitative mass spectrometry method to compare the extent of SNF2h-nucleosome cross-linking in the presence of different nucleotides (Figure 2B). In previous work we have shown that ADP•BeF₃ mimics an activated state of the SNF2h-nucleosome complex (Leonard and Narlikar, 2015; Racki et al., 2014; Sinha et al., 2017). We therefore, focused on domain level interactions that were enriched at least two-fold in the presence of ADP•BeF₃ relative to ADP.

Cross-links between the H4 tail and RecA lobe 2 of SNF2h are strongly enhanced in the ADP•BeF₃ state (Figure 2B). This result is consistent with previous work showing that the H4 tail activates ISWI remodelers and promotes a restricted active site conformation in the presence of ADP•BeF₃ (Clapier et al., 2001; Hamiche et al., 2001; Racki et al., 2014). We also found that the ADP•BeF₃ state promotes specific cross-links between the H2A/H2B acidic patch and lysines in the extended AutoN region, RecA lobe 1, the NegC region, and the DNA binding HAND-SANT-
SLIDE (HSS) region of SNF2h (Figure 2-Supplement 4). While cross-links between the acidic patch and the AutoN, HSS and NegC regions are compatible with structural constraints from previous studies, the cross-links with the RecA lobe 1 are not easily explained. Multiple studies of ISWI remodelers suggest that their RecA lobes bind at a location two DNA helical turns from the nucleosome dyad (SHL±2), quite far from the H2A/H2B acidic patch (Dang and Bartholomew, 2007; Kagalwala et al., 2004; Schwanbeck et al., 2004; Zofall et al., 2006). We hypothesize that the RecA-acidic patch cross-links arise from a population of higher-order SNF2h-nucleosome aggregates in our mass-spec samples and therefore focus below on cross-links to the remaining three regions.

To test the functional significance of cross-links to AutoN, we mutated the lysines that crosslink to the acidic patch in the ADP•BeF$_\alpha$ state. However, none of the mutants significantly altered remodeling by SNF2h (Figure 2-Supplement 1). These results suggest that the lysines in the AutoN region are in proximity to the acidic patch but do not make mechanistically significant interactions. We speculated, however, that other residues in AutoN might make functional interactions with the acidic patch. Previous work has indicated that many nucleosome binding proteins recognize the acidic patch via arginine residues. Near the AutoN lysines that crosslink to the acidic patch resides a key autoinhibitory motif specific to ISWI family remodelers that contains two arginine residues (Figure 2A, Figure 2-Supplement 4, R142 and R144). Autoinhibition by these arginine residues is relieved by a basic patch on the H4 tail, a nucleosomal epitope essential for maximal remodeling by ISWI-family remodelers (Clapier and Cairns, 2012; Clapier et al., 2002). We tested whether R142 and R144 functionally cooperate with the acidic patch by generating a mutated version of SNF2h with the two critical arginines of AutoN mutated to alanine (2RA), and measured the remodeling activity of this mutant. Consistent with previous reports, 2RA SNF2h remodells WT nucleosomes ~2-fold faster than WT SNF2h (Clapier and Cairns, 2012; Yan et al., 2016). However, 2RA SNF2h remodells APM nucleosomes ~50-fold faster than WT SNF2h (Figure 2F). This corresponds to a ~25-fold reduced dependency on the acidic patch for remodeling with 2RA SNF2h (Figure 2F). The same trend was also seen by an ensemble FRET remodeling assay (Figure 2-Supplement 2).

Together, these data suggest that the acidic patch contributes to relief of autoinhibition by R142 and R144 in AutoN. Since arginine residues are known to mediate this interaction in other systems, binding of either of the two arginines in AutoN to the acidic patch could provide a physical explanation for acidic patch recognition. However, given that neither carbodiimide chemistry nor any other commonly used cross-linking chemistry labels arginine residues, we
cannot determine whether R142 and R144 make direct contacts with the acidic patch based on our mass-spectrometry data. We were unable to observe detectable binding between an AutoN peptide containing the 2R residues and the nucleosomal acidic patch through pull down assays (data not shown), suggesting that either these residues do not physically interact with the acidic patch or the surrounding regions of the SNF2h protein are required for stable binding.

To investigate the functional significance of cross-links to the NegC region of SNF2h, we determined the effect of replacing a stretch of 32 residues in NegC with a flexible serine-glycine linker (mNegC). NegC is another autoinhibitory region of SNF2h that imposes flanking DNA length sensitivity on SNF2h by specifically slowing down remodeling of nucleosomes without flanking DNA (Leonard and Narlikar, 2015). Consistent with previous work, mNegC SNF2h slides WT 0/60 nucleosomes ~1.2-fold faster than WT SNF2h (Figures 2E and F) (Leonard and Narlikar, 2015). However, mNegC SNF2h slides APM nucleosomes ~100-fold faster than WT SNF2h (Figure 2F). As a result, sliding of APM nucleosomes by mNegC SNF2h is only ~2-fold slower than WT nucleosomes. Thus the mNegC mutation almost completely rescues the defect of the acidic patch mutation. These results suggest that residues in NegC also link activation of nucleosome sliding to acidic patch recognition.

The third category of cross-links entailed lysine residues in the HSS regions. Mutants in these lysines greatly reduced remodeling and, in contrast to the 2RA and mNegC SNF2h mutants, did not rescue the defects caused by the acidic patch mutations (data not shown). It is possible that the defects caused by these lysine mutations reflect direct contacts between the HSS residues and the acidic patch. However, because the HSS also contacts flanking DNA, the reduction in sliding rate could also arise from defects in binding DNA.

To better understand which crosslinks are dependent on H2A acidic patch binding, we also performed SNF2h-nucleosome cross-linking reactions with ADP•BeF$_x$ in the presence of the LANA peptide which competes for acidic patch binding (Figure 1E). Addition of the LANA peptide substantially reduced crosslinks between the H2A acidic patch and both AutoN and NegC (Figure 2-Supplement 5). In contrast, crosslinks between the HSS and the acidic patch were less strongly affected by LANA addition (Figure 2-Supplement 5). This suggests that HSS positioning near the acidic patch in the ADP•BeF$_x$ state is not strictly dependent on direct binding to the region of the H2A acidic patch contacted by the LANA peptide. As a result, we cannot unambiguously interpret HSS-acidic patch crosslinks as reflecting mechanistically significant interactions. Together, these data suggest that acidic patch recognition is strongly
linked to relief of autoinhibition by NegC and AutoN for nucleosome sliding by SNF2h.
Furthermore, activation of SNF2h involves conformational changes that bring both of these
autoinhibitory domains and the HSS in closer proximity to the acidic patch.

The acidic patch is required to promote the translocation phase of the SNF2h reaction

To gain additional insight into which steps in the remodeling cycle involve interaction with the
acidic patch, we turned to a single-molecule FRET assay (smFRET; Figure 3A). This assay is
analogous to the ensemble FRET remodeling assay described in Figure 2-Supplement 2, in that
the activity of SNF2h in sliding nucleosomes away from DNA ends increases the distance
between a donor and acceptor dye pair, and thus decreases the measured FRET efficiency.
However, with smFRET we can follow the remodeling of individual, surface-immobilized
nucleosomes, and thereby gain insights into the activity of SNF2h reaction steps that are
obscured in asynchronous, population-averaged ensemble assays.

smFRET has previously been used to study remodeling by several ISWI family members, as
well as by remodeling complexes from the SWI/SNF and INO80 families (Blosser et al., 2009;
Deindl et al., 2013; Harada et al., 2016; Hwang et al., 2014; Zhou et al., 2018). A key insight
revealed by these smFRET studies is that ISWI family remodelers do not slide nucleosomes in
a continuous manner, such that FRET decreases linearly over time, but rather in a series of
alternating phases: a “pause” phase, in which FRET (and therefore nucleosome position)
remain constant, and a “translocation” phase, in which the nucleosome is moved relative to the
DNA end. These repeating pause phases, which so far appear to be specific to ISWI, are
essential to our understanding of the mechanism of action of ISWI remodelers, because the
overall remodeling rates observed in ensemble assays are dominated by the durations of the
pause phases, not the translocation rate itself. Moreover, substrate cues such as the H4 tail and
flanking DNA have been shown to be sensed in the pause phase, not the translocation phase,
for ISWI family remodelers (Hwang et al., 2014). That is, shorter flanking DNA or mutation of the
H4 tail increases the durations of the pauses, thereby decreasing the overall remodeling rate,
while having no effect on the actual sliding rate of the nucleosome. These results suggest a
separation of regulation and activity in ISWI remodelers: translocation events are regularly
interrupted by pauses that allow for the periodic interrogation of substrate cues. This pausing
behavior also explains the ability of ISWI remodelers such as ACF to keep nucleosomes
centered: if the nucleosome is translocated off-center, the interruption of this translocation by a
new pause can trigger translocation in the opposite direction, restoring the nucleosome to a centered position.

As shown in Figure 3B, we find that SNF2h alone, like the ACF complex and the yeast ISWI enzymes, shares this alternating pause and translocation behavior at the single-molecule level.

We first wished to ascertain whether the acidic patch, like other nucleosomal epitopes recognized by ISWI remodelers, affects the regulatory pause phase. Because the rate of remodeling of APM nucleosomes by SNF2h is significantly slower (on the order of hours) than the rate of dye photobleaching (on the order of minutes; Figure 3-Supplement 1B), we assembled nucleosomes with a single point mutation in the acidic patch (E64R nucleosomes). We used the E64R mutation rather than the E64A mutation because the defect caused by this mutation was better rescued by the 2RA mutation in SNF2h than the defect in E64A (Figure 1-Supplement 3). The single-point mutation (E64R) is less deleterious than mutating all 4-acidic residues (APM), and SNF2h remodels E64R nucleosomes ~40-fold more slowly than WT nucleosomes as opposed to 200-fold more slowly with APM nucleosomes (Figure 1-Supplement 3). However, this remodeling rate is still very slow relative to the timescales typically measured by smFRET, which posed two additional challenges: an increase in the number of noise events (dye blinking, intensity fluctuations, etc) per remodeling trajectory, and an increase in the amount of data to be analyzed. These challenges were addressed through the use of custom in-house smFRET analysis software (“Traces”, https://github.com/stephlj/Traces) (Zhou et al. 2018, Johnson et al. 2018) to streamline the analysis of large data sets, and the adaptation of a computationally fast, versatile, open-source hidden Markov model (HMM) library called pyhsmm to quantify the durations of the pauses (see Methods).

As shown in the example trajectories in Figure 3C, remodeling of E64R nucleosomes by SNF2h proceeds through the same alternating pause and translocation phases as does remodeling of WT nucleosomes. However, the pauses are noticeably longer with E64R nucleosomes, by at least a factor of 2 (Figure 3D), indicating that the acidic patch epitope, like other substrate cues, is indeed sensed during the regulatory pause phase. We note that remodeling of E64R nucleosomes in ensemble assays is significantly slower than the photobleaching rate (Figure 3-Supplement 1B), so that by smFRET we only detect the fraction of remodeling events that are faster than photobleaching. As a result, the remodeling rate obtained for the E64R nucleosomes by smFRET represents an upper bound on the true remodeling rate (i.e. the E64R nucleosomes appear to remodel faster by smFRET than by ensemble assays (Figure 3-Supplement 1B)).
Given the ability of the 2RA mutation of the AutoN motif to partially rescue remodeling defects in APM nucleosomes in ensemble assays (Figure 2C), we next asked whether this rescue is due to a restoration of wild-type pause durations. In agreement with our ensemble results (Figure 2C), 2RA SNF2h remodels WT nucleosomes slightly faster than SNF2h by reducing pause durations (Figure 3D). The reduction is minor (~1.3 fold), consistent with a previous smFRET study of the effect of the 2RA mutation on pause durations for the ACF complex (Hwang et al., 2014). Furthermore, 2RA rescues the deleterious effect of the E64R acidic patch mutation, by nearly restoring wild-type pause durations (Figure 3D). The effects of the E64R nucleosomal mutation and the 2RA SNF2h mutation on pause durations therefore mirror the effects on overall remodeling rates of APM nucleosomes: 2RA remodels E64R nucleosomes nearly as fast as SNF2h remodels WT nucleosomes, but not as fast as 2RA remodels WT nucleosomes (Figure 1-Supplement 3). These results are consistent with a model in which relief of autoinhibition of the AutoN motif of SNF2h through direct or indirect interactions with the acidic patch enable pause exit (that is, promote the translocation phase).

The example traces in Figure 3C suggest an additional defect in remodeling of E64R nucleosomes: a reduction in the distance the nucleosome is moved during each translocation event (which we called the step size). Note that after two translocation events, WT nucleosomes with SNF2h or 2RA are moved from ~0.95 FRET to ~0.4 FRET (Figure 3C top and second from bottom), whereas after two translocation events the nucleosome in the E64R/SNF2h example trace has moved from ~0.95 FRET to ~0.75 FRET. Step size, like pause duration, plays an important role in regulating ISWI remodeler activity: since the pause durations dominate the overall remodeling rate, a smaller step size, which means more pauses per unit distance that the nucleosome is translocated, will mean a significant reduction in the overall remodeling rate (as observed in ensemble assays).

Step sizes can be quantified by converting the change in FRET between subsequent pauses to a change in the number of base pairs of DNA between the Cy5-labeled DNA end and the edge of the nucleosome. We accomplish this conversion by means of a calibration curve, described previously (Zhou et al., 2018). Like other ISWI family remodelers, SNF2h moves WT nucleosomes with an initial large step (~8 bp) followed by a smaller (~5 bp) step (Figure 3E, Figure 3-Supplement 3; (Blosser et al., 2009; Deindl et al., 2013)). However, SNF2h moves E64R nucleosomes a shorter distance in each translocation phase, as indicated by the leftward shift of the cumulative distributions in the red curves of Figure 3E, relative to the black curves (on average, E64R nucleosomes move about 6 bp in the first translocation and about 4 bp in the
Mutation of AutoN has little effect on the step size in the context of WT nucleosomes, but largely restores the step size in the context of E64R nucleosomes to that of SNF2h with WT nucleosomes (Figure 3E, magenta and blue curves). Given that ISWI family remodelers have been shown to translocate a nucleosome in elementary steps of 1-2 bp (Deindl et al., 2013), our results suggest that with E64R nucleosomes, SNF2h takes fewer of these elementary steps in succession during the translocation phase, before entering a new pause phase.

The major contributions to the remodeling defects observed with SNF2h and APM nucleosomes can therefore be attributed to two effects: first, an increase in pause duration, and second, a decrease in the distance travelled per translocation event, meaning that there are more (and longer) pauses in APM remodeling reactions per unit distance that the nucleosome is moved. Both of these effects are rescued by the 2RA mutation of the AutoN motif of SNF2h. We propose that the acidic patch is important for relieving autoinhibition by AutoN and thereby promoting exit from the pause phase. Further, we propose that the acidic patch is also involved in keeping AutoN out of the active site until the nucleosome has been translocated to the full extent (~8 bp initially, ~5 bp subsequently) and a new pause phase is entered (Figure 5).

The acidic patch is used by both ISWI and INO80 complexes

The ISWI ATPase forms complexes with several accessory proteins that regulate its localization and activity (He et al., 2006; Oppikofer et al., 2017; Tsukiyama et al., 1995; Varga-Weisz et al., 1997). ACF is one of the best studied of these complexes. ACF is a heterodimer of the ISWI ATPase subunit (SNF2h in humans) and the accessory subunit Acf1, and is implicated in gene repression, DNA replication, and DNA repair (Collins et al., 2002; Fyodorov et al., 2004; Lan et al., 2010). Biochemically, human ACF has the same core activity as SNF2h, but displays greater nucleosome affinity, enhanced sliding rates, and better kinetic discrimination of flanking DNA length (He et al., 2006; Yang et al., 2006). Despite these similarities, recent evidence suggests that ACF has some mechanistic differences from SNF2h on its own. For instance, in the context of ACF, AutoN regulates flanking DNA length sensing through interaction of an Acf1-specific domain called WAC (Hwang et al., 2014). SNF2h alone has no comparable domain. Furthermore, recent work has suggested that mutating the nucleosomal acidic patch causes a smaller defect in remodeling by ACF compared to SNF2h (Dann et al., 2017). Given this difference, we asked whether ACF requires the acidic patch for remodeling beyond binding. At saturating concentrations of ACF, where binding does not contribute to the overall remodeling
rate, ACF slides APM nucleosomes 10-fold more slowly than WT (Figure 4A), indicating that
ACF also uses the acidic patch in a step after binding. However, consistent with previous work,
ACF is less dependent on the acidic patch compared to SNF2h alone (Dann et al., 2017).

We next asked whether ISWI-family complexes uniquely use the acidic patch, or whether it is
also used by other chromatin remodeling enzymes. Recently, it was shown that while some
CHD family remodelers slide nucleosomes largely independent of the acidic patch, others are
dependent on it (Dann et al., 2017; Levendosky et al., 2016). It has been further shown that
SWI/SNF family enzymes also require the acidic patch for maximal activity (Dann et al., 2017).
These observations raise the possibility that acidic patch is a feature used by most remodeling
enzymes. To address this issue we determined if the acidic patch is required by yeast INO80,
which is in a distinct family from the CHD, ISWI and SWI/SNF families. INO80, like ACF, slides
nucleosomes preferentially in the direction of longer flanking DNA and can also create evenly
spaced nucleosome arrays (Udugama et al., 2011). This sliding activity is thought to be
important for positioning the +1 nucleosome at transcription start sites (Krietenstein et al.,
2016). Interestingly, we find that INO80 slides APM nucleosomes ~200-fold more slowly than
WT nucleosomes at saturating concentrations (Figure 4B). This result indicates that INO80 also
uses the acidic patch post-binding, but is more dependent on the acidic patch than the ACF
complex.

**Discussion**

In this study, we investigate the role of the highly conserved H2A acidic patch in chromatin
remodeling by ISWI enzymes. We find that the acidic patch is used post-binding in order to
activate remodeling by both INO80 and ISWI family remodelers. Furthermore, using a
combination of ensemble and single molecule methods, we show that the acidic patch is used
by SNF2h to relieve autoinhibition by the conserved AutoN and NegC motifs. Below we explore
the mechanistic and regulatory implications of these results.

ATP-dependent chromatin remodeling enzymes carry out specialized reactions on a complex
substrate. Understanding how recognition of this substrate is coupled to activity can provide a
means to understanding common principles underlying ATP-dependent chromatin remodeling
mechanisms. Owing to decades of study, ISWI enzymes provide a useful model system to
address this question. On the basis of crosslinking and footprinting studies, the ATPase domain
of ISWI enzymes is thought to bind and translocate DNA two helical turns from the nucleosomal
dyad (SHL±2) (Dang and Bartholomew, 2007; Kagalwala et al., 2004; Schwanbeck et al., 2004; Zofall et al., 2006). Work from several groups has also shown that for ISWI remodelers, recognition of both the H4 tail and flanking DNA enhances remodeling activity post-binding (Clapier et al., 2001; Hamiche et al., 2001; Yang et al., 2006). While the mechanisms by which these nucleosome cues activate remodeling are not well understood, the ISWI domains that recognize these cues are known. The C-terminal HAND-SANT-SLIDE (HSS) domain mediates flanking DNA recognition, while the H4 tail appears to be directly recognized by the second RecA lobe within the ATPase domain (Dang and Bartholomew, 2007; Yan et al., 2016). The acidic patch resides on a surface near SHL±6, far from where the ATPase domain engages the nucleosome. How then might the acidic patch be recognized and used by ISWI remodelers?

Our results suggest that two known autoinhibitory regions of ISWI enzymes, the AutoN region and the NegC region, functionally interact with the acidic patch, because mutating these regions dramatically reduces the dependence of SNF2h on the acidic patch for sliding. This suggests that a large role of the acidic patch is to relieve autoinhibition by both AutoN and NegC. While we do not have evidence for a direct interaction between the acidic patch and the two arginines in AutoN, our cross-linking mass spectrometry data suggests that activation of the enzyme places residues within both AutoN and NegC near this location. Our smFRET work here and previous smFRET work suggests that the AutoN region inhibits the transition from the pause phase to the translocation phase (Hwang et al., 2014). It has been shown that flanking DNA and the H4 tail are both sensed in the pause phase (Hwang et al., 2014). Further, previous ensemble work has suggested that the NegC region inhibits the transition between a flanking DNA sensing state of SNF2h and a translocation competent state of SNF2h (Leonard and Narlikar, 2015). We therefore propose that the acidic patch helps promote the translocation competent state of SNF2h by providing an alternative binding site for NegC and AutoN (Figure 5).

In addition to an increase in pause durations with the acidic patch mutations, the amount of DNA translocated within a translocation phase is reduced compared to WT nucleosomes. We hypothesize that translocation is interrupted by premature reversion of the enzyme to the autoinhibited state in the absence of stabilizing interactions with the acidic patch (Figure 5). As a result, more pauses are encountered per distance translocated. Our results lead to a model in which the acidic patch provides a binding surface for NegC and AutoN that sequesters these regions from inhibiting SNF2h (Figure 5). Combined with previous work, our results underscore how the strong coupling of relief of autoinhibition to recognition of two conserved nucleosome
cues (the H4 tail and the acidic patch) make this motor exquisitely specific for its complex substrate.

The acidic patch increases $K_{\text{m, app}}$ for SNF2h by ~5-fold, suggesting that interactions with the acidic patch also stabilize SNF2h binding. Thus the acidic patch appears to be used in at least two distinct steps of the SNF2h remodeling reaction. What residues in SNF2h might be interacting with the acidic patch in these two steps? In contrast to the rescuing effects of AutoN mutations on maximal activity, AutoN mutations additively increase $K_{\text{m, app}}$ beyond solely mutating the acidic patch (Figure 2–Supplement 3). This suggests that AutoN and the acidic patch do not cooperate in the ground state. It is thus possible that different SNF2h domains contact the acidic patch in different steps of the remodeling reaction. The different set of cross-links observed between the acidic patch and SNF2h regions in the ADP vs. ADP•BeF$_x$ state are consistent with such a possibility.

Based on previous work, we have hypothesized that SNF2h undergoes a large conformational change prior to adopting a translocation competent state that repositions the C-terminus from binding flanking DNA towards binding the nucleosome core (Leonard and Narlikar, 2015). Our crosslinking-MS data provides additional insights into the structural rearrangements that accompany enzyme activation. We find that crosslinks between the H2A/H2B acidic patch and the SANT domain increase in the ADP•BeF$_x$-bound state compared to the ADP-bound state (Figure 2B), consistent with the HSS binding to the nucleosome core. Interestingly, these crosslinks are not substantially reduced by LANA peptide binding, suggesting that the location of the HSS in the ADP•BeF$_x$-bound state is not strongly dependent on direct contacts with the residues contacted by the LANA peptide (Figure 2-Supplement 5). This result raises the possibility that the SANT domain and LANA peptide occupy adjacent regions on the nucleosome core. The SANT domain may then cross-link to the acidic patch when these sites are transiently exposed due to dynamics in LANA peptide binding. In contrast, NegC and AutoN crosslinks to the acidic patch are substantially reduced by the LANA peptide, suggesting LANA binding directly or indirectly displaces these regions from the nucleosome. Overall our results suggest that the key accessory regions of SNF2h namely, HSS, NegC and AutoN, are all positioned near the acidic patch in the activated state. In agreement with this observation, crosslinks between the C-terminus of SNF2h and NegC increase in the ADP•BeF$_x$ state (Figure 2B). The close positioning of multiple SNF2h accessory domains near the acidic patch raises the possibility that contacts between accessory domains may play a role in promoting the translocation-competent state. While substantial future work would be needed to test this
possibility, it is analogous to recent observations with the yeast CHD1 remodeling motor that
contacts between its N-terminus and the C-terminal DNA binding domain regulate the sliding
reaction (Sundaramoorthy et al., 2017).

Most chromatin remodeling ATPases also form large multi-subunit complexes, which regulate
the activity and specificity of the remodeling reaction. Here we find that the human ISWI
complex, ACF, requires the acidic patch for maximal activity but shows a ~20-fold smaller defect
upon mutation of the acidic patch than observed with SNF2h alone. This result is qualitatively
consistent with recent studies showing that the acidic patch has a smaller role in the activity of
ACF vs. SNF2h (Dann et al., 2017). Our results here provide a mechanistic framework for
understanding these recent observations. In particular, our results suggest that the accessory
protein Acf1 alters the mechanism of relief of autoinhibition by the acidic patch in a manner that
makes the reaction less dependent on the acidic patch, perhaps by providing an alternative
binding partner for the AutoN and NegC domains. Such a domain would be analogous to the
WAC motif of Acf1, which provides an alternative binding partner for the H4 tail (Hwang et al.,
2014). In contrast to ACF, we find that the yeast INO80 complex, a large multisubunit complex,
is as dependent on the acidic patch for nucleosome sliding as the isolated SNF2h ATPase.
Importantly, INO80 family remodelers are insensitive to the presence of the H4 tail and no
AutoN-like or NegC like motif has been identified in the ATPase subunit of INO80 (Udugama et
al., 2011). The acidic patch must then activate INO80 through a mechanism distinct from ISWI
complexes. Determining how the acidic patch is used by INO80 ATPases, and what roles the
ATPase and accessory subunits play in this mechanism, are important areas of future study.

Combined with previous results (Dann et al., 2017), our results suggest that several families of
chromatin remodelers require the acidic patch for remodeling. However, it is possible that this
surface is not a universal requirement for chromatin remodeling, as yeast CHD1 can remodel
nucleosomes largely independent of the acidic patch (Levendosky et al., 2016). CHD1 instead
uses an unidentified aspect of the histone H2A/H2B dimer to promote remodeling (Levendosky
et al., 2016). Yeast CHD1 and ISWI family remodelers have been thought to share a common
remodeling mechanism, as these families share biochemical activities, like nucleosome sliding
and spacing, and substrate cues required for maximal remodeling activity, such as the H4 tail
and flanking DNA (Ferreira et al., 2007; Stockdale et al., 2006). However, the enormous
difference in dependence on the acidic patch between yeast CHD1 and ISWI enzymes raise the
possibility that these families remodel nucleosomes through distinct mechanisms (Levendosky
et al., 2016). Importantly, while yeast CHD1 shares domains with ISWI remodelers such as the
SANT-SLIDE domains at the C-terminus and a version of the NegC region, called the C-terminal bridge (Hauk et al., 2010; Ryan et al., 2011), CHD family remodelers do not appear to possess an AutoN motif. Instead, remodelers like CHD1 have an N-terminal double chromodomain which has an analogous role as an autoinhibitory domain that is relieved by H4 tail binding (Hauk et al., 2010).

At a primary level, the requirement of the acidic patch provides a powerful means for remodelers to sense and respond to chromatin structure and nucleosome content. Thus, nucleosomes lacking histone H2A-H2B dimers or containing a modified acidic patch through histone variants or covalent modifications may be recognized and remodeled differently than canonical nucleosomes by different remodelers (Dann et al., 2017). Consistent with this possibility, nucleosomes containing histone H2Az, which have an extended acidic patch, are remodeled ~2-fold faster by ISWI complexes than canonical nucleosomes (Goldman et al., 2010). Analogously, recent work has shown that find that INO80 preferentially slides H2AZ nucleosomes over H2A nucleosomes (Brahma et al., 2017). Finally, given the growing list of factors that recognize the acidic patch, it is likely that remodelers and other chromatin binding proteins compete for access to the acidic patch. Indeed, binding by the LANA peptide to the acidic patch competes for remodeling by SNF2h. Sensitivity to the acidic patch could be a general mechanism to regulate the outcome of chromatin remodeling at loci where remodelers and other factors are jostling for access to their chromatin substrates.

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Methods

Expression and purification of chromatin remodeling enzymes

SNF2h was purified from *E. coli* as described previously with minor modifications (Leonard and Narlikar, 2015). DNA was precipitated following cell lysis by addition of 5% w/v Polyethylenimine (P3143, Sigma-Aldrich, St. Louis, MO) pH 7.9 dropwise to a final concentration of 0.1% and clarified by centrifugation. Following Cobalt affinity purification, the 6xHis tag was cleaved overnight with TEV protease and dialysed into SEC Buffer. TEV-cleaved SNF2h was then purified by anion exchange chromatography using a HiTrap Q column and Size Exclusion Chromatography (GE Life Sciences, Pittsburgh, MA). SNF2h concentration was determined by SDS-PAGE with BSA protein standards and staining with SYPRO Red (Thermo Fisher, Waltham, MA).

Human ACF was expressed and purified recombinantly from Sf9 insect cells by FLAG immunoaffinity purification as described previously with minor modifications (Aalfs et al., 2001). SNF2h-FLAG and Acf1 were expressed separately via infection with baculovirus. Nuclear extracts from each construct were generated and mixed together at a 10:1 Acf1:SNF2h-FLAG volume ratio to ensure full assembly of the complex. This mixture was then bound to FLAG M2-affinity resin (Sigma-Aldrich, St. Louis, MO), washed with increasing KCl concentrations, and eluted with buffer with 100mM KCl and 1mg/mL FLAG Peptide. ACF concentration was determined by SDS-PAGE with BSA standards and based on the intensity of the Acf1 band.

INO80 was purified by FLAG immunoprecipitation based on previously published methods (Shen, 2004; Zhou et al., 2018). Briefly, *S. cerevisae* with endogenously flag tagged INO80 was grown in YEPD at 30°C to saturation. Cells were pelleted by centrifugation for 10 min at 5000 rpm, resuspended with buffer H0.3, and pelleted again. Pelleted cells were then extruded through a 60mL syringe into liquid nitrogen to create “noodles”. Cell “noodles” were then lysed using a Tissue Lyser II (Qiagen, Hilden Germany) cooled in liquid nitrogen. Frozen lysate powder was resuspended in equal volume of H0.3 and spun in an SW28 rotor for 2 hr at 25,000 rpm at 4°C. Clarified lysate was mixed with equal volume buffer H0.3 and applied to FLAG M2-affinity resin (1 mL bead slurry per 40 mL of cleared lysate) equilibrated with H0.3 and incubated for 3 hours at 4°C. Resin was washed with 3x50 mL buffer H0.5 followed by 3x10 mL washes with buffer H0.1 and eluted with H0.1 supplemented with 1mg/mL FLAG peptide. Eluate was concentrated, aliquoted, flash frozen in liquid nitrogen, and stored at -80°C. INO80
concentration was determined by SDS-PAGE with BSA standards, based on the intensity of the Ino80-flag band.

Nucleosome labeling and reconstitution

Recombinant *Xenopus laevis* histones were expressed and purified from *E. coli* as previously described (Luger et al., 1999). Histone H2A E61A, E64A, D90A, D92A expression plasmid was a generous gift from the Tan lab at Penn State. Purified histone H2A E64R was provided by the Wolberger lab. Histone octamer was reconstituted as previously described (Luger et al., 1999; Zhou and Narlikar, 2016), except for smFRET nucleosomes where a 2:1 unlabeled:labeled H3 mixture was used during octamer assembly to generate nucleosomes with mostly one H3 or neither H3 labeled. Histone H3 with a cysteine introduced at position 33 was labeled with either Cy3 (for smFRET) or Cy5 (for ensemble assays) prior to histone octamer assembly via cysteine-maleimide chemistry. Cy3-labeled (for ensemble assays) and Cyanine 5 SE-labeled and biotinylated DNAs (for smFRET) were generated by PCR with HPLC-purified, labeled primers (Cy5 primers: TriLink Biotechnologies, San Diego, CA; Cy3 and biotinylated primers: IDT, Coralville, IA) and purified by PAGE. The strong, synthetic 601 nucleosome positioning sequence (Lowary and Widom, 1998) was used to assemble all nucleosomes in this study, with an arbitrary sequence for DNA flanking the 601 positioning sequence (Figure 1-Supplement 1). These DNAs were assembled with either wild-type or APM octamers by salt gradient dialysis and purified by glycerol gradient centrifugation (Zhou and Narlikar, 2016).

Native gel remodeling assay

All remodeling reactions were performed under single turnover conditions (enzyme in excess of nucleosomes). Reactions with SNF2h were performed at 20°C with 20nM nucleosomes, 12.5mM HEPES pH 7.5, 2mM Tris pH 7.5, 70mM KCl, 5 mM ATP•MgCl2, 3mM MgCl2,0.02% NP40, and ~3%(v/v) glycerol. Reactions with ACF and INO80 were performed as above at 30°C and with minor changes in buffer composition (ACF: 10 nM nucleosomes, 12.5 mM HEPES pH 7.9, 2 mM Tris pH 7.5, 60 mM KCl, 2 mM ATP•MgCl2, 3 mM MgCl2, 0.02% NP40, 0.3 mg/mL FLAG peptide, and ~9% glycerol; INO80: 40 mM Tris pH 7.5, 60 mM KCl, 2 mM ATP•MgCl2, 1.1 mM MgCl2, 0.02% NP40, 0.5 mg/mL FLAG peptide, and 1% glycerol). Reactions were started with addition of enzyme and time points were quenched with excess ADP and plasmid DNA. Time points were then resolved by native PAGE (6% acrylamide, 0.5XTBE) and scanned on a Typhoon variable mode imager (GE Life Sciences, Pittsburgh, PA) by scanning for fluorescent...
labels. Gels were then quantified by densitometry using ImageJ. The fraction of nucleosomes end-positioned (i.e. unremodeled) at a given time point was determined by the ratio of fast-migrating nucleosomes to the total nucleosome intensity. This was fit to a single exponential decay using Prism 6 (GraphPad, La Jolla, CA) (Equation 1),

\[ y = (y_0 - p) e^{-k_{obs} t} + p \]  

where \( y_0 \) is the initial fraction end-positioned, \( k_{obs} \) is the observed rate constant, and \( p \) is the fraction end-positioned at plateau. Reactions in a given concentration series were fit constrained to a common \( y_0 \) and \( p \). Concentration series were fit to a cooperative binding model (Equation 2),

\[ k_{obs} = k_{max} \frac{X}{(K_m^{app})^h + X^h} \]  

where \( X \) is the concentration of SNF2h, \( h \) is the hill coefficient, \( K_m^{app} \) is the apparent \( K_m \), and \( k_{max} \) is the saturating rate constant. Competition assays were performed as described above with varying concentrations of LANA peptide and fit to a single exponential decay. This was then fit to a simple competition binding model (Equation 3),

\[ k_{obs} = \frac{k_0}{1 + \frac{X}{K_I}} \]  

where \( k_0 \) is the rate constant without peptide, \( X \) is the concentration of peptide, and \( K_I \) is the inhibition constant.

**ATPase assay**

ATPase reactions were performed under multiple turnover conditions (nucleosomes in excess of enzymes). Reactions were performed with 25nM SNF2h, 12.5mM HEPES pH 7.5, 2mM Tris pH 7.5, 70mM KCl, 7.5µM ATP•MgCl₂, 3mM MgCl₂, 0.02% NP40, ~3%(v/v) glycerol, and trace amounts of \( \gamma^{32P}-ATP \). Reactions were started with addition of enzyme, and 2.5 µL time points were quenched with an equal volume of 50 mM Tris pH 7.5, 3% SDS, and 100 mM EDTA. Inorganic phosphate was resolved from ATP on a PEI-cellulose TLC plate (Select Scientific) with 0.5 M LiCl/1M formic acid mobile phase. Plates were dried, exposed to a phosphorscreen overnight, and scanned on a Typhoon variable mode imager. Rate constants were determined by fitting a line through the first 10% of inorganic phosphate generated using Prism.
**Ensemble FRET remodeling assay**

Ensemble FRET remodeling assays were performed under the same conditions as gel remodeling assays. Reactions were initiated by addition of enzyme and then measured in a K2 fluorometer (ISS) equipped with a 550 nm short pass excitation filter and a 535 nm long pass emission filter. Reactions were excited at 515 nm and emission was measured at 665 nm. The resulting curves were fit to a two-phase exponential decay (Equation 4),

\[ y = p + (y_0 - p)(f_{fast} e^{-k_{fast}t}) + (1 - f_{fast}) e^{-k_{slow}t} \]  

where \( f_{fast} \) is the fraction in the fast phase and \( k_{fast} \) and \( k_{slow} \) are the remodeling rates of the fast and slow phase respectively.

**Crosslinking mass spectrometry**

Crosslinking mass spectrometry samples were prepared by incubating 72µg of mononucleosomes without flanking DNA at 9µM final concentration with 24µM SNF2h in buffer containing either ADP or ADP•BeFx (15 mM HEPES pH 7.5, 140 mM KCl, 0.5mM ADP•Mg, 0.5mM MgCl₂, ±0.5mM BeFx 1:5 BeCl₂:NaF) for 10 min at 30°C. The crosslinking reaction with the LANA peptide was performed the same as the ADP•BeFx condition with 30µM peptide added. The samples were then reacted with 1 mM EDC and 20 µM N-hydroxysulfosuccinimide (added as a 10x stock in water) for 60 minutes at room temperature. Crosslinking reactions were then quenched by adding 10 mM Tris pH 7.5 and samples were acetone precipitated and washed once with cold acetone. The pellet was resuspended in 8M Urea, 5 mM TCEP, 100 mM ammonium bicarbonate and heated at 56°C for 25 minutes, followed by alkylation with 10 mM iodoacetamide for 40 min at room temperature. The sample was diluted 5-fold with 100 mM ammonium bicarbonate and digested with 1:25 trypsin for 4 hours at 37°C followed by addition of a second aliquot of trypsin and overnight digestion.

Crosslinked peptides were desalted using 100 µl OMIN C18 tips (Agilent), fractionated by size-exclusion chromatography (SEC), and analyzed by LC-MS similarly to a previously described method (Zhou et al., 2017). Briefly, trypsin digests were acidified to 0.2% TFA, desalted, and run over a Superdex Peptide PC 3.2/300 SEC column (GE Healthcare). SEC fractions eluting between 0.9 ml and 1.4 ml were dried and resuspended in 0.1% formic acid for LC-MS. Each fraction was separated over a 15 cm x 75 µm ID PepMap C18 column (Thermo) using a NanoAcquity UPLC system (Waters) and analyzed by a Fusion Lumos mass spectrometer.
Precursor ions were measured from 375-1500 m/z in the Orbitrap analyzer (resolution: 120,000; AGC: 4.0e5). Ions charged 3+ to 8+ were isolated in the quadrupole (selection window: 1.6 m/z units; dynamic exclusion window: 30 sec; MIPS Peptide filter enabled), fragmented by HCD (Normalized Collision Energy: 28%) and measured in the Orbitrap (resolution: 30,000; AGC: 5.0e4). The cycle time was set to 3 seconds.

Peaklists were generated using PAVA (UCSF) and searched for crosslinked peptides with Protein Prospector 5.19.22 (Trnka et al., 2014) against a target database containing human SNF2h plus the four core histone sequences from *X. laevis* concatenated with a decoy database containing 10 randomized copies of each target sequence (total database size 55 sequences). Loss of the initiator methionine and carbamidomethylation of cysteine. Methionine oxidation, peptide N-terminal glutamine to pyroglutamate formation, acetylation at the protein N-terminus, and mis-annotation of the monoisotopic peak (1Da neutral loss) were treated as variable modifications. EDC was designated as a heterobifunctional crosslinking reagent with specificity of aspartate, glutamate, and the protein C-terminus on one side and lysine and the protein N-terminus on the other with a bridge mass corresponding to loss of H2O. A mass modification range of 400-5000 Da was specified on these residues and 85 product ion peaks from the peaklist were used in the search. Precursor and product ion tolerances were 8 and 25 ppm respectively.

Crosslinked spectral matches (CSMs) were initially classified as in (Zhou et al., 2017). The dataset was then aggregated into unique crosslinked residue-pair level data with a corresponding spectral count value. Due to the prevalence of multiple, closely spaced Asp and Glu residues in a typical tryptic peptide, site-localization of EDC crosslinks is more challenging than with homobifunctional lysine-directed reagents. To address this, when the site-localization was judged to be ambiguous, all possible residue-pairs were kept with an annotation noting the ambiguity. When calculating spectral counts, fractional spectral counts were assigned to these ambiguous site localizations so that a given spectrum was awarded exactly 1 spectral count. For instance, a product ion spectrum matching equally well to both K91.H4-D65.H2B or K91.H4-E68.H2B contributes 0.5 towards the spectral counts of each residue-pair. Decoy CSMs were retained throughout this aggregation and spectral counting process. A linear SVM model, built on five features of the Protein Prospector search output (score difference, % of product ion signals matched, precursor charge, rank of peptide 1, and rank of peptide 2) was constructed to sort crosslinked residue pairs into decoy and target classes. Crosslinked residue-pairs with an SVM score greater than 1, score difference greater than 5, and at least one spectral count are
reported. The final residue-pair level data set is reported at specificity of 99.7% corresponding to 0.05% FDR. The number of unique crosslinks in the ADP condition was 974, while 1470 crosslinks were unique to the ADP•BeFx condition and 707 crosslinks were common to both conditions.

To determine which protein domains are involved in SNF2h mediated nucleosome sliding, residue level crosslink spectral counts were aggregated into domain level counts. Each domain pair was assigned a minimum spectral count of 1 to avoid dividing by zero, and the log2 ratio of spectral counts for each domain pair in the ADP•BeFx condition to the ADP condition were calculated. Domain pairs with a Log2 ratio of exactly 0 were treated as NA values. The remaining data were normalized such that the median value was set at 0. Hence, most domain-domain interactions were assumed to not change substantially between conditions.


The sequence of SNF2h was identical to the Human entry in Uniprot (O60264) with an additional two amino acids at the N-terminus to match the construct used. All residue numbers are therefore shifted from the Uniprot entry by 2 aa. SNF2h domains were defined as follows: Snf2h1: 1-83, AutoN: 84-160, Snf2h2: 161-183, RecA1: 184-402, RecA2: 403-641, NegC: 642-703, Snf2h4: 704-736, HAND: 737-839, SANT: 840-894, SLIDE: 895-1013, Snf2h5: 1014-1054.

Annotated Mass Spectra are available using MS-Viewer at: [http://msviewer.ucsf.edu/prospector/cgi-bin/msform.cgi?form=msviewer](http://msviewer.ucsf.edu/prospector/cgi-bin/msform.cgi?form=msviewer)

The ADP data set is accessed with search key: 2x0kr2kzq1

The ADP•BeFx data set is accessed with search key: fjamygr8pl

The ADP•BeFx in the presence of LANA peptide is accessed with search key: c5o2mcxwum.

Raw mass spectrometry data is available in the MassIVE repository at UCSD with accession key: MSV000082136

**Single molecule FRET measurements.**
smFRET experiments were performed as previously described in (Zhou et al., 2018) with modifications to the reaction buffers as described below.

**Sample preparation and imaging.** Briefly, as in (Zhou et al., 2018), quartz slides were PEGylated and then incubated with neutravidin (A2666, ThermoFisher Scientific, Waltham, MA) to mediate attachment of biotinylated nucleosomes. After removal of unbound neutravidin, biotinylated nucleosomes at 12.5 pM in a modified Wash Buffer (12 mM HEPES-KOH, pH 7.5 at 22°C, 60 mM KCl, 1.4 mM MgCl₂, 10% glycerol, 0.1 mM EDTA, 0.02% Igepal, 1% [w/v] glucose, and 0.1 mg/mL acetylated BSA) were added and incubated for 10 minutes. Unbound nucleosomes were removed by washing with Wash Buffer. All incubations and experiments were performed at 20°C. Nucleosomes were imaged on a custom-built prism-based TIRF setup.

Immediately prior to data acquisition, the sample chamber was flushed with a modified imaging buffer (53 mM HEPES-KOH, pH 7.5 at 22°C, 9.1 mM Tris-acetate, pH 7.5 at 22°C [contributed by the Trolox], 63 mM KCl, 1.41 mM MgCl₂, 10% glycerol, 0.1 mM EDTA, 0.02% Igepal, 1% [w/v] glucose, 0.1 mg/mL acetylated BSA, 2 mM Trolox [Sigma 238813, made as an 11 mM stock in Tris-acetate, pH’d to 7.5 with 1 M NaOH, and stored at 4°C], 0.03 mM β-mercaptoethanol, 2 U/μL catalase [=0.2 mg/mL; Sigma E3289], and 0.08 U/μL glucose oxidase [0.8 mg/mL, Sigma G2133; made with the catalase as a 100x stock in SPB, and stored at 4°C for not more than one week]). Images were collected using Micro-Manager (www.micro-manager.org, San Francisco, CA) (Edelstein et al., 2010) at 7.4 Hz, with an exposure time of 100 ms. To start each reaction, saturating enzyme (400 nM for WT nucleosomes; 2 μM for E64R nucleosomes) and saturating ATP (1 mM) in 300 μL imaging buffer were added via an automated syringe pump (J-KEM Scientific, St. Louis, MO).

**Data analysis.** The number of remodeling events per smFRET experiment was roughly an order of magnitude lower with E64R nucleosomes compared to WT nucleosomes, necessitating significantly larger data sets for the E64R nucleosomes compared to WT. To streamline the data analysis process with these data sets, as well as to improve the overall quality of the data, we made use of the custom in-house software we have developed for smFRET image analysis, called Traces, available for download at [https://github.com/stephlj/Traces](https://github.com/stephlj/Traces) (Zhou et al., 2018, Johnson et al. 2018). In addition, the long pauses exhibited by SNF2h remodeling E64R nucleosomes are subject to an increased number of artifacts, such as dye blinking or slight fluctuations in the noise, which can complicate quantification of smFRET trajectories. We therefore used the python-based HMM library pyhsmm ([https://github.com/mattjj/pyhsmm](https://github.com/mattjj/pyhsmm)),
which we adapted for the analysis of smFRET data as part of the Traces package, to quantify pause durations. This particular HMM package, which fits a discrete state HMM to each trajectory generated by Traces, enables a reduction in the likelihood of the HMM identifying artifacts as real transitions, and also reduces analysis time.

As described in Figure 3–Supplement 1, and consistent with previous smFRET studies with nucleosomes (Blosser et al., 2009; Deindl et al., 2013; Hwang et al., 2014), we observe two predominant clusters of FRET values, at 0.57 and 0.95 FRET, in the absence of remodeler. These FRET states correspond to two of the four populations of nucleosomes that result from mixing unlabeled H3 with Cy3-labeled H3 during octamer formation: some nucleosomes will have a Cy3 label on the H3 proximal to the Cy5-labeled DNA end, resulting in the higher FRET state, and some will have a Cy3 label on the H3 distal to the Cy5-labeled DNA end, resulting in the mid-FRET state. There will also be a population of nucleosomes with both H3 histones unlabeled, which show no FRET; and a population with both copies of H3 labeled, which are distinguishable by two-step photobleaching of the Cy3 dye, and are excluded from all analyses (i.e. both calibration curve data (Zhou et al., 2018) and remodeling data). We also excluded any trajectories to which pyhsmm fit an initial FRET value lower than 0.775 FRET, since nucleosomes with distally labeled H3's do not provide as great a dynamic range for monitoring nucleosome remodeling. We excluded any part of a trajectory including and subsequent to backtracking events (where the nucleosome was moved away from the center, that is, where FRET increased instead of decreasing). Each data set (e.g., E64R/SNF2h) consists of at least 100 trajectories collated from at least 3 (typically 5 to 7) different experiments. All errors were bootstrapped over trajectories (that is, for each data set, the ≥100 trajectories were resampled with replacement, and reported values, such as the means of each pause duration, were recalculated for each bootstrapped sample). Reported errors are standard deviations of the bootstrapped values. A similar procedure was used to obtain the errors on the cdfs in Figure 3E and Figure 3-Supplement 3; the shaded regions represent +/- the standard deviation of the set of bootstrapped cdfs.
Figure 1 The acidic patch is an important epitope for remodeling post-binding.
A. Charge profile of the nucleosome (left) and magnification of the acidic patch region (right) (PDBID: 1KX5, charge profile generated using ABPS and UCSF Chimera (Pettersen et al., 2004)). Residues of the acidic patch mutated in this study are shown in red. B. Example of a Cy5-fluorescent scan of a gel-based remodeling assay with 0/60 WT and acidic patch mutant (APM) nucleosomes (1μM SNF2h, 20nM nucleosomes, saturating ATP). C. Quantification of the gel in B and fits to a single exponential decay. Inset is zoomed to show faster time points with WT nucleosomes. D. Gel remodeling rate as a function of enzyme concentration, plotted on a log(2) scale. APM nucleosomes are remodeled substantially slower even at saturating concentrations of enzyme. Data were fit to a cooperative binding model (WT nucleosomes: $K_{M}^{app} = 61$ nM, $k_{max} =2.5$ min$^{-1}$, $h=1.5$; APM nucleosomes: $K_{M}^{app} = 280$ nM, $k_{max} =.012$ min$^{-1}$, $h=1.2$). E. Remodeling is inhibited by competition for the acidic patch. WT nucleosomes were remodeled with sub-saturating SNF2h and in the presence of KSHV LANA peptide. Remodeling is inhibited by the peptide but not when the arginine anchor is mutated to alanine. F. Inhibition curve with the LANA peptide ($K_i=1.21$ µM). Error bars represent standard errors on the mean for 3 replicates, except for the no-peptide condition in F, which had 2 replicates. G. ATPase activity of WT SNF2h. Nucleosomes were in excess of SNF2h and at saturating concentrations. APM nucleosomes stimulated ATPase activity 4-fold weaker than WT.

Figure 1-Source Data 1 Values used to obtain plots

Figure 2 Dependence on the nucleosome acidic patch is linked to relief of autoinhibition
A. Domain architecture of SNF2h. The 2 critical arginines (R142, R144) of AutoN are highlighted in blue, while the NegC region replaced with a flexible GGS linker (Leonard and Narlikar, 2015) is highlighted in brown. Intervening sequences with no known domain annotations are numbered as in B. B. Direct domain-domain interactions of SNF2h nucleosomes probed by crosslinking mass spectrometry. SNF2h mononucleosomes were crosslinked with the zero-length reagent EDC in the presence of either the ATP transition state analog ADP•BeF$_x$ or ADP. Residue-residue crosslink data were aggregated into domain-domain level spectral counts. The fold change between ADP•BeF$_x$ and ADP conditions is displayed as the log2 of the spectral count ratio. The data were centered to the median value and the color scale depicts the interactions most enriched in ADP•BeF$_x$ in magenta, and ADP in blue. Light grey tiles indicate no crosslinks observed in either condition. Domains are listed from N- to C-term within each protein. The distribution of fold changes in domain-domain spectral counts is plotted as a
histogram below. It should also be noted that due to the structure of the histone octamer and the 2:1 stoichiometry of the SNF2h-Nucleosome complex, we cannot distinguish between intramolecular and intermolecular SNF2h-SNF2h or Histone-Histone crosslinks. C and D. Mutation of AutoN or NegC increases remodeling rate with WT and APM nucleosomes. Cy5 fluorescent scans of native gel remodeling assays. E. Saturating remodeling rates ($k_{\text{max}}$) for WT and APM nucleosomes with WT, 2RA, and mNegC SNF2h. Mutation of the acidic patch has a substantially lower effect on the $k_{\text{max}}$ for 2RA and mNegC SNF2h than for WT SNF2h (8-fold and 2-fold respectively vs. 200-fold). Error bars represent standard errors on the mean for N=3 replicates.

**Figure 2-Source Data 1 Values used to obtain plots**

**Figure 3. The acidic patch interacts antagonistically with AutoN to promote pause exit and persistent translocation**

A. Schematic of the smFRET setup. Nucleosomes labeled on histone H3 with a Cy3 donor dye and on one end of the DNA with a Cy5 acceptor dye are immobilized on the surface of a slide and imaged with a prism-based TIRF microscope. The end-positioned nucleosomes used here have an initial high FRET efficiency (see also Figure 3-Supplement 2). As remodeling proceeds and the nucleosome is moved away from the DNA end, the FRET efficiency is reduced. B. Example time-course of remodeling of a single surface-attached, WT nucleosome in the presence of saturating SNF2h and ATP (400 nM and 1 mM, respectively). Vertical yellow line indicates addition of enzyme and ATP; horizontal pink, green, and black lines are the output of an HMM fit used to quantify pause durations and locations (see Methods). Note the alternating “pause” and “translocation” phases of the remodeling reaction; in keeping with the ISWI literature, we call the first pause the “wait” pause, the second pause “p1”, the third “p2”, etc. Intensity and FRET data here and in C have been smoothed with a 0.95-second median filter for visualization only. C. Example time-courses of remodeling of, from top to bottom, WT nucleosomes by SNF2h, APM nucleosomes by SNF2h, WT nucleosomes by 2RA SNF2h, and APM nucleosomes by 2RA SNF2h. Saturating enzyme (400 nM enzyme with WT nucleosomes; 2 μM enzyme with APM nucleosomes) and ATP (1 mM) were used in all cases. Additional examples are shown in Figure 3-supplement 6. D. Quantification of the first three pause durations. *Indicates a lower limit; as shown in Figure 3-Supplement 2B, remodeling of APM nucleosomes by SNF2h is too slow to capture by smFRET, due to the competition between photobleaching of the dyes and remodeling by SNF2h. Errors were bootstrapped (see Methods). E. Empirical cumulative distribution functions (cdfs) of the change in nucleosome
position during the first translocation event (top panel) and the second translocation event (bottom panel) for different combinations of nucleosome constructs and enzymes. Roughly 50% of the initial translocation events by SNF2h on WT nucleosomes move the nucleosome 7 bp or fewer (black dashed lines); in contrast, nearly 80% of initial translocation events by SNF2h on E64R nucleosomes move the nucleosome 7 bp or fewer. Similarly, during the second translocation event, SNF2h moves WT nucleosomes roughly 5 bp or fewer 50% of the time, whereas again nearly 80% of second translocation events for SNF2h with E64R nucleosomes result in step sizes 5 bp or fewer. Shaded areas represent bootstrapped error estimates (see Methods). See also Figure 3-Supplement 6 for additional representations of these data and a further discussion of step sizes.

Figure 3-Source Data 1 Values used to obtain plots

Figure 4 The acidic patch is used by ACF and INO80
A. Gel remodeling assay with human ACF and 0/60 nucleosomes. Saturating concentrations of enzyme and ATP were used. B. Gel remodeling with the yeast INO80 complex and 0/60 nucleosomes. Reactions were performed with saturating enzyme and ATP. Error bars represent standard error of the mean for N=3 replicates.

Figure 4-Source Data 1 Values used to obtain plots

Figure 5 Model for nucleosome remodeling by SNF2h
After binding the nucleosome, SNF2h is in equilibrium between an active and autoinhibited state. In the autoinhibited state AutoN and NegC hold the remodeler in an inactive state. The active state is promoted by AutoN and NegC binding near the acidic patch and by H4 tail binding the ATPase domain resulting in conformational changes that bring HSS in close proximity to the acidic patch. From this active state, SNF2h can translocate DNA around the octamer. Although SNF2h remodels nucleosomes as a dimer at saturating enzyme concentrations (Blosser et al., 2009; Racki et al., 2009), in this model we display only one of the protomers for simplicity.

Figure 1-figure supplement 1. Sequences of DNA constructs used in this work
601 nucleosome positioning sequence is underlined; Pst18 restriction site (where present) is in blue. The sequence of the 3/78 construct is the same as the “n=3” construct in Blosser et al., and the 3/78 construct in Zhou et al. 2018 (Blosser et al., 2009; Zhou et al., 2018).

Figure 1 Supplement 1-Source Data 1 Values used to obtain plots
Figure 1 – figure supplement 2. Representative ATPase assay fits.

Saturating concentrations of nucleosomes (200nM WT nucleosomes, 187 nM APM nucleosomes) were incubated with 12.5 nM enzyme and 7.5 µM ATP•Mg, and trace amounts of γ-P\textsuperscript{32}ATP. Reactions were analyzed by thin layer chromatography followed by radiography. Initial rates were determined by a linear fit to the first 10% of ATP hydrolyzed in the reaction.

Figure 1 Supplement 2-Source Data 1 Values used to obtain plots

Figure 1 – figure supplement 3. Maximal remodeling rates for various acidic patch mutations.

0/60 Cy3-DNA labeled nucleosome constructs were remodeled with saturating concentrations of either WT or 2RA SNF2h (N=3). The mean and standard error are shown for the maximal remodeling rates. Residues in the “arginine anchor” binding site of the nucleosome (E61, D90, E92) produce the same defect when mutated individually or in combination, suggesting a cooperative role for these residues in SNF2h remodeling. Mutating residues in the acidic patch but outside the arginine anchor site (E64) has a weaker effect than mutating the arginine anchor residues. However, all acidic patch mutations have a weaker defect in the context of 2RA mutant SNF2h.

Figure 1 Supplement 3-Source Data 1 Values used to obtain plots

Figure 2 – figure supplement 1. Mutations to the N-terminus have minimal effects on remodeling

A and B. Left. Cy3-fluorescence scan of native gel remodeling assays with WT nucleosomes and either WT or mutant SNF2h. Saturating concentrations of SNF2h were used for all constructs. Right. Quantifications of the gel images shown at the left. The residues chosen are lysines in AutoN that crosslinked to the acidic patch in only the ADP•BeF\textsubscript{x} state. In all cases, the mutations showed no appreciable effect on remodeling (<2-fold difference).

Figure 2 Supplement 1-Source Data 1 Values used to obtain plots

Figure 2 - figure supplement 2. Ensemble FRET remodeling shows similar effects as the gel-based assay

A. FRET remodeling assay. Saturating concentrations of enzyme were added to 8 nM FRET-labeled nucleosomes. Cy5 fluorescence was monitored over time and fit to a two-phase exponential decay. Reactions were normalized to the initial (maximal) and plateau (minimal) fluorescence from the fit. B. Remodeling rates of the two phases. Error bars represent standard
error of the mean for N=3 replicates.

**Figure 2 Supplement 2-Source Data 1 Values used to obtain plots**

**Figure 2 – figure supplement 3. Saturation Kinetics Experiments**
Fits of the dependence of remodeling rates on enzyme concentration for each enzyme-substrate pair. Previous work indicates that SNF2h cooperatively remolds nucleosomes as a dimer, suggesting the data should fit to a hill coefficient greater than 1 (Racki et al., 2009). However, we cannot rule out that mutations may affect the cooperative association of SNF2h. To address this, reactions were fit with either a floating hill coefficient (top) or constrained to hill coefficient of 1 (middle) or 2 (bottom). Both $K_{\text{m,app}}$ and $k_{\text{max}}$ change less than 2-fold regardless of whether the data are fit to a fixed hill coefficient or fit to a floating coefficient.

**Figure 2 Supplement 3-Source Data 1 Values used to obtain plots**

**Figure 2 – figure supplement 4. Acidic Patch-SNF2h Crosslinks Unique to the ADP•BeF\(_x\) condition**
Residue level cross-links between SNF2h and the H2A/H2B acidic patch. Each dot represents a pair of amino acid residues that were identified in either the ADP-BeFx experiment or the ADP experiment (right two panels, black). The left panel displays only the cross-links uniquely identified in the presence of ADP-BeFx (magenta). The area of each dot is scaled to the number of spectral counts identified. Domain boundaries are indicated by bold grey lines and labeled for SNF2h. Only cross-links to the H2A and H2B acidic patches (defined as H2A:44-100 and H2B:100-122) are plotted here for clarity. Crosslinks between H2A acidic patch and SNF2h AutoN domain are only observed with ADP-BeFx. Likewise crosslinks between the H2B acidic patch and the SNF2h NegC domain are only observed with ADP-BeFx. Positions of the 2RA mutants (R142, R144) are marked by horizontal red lines.

**Figure 2 – figure supplement 5. Comparison of crosslinks in the ADP•BeFx state with and without LANA peptide**
A. Domain diagram of SNF2h B. Direct domain-domain interactions of SNF2h and Nucleosomes with ADP•BeFx in the presence or absence of the LANA peptide. The LANA peptide is expected to compete with domains on SNF2h that bind the acidic patch resulting in reduced crosslinking to this region of the nucleosome. Residue-residue crosslink data were aggregated into domain-domain level spectral counts and plotted as in Figure 2B. The fold change between ADP•BeF\(_x\) with and without LANA peptide is displayed as the log2 of the spectral count ratio with the data centered to the median fold change. The domain interactions most enriched in the presence of
the LANA peptide are colored in brown, while the interactions promoted in the absence of the peptide are colored in magenta. Light grey tiles indicate no crosslinks observed in either condition. Crosslinks between the H2A acidic patch and both the RecA lobes and the autoinhibitory domains are strongly reduced, while crosslinks to the HSS domain are less affected. The distribution of fold changes in domain-domain spectral counts is plotted as a histogram to the right.

**Figure 3 – figure supplement 1. smFRET controls**

A. Kernel density estimation plots (KDE, blue) and empirical cumulative density functions (cdfs, orange) of initial FRET values for 3/78 nucleosomes alone (in the presence of 1 mM ATP, but in the absence of remodeler; previous work with ISWI family remodelers has shown the addition of remodeler does not affect these initial FRET (Blosser et al., 2009; Deindl et al., 2013)). Left panel, wild-type H2A-containing nucleosomes; right panel, H2A/E64R-containing nucleosomes. N indicates the number of nucleosomes included in the cdf. KDEs are more intuitive—the y-axis is analogous to the frequency axis of a histogram—but cdfs have the advantage of not requiring any smoothing or binning. Note that here nucleosomes were imaged at a significantly higher laser power (20 mW) than for measuring remodeling (11.5 mW), to ensure that the majority of both donor and acceptor dyes photobleached within the 5 minute imaging interval, so that dyes with two-step photobleaching events could be excluded. Only nucleosomes with an initial FRET value greater than 0.775 (black dashed lines) were included for further analysis. KDEs have Gaussian kernels with bandwidths ~0.01. B. Our ability to measure remodeling of E64R nucleosomes by SNF2h is limited by the photobleaching rate. Previous work has shown remodeling in ensemble FRET assays to be comparable to remodeling in surface-immobilized smFRET assays (Blosser et al., 2009; Hwang et al., 2014), as well as to that of surface-attached dinucleosome constructs (Hwang et al., 2014). However, as shown here, remodeling of surface-attached E64R nucleosomes (red data) appears to go to completion in about 4 minutes (inset), whereas remodeling of these same nucleosomes in ensemble FRET assays (blue data) takes over two hours to go to completion. This discrepancy is not due to an effect of the surface immobilization, but rather to the competition between remodeling and photobleaching. In the presence of SNF2h alone, without ATP (black data), nucleosomes photobleach on roughly the same timescale as the apparent remodeling of surface-attached E64R nucleosomes. Thus we are able to put only a lower bound on pause durations for remodeling of E64R nucleosomes, because slower remodeling events are undetected due to the relatively fast photobleaching of the dyes. The smFRET data shown here corresponds to the
summed Cy5 intensities of all individual nucleosomes in the indicated data set, binned in 1-second intervals to simulate an ensemble measurement.

Figure 3-figure supplement 2. Additional example traces of SNF2h or SNF2h/2RA remodeling single nucleosomes, plotted as in Figure 3C.

Figure 3-figure supplement 3. The step size of WT SNF2h with WT nucleosomes is comparable to step sizes previously described for ISWI family remodelers.

A. Three different representations of the change in nucleosome position (that is, the step size) during the first (top) or second (bottom) translocation event observed by smFRET, for the ≥100 remodeling trajectories collected with WT nucleosomes and WT SNF2h. The three representations are: histograms in black; kernel density estimations (KDEs) with two different bandwidths (0.6, solid blue lines; 0.4, dashed blue lines) in blue; and cumulative distribution functions (cdfs) in orange. Vertical black dashed lines indicate the mean value, with the value of this mean step size ± S.E.M. given on the KDE/cdf plot. The histograms in black illustrate how the step sizes for SNF2h generally form a cluster, consistent with previous reports for other ISWI-family remodelers (Blosser et al., 2009; Deindl et al., 2013; Hwang et al., 2014). However, the other two representations of the same data reveal additional structure that is difficult to discern from the histograms alone. For example, the purple box over the top KDE indicates steps that likely represent nucleosomes that skipped the p1 pause, or whose p1 pause was too short to detect, such that the nucleosome was moved ~8 bp + 5 bp = 13 bp during the first translocation. For WT ISWI-family remodelers remodeling WT nucleosomes, the initial step has been reported to range from 6.9 ± 0.1 bp to 7.4 ± 0.1 bp, and the second step from 3.0±0.1 to 3.7±0.2 (Blosser et al., 2009; Deindl et al., 2013; Hwang et al., 2014). The difference of ~ 1 bp in the size of the second step between our work and that in the literature likely results from the different approaches used to obtain the mean step size. B. KDE and cdf plots as in A, but for the other three combinations of nucleosomes and SNF2h constructs shown in Figure 3. Due to their lack of binning or smoothing parameters, cdfs represent the best quantitative comparison between data sets. However, for assessing the step size within a single data set, KDEs can provide a more intuitive picture. Cdfs here are the same as in Figure 3E (though shown here without the error estimate).

Supplementary File 1 - Crosslinked residue pairs identified from EDC treatment of SNF2h-nucleosomes in the presence of: ADP, ADP-BeFx, or ADP-BeFx and LANA peptide.
References


Lan, L., Ui, A., Nakajima, S., Hatakeyama, K., Hoshi, M., Watanabe, R., Janicki, S.M., Ogiwara, H., Kohno, T., Kanno, S.-I., et al. (2010). The ACF1 complex is required for DNA double-strand...


within the INO80 Chromatin Remodeling Complex Implicates the Yeast Rvbs as Protein Assembly Chaperones. Cell Rep 19, 2033–2044.

A. Microscope setup: Totally internally reflected 532 nm laser.
- Cy3 donor on H3
- Cy5 acceptor on DNA

B. FRET intensity plots:
- Cy3 or Cy5 intensity (a.u.) vs. Time (sec)
- Waveform plots with markers p1 and p2

C. FRET duration plots:
- WT/SNF2h
- E64R/SNF2h
- WT/2RA
- E64R/2RA

D. Cumulative probability bars:
- WT/SNF2h, E64R/SNF2h, WT/2RA, E64R/2RA

E. Cumulative probability plots:
-wait→p1
-p1→p2
A C F

Acf1

Snf2h

ACF

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WT Nucleosomes

APM Nucleosomes

\[ k_{obs} \text{ (min}^{-1} \text{)} \]

Nucleosome: WT  AP M

10-Fold

B

INO80

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WT Nucleosomes

APM Nucleosomes

\[ k_{obs} \text{ (min}^{-1} \text{)} \]

Nucleosome: WT  AP M

200-Fold
0/60:
5′- CTGGAGAATCCCGGCTCAATGGTCTAGACAGCTCTAGCACCGCTCTAAACGCACGTACGCGCTGTCCCCCGCGTTTTAACCGCCAAGGGG... 
3′- GACCTGTCATTCGCCCAAGCTCCGGAGTTAACCAGCATCTGTCGAGATCGTGGCGAATTTGCGTGCATGCGCGACAGGGGGCGCAAAATTGGCGGTTCCC... 
  ...ATTACTCCCCATCTCCAGGCAAGTGCAGATATATACATCCTGTGCATCTATTGAACAGCGACCTTGCCGGTGCCAGTCGGATAGTGTTCCGAGCTCCC... 
  ...TAATGAGGGATCAGAGGTCGGCTCAGCTCTATATATGAGCACCGTAGATAACTTGTCGCTGGAACGGCCACGGTCAGCCTATCAGAAGGCTCC... 
  ...CTCCCACTCTAGAGGATCCTGGGTACGGGTACC-3′ 
  ...GAGGGTGAGATCTCCTAGGGGCCCATGG/5Bioteg/-5′

3/78:
5′- /5Cy5/GCCCTGGAGAATCCCGGCTCAATGGTCTAGACAGCTCTAGCACCGCTCTAAACGCACGTACGCGCTGTCCCCCGCGTTTTAACCG... 
3′- CCGGACCTGTCATTCGCCCAAGCTCCGGAGTTAACCAGCATCTGTCGAGATCGTGGCGAATTTGCGTGCATGCGCGACAGGGGGCGCAAAATTGGCGGTTCCC... 
  ...CCAAGGGGATTACTCCCCCTAGCTCCACGCGCTCTAGATATATACATCCTGTGCATCTATTGAACAGCGACCTTGCCGGTGCCAGTCGGATAGTGTTCCGAG... 
  ...GTTCCCCTAATGAGGGATCAGAGGTCGGCTCAGCTCTATATATGAGCACCGTAGATAACTTGTCGCTGGAACGGCCACGGTCAGCCTATCAGAAGGCTC... 
  ...CTCCCACTCTAGAGGATCCTGGGTACGGGTACC-3′ 
  ...GAGGGTGAGATCTCCTAGGGGCCCATGG/5Bioteg/-5′
Fraction ATP hydrolyzed

Time (min)

WT SNF2h Alone
WT SNF2h WT Nucleosomes
WT SNF2h APM Nucleosomes
Nucleosome: WT E61A E64A E64R D90A D90R/E92A APM

SNF2h: WT 2RA
**A**

[Graph showing time (min) on the x-axis and normalized cy5 Fluorescence on the y-axis. The graph includes four lines representing different nucleosome conditions: WT Nucs WT SNF2h, APM Nucs WT SNF2h, WT Nucs 2RA SNF2h, APM Nucs 2RA SNF2h.]

**B**

[Bar graph showing observed rate constants $k_{obs}$ for WT and APM nucleosomes with and without SNF2h. The graph includes two bars for each nucleosome condition: $k_{fast}$ and $k_{slow}$.]
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A

WT (N=329)

E64R (N=377)

B

Normalized Intensity

Normalized Intensity

Time (min)

Time (min)

E64R/SNF2h, ensemble
E64R/SNF2h, smFRET
WT/SNF2h, no ATP, smFRET

o E64R/SNF2h, ensemble
o E64R/SNF2h, smFRET
o WT/SNF2h, no ATP, smFRET
WT/SNF2h

wait → p1

Density

Cumulative probability

Change in nucleosome position (bp)

p1 → p2

E64R/SNF2h

wait → p1

Density

Cumulative probability

Change in nucleosome position (bp)

p1 → p2

WT/2RA

Density

Cumulative probability

Change in nucleosome position (bp)

E64R/2RA

Density

Cumulative probability

Change in nucleosome position (bp)

5.1±0.2 bp

5.6±0.3 bp

7.8±0.3 bp

4.4±0.5 bp

5.6±0.3 bp

7.5±0.3 bp

5.3±0.3 bp

8.7±0.3 bp

5.1±0.2 bp