

# ATP-dependent chromatin remodeling: genetics, genomics and mechanisms

Diana C Hargreaves<sup>1</sup>, Gerald R Crabtree<sup>1</sup>

<sup>1</sup>Howard Hughes Medical Institute, Beckman Center B211, 279 Campus Drive, Mailcode 5323, Stanford University School of Medicine, Stanford, CA 94305-5323, USA

**Macromolecular assemblies that regulate chromatin structure using the energy of ATP hydrolysis have critical roles in development, cancer, and stem cell biology. The ATPases of this family are encoded by 27 human genes and are usually associated with several other proteins that are stable, non-exchangeable subunits. One fundamental mechanism used by these complexes is thought to be the movement or exchange of nucleosomes to regulate transcription. However, recent genetic studies indicate that chromatin remodelers may also be involved in regulating other aspects of chromatin structure during many cellular processes. The SWI/SNF family in particular appears to have undergone a substantial change in subunit composition and mechanism coincident with the evolutionary advent of multicellularity and the appearance of linking histones. The differential usage of this greater diversity of mammalian BAF subunits is essential for the development of specific cell fates, including the progression from pluripotency to multipotency to committed neurons. Recent human genetic screens have revealed that BRG1, ARID1A, BAF155, and hSNF5 are frequently mutated in tumors, indicating that BAF complexes also play a critical role in the initiation or progression of cancer. The mechanistic bases underlying the genetic requirements for BAF and other chromatin remodelers in development and cancer are relatively unexplored and will be a focus of this review.**

**Keywords:** chromatin; genetics; biochemistry; BAF; SWI/SNF; development

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## Introduction

In eukaryotic cells, genetic information encoded in over 1 meter of DNA is packaged into chromatin and compartmentalized in the nucleus. The basic unit of chromatin is the nucleosome [1], which consists of 146 base-pairs of duplex DNA wrapped around a histone octamer comprised of two of each of the conventional histone proteins: H2A, H2B, H3 and H4. Highly related histone variants are also incorporated throughout the genome for regulatory purposes [2]. A fifth histone protein, H1, promotes higher order chromatin structures by encouraging compaction of neighboring nucleosomes from “beads on a string” to the 30 nm fiber. H1 is more abundant in heterochromatin than in euchromatin, leading to the greater compaction and more condensed appearance of heterochromatin. As the 30 nm fiber accounts for only ~25-fold

of a 5 000-fold DNA-to-nucleus compaction ratio, several other as yet unknown mechanisms must contribute to higher order compaction and nuclear organization.

Despite this intricate packaging, DNA must be accessible for critical cellular processes such as transcription, replication, recombination, and repair. DNA accessibility is facilitated by two classes of enzymes, ATP-dependent nucleosome remodelers and histone modifying enzymes. Histone modifying enzymes post-translationally modify the N-terminal tails of histone proteins to alter the structure of chromatin and provide binding sites for regulatory proteins. Many chromatin-associated proteins contain protein domains that bind these moieties such as the bromodomain that recognizes acetylated residues. Through direct interactions with histone tails, these proteins are targeted to specific sites on chromatin, such as transcriptionally active regions abundant in H3K4me3, or repressed regions marked with H3K27me3.

Chromatin remodeling complexes (CRCs) in contrast utilize the energy of ATP to disrupt nucleosome DNA contacts, move nucleosomes along DNA, and remove

Correspondence: Gerald R Crabtree  
E-mail: crabtree@stanford.edu

or exchange nucleosomes. They thus make DNA /chromatin available to proteins that need to access DNA or histones directly during cellular processes. The essential role of these enzymes is reflected in the fact that many of them are required for diverse but specific aspects of embryonic development including pluripotency, cardiac development, dendritic morphogenesis and self-renewal of neural stem cells. In the adult, deletion or mutation of these proteins often leads to apoptosis or tumorigenesis as a consequence of dysregulated cell cycle control. Here we review the four major families of CRCs, highlighting their specialized functions.

### Early genetics

The first chromatin remodeling proteins were identified in two independent screens for mutants affecting mating-type switching and growth on sucrose, and were named SWI or SNF for switching defective or sucrose nonfermenting [3, 4]. Along with several cytoplasmic signaling proteins, these screens returned what were subsequently found to be regulators of chromatin structure. The *swi2* and *snf2* mutations could be suppressed by mutations in *H2A* and *H2B*, and the mutants exhibit altered nuclease sensitivity at the *suc2* gene, indicating that SWI/SNF proteins alter chromatin structure to enable transcription [5].

The yeast proteins SWI1/ADR6, SWI2/SNF2, SWI3, SNF5, and SNF6 are co-associated in a large multi-subunit complex called SWI/SNF [6, 7]. Inactivation of any single subunit similarly affects transcription due to dissolution of the complex, although these proteins are not essential for yeast viability [4, 8, 9]. SWI/SNF proteins are required for transcription by sequence-specific transcription factors, including yeast GAL4 and the glucocorticoid receptor expressed in yeast [4, 10, 11]. In addition, LexA fusions of SNF2, SNF5, and SNF6 can activate transcription when bound to DNA [6, 10]. The SWI/SNF complex was thus proposed to be a general activator of transcription, working in coordination with sequence-specific transactivators and the histone acetylase GCN5 [12].

The SWI/SNF family is evolutionarily conserved, and homologous proteins were subsequently identified in flies, plants, and mammals. The *Drosophila* homologue of SWI2/SNF2, Brahma (BRM), was identified in a screen for genes that suppress the body segment defects caused by mutations in Polycomb [13]. BRM mutants cause homeotic transformations consistent with the reduced transcription of homeotic genes [14]. The demonstration of antagonism between BRM and Polycomb, a known chromatin regulator, provided complementary support in flies for the role of SWI/SNF as a regulator

of chromatin structure [15]. It was subsequently found that BRM localizes to a fraction of actively transcribed genes on polytene chromosomes and is required for Pol II localization to these loci [16, 17]. However, in contrast to the genetic studies in yeast, mutants for the fly homologue of SNF5 do not phenocopy the BRM mutant [18, 19].

### The ATPase homology

The yeast SWI2/SNF2 gene is homologous to a number of other ATP-binding helicases of the DEAD/H family [20]. The sequence similarity includes the catalytic ATPase domain and seven characteristic protein motifs [21]. Performing a homology search in the human genome against the human homologue of SWI2/SNF2, BRG1, reveals a high degree of homology with 26 other ATPase domain-containing proteins (Table 1). These include proteins involved in transcriptional regulation (BTAF1), DNA methylation/gene silencing (LSH), DNA repair (CSB, HARP, HIP116), DNA recombination (RAD54, RAD54b), chromosome stability (ATRX), proteins of unknown function (HuF2, HEL1), as well as other chromatin remodelers. Despite the similarity in their core enzymatic domains, these proteins are genetically nonredundant *in vivo* indicating that they have specialized functions. This nonredundancy is not simply due to exclusive expression patterns as coexpressed ATPases exhibit very different phenotypes upon deletion in the same cell type.

The chromatin remodelers can be further subdivided into four families: BRG1 and hBRM are subunits of the SWI/SNF family, hINO80, hDomino, and SRCAP are ATPases of the INO80/SWR1 family, hSNF2H and hSNF2L belong to the ISWI family, and CHD1-9 serve the CHD family. These ATPases share affinity for the nucleosome and display DNA- and nucleosome-dependent ATPase activity *in vitro* [90]. Most chromatin remodelers form large multi-subunit complexes, which most likely alters the activity of the core ATPase *in vivo*. The accessory subunits commonly contain interaction domains that may directly regulate the enzymatic activity of the complex, facilitate binding to transcription factors and other chromatin modifying enzymes, and target the complex to DNA and/or modified histones. Among others, Actin-related proteins (Arps) are frequently associated with chromatin remodelers and genetic studies in yeast and mice have demonstrated an unequivocal requirement for these proteins [91].  $\beta$ -actin is also an intrinsic subunit of the INO80, SWR1, and mammalian SWI/SNF or BAF complexes; however, its function has been difficult to discern because actin is essential for viability. It is most

**Table 1** ATPases of the DEAD/H helicase family

| ATPase      |  | Function  | References |
|-------------|--|---|------------|
| hBRG1       |  | Homozygous mutant embryos die during the periimplantation stage; Conditionally inactivated mutants display defects in ES cell pluripotency and T cell development; tumor suppressor, frequently lost or silenced in tumor cell lines and primary human tumors | [22-26]    |
| hBRM        |  | Homozygous mutant mice are slightly larger than normal; frequently lost or silenced in tumor cell lines and primary human tumors  | [26-28]    |
| hSNF2L      |  | Regulates homeotic gene transcription; an associated subunit, BPTF, is essential for early embryonic development  | [29-31]    |
| hSNF2H      |  | Homozygous mutant embryos die during the periimplantation stage; involved in chromatin assembly and replication through pericentric heterochromatin; mediates phosphorylation of H2A.X(Tyr142) following DNA damage; loads cohesin onto chromatin             | [32-37]    |
| CHD1        |  | Essential for ES cell pluripotency; regulates transcriptional elongation and pre-mRNA splicing  | [38, 39]   |
| CHD2        |  | Homozygous mutant mice exhibit growth delays late in embryogenesis and perinatal lethality; heterozygotes exhibit kidney abnormalities  | [40]       |
| CHD3/Mi2-α  |  | Autoantigen in dermatomyositis  | [41-43]    |
| CHD4/Mi-2β  |  | Autoantigen in dermatomyositis; regulates erythroid and T cell development via interactions with Ikaros; regulates B cell development via interactions with Bcl-6   | [41-46]    |
| CHD5        |  | Frequently deleted in human neuroblastomas  | [47, 48]   |
| CHD6        |  | Unknown   |            |
| CHD7        |  | Mutated in human CHARGE syndrome; Homozygous mutant mice exhibit developmental defects similar to those present in CHARGE syndrome patients; Interacts with hBRG1 and PBRM1 in human neural crest-like cells  | [49-51]    |
| CHD8        |  | Homozygous mutants die early in embryogenesis due to a failure in CHD8-mediated repression of p53-dependent apoptosis   | [52]       |
| CHD9        |  | Expressed in bone marrow stromal progenitor cells   | [53, 54]   |
| HELLS/LSH   |  | Homozygous mutant mice die shortly after birth; Lsh <sup>-/-</sup> cells exhibit defects in <i>de novo</i> DNA methylation by Dnmt3a and Dnmt3b   | [55-60]    |
| hHEL1       |  | Unknown   |            |
| hINO80      |  | Implicated in the DNA damage response, checkpoint pathways, and DNA replication   | [61]       |
| BTA1/Mot1   |  | Regulation of pre-initiation complex assembly   | [62]       |
| ERCC6/CSB   |  | Mutated in 80% of human Cockayne Syndrome cases; involved in transcription coupled repair, nucleotide excision repair, and base excision repair   | [63]       |
| RAD54B      |  | Homologous recombination  | [64-66]    |
| hRAD54      |  | Homologous recombination  | [67-69]    |
| ATRX        |  | Mutated in α-thalassaemia; binds to telomeres, subtelomeric regions, and GC-rich tandem repeats to facilitate H3.3 exchange   | [70-72]    |
| hDomino     |  | Homozygous mutant mice die in embryogenesis due to defects in hematopoiesis; catalyzes H2A.Z deposition; prevents cell cycle arrest and senescence; facilitates DNA repair  | [73-78]    |
| TTF2/HuF2   |  | Unknown   |            |
| hHARP       |  | Recovery from replication fork arrest during S phase  | [79-83]    |
| HLTF/HIP116 |  | Ubiquitinates PCNA to facilitate recovery of stalled replication forks; frequently silenced in colon and digestive tract cancers  | [84-86]    |
| SRCAP       |  | Catalyzes H2A.Z deposition  | [87, 88]   |
| RAD54L2     |  | Interacts with sumoylated transcription factors to regulate transcription of target genes   | [89]       |

likely the contribution of these additional subunits that lends functional specificity to highly similar ATPases.

### Mechanistic features of DNA-dependent ATPases

Until recently, the mechanisms of action of most of these complexes were thought to be relatively well understood. However, recent genomic and genetic data in mammals have raised questions about the mechanisms by which these complexes produce their biologic activities. Early studies done primarily on nucleosomal templates suggested that these complexes might mediate their biologic actions by simply moving or exchanging nucleosomes. Indeed for the SWR1 complex, genetic studies have nicely documented the importance of the nucleosome exchange reaction first observed *in vitro* [92]. However, many investigators including ourselves have failed to find substantial changes in nucleosome positioning after acute conditional mutation of proteins such as Brg1, despite rapid and extensive changes in the regulation of target genes. These studies have suggested that the biologic actions of CRCs may be achieved not only by nucleosome movement but also additional mechanisms. Genome-wide studies of Brg1 have shown that BAF complexes generally do not bind at promoters, which are less compact than other genomic regions. Hence, these complexes may primarily occupy nucleosomal assemblies that are more complex than a simple “beads on a string” template where nucleosomes are not free to slide on DNA as observed *in vitro*. In addition, despite the definitive requirement for complex subunits *in vivo*, they are not required for the remodeling activity of the ATPase *in vitro*, further cautioning that *in vitro* mechanistic studies may be only partially informative. Thus, there is an urgent need to develop new and better mechanistic approaches to study the complexes upon chromatin templates with all of the histone modifications and complexity of the actual genomic sites where they carry out their functions. With these caveats, we will review what is known of the mechanism underlying the biologic actions of ATP-dependent chromatin remodelers.

Structural studies have focused on understanding how the hydrolysis of ATP catalyzes the movement of DNA around nucleosomes. Although the electrostatic interactions governing the DNA-nucleosome association are energetically unfavorable to unwrapping, it has been shown that CRCs can disrupt these contacts in order to promote nucleosome sliding, DNA exposure, and nucleosome exchange on DNA templates *in vitro*. An early hypothesis supported a mechanism of twist diffusion, whereby DNA twists around the nucleosome to accommodate the gain of a base pair from the linker DNA. The twist is

propagated through the rest of the DNA:histone contacts, resulting in the advance of the nucleosome along DNA. However, this model was rejected on the basis that large impediments to DNA twisting, such as DNA hairpins or biotin crosslinks, produced no defect in nucleosome sliding [93, 94]. A more favored model is the ‘loop recapture’ model, which argues for the generation of a loop of DNA created by new histone contacts with neighboring linker DNA [94]. As in the twist diffusion model, this loop is propagated around the nucleosome to advance the nucleosome along DNA. Although energetically more costly, the loop recapture model explains how DNA might move around the nucleosome without changes to rotational phasing.

The initial creation of the DNA loop or bulge may be the result of the translocase activity that has been ascribed to the SWI/SNF, ISWI, and ACF ATPases [95-98]. In the case of SWI/SNF, Saha and colleagues postulate that the ATPase binds to a specific location on the nucleosome, from which it utilizes its 3' → 5' translocase activity to draw DNA from one entry/exit and pump it to the other in a directional wave [99]. DNA footprinting and crosslinking experiments have placed the ATPase at a site of weak DNA:histone contact, where torsional strain might be tolerated for the propagation of the loop [96, 99-101]. In addition, electron micrographic reconstructions of the larger RSC and SWI/SNF complexes indicate that these CRCs form multi-lobed C-shaped structures that cradle the nucleosome in a central cavity with its entry and exit points exposed [101-103]. Binding of the complex to the nucleosome creates significant rearrangement of the DNA with respect to the histone octamer even in the absence of ATP hydrolysis, which may facilitate the creation of a DNA bulge required for ATP-dependent translocation [104].

In contrast to SWI/SNF, the smaller ISWI CRCs make limited contacts with the nucleosome and the extranucleosomal DNA [100, 105-107]. These complexes bind their substrate as a dimeric motor to facilitate the bidirectional and processive translocation of DNA over the nucleosome [95, 108, 109]. This is consistent with the role of these remodelers in nucleosome spacing, and their ability to sample DNA linker lengths to position nucleosomes equidistant from either end [94, 107, 110, 111]. In this case, ATP hydrolysis by one of the two ATPases results in loosened DNA-histone contacts that may act similarly to the ATP-independent conformational change upon SWI/SNF binding to promote DNA translocation following ATP hydrolysis by the second ATPase [112].

Although the *in vitro* translocase activity of the SWI/SNF and ISWI ATPases is remarkably similar, their as-

sociated subunits have a profound impact on the resulting function of the remodeling complex. For example, subunit composition determines the stoichiometry of the complex to the nucleosome, 1:1 for SWI/SNF versus 2:1 for ISWI. In addition, the Dbp4 subunit of ISWI makes significant contacts with extranucleosomal DNA, thus rendering ISWI activity dependent on linker length [105]. These biochemical differences correspond to functional specialization *in vivo*. Namely, nucleosome sliding and perturbation by SWI/SNF promotes access to DNA, while nucleosome spacing by ISWI facilitates chromatin formation and gene silencing.

Structural studies may also shed light on template selection *in vivo*. For example, it has been determined by mutational analysis that the catalytic activity of ISWI is uniquely affected by a basic patch of residues (K<sub>16</sub>R<sub>17</sub>H<sub>18</sub>R<sub>19</sub>) on the H4 tail [113, 114]. Similarly, acetylation of H4K16 reduces remodeling by the ISWI remodeler ACF [115]. This suggests that ISWI is targeted away from transcriptionally active, H4K16 acetylated chromatin and areas where even spacing of nucleosomes occludes either the H4 tail or linker DNA. On a related note, the extensive contact between SWI/SNF and RSC and their nucleosome substrate would seem to preclude the binding of these complexes to condensed chromatin. This is in agreement with reports suggesting that H1 incorporation renders chromatin resistant to remodeling [116-118].

Bioinformatic analysis of chromatin remodelers suggests that the core ATPase domain is remarkably similar to that of the DEAD/H helicases, indicating that they most likely utilize ATP to facilitate movement of DNA by similar mechanisms. Indeed, these enzymes and the related RAD54 protein share an ability to translocate DNA similar to the DEAD/H helicases, despite no evidence for helicase activity of SWI/SNF proteins [119]. Despite these similarities, these proteins are genetically nonredundant, demonstrating functional specificity [120]. Extensive structural analyses of the SWI/SNF and ISWI complexes have demonstrated differences in the reaction product, substrate selection, enzymatic activity, and targeting of these complexes. However, as alluded to above, our understanding of the relative ability of CRCs to remodel chromatinized templates may be misguided due to the complexity of reproducing physiological templates *in vitro*. It is possible that nucleosome movement is only one aspect of CRC function and that studies utilizing native chromatin templates will reveal other activities.

### Chromatin regulators as macromolecular machines

Given the importance of associated subunits, it is

appropriate to consider what defines a macromolecular complex and why nature would choose to partner proteins in this manner. Perhaps the most well defined macromolecular complex is the ribosome, which we can use as a standard for what constitutes a subunit of a complex as opposed to an associated protein. By this criteria, a subunit can be defined as a protein that forms a stable, interlocking association with the complex which is resistant to exchange with free subunits and can only be disrupted by denaturation. Subunits exhibit biochemical dedication to the complex as assessed by glycerol sedimentation, which is often 50% or more of the total protein in case of CRCs. Certain subunits can be found in more than one complex. For example, virtually all BAF53b and BAF45 migrate with the ATPase Brg1 in mammalian BAF complexes. However, Polybromo migrates with Brg1 and with a large unidentified complex, yet Polybromo is a stable subunit of BAF complexes [121]. Lastly, genetic deletion of subunits often results in similar phenotypes. However, different phenotypes can emerge if a subunit is employed in another complex or is only required for a subset of complex functions.

Why would a DNA-dependent ATPase be driven evolutionarily to function with other subunits, rather than simply use the activities of these other proteins in solution? One answer seems to lie in the rapid coupling of reactions. The probability of an effective collision between two molecules is a 3<sup>rd</sup> order function of proximity. Hence complexes are probably assembled from active subunits during the course of evolution to provide proximity for coupled reactions. This means that if a chromatin regulatory complex carries out steps 1, 2, 3, 4, 5, 6 and 7 on its surface, then the genetic lack of a subunit might result in loss of reactions 5-7 while another mutation would result in the loss of reactions 1-7 thereby giving a different phenotype to two genes that are entirely dedicated, for example BRM and OSA in flies [18]. Unfortunately, we have very little understanding of the other coupled reactions that almost certainly are serially and stereospecifically organized on the surface of chromatin regulators. The development of small molecule inhibitors to parse the sequence of steps is likely to be invaluable to understanding the nature of these coupled mechanisms.

A second evolutionary force driving the formation of macromolecular complexes is the opportunity for diversification of function afforded through combinatorial assembly of the complexes. Combinatorial assembly seems to be a feature of mammalian complexes and arises from the fact that subunits are encoded by gene families. These family members provide mechanistic variation leading to the functional specialization of a specific complex assembly. As we will discuss later, this latter point is an

intriguing property of the mammalian BAF complex and probably most other chromatin remodeling complexes, which has dramatically broadened its role from yeast to mammals. Here again, small molecule inhibitors interfering with the function of specific subunits or emerging from unique interfaces between subunits could be vital to understanding how the subunits work as a complex. Of note, the development of small molecule inhibitors targeting the bromodomain of BRD4 was recently reported, which could be modified to target any of the several bromodomains present in subunits of the BAF complex [122, 123].

### Actin and actin related proteins in chromatin remodeling

The long debate over the existence of actin in the nucleus centered on the fact that actin filaments could never be detected in the nucleus despite concerted efforts. However, the discovery that monomeric actin is an intrinsic subunit of the BAF complexes resolved this argument and also explained why filaments were not visible [124].  $\beta$ -actin is present at what appears to be a 1:1 molar ratio with Brg1 and binds directly to the Brg1 ATPase [125]. Actin is not exchangeable *in vitro* and remains associated with BAF complexes even in 3 M urea. How monomeric actin might be contributing to chromatin remodeling by the BAF complex is unknown, but it is intriguing to speculate that the ATPase activity of  $\beta$ -actin is involved analogous to its regulation of myosin. In association with myosin, actin acts as an exchange factor to remove ADP from the active site of myosin, which initiates the myosin power stroke and thereby increases the rate of ATP hydrolysis by myosin [126]. Thus one possible role of actin in the BAF complex is to act as an exchange factor to promote a similar conformational twist of the Brg/Brm ATPase. Actin appears to be necessary for DNA-dependent ATPase activity of BRG1 as removal of actin from its binding site on BRG1 by treatment of BAF complexes with 5 M urea followed by renaturation leads to impairment of ATPase activity [124]. However, it is not clear that the BRG1 protein was fully renatured in these experiments. Actin was also found to be a component of the SWR1 complex and recent evidence indicates that it also is required for optimum ATPase activity of the SWR1 complex [92].

Arps are highly similar to actin, particularly over the ATP/ADP-binding pocket of the actin ATPase domain known as the 'actin fold'. However, unlike actin, Arps cannot polymerize and do not have ATPase activity, with the possible exception of yeast Arp4. Arps have recently been classified into 11 subfamilies with Arp4-Arp9 being

predominantly localized in the nucleus [127]. These proteins are dedicated, conserved subunits of the SWI/SNF and INO80 CRCs that associate directly with the ATPase via the conserved N-terminal HSA domain [128]. Arp7 and Arp9 appear to be specific to fungi and may be functionally similar to Arp4 and actin, which are present in the orthologous CRCs of higher eukaryotes [127]. Due to a unique insertion, Arp4 has the ability to bind histones, modified histones, and nucleosomes, which is critical to the function of the yeast NuA4, INO80, and SWR1 complexes in DNA repair [129]. In addition, Arp4 is the only known Arp with ATP-binding activity, which contributes to NuA4 function through Arp4 association and disassociation from the complex [130]. Arp6 is a unique subunit of SWR1, which regulates binding to H2AZ prior to H2AZ exchange by the SWR1 complex [131]. Finally, Arp5 and Arp8 have been shown to be essential subunits of  $\gamma$ INO80 as deletion of Arp5 or Arp8 phenocopies Ino80 deletion in yeast [132].

In the mammalian BAF complexes, the two Arp4 homologues, BAF53a and BAF53b, are used sequentially in the development of the nervous system. BAF53a is present in neural stem cells lining the ventricles and is rapidly replaced by BAF53b at cell cycle exit [121]. BAF53b is a dedicated subunit of the neuron-specific nBAF complex and its deletion leads to death shortly after birth due to a failure of dendritic morphogenesis in post-mitotic neurons [133]. Similarly, knockdown of the *Drosophila* homologue of BAF53a/b, BAP55, in class I dendrite arborization neurons during embryogenesis results in aberrantly oriented dendrites and reduced arborization while deletion of BAP55 leads to a highly specific mistargeting of olfactory projection neurons [134, 135]. Because BAF53a and b do not possess ATPase activity and are not required for the enzymatic activity of Brg *in vitro* [124, 133], they must contribute to complex function by other means, perhaps through complex targeting to histones or nucleosomes. Alternatively, they might function like Arp2/3 in generation of dimers that control actin dynamics [136].

### Chromatin remodeling complexes by family

#### *SWI/SNF family*

**SWI/SNF** Yeast SWI2/SNF2 is incorporated into a 1.14 MDa multi-subunit complex of 8-11 subunits. When we purified a family of related mammalian complexes in HeLa cells we called them mSWI/SNF because we found that four subunits were clearly homologous to ones present in the yeast SWI/SNF complex [137-139]. However, we and other laboratories have characterized the complexes from non-transformed cells and find that

the mammalian complexes are less similar to ySWI/SNF than we initially thought [120, 121, 124]. Recent studies have shown that many BAF subunits are bona fide tumor suppressors, which are generally inactivated in malignant cell lines and can be deleted with no apparent phenotype [140]. Hence, we and other investigators who took the same tack were most likely studying partially assembled complexes. Hence we have used BAF for Brg/Brm-associated factors for the mammalian complexes to discourage extrapolation, which has often led to incorrect conclusions. Because SWI, SNF, and BAF were not acceptable to the HUGO nomenclature committee, we suggested an alternate nomenclature of SMARC (SWI/SNF related, actin containing, regulators of chromatin). Unfortunately, this nomenclature has been extended by HUGO to proteins that do not have these characteristics. At present all of these nomenclatures are in use and Table 2 provides a translation for the reader.

Mammalian BAF and fly BAP (for Brahma-associated factors) complexes have lost, gained, and poached SWI/SNF subunits in response to the changes in epigenetic regulation that accompanied the evolution of multicellularity. The BAF complex is composed of five ySWI/SNF orthologues (BRG1/hBRM, BAF155/170, BAF60, BAF53a/b, and BAF47) and several additional unique subunits (BAF250a/BAF250b, BAF200, BAF45a/b/c/d, Brd9, and Brd7), including two subunits, BAF57 and actin, which are related to Nhp10 and actin found in the yeast INO80 and SWR1 complexes. Five yeast subunits have been discarded, resulting in a mammalian complex of about 2 MDa, which is larger than the calculated molecular weight of the known subunits, indicating that several subunits have yet to be identified. In addition to the amino terminal DEXDc and HELICc subdomains shared with other DEAD/H helicases, Brg1 and Brm have a C-terminal bromodomain. The bromodomain was first identified in BRM, the *Drosophila* homologue of ySWI2/SNF2, and binds acetylated residues on histone tails [14]. Additional domains present in the accessory subunits presumably facilitate interactions with proteins (LXXLL, BAH, SANT, SWIRM, SWIB), DNA (ARID, HMG, Zn finger, Leucine zipper), and modified histones (Bromodomain, Chromodomain, PHD domain), although this has yet to be confirmed genetically. The bromodomain of Brg1 is not required for its *in vivo* function, as a mutant lacking the bromodomain fully rescued the knockout phenotype in both flies and mice [90, 141]. However, additional bromodomains in Brd7 and Brd9 might function redundantly in recruiting the complexes to specific genetic loci. Finally, while ySWI/SNF lacks actin, the mammalian BAF complex has approximately one actin molecule per complex, which may enhance the ATPase

activity of BRG1/hBRM, as in the case of SWR1 [124].

Perhaps even more important than gain or loss of subunits in the evolution of SWI/SNF is the expansion of gene families that encode homologous BAF subunits. Whereas ySWI/SNF is monomorphic, BAF complexes are arrayed from several possible options in each of the following gene families: Brg1/Brm, BAF250a/b, BAF155/BAF170, BAF60a/b/c, BAF45a/b/c/d, BAF57, BAF53a/b, BAF47, and actin. Genetic studies indicate that subunit exchange helps drive the transition from pluripotency to multipotency to the committed post-mitotic neuron. Pluripotent embryonic stem (ES) cells express BAF155 but not BAF170, Brg1 but not Brm, and BAF53a but not BAF53b [23, 24, 142, 143]. Consistent with a requirement for this particular assembly for pluripotency, deletion of Brg1 results in early embryonic lethality during implantation [22], while Brm-deficient mice are viable, though a little larger than normal [28]. Differentiation of ES cells into neural progenitors leads to the activation of Brm and BAF170 and repression of BAF60b [121]. The final step to post-mitotic neurons is accompanied by the repression of BAF53a and the activation of BAF53b, BAF45b and BAF45c, which assemble in nBAF complexes, apparently unique to the nervous system [121]. The mechanism underlying the switch in subunits during the development of the vertebrate nervous system appears to be a triple negative genetic circuit in which REST represses miR-9/9\* and miR-124, which in turn repress BAF53a leading to cell cycle exit, the activation of the alternative BAF53b subunit, and neural differentiation [144]. This npBAF to nBAF switch is essential for the development of the vertebrate nervous system, since mutations affecting either state are lethal. Another example of selective assembly occurs in the developing heart, where BAF60c designates the region of the embryo with cardiogenic potential [145]. Complexes containing BAF60c are uniquely required for heart development, and can directly facilitate the formation of heart tissue from mesoderm in the presence of tissue-specific factors [145, 146]. In each of these cases, the expression of homologous subunits could not compensate for the loss of the correct subunit, arguing for the exquisite specificity of each composite in instructing cell fate decisions.

The unique composition of BAF at each developmental stage correlates with a specific gene expression program that is required for maintaining cell state, although the mechanistic basis for this maintenance is unknown. In ES cells, Brg1 binding was mapped by chromatin immunoprecipitation in combination with high resolution sequencing [147]. Brg1 binds to about 6 000 sites in large islands of several hundred bases. About 300 000

**Table 2** The SWI/SNF family

| Subfamily       |                                      | SWI/SNF                                   |  |                    |   |   |  |
|-----------------|--------------------------------------|---|--|--------------------|---|---|--|
| Species         | Yeast                                | Fly                                       | Human  | Mouse              | Mouse   | Mouse   |  |
| Complex         | SWI/SNF                              | BAP                                       | BAF  | esBAF              | npBAF   | nBAF  |  |
|                 | Swi2/Snf2                            | BRM                                       | BRG1/SMARCA4 or hBRM/SMARCA2                       | BRG1/SMARCA4       | BRG1/SMARCA4                                      | BRG1/SMARCA4                                      |  |
|                 |                                      | OSA/eyelid or BAP170                      | BAF250a/ARID1A or BAF250b/ARID1B or BAF200/ARID2*  | BAF250a/ARID1A     | BAF250a/ARID1A or BAF250b/ARID1B or BAF200/ARID2* | BAF250a/ARID1A or BAF250b/ARID1B or BAF200/ARID2* |  |
|                 | Rsc1, Rsc2, Rsc4                     | Polybromo                                 | BAF180*  |                    |   |   |  |
|                 | Rsc8                                 | MOR/BAP155                                | BAF155/SMARCC1 and/or BAF170/SMARCC2               | BAF155/SMARCC1     | BAF155/SMARCC1 and/or BAF170/SMARCC2              | BAF155/SMARCC1 and/or BAF170/SMARCC2              |  |
|                 | Rsc6                                 | BAP60                                     | BAF60a/SMARCD1 or BAF60b/SMARCD2 or BAF60c/SMARCD3 | BAF60a/SMARCD1     | BAF60a/SMARCD1                                    | BAF60a/SMARCD1                                    |  |
|                 | Arp7, Arp9                           | BAF55 or BAP47                            | BAF53a or BAF53b                                   | BAF53a             | BAF53a  | BAF53b  |  |
|                 | Sfh1                                 | SNR1/BAP45                                | BAF47/hSNF5/SMARCB1                                | BAF47/SNF5/SMARCB1 | BAF47/SNF5/SMARCB1                                | BAF47/SNF5/SMARCB1                                |  |
|                 |                                      | SAYP                                      | BAF45a or BAF45b or BAF45c or BAF45d               | BAF45d             | BAF45a or BAF45d                                  | BAF45b or BAF45c or BAF45d                        |  |
|                 |                                      | BAP111/dalao                              | BAF57/SMARCE1                                      | BAF57/SMARCE1      | BAF57/SMARCE1                                     | BAF57/SMARCE1                                     |  |
|                 |                                      | $\beta$ -actin                            | $\beta$ -actin                                     | $\beta$ -actin     | $\beta$ -actin                                    | $\beta$ -actin                                    |  |
| Unique Subunits | Swi1/Adr6, Swp82, Taf14, Snf6, Snf11 | Rsc3, 5, 7, 9, 10, 30, Htl1, Lbd7, Rtt102 | Brd7*, Brd9  |                    |   |   |  |

\*ARID2, BAF180, and Brd7 are proposed subunits of the PBAF complex. Despite coassociation with BRG1, these proteins are not dedicated subunits of the BAF complex as assessed by glycerol gradient centrifugation ([121], Hargreaves DC and Crabtree GR, unpublished observations).

molecules are present on average in each ES cell, indicating that the peaks are likely the result of the binding of several complexes, rather than an individual complex. In contrast to the expectation that Brg1 would function primarily at promoters, only 12% of peaks were within 500 bp of the transcription initiation site [147]. Brg1 occupancy is more highly correlated with H3K4me1, a modification enriched at enhancers and regulatory elements, than H3K4me3, a mark of active transcription [147, 148]. Consistent with this, *pou5f1* and *nanog* show peaks of Brg1 binding at each of the mapped enhancers critical to the positive feedback loop of pluripotency [149, 150]. Rapid deletion of Brg1 in ES cells followed by analysis of RNA by transcript arrays led to the unexpected discovery that Brg1 represses most of its direct targets. As Brg1 binding is non-overlapping with Suz12, a Polycomb group protein, BAF must mediate repression via a distinct mechanism(s). Interestingly, Brg1-dependent repression occurs in functional coordination with pluripotency genes such as Oct4 and Sox2, to which it binds in solution and on chromatin. Genome-wide, Brg1 binding sites in ES cells physically overlap with the binding sites of Oct4, Sox2, Nanog, Smad1, and STAT3, all critical regulators of pluripotency, at far higher levels than predicted [147]. Proteomic studies in ES cells also revealed that the esBAF complex binds to many of the pluripotency factors suggesting that the specialized subunit composition of esBAF complexes is tailored to the needs of a pluripotent cell.

The possibility that BAF complexes mediate repression at a distance originated from studies of the role of Brg1 and BAF complexes in lymphocytes indicating that these complexes act on a distal silencer to repress transcription of the *Cd4* gene [151]. Deletion of *Brg1* or the silencer gives similar phenotypes of premature derepression of CD4 expression [151]. These studies and the fact that Brg1 most commonly binds to intragenic regions suggest that Brg1 acts from a distance to regulate transcription, perhaps by creating long-range interactions. Indeed, Brg1-dependent looping has been observed at both the  $\alpha$ - and  $\beta$ -globin loci and is required for developmentally regulated transcription [152, 153]. Thus, long-range interactions from distal regulatory sites, and not nucleosome sliding, may more accurately account for BAF's observed phenotypes. In the example of the CD4 silencer in T cells, BAF is thought to prevent accumulation of H1 linker histones, allowing the corepressor Runx1 to bind and repress CD4 [154].

Finally, the antagonism between BRM and Polycomb first observed in *Drosophila* also warrants revisiting. Brahma was identified in a screen for mutants that suppress the body segment phenotypes resulting from

mutations in Polycomb. *In vitro*, Polycomb can inhibit remodeling by Brm. However, as Brahma is required for homeotic gene transcription and, by some accounts, transcription of all *Drosophila* genes, it is unclear how direct this antagonism is. In mouse ES cells, Brg1 binding was anti-correlated with Polycomb and H3K27me3, indicating that co-occupancy of Brg1 and Polycomb is actively avoided [147]. Recently, a more direct antagonism was demonstrated at the *p16<sup>ink4a</sup>/p19<sup>Arf</sup>* locus where expression of BAF47 in a BAF47-deficient tumor cell line resulted in Polycomb eviction from the locus, re-expression of the gene, and growth arrest [155]. It has subsequently been shown that Polycomb represses many genes upon loss of BAF47 in tumorigenesis and that the double mutant with Ezh2, a subunit of the Polycomb Repressive Complex 2 (PRC2), expresses wild-type levels of these genes and survives tumor-free [156]. Interestingly, many of the genes affected by this antagonism are part of a stem cell-associated gene set, suggesting that this antagonism may also be critical during normal development.

As mammalian genetic studies come to light, several longstanding tenants concerning the biologic role of SWI/SNF are being challenged. Clearly, extrapolation of findings from the ySWI/SNF complex to the mammalian BAF complex is risky and direct genetic and biochemical analyses of the mammalian complex have led to surprising conclusions. Yeast SWI/SNF binds almost exclusively to promoters and activates its direct targets concomitant with nucleosome displacement. In contrast, BAF complexes are most often found in intergenic regions where they both activate and repress genes and the role of nucleosome movement seems less clear. Furthermore, BAF complexes are polymorphic and hence possess greater functional diversity through combinatorial assembly. By exchanging subunits, BAF can reinterpret the genome, in effect increasing the total number of potential BAF targets in an organism. To date, this mechanism has only been observed in vertebrates, possibly as a means of increasing effective gene number or patterns of gene regulation during development. Although a similar exchange of subunits has not been described for other CRCs, it seems likely that they also take advantage of the diversification of function afforded by combinatorial assembly.

**RSC** A complex related to ySWI/SNF was identified called RSC for Remodel the Structure of Chromatin based on shared homology between the yeast ATPases STH1 and SWI2/SNF2 [157]. RSC shares two identical subunits with ySWI/SNF, Arp7 and Arp9, and several homologous subunits [157] (Table 2). There is also evidence of an additional biochemically distinct SWI/

SNF-related complex in *Drosophila* and mammals that nevertheless shares many subunits with the BAF and BAP complexes. These are the PBAP and PBAF complexes, named after their signature subunit, Polybromo (BAF180), a large protein containing several bromodomains, two BAH protein interaction domains, an HMG domain, and a C2H2 zinc finger domain. BAF180 is structurally related to the yeast Rsc1, Rsc2, and Rsc4 proteins, which are unique to the RSC complex, and thus, PBAP and PBAF are presumed to be evolutionary counterparts of RSC [158]. BAF200 is also present only in PBAF, while hBRM and BAF250 are unique to BAF [159]. It is worth noting that similarly to the case with ySWI/SNF and BAF, there are significant biochemical differences between RSC and PBAF that suggest these complexes may not be functional orthologues.

Genetic studies in yeast indicate that ySWI/SNF and RSC are nonredundant and functionally specialized. Both complexes appear to regulate gene expression, but target non-overlapping sets of genes [8, 9, 160]. STH1, SFH1, RSC3, and RSC9, all components of the RSC complex, are required for cell cycle progression through G2/M, while ySWI/SNF is not essential for viability [161-163]. Related to this, RSC has been implicated in sister chromatid cohesion, chromosome segregation, and ploidy maintenance [164-167]. *In vitro*, PBAF was implicated in ligand-dependent transactivation by nuclear hormone receptors [168]. Consistent with this, deletion of BAF180 in mice resulted in severe hypoplastic ventricle development and trophoblast placental defects due to reduced retinoic acid-induced gene expression [169]. However, flies deficient in Polybromo and/or BAP170 have no defects in ecdysone-induced transcription suggesting that the role of PBAF in nuclear receptor-dependent activation is not evolutionarily conserved [170]. Whether PBAP or PBAF acts more broadly to influence cell cycle is not known, although there is evidence that BAF180 localizes to kinetochores during mitosis [158]. Of note, BAF180 predominantly co-migrates on glycerol gradients with a very large 4 MDa complex distinct from the other BAF subunits, which could account for some of its activities, for example at the kinetochore [121].

### *INO80 family*

Continuing in the theme of unexpected roles for chromatin remodelers, we next discuss the INO80 family. The INO80 family includes the yeast INO80 complex and its orthologues Pho-dINO80 (*Drosophila*) and INO80 (human), the yeast SWR1 complex and its orthologue SRCAP (human), and the yeast NuA4 complex and its orthologues Tip60 (*Drosophila*) and TRAAP/Tip60 (humans) (Table 3). The core ATPase subunits of the INO80

family are characterized by a split ATPase domain. This unique module retains ATPase activity, and acts as a scaffold for the association of two RuvB-like proteins, Rvb1 and Rvb2. RuvB is a bacterial ATP-dependent helicase that forms a double hexamer around Holliday junctions to promote their migration during homologous recombination [171]. Rvb1 and Rvb2 are present at 6:1 stoichiometry, suggesting that they form hexamer structures similar to the bacterial RuvB protein. Unlike CRCs of other families, the INO80 complex exhibits DNA helicase activity *in vitro* and binds specialized DNA structures *in vitro* that resemble Holliday junctions and replication forks [172, 173]. However, it should be noted that the INO80 family does have affinity for nucleosomes, in particular the H2A variants H2AZ and H2AX. INO80 and SWR1 can slide nucleosomes *in vitro* on a reconstituted chromatin template and even evict histones from DNA [132, 174, 175].

The INO80 complexes contain  $\beta$ -actin and several Arps, which are shared with the mammalian BAF and fly BAP complexes. The yINO80 complex contains Arp4, Arp5, Arp8, and actin, of which Arp5, Arp8, and actin are conserved in flies and mammals. Arp4 and actin are also present in the yeast SWR1 and NuA4 complexes. The Arp4 mammalian homologue, BAF53a, is present in the INO80, SRCAP, and Tip60 complexes of the INO80 family, and as mentioned previously, the BAF and PBAF complexes. In flies, the SWR1 complex contains BAP55, an Arp that is also present in the BAP and PBAP complexes. This conservation argues for a critical actin- or Arp4-dependent role shared by INO80 and BAF complexes, possibly in targeting these complexes to structural elements. The fact that ySWI/SNF has neither Arp4 nor actin implies that the BAP and BAF complexes may have acquired mechanistic features of the INO80 family that are not operational in ySWI/SNF.

***INO80 and SWR1*** Yeast Ino80, the Swi2/Snf2-related ATPase of the INO80 complex, was identified in a screen for regulators of phospholipid biosynthesis [176]. It was subsequently found that yIno80 associates with 14 other proteins to form a complex of 1.2 MDa [172, 176]. As outlined above, Rvb1, Rvb2, Arp4, Arp5, Arp8, and actin are among these subunits, as well as Taf14, Ies1-6, and Nhp10. As evidenced by its role in phospholipid biosynthesis, yINO80 regulates gene expression downstream of several promoters, both positively and negatively [177].

Swr1 was identified based on its homology to Ino80 and was subsequently found to catalyze the exchange of H2A for Htz1 [92, 178]. Htz1 and H2B copurify with Swr1, and genome-wide transcription profiles reveal a ~40% overlap in genes regulated by Htz1 and Swr1 [92].

**Table 3** The INO80 family

| Subfamily       | INO80                  |                      |   |                | SWR1             |
|-----------------|------------------------|----------------------|---|----------------|------------------|
| Species         | Yeast                  | Fly                  | Human                                   | Human          | Yeast            |
| Complex         | INO80                  | Pho-dINO80           | INO80                                   | SRCAP          | SWR1             |
|                 | Ino80                  | dIno80               | hIno80                                  | SRCAP          | Swr1             |
|                 | Rvb1, Rvb2             | Reptin, Pontin       | Tip49a, Tip49b                          | Tip49a, Tip49b | Rvb1, Rvb2       |
|                 | Arp4, Arp5, Arp8, Act1 | dArp5, dArp8, dActin | Baf53a, Arp5, Arp8                      | Baf53a, Arp6   | Arp4, Arp6, Act1 |
|                 | Taf14                  |                      |   | GAS41          | Yaf9             |
|                 | Ies2                   |                      | hIes2                                   |                |                  |
|                 | Ies6                   |                      | hIes6                                   |                |                  |
|                 |                        |                      |   | DMAP1          | Swc4/Eaf2        |
|                 |                        |                      |   | YL-1           | Swc2/Vps72       |
|                 |                        |                      |   | dBrd8          | Bdf1             |
|                 |                        |                      | H2AZ, H2B                               | H2AZ, H2B      |                  |
|                 |                        |                      | Znf-HIT1                                | Swc6/Vps71     |                  |
|                 |                        |                      | dTral                                   | Tral           |                  |
|                 |                        |                      | dTIP60                                  | Tip60          |                  |
|                 |                        |                      | dMRG15                                  | MRG15, MRGX    |                  |
|                 |                        |                      | dEaf6                                   | FLJ11730       |                  |
|                 |                        |                      | dMRG15                                  | MRG15          |                  |
|                 |                        |                      | E(pc)                                   | EPC1, EPC-like |                  |
|                 |                        |                      | dING3                                   | ING3           |                  |
| Unique Subunits | Ies1, Ies3-5, Nhp10    | Pho                  | Amida, NFRKB, MCRS1, FLJ90652, FLJ20309 |                | Swc3,5,7         |
|                 |                        |                      |   |                | Eaf5, Eaf1/Vid21 |

Rvb1, Rvb2, Arp4, and actin are shared subunits with INO80. The remaining subunits are Arp6, Yaf9, Swc2-7, and Bdf1, Swc2 being a potential histone chaperone-like subunit based on its affinity for Htz1 [179]. The role of Bdf1, a bromodomain-containing protein, is not known, but this may suggest that H2AZ exchange is regulated by Bdf1-dependent binding to acetylated residues on histone tails. The role of SWR1 in H2A variant exchange is conserved throughout evolution: the *Drosophila* Tip60 complex catalyzes the exchange of H2A for H2Av (a variant with similarities to both H2AZ and H2AX) [180] and mammalian SRCAP complex catalyzes the exchange of H2A for H2AZ [87, 88].

Yeast *ino80* and *swr1* mutants are hypersensitive to DNA damage-inducing agents, but the transcriptional response induced by damage is unaffected [181, 182]. This implies a direct role for these complexes in DNA repair, perhaps at the site of damage. Upon DNA damage, H2AX is phosphorylated at double-stranded breaks (DSBs) by the Mec1/Tel1 kinases (ATM/ATR in mammals), leading to the recruitment of several DNA repair proteins. The lesion can then be repaired by homologous recombination (HR) or non-homologous end joining (NHEJ) depending on which downstream pathway is initiated. *ino80* mutants have defects in both the HR and NHEJ pathways, while *swr1* mutants have defects in NHEJ alone [175]. Both the INO80 and SWR1 complexes are recruited to DSBs via direct binding of complex subunits to phosphorylated H2AX, or  $\gamma$ -H2AX. Arp4, a shared subunit of INO80 and SWR1, mediates SWR1 recruitment to DSBs, while Nph10 (and Ies3) is required for INO80 recruitment, perhaps in association with Arp4 [129, 182]. Based on its ability to exchange histones, it was initially proposed that SWR1 is recruited to DSBs to exchange  $\gamma$ -H2AX for an unmodified histone. However, it is INO80 that facilitates nucleosome eviction at DSBs, allowing the recruitment of repair factors. Specifically, *ino80* and *arp8* mutants are defective in histone eviction, which leads to impaired recruitment of Mec1 and Rad51 and defects in ssDNA resection by the nuclease Mre11, both prerequisites for repair by HR [174]. INO80 and an associated protein YY1 also promote HR repair in human cells, indicating that the role of INO80 in repair is conserved [173]. Although SWR1 does not appear to exchange histones at the site of damage, it is required for efficient recruitment of Mec1 and Ku80 to DNA ends during NHEJ [175].

Following the initiation of repair at DSBs, checkpoint pathways are activated downstream of Mec1/Tel1 to coordinate the cell cycle with DNA repair. Interestingly, *ino80*, *ies4*, *arp5* and *arp8* mutants have defects in checkpoint recovery and reduced viability in the face of repli-

cative stress. It was shown that Mec1/Tel1 phosphorylate Ies4 of the INO80 complex, which acts redundantly with checkpoint factor Tof1 to regulate the cell cycle following release from replicative stress [183]. In parallel, INO80 was shown to promote recovery of stalled replication forks during the S-phase checkpoint response [184]. Specifically, INO80 binds replication forks during S phase and recruits Rad18 and Rad51 to stalled replication forks in order that they might be processed [184, 185]. In contrast, SWR1 is not present at stalled replicative forks and the corresponding mutant does not exhibit decreased viability in response to replicative stress [186, 187].

Finally, INO80 and SWR1 have been identified in genetic screens for proteins that regulate telomere length [187, 188]. *Ino80*, *Nph10*, *Ies3*, unique subunits of the INO80 complex, and Arp4, a common subunit of both INO80 and SWR1, localize to telomeres. Mechanistically, it is possible that INO80 is recruited to telomeres because they share several features with DSBs, including the incorporation of  $\gamma$ -H2AX and the recruitment of Mec1/Tel1. These features appear to regulate telomere length during normal replication and intensify during telomere shortening to “repair” the DNA end. *Ies3*, which is not involved in DSB repair, may mediate the specialized function of the INO80 complex at telomeres following  $\gamma$ -H2AX-dependent recruitment. Indeed, *Ies3* associates directly with the telomerase complex and deletion of *Ies3* specifically results in impaired growth and telomeric instability [188].

**NuA4/Tip60** As mentioned above, the *Drosophila* Tip60 complex catalyzes the exchange of H2A for H2Av, similar to the yeast SWR1 and human SRCAP complexes [180]. However, the dTip60 complex harbors an additional seven subunits, including the eponymous subunit Tip60, a histone acetyltransferase (HAT). In fact, based on a number of shared subunits, the Tip60 complex seems to be the evolutionary product of the combined SWR1 and NuA4 yeast complexes. The related Tip60/TRRAP complex in humans also has both chromatin remodeling and HAT activity, the key enzymatic subunits being the Swr1-related ATPase, p400, and Tip60 [189-192]. Intriguingly, Arp4 and/or actin are common to the entire INO80 family in all organisms.

Tip60 has been shown to regulate the DNA damage response through the acetylation of histones and other regulatory proteins. In all organisms, Tip60 is recruited to DSBs, possibly through an Arp4-dependent mechanism similar to that shown for SWR1. In yeast, NuA4 acetylates H4 at breaks following recruitment by Arp4 [129]. *Drosophila* Tip60 acetylates phosphorylated

H2Av at DSBs and facilitates its exchange with unmodified H2Av [180]. Likewise, the acetylation of  $\gamma$ -H2AX by human TIP60 after ionizing radiation is required for its subsequent ubiquitination and removal [193]. Tip60 also acetylates ATM upon recruitment to DSBs, which activates this kinase to initiate the repair process. Interestingly, Tip60 enzymatic activity is dependent on an interaction between the Tip60 chromodomain and H3K9me3 at DSBs and disruption of this interaction or depletion of H3K9me3 results in increased sensitivity to ionizing radiation and genomic instability [194]. In an unrelated fashion, Tip60 was found to acetylate lysine 120 on p53 upon increasing levels of DNA damage [195, 196]. p53 acetylation by Tip60 leads to p53's selective activation of pro-apoptotic genes, shifting the balance from cell cycle arrest to apoptosis. Finally, Tip60 appears to play a regulatory role in oncogene-induced DNA damage response as haplo-insufficiency of Tip60 accelerates Myc-induced tumor progression [197].

In addition to its role in the DNA damage response, Tip60 appears to regulate several other pathways critical for cell viability. Along with Brg1, Tip60 was identified in a screen for factors affecting ES cell self-renewal and morphology [198]. Using promoter ChIP-on-chip analysis, Fazio and colleagues found that the p400-Tip60 complex is bound to about 55% of all promoter regions, including active and silent genes, where it acetylates H4. Knockdown of Tip60 resulted in the upregulation of many developmental genes coregulated by Nanog and dysregulation of the ES cell cycle. Perhaps as a result, Tip60 ablation causes embryonic lethality in mice and flies [199, 200].

In summary, the INO80 family of CRCs function in a diverse array of cellular processes, including DNA repair, cell cycle checkpoint, and telomeric stability. Some intriguing features of these complexes include the special affinity for the histone variants H2AZ and H2AX and the recruitment of these complexes to specialized DNA structures, such as DSBs, Holliday junctions, telomeric ends, and replication forks. The unique split ATPase domain of Ino80 and Swr1 and their association with the RuvB-like proteins, Rvb1 and Rvb2, may account for the specialized function of these complexes. Of note, RAD54, a related ATPase, can bind Holliday junction-like structures and promote bidirectional branch migration in an ATP-dependent manner [201]. Alternatively, their recruitment may reflect the incorporation of H2A variants at critical sites. It is clear that the associated Arps are absolutely required for the function of these complexes, which may reflect a critical role for Arps and actin in targeting and/or regulation of these complexes.

### ISWI family

The CRCs of the imitation SWI (ISWI) family were first identified in *in vitro* assays for nucleosome remodeling activities in *Drosophila* embryo extracts [202]. These complexes are characterized by the DEAD/H-related ATPase ISWI, which in addition to the conserved ATPase domain contains a SANT and a SLIDE domain. ISWI is incorporated into several distinct complexes in *Drosophila*, including the ACF, CHRAC, and NURF complexes (Table 4). ACF and CHRAC are highly similar as they both contain the large Acf1 subunit in addition to ISWI [203, 204]. Acf1 has a bromodomain, 2 PHD fingers, and a putative heterochromatin targeting domain. NURF is characterized by the NURF301 subunit, which is structurally related to Acf1 apart from four LXXLL motifs, which allow interactions with nuclear hormone receptors. Homologous complexes have been identified in yeast and humans. The related human ATPases hSNF2H and hSNF2L are accompanied by hACF and BPTF in ACF/CHRAC and NURF complexes, respectively (Table 4).

Initial identification of these complexes indicated a role in transcriptional activation as they promote transcription from *in vitro* templates [202, 205, 206]. *In vivo*, NURF301-deficient flies and BPTF morpholino frogs exhibit developmental defects consistent with reduced expression of homeotic genes [207, 208]. NURF301 is required for transcription of the GAGA gene targets *ultrabithorax*, *engrailed*, *hsp70*, and *hsp26*, and the *wingless* target *senseless* through direct association with cognate transcription factors [205, 208]. BPTF is similarly required for homeotic gene transcription in *Xenopus* via an interaction between the PHD domain of BPTF and H3K4me3 at HOX genes [31]. In mice, deletion of SNF2h or BPTF is embryonic lethal. ES cell lines could not be derived from SNF2h-deficient mice due to the requirement for SNF2h in survival of the inner cell mass *ex vivo* [32]. BPTF is required at a later stage in the differentiation to the endoderm, mesoderm, and ectoderm lineages, possibly in conjunction with the Smad pathway [30].

The abovementioned studies highlight the somewhat predicted role of ISWI in regulating gene expression in euchromatic regions. However, ISWI is also enriched in heterochromatic regions, suggesting a role for ISWI in initiation or maintenance of heterochromatin formation [209]. ISWI is required for fly embryogenesis, so flies expressing an ATPase-dead dominant negative mutant of ISWI, ISWI<sup>K159R</sup>, were analyzed. Expression of ISWI<sup>K159R</sup> results in decondensation of the male X chromosome, presumably reflecting a loss of chromatin compaction. Interestingly, mutations in the dosage compensation complex that acetylates H4K16, cause the male X to appear

**Table 4** The ISWI family

| Subfamily       | ACF/CHRAC |            |       |      |         |                   |                   |        | NURF       |        |
|-----------------|-----------|------------|-------|------|---------|-------------------|-------------------|--------|------------|--------|
| Species         | Yeast     | Yeast      | Yeast | Fly  | Fly     | Human             | Human             | Human  | Fly        | Human  |
| Complex         | ISW1a     | ISW1b      | ISW2  | ACF  | CHRAC   | ACF               | CHRAC             | WICH   | NURF       | NURF   |
|                 | Isw1      | Isw1       | Isw2  | ISWI | ISWI    | hSNF2H            | hSNF2H            | hSNF2H | ISWI       | hSNF2L |
|                 |           |            | Itc1  | ACF1 | ACF1    | WCRF180/<br>hACF1 | WCRF180/<br>hACF1 | WSTF   | NURF301    | BPTF   |
|                 |           |            | Dpb4  |      | CHRAC14 |                   | hCHRAC17          |        | NURF55/p55 | RbAp46 |
|                 |           |            | Dls1  |      | CHRAC16 |                   | hCHRAC15          |        |            | RbAp48 |
| Unique Subunits | Ioc3      | Ioc2, Ioc4 |       |      |         |                   |                   |        | NURF38     |        |

more condensed, consistent with increased ISWI activity in the absence of H4 acetylation. NURF301 mutant flies are viable and recapitulate the male X phenotype, while ACF mutant flies are semi-embryonic lethal, providing a clear example of functional specialization of the ISWI complexes by associated subunits [210, 211]. In the 30% of ACF-deficient flies that do survive, normal diversification of euchromatin versus heterochromatin during blastoderm development is perturbed [210]. In addition, ACF mutant flies display defects in heterochromatic silencing as assessed by defects in pericentric position effect variegation and Polycomb-mediated repression [35]. The phenotypes observed upon loss of ISWI, ACF, and NURF301 may correlate with the unique ability of ISWI to catalyze the formation of regularly spaced nucleosomal arrays *in vitro*, as regular spacing of nucleosomes in native chromatin may result in greater compaction. However, closer examination revealed that nucleosome spacing is normal ISWI mutant flies despite global perturbations in chromatin compaction [212, 213]. The authors instead ascribed the defect to a loss of H1 on chromatin as H1 mutant flies have a similar defect in compaction.

A specific role for the mammalian WICH complex in DNA replication of condensed regions has also been observed. hSNF2h, ACF1, and the ACF1 homologue WSTF are particularly enriched in replicating pericentric heterochromatin, suggesting that they might have a role in the replication of condensed chromatin [33, 214]. Indeed, knockdown of these proteins led to slower S phase progression that could be ameliorated by treatment with 5-aza-2-deoxycytidine, a DNA methylase inhibitor that results in chromatin decondensation [33]. Further exploration revealed that WSTF is directly recruited to replication foci by PCNA and that knockdown of WSTF results in greater compaction and increased HP1 association of newly replicated DNA [34]. Finally, WSTF binds to  $\gamma$ -H2AX following DNA damage and stabilizes  $\gamma$ -H2AX by phosphorylating Tyr142 [36].

The ISWI family has been implicated in a variety of cellular functions, including transcriptional repression, heterochromatin formation, DNA replication, and ES cell pluripotency. Like Brg1 and Brm, ISWI can assemble with different homologous subunits such as ACF1 or NURF301 to form distinct functional complexes. It is tempting to speculate that combinatorial assembly is an important regulatory mechanism that governs the function of ISWI in developmental or cellular transitions as it does for the BAF complex. Consistent with this hypothesis, ACF1 expression appears to be developmentally regulated in *Drosophila* and constitutive ACF1 expression is lethal [210]. Careful genetic analysis of this type may reveal that subunit switch is a common theme among chromatin remodelers. Finally, it will be important to determine how the unique ability of the ISWI ATPase to order nucleosomes *in vitro* relates to the phenotypes observed *in vivo*.

#### CHD family

The ATPases of the CHD family are characterized by N-terminal tandem chromodomains in addition to the conserved DEAD/H-related ATPase domain. There are nine known family members that have been further categorized into three families based on similarities in domain structure. CHD1 and CHD2 have a C-terminal DNA binding domain, CHD3 and CHD4 lack the DNA binding domain but have N-terminal paired PHD fingers, and CHD5-9 have additional functional domains in their C termini. Of these nine members, CHD1 is the only family member in yeast. Here, we will review the known functions of a few of the well-characterized family members, beginning with CHD1.

**CHD1/2** Biochemical characterization of CHD1 demonstrated that CHD1 exists as a monomer or dimer, unlike the enzymatic subunits of other CRCs [215, 216]. yChd1 has also been shown to interact with the multi-subunit

SAGA and SLIK complexes, two highly related yeast HAT complexes that preferentially acetylate H3 and H2B [216]. CHD1 is targeted to sites of active transcription through PHD-mediated recognition of H3K4me3 [217, 218] and associates with other preinitiation factors to facilitate transcriptional elongation and splicing [39]. Consistent with these findings, genome-wide binding of Chd1 correlates globally with H3K4me3 and Pol II in ES cells [38]. Knockdown of Chd1 in ES cells results in the loss of endoderm lineage potential and the appearance of heterochromatic foci, suggesting that Chd1 is required to maintain an open chromatin state for pluripotency [38]. Interestingly, *Drosophila* CHD1 is required for the deposition of histone variant H3.3 into decondensing sperm chromatin during embryogenesis [219]. Although it is not known whether the related CHD2 protein also functions in transcription or nucleosome assembly, Chd2-deficient mice exhibit decreased neonatal viability and organ failure [40].

**CHD3/4** CHD3 and CHD4, or Mi-2 $\alpha$  and Mi-2 $\beta$ , are incorporated into a large protein complex with both histone deacetylase and remodeling activities called NURD (Table 5). The other subunits of NURD include HDAC1, HDAC2, RbAp48, RbAp46, MTA1, MTA2, MTA3, MBD2, and MBD3. The deacetylase activity of HDAC1/2 on nucleosomal templates is stimulated by ATP, suggesting that the chromatin remodeling activity of NURD facilitates access to acetylated histones [220, 221]. It is thus tempting to assume that NURD represses transcription by binding to methylated DNA in an MBD2/3-dependent manner, remodeling the surrounding chromatin, and removing active histone marks. However, it is unclear how these accessory proteins contribute to the considerably more complex NURD-dependent processes observed *in vivo*.

Our limited genetic knowledge suggests that NURD associates with transcription factors to regulate key developmental transitions. The role of Mi-2 $\beta$  in T lymphocytes was extensively analyzed after it was found to associate with Ikaros, a transcriptional repressor important in the development of T cells [222]. Like Ikaros, Mi-2 $\beta$  is required for thymocyte development and for proliferative expansion of mature peripheral T cells. Interestingly, Mi-2 $\beta$  also promotes CD4 expression during T cell development by antagonizing Ikaros [46, 223]. This is in contrast to Brg, which represses CD4 expression in developing T lymphocytes [151]. Through association with another repressor, Bcl-6, MTA-3/NURD also regulates cell fate decisions in the B lymphocyte lineage [44]. The role of NURD in embryonic development has not been reported. MBD3 was shown to be critical for maintaining

**Table 5** The NURD complex

| Subfamily | NURD      |   |
|-----------|-----------|---|
| Species   | Fly       | Human                                   |
| Complex   | Mi-2/NuRD | NuRD                                    |
|           | dMi-2     | CHD3/Mi-2 $\alpha$<br>CHD4/Mi-2 $\beta$ |
|           | dMBD2/3   | MBD2, MBD3                              |
|           | dMTA      | MTA1,2,3                                |
|           | dRPD3     | HDAC1,2                                 |
|           | p55       | RbAp46,48                               |
|           | p66/68    | p66 $\alpha$ , $\beta$                  |

pluripotency in ES cells through suppression of the trophoctoderm lineage, but it is unclear whether this transcriptional repression is mediated by Mi-2/NURD [224, 225]. Of note, dMi-2 is essential for the development of germ cells in flies [226].

Similarly to INO80, SWR1 and TIP60, NURD is recruited to sites of DNA damage, although this recruitment appears to be mediated by the DNA repair protein PARP rather than  $\gamma$ -H2AX itself [227, 228]. Following recruitment, Mi-2 $\beta$  is phosphorylated by ATM and facilitates the RNF8/RNF168-dependent ubiquitination of  $\gamma$ -H2AX and subsequent RNF8 and BRCA1 accumulation at DSBs [229, 230]. Loss of Mi-2 $\beta$ , in turn, results in aberrant DSB repair and G2/M checkpoint activation, leading to increased sensitivity to DNA damage. Interestingly, loss of Mi-2 $\beta$  in the absence of induced DNA damage led to spontaneous DNA damage, increased levels of p53 and p21, reduced cell proliferation, and apoptosis [230, 231]. Consistent with these findings, it was shown that NURD subunits are lost during premature and normal aging, leading to changes in higher order chromatin structure and spontaneous DNA damage [232]. These data indicate that NURD acts as a gatekeeper of genomic stability.

**CHD5** CHD5 was recently found to be the tumor suppressor gene located in 1p36, a region that is commonly deleted in malignancies of epithelial, neural, and hematopoietic origin [233]. CHD5 is required for p16<sup>Ink4a</sup>/p19<sup>Arf</sup> expression and CHD5 knockdown results in dysregulation of the p19<sup>Arf</sup>/p53- and p16<sup>Ink4a</sup>/Rb-mediated tumor suppressive pathways. Interestingly, loss of heterozygosity (LOH) was not observed in most tumor cell lines, indicating that heterozygosity predisposes to tumor formation. The frequency of this deletion in a diversity of cancers indicates that the requirement for CHD5 in transactivation of Ink4a/Arf may be relatively

universal.

**CHD7** The final Chd family member of note is CHD7. Recently, haploinsufficiency of CHD7 was found to account for 10 of 17 cases of the human CHARGE syndrome, features of which include coloboma of the eye, heart defects, atresia of chonae, renal anomalies, genital anomalies, and ear abnormalities [49]. Mice deficient in CHD7 recapitulate many of the human defects as a result of dysregulated transcription of key developmental genes [50, 234, 235]. Interestingly, CHD7 was found to collaborate with BRG1 in neural crest-like cells at distal enhancer elements to regulate the neural crest transcriptional program [51]. Although this regulation was initially ascribed to the PBAF complex based on the association of CHD7 with BAF180, it is more likely mediated by nBAF that is present in post-mitotic neurons and neural crest cells [121]. Genome-wide binding studies further demonstrated that CHD7 colocalizes with Brg1 and H3K4me1 at distal enhancer sites in mouse ES cells, although SWI/SNF clearly has a more extensive role in defining pluripotency as CHD7-deficient mice survive as late as E10.5, when they presumably die due to defects in cardiac development [51, 236, 237].

### Chromatin remodeling complexes and cancer

Several CRCs, most prominently the BAF complex, have been implicated in cancer initiation or progression. Early studies demonstrated that many cell lines have lost both BRG1 and hBRM expression and that introduction of BRG1 or hBRM results in slower or arrested growth [238]. Other cell lines such as HeLa have partially assembled complexes whose deletion often does not affect the cells and hence these partial complexes seem to be inactive. However, the first definitive evidence that BAF complexes are tumor suppressive came from Versteeg and colleagues who found that BAF47 (SNF5) underwent LOH in nearly all cases of pediatric rhabdoid sarcoma, in which the other allele was mutated or silenced by methylation [239]. These are extraordinarily aggressive tumors of young children that appear to have a short interval between loss of the second allele and malignancy. This disease can be recapitulated in *Snf5*<sup>+/-</sup> mice, which give rise to tumors that are histologically similar to human rhabdoid tumors following LOH [240]. Furthermore, conditional inactivation of *Baf47* in mice leads to lymphomas with 100% penetrance in a median onset time of 11 weeks [241]. Thus BAF47 is a *bona fide* tumor suppressor that when lost gives rise to tumors more quickly than in any other tumorigenic mouse model. Biallelic loss of *BRG1* also occurs in prostate, lung, breast

and pancreatic cancer cell lines, and has been shown to result in pediatric rhabdoid sarcomas [27, 242, 243]. *hBRM* and *BRG1* were lost in 26% and 23% of small-cell lung cancer cell lines and 76% and 77% of non-small-cell lung cancer cell lines, respectively [244]. Consistent with these data, about 10% of *Brg1*<sup>+/-</sup> mice spontaneously acquire tumors within one year [22, 245]. Among the other BAF subunits, *ARID1A* (BAF250a) was found to be deleted in 10% of breast carcinomas, 30% of renal carcinomas, and 57% of primary ovarian clear cell carcinomas [246, 247]. *PBRM1* was deleted in 48% of breast cancer cell lines tested [248], and *BRD7* was frequently deleted in a subset of p53 intact human breast tumors [249]. Collectively, these data argue that BAF and possibly PBAF complexes play a critical role in suppressing tumorigenesis.

Efforts are now focused on how loss of BRG1 might promote cell growth or dysregulation of the cell cycle. It was shown, for example, that Rb is unable to induce G1 arrest in BRG1/BRM-deficient cell lines, while expression of BRG1 or BRM in these cells restores growth inhibition. BRG1 and Rb physically interact and regulate cell cycle in conjunction with cyclin proteins [238, 250]. During G1, Rb-SWI/SNF associates with HDACs to repress the cyclin A and E genes. Phosphorylation of Rb by cyclin D/CDK disrupts the interaction between Rb-SWI/SNF and HDACs, resulting in cyclin E upregulation and progression into S phase. The Rb-SWI/SNF complex remains intact and repressive for the cyclin A gene until cyclin E/CDK2 accumulates and phosphorylates BRG1/BRM and BAF155, leading to disruption of the Rb-SWI/SNF complex and exit from S phase [251]. In addition, BRG1 promotes hypophosphorylation of Rb through enhancing the expression of p21 [252]. In other work, it was shown that SWI/SNF co-associates with p53 and BRCA1, and that SWI/SNF is required for p53- and BRCA1-mediated transcription [253-255]. Similarly, BAF180 and BRD7 regulate transcription of a subset of p53-dependent targets induced during oncogene-induced and replicative senescence [249, 256].

Despite considerable progress on the mechanism of tumorigenesis, a comprehensive understanding of the BAF complex in cancer is lacking. Why deletion of dedicated subunits does not result in similar tumorigenic phenotypes is unknown. One might imagine that loss of BRG1, BAF250a and BAF47 would result in tumorigenesis with similar kinetics and penetrance, but their phenotypes are distinct and tissue specific. The answer may lie in the combinatorial diversity of the BAF complex, which underlies important developmental transitions and appears to have an instructive role in certain cell fates [146]. Differential subunit usage could contribute to tissue- or

gene-specific effects, which might result in distinct tumorigenic phenotypes upon loss of a given subunit. For example, incorporation of BAF250a or BAF250b is anti- and pro-proliferative, respectively, and it is BAF250a that is lost in human carcinomas [247, 257]. If oncogenesis does indeed arise from differential subunit usage, the unique complex interface of malignancy could be specifically targeted using small molecule inhibitors. This theory does not address the specific role of BAF47, which is monomorphic. In this case, it is possible that BAF47 is not required for all of the functions of the BAF complex or that it assembles into specific complexes selectively. Alternatively, loss of BAF47 may result in formation of partial complexes with detrimental and nonphysiological activities. At the molecular level, loss of BAF47 results in upregulation of the Polycomb protein Ezh2 and subsequent repression of *p16<sup>ink4a</sup>/p19<sup>Arf</sup>* [155, 156]. It remains to be seen whether BRG1/hBRM and other subunits of the SWI/SNF complex similarly antagonize Polycomb to suppress tumorigenesis.

The loss or inactivation of CRC-associated proteins is frequently observed in cancer, and many such as BAF47, BAF250, BRG1, and CHD5 are *bona fide* tumor suppressors. Others appear to play a more supporting role, and are among the class often referred to as passengers in the process of oncogenesis. Mechanistically, CRCs may directly repress growth by interacting with other key tumor suppressors, or they may act independently to maintain genomic integrity. Certainly, members of the BAF complex have been implicated in growth restriction and cell cycle arrest mediated by Rb, p53, BRCA1, and *p16<sup>ink4a</sup>/p19<sup>Arf</sup>*. However, given the unprecedented rapid onset and penetrance of tumors following loss of BAF47, it is likely that interactions with known tumor suppressors will not fully explain its tumor suppressive function. The extensive role of related CRCs in DNA repair, DNA replication, and genomic integrity suggests the intriguing possibility that BAF may also contribute to these surveillance mechanisms. Further mechanistic studies are imperative to understand whether the loss of chromatin remodeling activity is related to a breakdown in these basic cellular processes.

## Outlook

The first CRC,  $\gamma$ SWI/SNF, was isolated for its role in transcriptional activation of signal-dependent gene targets. Subsequent work in yeast and *Drosophila* contributed to the strong perception that SWI/SNF and related CRCs are recruited by signal-dependent transcription factors to disrupt nucleosome:DNA contacts to enable transcription. Certainly, it has been well documented that

ATP-dependent remodelers are required for transcription and can catalyze nucleosome sliding and exchange on DNA templates *in vitro*. However, these machines are absolutely critical for many other cellular processes such as DNA replication, recombination, and repair. It is not clear that nucleosomal movement can account for all of the biologic activities of CRCs observed *in vivo* despite clear affinity of the complexes for nucleosomes. CRCs may bind nucleosomal assemblies where they conduct a range of activities, including, but not limited to, nucleosome remodeling. There is a clear need for experimental model systems that faithfully recapitulate the native state of chromatin with its higher order structure and local histone modifications in order to define and characterize these as yet undefined mechanisms.

This unique class of machines is both omnipresent and hard to define. Despite the similarity between their core enzymatic subunits, their common affinity for nucleosomes, and their common ability to disrupt nucleosomal templates *in vitro*, there is little functional similarity between members *in vivo* and as a result, very little predictive power for members of unknown function. In many cases, additional associated subunits radically alter the activity, mechanism, or targeting of the complex to lend it specific function, as in the case of BAF53b in post-mitotic neurons. The utilization of genetic models and genome-wide binding studies to parse out the role of each subunit may reveal that subunit usage is a common theme among CRCs to generate specificity. Capitalizing on this knowledge, we may reveal new therapeutic targets for the treatment of cancer and other CRC-associated diseases.

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