The SAGA chromatin-modifying complex: the sum of its parts is greater than the whole

Jelly H.M. Soffers and Jerry L. Workman
Stowers Institute for Medical Research, Kansas City, Missouri 64110, USA

There are many large protein complexes involved in transcription in a chromatin context. However, recent studies on the SAGA coactivator complex are generating new paradigms for how the components of these complexes function, both independently and in concert. This review highlights the initial discovery of the canonical SAGA complex 23 years ago, our evolving understanding of its modular structure and the relevance of its modular nature for its coactivator function in gene regulation.

Chromatin-modifying and remodeling complexes are important for epigenetic regulation of gene expression. Studies into their exact molecular mode of action have inspired different research areas. Chromatin and structural biology work has helped us understand how these complexes interact with chromatin and their enzymatic substrates. Translational research has profiled epigenetic landscapes in homeostatic, differentiation, and pathological conditions to better understand how chromatin modifications regulate gene expression. However, a detailed understanding of the molecular basis of the gene regulatory actions of chromatin modifiers ultimately starts with a profound understanding of subunit functions, mode of assembly, and complex structure, which together dictate their capacity to alter chromatin structure. This review focuses on one of the first identified chromatin-modifying complexes, the Spt/Ada/Gcn5 acetyltransferase [SAGA] complex. We review its initial discovery, studies that have shaped our view on the internal organization of its subunits into modules, and the latest structural work that visualizes the modules and provides insights into their function. Finally, we discuss how modules also function as noncanonical complexes.

Discovery of the yeast SAGA complex

Studies designed to investigate the molecular mechanisms of eukaryotic gene regulation in yeast led to the discovery of the first chromatin-modifying and remodeling complexes. One set of genetic experiments took advantage of the Ty transposable elements. Ty element insertion mutations interfere with transcription because of Ty-encoded transcription signals. Suppressor of Ty [Spt] mutations restore transcription as they disrupt the genes encoding factors that recognize the Ty transcription signals [Winston et al. 1987; Fassler and Winston 1988]. As such, Spt mutations uncovered proteins that promote eukaryotic gene expression and identified the TATA-binding protein (TBP)-related set of Spt proteins that includes Spt3 [Eisenmann et al. 1992; Laprade et al. 2007], Spt7 [Gansheroff et al. 1995], Spt8 [Eisenmann et al. 1994], and Spt20 [Marcus et al. 1996; Roberts and Winston 1996], also (for review, see Winston and Carlson 1992).

Another set of genetic experiments identified proteins that interacted with transcription activator proteins. Activators bind a specific promoter DNA sequence, recruit components of the transcription machinery and stimulate transcription [Hall and Struhl 2002; Carpenter et al. 2005; Ferreira et al. 2005]. Natural activators in yeast are Gal4 and Gcn4. An unusually potent artificial hybrid Gal4-VP16 activator is formed by the DNA-binding portion of Gal4 fused to a highly acidic portion of the herpes simplex virus protein VP16 that interacts with DNA-binding proteins [Sadowski et al. 1988]. Gal4-VP16 overexpression is lethal and may be due to sequestration of general transcription factors away from productive transcription complexes, perhaps via concentration-dependent liquid phase separation of abundant Gal4-VP16 and its interactors [Guarente 1995; Bolognesi et al. 2016; Boija et al. 2018]. Like the Spt mutations, mutations that suppress Gal4-VP16 toxicity identified proteins that promote gene expression. These proteins were termed transcriptional adaptor [Ada] proteins, because they are thought to form a physical bridge between the upstream DNA-bound activators and the transcriptional machinery at a promoter [Guarente 1995]. Ada1–5 interact with Gcn4 and Gal4-
VP16 in yeast (Marcus et al. 1994). Ada5 is identical to Spt20 (Marcus et al. 1996; Roberts and Winston 1996). Ada4 is currently better known as Gcn5, a nuclear histone acetyltransferase [HAT] enzyme (Brownell et al. 1996; Wang et al. 1997). Ada2, Ada3, and Gcn5 not only interact with activators, but also with each other, and Ada2 also interacts indirectly with TBP (Georgakopoulos and Thir- eos 1992; Marcus et al. 1994; Silverman et al. 1994; Hori- uchi et al. 1995; Candau and Berger 1996; Saleh et al. 1997). These data suggested that Ada proteins are part of activator complexes that mediate interactions between acidic activators and TBP.

The roles for the Spt and Ada proteins in transcription regulation converged with the discovery of the Spt/Ada/ Gcn5 acetyltransferase (SAGA) complex. A collaboration of biochemists and geneticists discovered that the adaptor Gcn5 functions as a catalytic subunit in a high-molecular- mass native HAT complex that acetylated nucleosomal histones at H3 and H2B (Grant et al. 1997). This complex contained Ada2, Ada3, Ada5/Spt20, as well as Spt3, Spt7, and Spt8. The function of Gcn5 as a histone acetyltrans- ferase within the SAGA adaptor complex indicated the importance of histone acetylation during steps in transcription activation, mediated by interactions with transcription activators and general transcription factors. Concomitantly with its discovery, Roberts and Winston (1997) demonstrated that SAGA genetically interacted with the Switch/ Sucrose nonfermentable [Swi/Snf] chromatin remodeling and Mediator complexes. This formed the first indication that SAGA was important for additional aspects of chromatin remodeling and transcription regulation.

A comprehensive overview of SAGA subunits identified in yeast and higher eukaryotes

Much of the early work described above was performed using yeast as a model organism, but the SAGA complex is also conserved in higher eukaryotes. To date, 18 conserved subunits have been identified, as well as some species-specific associated factors [Table 1]. Concomitantly, additional coactivator functions of the SAGA complex emerged. This section summarizes the discovery of these additional subunits and the implications for gene regulation.

Gen5-associated adaptor proteins

The adaptor proteins are conserved in higher eukaryotes. In Drosophila and humans, two Ada2 paralogs exist: Ada2a and Ada2b (Barlev et al. 2003; Kusch et al. 2003; Guelman et al. 2006a). Ada2a incorporates into the Drosophila Ada2a-containing ATAC complex, also a Gcn5-type HAT complex (Guelman et al. 2006a; Suganuma et al.

| Orthologous SAGA complexes |
|-----------------------------|
| **S. cerevisiae** | **S. pombe** | **Drosophila melanogaster** | **Homo sapiens** |
| HAT module | Gcn5 | Gcn5 | KAT2 (GCN5) | KAT2A/KAT2B (GCN5-L/PCAF) |
| Ada2 | Ada2 | Ada2b | TADA2b |
| Ngg1 [Ada3] | Ngg1 [Ada3] | Ada3 | TADA3 |
| Sgf29 | Sgf29 | Sgf29 | Sgf29 |
| DUB module | Ubp8 | Ubp8 | dNonstop | USP22 (UBP22) |
| Sgf11 | Sgf11 | dSgf11 | ATXN7L3 |
| Sgf73 | Sgf73 | dATXN7 | ATXN7/ATXN7L1/L2 (SCA7) |
| Sus1 | Sus1 | dE(y)2 | ENY2 |
| Core module* | Taf5 | Taf5 | WDA/TAF5L | TAF5L |
| Taf6 | Taf6 | SAF6/TAF6L | TAF6L |
| Taf9 | Taf9 | TAF9 | TAF9/TAF9b |
| Taf10 | Taf10 | TAF10b | TAF10 |
| Taf12 | Taf12 | TAF12 | TAF12 |
| Spt7 | Spt7 | Spt7 | SUPT7L (STAF65G) |
| Hfi1 [Ada1] | Hfi1 [Ada1] | Ada1 | TADA1 |
| Spt20 | Spt20 | Spt20 | SUPT20H |
| Spt3 | Spt3 | Spt3 | SUPT3H |
| Spt8 | Spt8 | — | — |
| TF-binding module | Tra1 | Tra1 | Nipped-A [Tra1] | TRRAP |
| Species-specific subunits | Ppr5 | — | SF3B3 | SF3B [SAP130] |
| Cdh1 | — | — | — |

Table 1. Overview of orthologous subunits of the SAGA complex

Shown are the names of each subunit from orthologous SAGA complexes in *S. cerevisiae*, *S. pombe*, *Drosophila melanogaster*, and *Homo sapiens* according to the official nomenclature. Names in brackets are alternative, commonly used names. Paralogous subunits are separated with a slash and are indicated if they have been described as part of SAGA. In most cases, paralogous subunits are mutually exclusive within SAGA complexes. Adapted from Helmlinger and Tora (2017) with permission from Elsevier.

*The core module is composed of Spt and TAF subunits, as well as ADA1. Formerly, the TAFs have been singled out and described as the TAF module, and the remaining subunits as the Spt modules [see Fig. 2, below].
Ad2b is part of *Drosophila* and human SAGA (Martinez et al. 1998; Muratoglu et al. 2003; Qi et al. 2004; Gamper et al. 2009). In fact, two *Ada2b* isoforms exist in *Drosophila*; SAGA contains the long isoform of *Ada2b* (Kusch et al. 2003), and so does another Gcn5-type HAT complex named ADA (Softers et al. 2019). The *Drosophila* Chiffon Gcn5-type HAT incorporates the shorter *Ada2b* isoform (Torres-Zelada et al. 2019).

The Gcn5 family member proteins are PCAF [p300/ CBP-associated factor, also known as KAT2B] and two isoforms of GCN5 (Gen5-L and Gen5-S). The PCAF protein shares extensive similarity with yeast Gcn5, but it has an N-terminal extension that interacts with the transcriptional coactivator p300/CBP (Yang et al. 1996, Smith et al. 1998). The less abundant Gen5-S protein is similar in length to yeast, mouse, and *Drosophila* Gen5p (Candau and Berger 1996; Yang et al. 1996, Smith et al. 1998; Xu et al. 1998), whereas the predominantly expressed Gen5-L protein contains an extended N-terminal domain akin to PCAF (Smith et al. 1998). The human Ada2 and Gen5-S proteins correspond to functional homologs of the yeast adaptors (Candau and Berger 1996). Gen5-L was identified as part of the human SAGA complex (Smith et al. 1998).

SAGA factor 29 (Sgf29) is a conserved SAGA subunit (Sanders et al. 2002, Vermeulen et al. 2010; Lee et al. 2011). It is a bone fide Ada factor that rescues Gal4-VP16-mediated toxicity, just like the other Ada proteins that had been identified previously [Lee et al. 2011]. This subunit contains a tandem tudor domain that recognizes H3K4me3 [Li et al. 2010; Bian et al. 2011].

### The SAGA complex contains TBP-associated factors

Transcription initiation requires the assembly of the pre-initiation complex (PIC), formed by polymerase II [Pol II] along with the general transcription factors, which include TFIIA, TFIIB, TFIID, TFIIE, TFIIF, and TFIH. PIC formation typically starts with the recognition of the core promoter by TFIID, which consists of TBP and up to 14 other proteins called TBP-associated factors [TAFs] (Thomas and Chiang 2006). In agreement with its widespread role in transcription initiation, the SAGA complex also contains TAF subunits. In fact, human SAGA was initially named STAGA because it contained TAF9 [TAF9b] (Ogryzko et al. 1998; Martinez et al. 2001).

SAGA-related complexes also contain TAFs. One such complex is the TBP-free TAF-containing complex (TFTC) (Wieczorek et al. 1998). TFTC acetylates similar lysine residues to SAGA, and is in fact very similar in structure and subunit composition to mammalian SAGA (Brand et al. 1999a, b; Cavusoglu et al. 2003). Furthermore, PCAF also associates with multiple TAFs [Kotani et al. 1998].

The TAFs that form an integral part of yeast SAGA are Tafl4, Tafl6, Tafl9, Tafl10, and Tafl12 (Grant et al. 1998a). Although both yeast and metazoan incorporate TAFs, metazoans have evolved several paralogs not found in yeast. Whereas the TAFs that are part of yeast SAGA are all shared with TFIID, in *Drosophila* SAF6 is the SAGA-specific paralog of TAF6, TAF10b is the paralog of TAF10 [Georgieva et al. 2000; Weake et al. 2009], and WDA is the paralog of TAF5 [Guelman et al. 2006b]. Similarly, human SAGA incorporates TAF5L, TAF6L, and TAF9B, whereas TAF5, TAF6, and TAF9 are found in TFIID [Helmlinger and Tora 2017].

### The SAGA complex contains the activator-interacting protein Tra1

Yeast Tra1 is related to the ataxia telangiectasia mutated [ATM] family of phosphatidylinositol 3 [PI 3] kinases [PIKKs] and is a conserved SAGA subunit (Grant et al. 1998b; McMahon et al. 1998; Saleh et al. 1998; Kusch et al. 2003). PIKKs contain numerous conserved domains, including HEAT [huntingtin, elongation factor 3 protein, phophatase 2A, and lipid kinase TOR] and FAT [focal adhesion translocase] domains [Knutson and Hahn 2011; Cheung and Diaz-Santin 2019]. The Tra1 pseudokinase domain is required for nuclear localization and incorporation into SAGA and other coactivator complexes [Berg et al. 2018; Elias-Villalobos et al. 2019]. The best characterized function of Tra1 in SAGA is its ability to associate with activators. VP16 interacts with the C-terminal half of Tra1 [Brown et al. 2001], whereas the N-terminal HEAT domain binds yeast Gal4 [Lin et al. 2012], Gcn4 and Rap1 [Han et al. 2014], human p53 [Li et al. 2004], and c-Myc [Park et al. 2001]. The multiple activator interaction domains may provide a simple mechanism to integrate signaling from multiple activators; for example, by different activators binding Tra1 simultaneously and/or competitively [Cheung and Diaz-Santin 2019]. In addition, the presence of multiple activator binding sites may serve to increase avidity by binding single activator with multiple contact points as described between activator Gcn4 and Mediator subunit Med15 [Tuttle et al. 2018; Cheung and Diaz-Santin 2019].

Tra1 is not the only subunit of SAGA that interacts with activators. Gcn5 interacts with Myc [Liu et al. 2003; Zhang et al. 2014] and Adal, Tafl6, Tafl9, and Tafl12 interact with Gcn4 and/or VP16 [Goodrich et al. 1993; Uesugi et al. 1997; Natarajan et al. 1998; Klein et al. 2003; Fishburn et al. 2005; Reeves and Hahn 2005]. These multiple interactions facilitate the recruitment of SAGA coactivation activities to an activator-bound promoter.

### The SAGA complex not only contains histone acetylation, but also ubiquitin protease activity

The next major discovery was that yeast SAGA contains not only HAT activity, but also histone deubiquitinase [DUB] activity. H2B lysine 123 undergoes ubiquitination, which serves as a transient signal that orchestrates a diverse set of events during transcription [Shilatifard 2006; Fleming et al. 2008; Cheon et al. 2020]. Four subunits have been identified in yeast SAGA that are required to cleave monoubiquitin from H2B. The ubiquitin protease 8 [Ubp8] is the catalytic protease subunit (Sanders et al. 2002, Henry et al. 2003; Daniel et al. 2004). SAGA-associated factor 73 [Sgf73] [McMahon et al. 2005] anchors the DUB subunits to the SAGA complex [Rodriguez-Navarro et al. 2004;
McMahon et al. 2005; Köhler et al. 2008) and zinc finger protein SAGA-associated factor 11 [Sgf11] and SI gene upstream of ySa1 [Sus1] are required for protease activity (Powell et al. 2004; Ingvarsdottir et al. 2005; Lee et al. 2005; Köhler et al. 2006). In humans, more variety exists in the subunits that encode DUB activity [Köhler et al. 2010; Atanassov et al. 2016]. Furthermore, ATAXIN-7, the homolog of yeast Sgf73, exists as three paralogous proteins in humans [Helminger and Tora 2017].

Sus1 is also a functional component of the nuclear pore-associated mRNA export machinery TREX-2 (Rodriguez-Navarro et al. 2004; García-Oliver et al. 2012). A physical interaction between SAGA and TREX-2 (Ellisdon et al. 2010) is essential for targeting the transcriptional machinery to the periphery of the nuclear pore complex and contributes to transcription-coupled mRNA export. Moreover, a functional interaction is required for maintaining the ubiquitination status linked to genome stability, since SAGA deubiquitination activity depends on TREX-2 integrity [Evangelista et al. 2018].

Species-specific subunits

The SAGA complex incorporates subunits that are important for mRNA splicing. These are associated with the U2 small nuclear ribonucleoprotein particle [U2 snRNP] that forms a core constituent of the spliceosome [Zhang et al. 2020]. Human SAGA contains spliceosome-associated protein 130 [SAP130], a subunit of splicing factor SF3B found in the U2 snRNP [Martinez et al. 2001]. This finding suggested a function for SAGA in transcription-coupled pre-mRNA splicing. Drosophila SAGA contains SF3B3 and SF3B5. Although SF3B5 does not appear to function in SAGA’s histone-modifying activities, it is still required for expression of a subset of SAGA-regulated genes [Stegeman et al. 2016]. Both factors associated with SAGA independent of RNA and interacted with Sgf29 and Spt7 [Stegeman et al. 2016]. S. cerevisiae SAGA interacts with Ppr5 via Spt3 and Spt7 in Shao et al. [2020]. Ppr5p is a spliceosomal RNA-dependent ATPase required for stable binding of U2 snRNP to the pre mRNA and subsequent prespliceosome assembly [Ruby et al. 1993; Xu and Query 2007]. Shao et al. [2020] proposed that the interaction with SAGA mediates a balance between transcription initiation/elongation and prespliceosome assembly.

Finally, the chromatin remodeling protein chromo-ATPase/helicase DNA-binding domain 1 (Chd1) associates with SAGA and SAGA-like complex (SLIK) in S. cerevisiae [Pray-Grant et al. 2005]. Chd1 has functions in chromatin remodeling, transcription regulation, and transcription elongation [Tran et al. 2000; Simic et al. 2003]. One of the two chromodomains of Chd1 specifically interacts with the H3K4me3, a mark that is associated with transcriptional activity [Pray-Grant et al. 2005]. Furthermore, the SLIK complex shows enhanced acetylation of a methylated substrate, and this activity was dependent on a functional methyl-binding chromodomain [Pray-Grant et al. 2005]. This suggest that there can be crosstalk between Gcn5-mediated acetylation and the recognition of promoter methylation marks.

SAGA structure meets function

SAGA subunits coactivate gene expression at multiple points during the process of transcription. How the subunits organize and how this potentiates transcription is discussed next.

Genetic and biochemical experiments identify critical core subunits

The first genetic and biochemical studies grouped SAGA subunits based on their mutant phenotypes and effects on the composition of the complex [Grant et al. 1998c]. Complete deletion of the adaptor proteins Gcn5, Ada2, or Ada3 cause the Ada− growth phenotype, which is characterized by the inability to grow on minimal medium and the inability to repress Gal4-VP16-mediated toxicity [Berger et al. 1992; Piña et al. 1993; Marcus et al. 1994; Eberharder et al. 1999; Lee et al. 2011]. Deletion of Spt3 or Spt8 causes the Spt− phenotype, which is characterized by the abnormal initiation of certain transcripts [Winston and Carlson 1992]. Deletion of the gene Spt7 or Spt20 causes both Ada− and Spt− phenotypes [Marcus et al. 1996; Roberts and Winston 1996] and their deletion is synthetically lethal in combination with mutations in genes that encode members of the Swi/Snf or Mediator complexes [Roberts and Winston 1997]. In contrast, null mutations in the SAGA genes Gcn5, Spt3, and Spt8 do not cause lethality in combination with mutations in these other transcriptional regulators, which sets Spt20 and Spt7 apart [Roberts and Winston 1997]. In addition, Spt20, Spt7, and also Ada1 deletion disrupts SAGA complex formation as well as function [Grant et al. 1997; Horiuchi et al. 1997; Roberts and Winston 1997; Sterner et al. 1999; Wu and Winston 2002]. Together, these studies demonstrate that Spt7, Spt20 and Ada1 are critical structural core components of the S. cerevisiae SAGA complex.

Architecture of the SAGA complex

The first structure of the SAGA complex was derived from electron microscopy and mapped subunit positions with different labeling techniques and mutant yeast strains [Wu et al. 2004]. This divided SAGA into five domains [Fig. 1A]. Domains I and V are involved in transcriptional regulatory functions as domain I consists mainly of the Tra1 protein, required for interaction with activators, and V contains the TBP-binding protein Spt3. Domain III contains the HAT Gcn5, and central domains II, III, and IV contain the TAF subunits and the structural components Spt7, Spt20, and Ada1. Later studies roughly mapped the DUB subunits a region corresponding to domain III, close to TAF subunits Ada1 and Spt7 [Fig. 1B, Setiaputra et al. 2015]. These models showed that the Spt subunits form a TBP-binding surface that is adjacent to Tra1 on one side and both enzymatic modules at the other lateral side of the TBP-binding surface [Wu et al. 2004; Setiaputra et al. 2015].

Tra1 was initially thought to be a critical structural core subunit, mainly because its large size (≈400 kDa) would
allow it to act as a scaffolding protein. However, the Ada1 and Spt20 mutants that lack Tra1 are still able to form partial SAGA complexes; thus, Tra1 cannot be a critical subunit that nucleates SAGA assembly (Wu and Winston 2002; Lee et al. 2011). In fact, Tra1 assumes a peripheral position and is not located centrally in the complex at all, nor does it extensively cross-link to different subunits (Figs. 1C,D, 3) [Han et al. 2014; Diaz-Santin et al. 2017; Sharov et al. 2017; Cheung and Diaz-Santin 2019]. The main connection forms between the Tra1 FAT domain and several regions of Spt20 and Taf12 (Fig. 3A,B, Han et al. 2014; Elias-Villalobos et al. 2019; Papai et al. 2020; Wang et al. 2020).

Further cross-linking mass spectrometry experiments corroborated these data and show that TAFs and Ada1 assume a central position and are interlinked with Spt subunits, and link the HAT and DUB module to the TAF/Ada1 center, whereas Tra1 does not form many interprotein connections (Fig. 1D, Han et al. 2014).

Figure 1. Schematic overview of the architecture of the SAGA complex based on electron microscopy, immunolabeling, and cross-linking mass spectrometry techniques. (A) The first S. cerevisiae SAGA structure divides it in five domains (adapted from Wu et al. 2004, with permission from Elsevier). Tra1 occupies domain I, and domains II–IV contain TAFs, core (Ada1, Spt7, and Spt20) and HAT subunits (Gcn5). Domain V is flexible and contains Spt3. (b) The subsequent SAGA structure locates the DUB module as bulging density in close proximity to TAF and Spt subunits. (Adapted from research originally published in Setiaputra et al. 2015. © the American Society for Biochemistry and Molecular Biology). This structure highly resembles the previous structure (Wu et al. 2004), when domain V is flipped toward domain I. The Spt subunits form a TBP-binding surface adjacent to Tra1, and the DUB and HAT modules locate to the right. Vertical dashes indicate the midplane. Subunit Spt3 and the DUB module are located on the backside. (C) Cryo-EM structure showing the narrow hinge region that forms the peripheral connection between Tra1 and the core modules (arrow; Tra1 cross-links to Spt20 and Taf12). The DUB (purple) and HAT module (yellow) associate with the structural core (green/white). The vertical stripes indicate the midplane. The DUB module anchors via the backside. Both enzymatic modules form a clamp that engages nucleosomes, and the HAT module can swivel ∼15° toward the DUB module (dashed arrow) (adapted from Cheung and Diaz-Santin 2019). (D) The architectural map determined by cross-linking mass spectrometry (adapted with permission from Han et al. 2014; © 2014 The Authors) reveals the close proximity of Tra1 and the Spt subunits, and the peripheral connections of Tra1 to the rest of the complex. The TAFs with Ada1 assume a central position and are interlinked with Spt subunits. The HAT and DUB module link to the TAF/Ada1 center, and the DUB module cross-links also to Spt20.

Modular divisions of the SAGA complex

A yeast combinatorial depletion study generated a comprehensive interaction matrix for all the SAGA subunits (Lee et al. 2011). The modular organization of the complex as well as the interrelationships between its subunits were determined with computational methods after the systematic purification of several tagged SAGA subunits in genetic deletion strains [Lee et al. 2011]. This model divided the SAGA subunits in four modules (Fig. 2A): the enzymatic HAT module (Gcn5, Ada3, Ada2, and Sgf29); the DUB module (Ubp8, Sgf73, Sgf11, and Sus1); the TAF module, composed of all of the SAGA TAF proteins (Taf6, Taf5, Taf12, Taf9, and Taf10); and the SPT module, consisting of all of SAGA’s SPT proteins (Spt7, Spt8, Spt3, and Spt20) together with Tra1 and Ada1.

Thus, the critical core subunits (Ada1, Spt7, and Spt20) that had been mapped to a central location within the complex (Wu et al. 2004; Han et al. 2014; Setiaputra et al. 2015) also reside together in the Spt module (Lee et al. 2011). However, the exact positions of Ada1, Spt7, and Spt20 raised the question of how they could form a central structural core. Spt7 and Spt20 cross-linked to different sets of subunits and not to each other (Fig. 1C; Han et al. 2014). Specifically, Spt7 cross-linked to Spt8, primarily between the C terminus of Spt7 and WD40 repeats at the center of Spt8 (Han et al. 2014). Spt7 also cross-linked to Gcn5 and Ada3, showing close positioning of Spt7 to the HAT module, in agreement with the models by Wu et al. (2004) and Setiaputra et al. (2015). In contrast, Spt20 cross-linked to neither Spt7, Spt8, nor the HAT module. Instead, Spt20 cross-linked to TAFs, the DUB module, and Spt3 (Han et al. 2014).

The latest high-resolution cryo-EM structures of SAGA from S. cerevisiae and P. pastoris corroborate these results and show how Ada1, Spt20, and Spt7 together can be part of one and the same structural core (Fig. 2B; Papai et al. 2020; Wang et al. 2020). These structures reveal that the TAFs, Ada1, and Spt subunits together form the structural core of the SAGA complex. Spt20 and Spt7 are found at opposing lateral sides and Ada1 at a central position (Figs. 2B, 3A). Moreover, they resolve questions on the subunit stoichiometry. Both SAGA and TFIID contain Taf5, Taf6, Taf9, Taf10, and Taf12, but TFIID contains two copies of these TAFs [Leurent et al. 2002]. The first structures could not yet conclusively resolve their stoichiometry [Wu et al. 2004; Setiaputra et al. 2015]. However, these cryo-EM studies reveal that each subunit of the core module is
present as one copy [Fig. 2B]. Finally, these studies categorize the SAGA modules as follows: Tra1 module, core module (Spt, TAFs, Ada1, and Spt20), HAT module (Gcn5, Ada2, Ada3, and Sgf29), and DUB (Sgf73, Sgf11, Sus1, and Ubp8) [Fig. 2B; Papai et al. 2020; Wang et al. 2020].

The histone fold gives rise to a histone-like octamer structure within SAGA’s core module

All of the SAGA core module subunits, except for Taf5 and Spt20, have a histone fold (HF) motif (Gangloff et al. 2001a,b). The HF is a protein–protein interaction motif that was originally described in the heterodimerization of histones H4 and H3, and also H2A and H2B and their assembly from a central H3-H4-heterotetramer into the histone octamer as it is found in nucleosomes (Luger et al. 1997). The motif is composed of three helices separated by two short strap loops and it assembles into heterodimers by interleaving the helices into the “handshake motif” and juxtaposing the strap loops into short parallel β-bridges (Arents et al. 1991).

TAFs similar to H3, H4, and H2B were identified in Drosophila (Goodrich et al. 1993; Weinzierl et al. 1993; Kokubo et al. 1994; Mengus et al. 1995), humans (Weinzierl et al. 1993; Hisatake et al. 1995; Mengus et al. 1995; Hoffmann et al. 1996; Ogryzko et al. 1998), and yeast (Poon et al. 1995; Reese et al. 2000) (also for review, see Burley and Roeder 1996). Soon, biochemical studies emerged that showed that the TAFs in TFIID could form a histone octamer-like structure, like histones do as part of the nucleosome (Hoffmann et al. 1996). The Drosophila H3–H4-resembling TAFs were found to dimerize via HF domains (Nakatani et al. 1996) and Xie et al. (1996) demonstrated with X-ray diffraction studies that a Drosophila [Taf6-9]x2 tetramer forms, and postulated that four copies of Taf12 will complete the octamer. In 2001, Selleck et al. (2001) crystalized an octamer composed of two copies of the tetramer formed by Taf6, Taf9, Taf4, and Taf12, and other studies show that this octamer is complemented by a hexamer of remaining TAF subunits in TFIID (Kolesnikova et al. 2018) in agreement with findings in insect cells (Biemiossek et al. 2013).

Whether histone-like TAFs arranged into an octamer core in SAGA remained largely unclear until the stoichiometry and structure of the SAGA core subunits were resolved. The SAGA core contains an octamer of TAF subunits formed by the following histone fold dimer pairs: Taf12–Ada1, Taf9–Taf6, Taf10–Spt7, and Spt3 [Fig. 2B; Papai et al. 2020; Wang et al. 2020]. The Spt3 dimer pair is not formed by two proteins, as one copy of Spt3 contains both halves of an interacting histone fold pair located at the N-terminal and C-terminal ends (Birck et al. 1998). This octamer configuration explains why previous EM studies located Spt7 in proximity to the TAFs and especially Taf10, and mapped Ada1 in proximity of Taf12 [Fig. 2B; Wu et al. 2004; Han et al. 2014].

In contrast to the canonical symmetric histone octamer that contains two copies of each histone, the octamer in SAGA is asymmetric and formed by one copy of each
HF protein [Papai et al. 2020; Wang et al. 2020]. In addition, it is not built up from a central histone fold domain, but from one end through interactions with Taf5 [Papai et al. 2020; Wang et al. 2020]. Taf5 lacks a HF domain but contains WD40 domain repeats [Durso et al. 2001]. WD40 repeat proteins act as scaffold proteins because of their β propeller structure [Xu and Min 2011]. In SAGA, the Taf5 WD40 propeller docks to two HF pairs (Taf6-9 and Taf10-Spt7) and binds Spt20 [Wang et al. 2020]. Spt20 also contains an extended loop that forms a wedge between the N-terminal and C-terminal Taf5 domains, thereby stabilizing them in a defined orientation. The N-terminal helical domain of Taf5 binds the HEAT repeat region of Taf6 [Papai et al. 2020; Wang et al. 2020]. This interaction is conserved, and influences the dimerization of human TAF6 and TAF9 [Scheer et al. 2012]. Conversely, Taf6 contributes one β strand to the Taf5 propeller, making them obligate heterodimers [Papai et al. 2020; Wang et al. 2020]. Work on human cells further demonstrated that TAF6’s HF dimer partner TAF9 also interacts with...
the WD40 domain of TAF5 in a discrete TAF5–TAF6–
TAF9 subcomplex, which further elaborates the elabo-
rate interactions between Taf5, Taf6, and Taf9 (Antonova
et al. 2018). Taken together, both HF and non-HF core sub-
units interact and define the architecture of the core.

The core structure is important for TBP binding

Recent work has addressed how TBP is loaded onto
promoters by TFIIID (Patel et al. 2020), but how SAGA
mechanistically contributed to TBP loading had remained
largely unclear. Now that the structure of the SAGA core
is resolved, several studies converge into a model where
SAGA tethers TBP to Spt subunits and loads it onto
DNA when certain conformational changes are induced,
as discussed next.

Genetic and biochemical studies have implicated all
SAGA Spt subunits in SAGA–TBP association, and Spt3
and Spt8 bind TBP directly (Eisenmann et al. 1992, 1994,
Marcus et al. 1996; Roberts and Winston 1996; Sterner
et al. 1999; Belotserkovskaya et al. 2000; Larschan and
Winston 2001; Warfield et al. 2004; Sermwittayawong
and Tan 2006; Labrade et al. 2007). Sermwittayawong
and Tan (2006) proposed a two-step model in which
SAGA, via Spt8, binds TBP to facilitate its subsequent
transfer to a TATA box. However, the role of Spt8 has re-
mained controversial. No Spt8 homologs have been de-
scribed in higher eukaryotes (Helmilinger and Tora
2017), suggesting that other subunits must sustain TBP
binding. Even in yeast, two SAGA-like complexes that
lack Spt8 are able to engage TBP and coactivate gene ex-
pression, indicating that Spt8 is not necessarily always re-
quired (Saleh et al. 1997; Grant et al. 1998b). In agreement
with a more modulatory role for Spt8 in TBP binding, im-
pairment of Spt8 barely impacts mRNA synthesis, where-
as Spt3 is absolutely required (Baptista et al. 2017). In the
model proposed by Ben-Shem and colleagues (Papai
et al. 2020), Spt8 engages TBP, and introduction of TFIIA
leads to transfer of TBP from Spt8 to Spt3. Spt8 binds the
N-terminal region of TBP (Wu and Winston 2002,
Han et al. 2014; Papai et al. 2020). The cryo-EM structures
show how multiple Spt3 helices bind the C-terminal stir-
rup of TBP (Papai et al. 2020). In SAGA, the Spt3 HFs are
tilted compared with its analogous H2A–H2B histone pair
in the nucleosome. This tilt prevents the participa-
tion of Taf10 α3 helix in the interaction, such that Spt3
is barely linked to Taf10 and able to engage TBP (Papai
et al. 2020). Spt20 may also aid the Spt3–TBP binding.
Spt20 cross-links to the N-terminal side of TBP near the
Spt8 interaction site, but also to both ends of the Spt3
C-terminal HF (Han et al. 2014), and Spt20 deletion leads
to a severe reduction in TBP binding at the
GAL 1 locus (Dudley et al. 1999). TBP–Spt3 binding induces a con-
formational change in Spt3 that allows DNA to pass between
Tra1 and the remainder of the SAGA complex, which fa-
cilitates TBP loading onto promoters (Papai et al. 2020).
Since Spt3 and TFIIH mutually exclusively bind TBP (War-
field et al. 2004), the actual release of TBP from SAGA is
important for further PIC assembly (Papai et al. 2020),
but how this is regulated is not currently understood. Bio-
chemical studies suggest that TBP/TFIIA binding directs
the dissociation of the TATA box from the surface of the
histone octamer (Godde et al. 1995). Histone acetylation
also weakens the interaction between nucleosomal
DNA and histone tails, since lysine acetylation affects
the higher-order folding of chromatin fibers and loosens
—but does not totally abolish—the contacts between
the DNA and the nucleosomes [Li et al. 2007]. Indeed, in
vitro binding of the TFIIA–TBP complex is stimulated by
Gcn5 HAT activity (Biswas et al. 2004). Thus, SAGA
HAT activity might support the role of TFIIA and help
to facilitate TBP release by promoting its binding to the
TATA box.

The core structure allows a flexible link of the enzymatic
modules

The yeast DUB module forms two functional lobes orga-
nized around the catalytic subunit. N-terminal residues
of Sgf73 are located between the two Ubp8 domains and
potentiate Ubp8 activity (Köhler et al. 2010; Samara
et al. 2010). Ubp8, the subunit containing DUB activity,
is inactive unless associated with the three other subunits
of the DUB module (Köhler et al. 2008; Lang et al. 2011).
In contrast to yeast, in humans and Drosophila the anchoring
subunit Ataxin-7 is completely dispensable for DUB
module enzymatic activity (Mohan et al. 2014). The DUB
module joins the SAGA core module close to the
Tra1 hinge region and subunit Sgf73 forms the anchor
(Köhler et al. 2010; Samara et al. 2010; Morgan et al.
2016; Wang et al. 2020). The C-terminal part of Sgf73 is re-
quired for insertion into the core module (Lee et al. 2009;
Kamata et al. 2013; Durand et al. 2014; Han et al. 2014)
and enters the core proximal to the Taf9 histone fold, tra-
verses an elongated domain of Ada1, and exits next to the
Spt20 SEP [shp1, eyc, and p47] domain (Fig. 3A–C; Papai
et al. 2020; Wang et al. 2020). Tra1 and Spt20 are critical
for the association of the DUB module in S. pombe (Elí-
as-Villalobos et al. 2019). Likely, Tra1 stabilizes the con-
figuration of Spt20 in such a way that the DUB module
can anchor. This finding further stresses that non-HF-con-
taining core subunits are also important for SAGA struc-
ture and explains why Ada1 and Spt20 are critical struc-
tural subunits, required for the integration of the
DUB activity.

Several studies investigated how the enzymatic mod-
ules contact the nucleosome, but exact interaction sites
remain hard to identify (Durand et al. 2014; Papai et al.
2020; Wang et al. 2020). The structure of the DUB module
bound to an ubiquitinated nucleosome superimposed
onto the S. cerevisiae SAGA structure shows that one nu-
cleosome fits between the HAT and DUB module (Wang
et al. 2020). The HAT module maps close to the DUB
module, and is the most mobile region of the complex
(Setiaputra et al. 2015; Papai et al. 2020; Wang et al.
2020). It adopts multiple conformations and orientations.
Accordingly, the bulk of this module is visible only as an
elongated featureless density. In contrast to the DUB
module, no parts of the HAT module are embedded in
the SAGA core. The HAT module docks to SAGA at
subunit Taf6, where two helical domains—attributable to Ada3—lie at the surface of the Taf6 HEAT repeat domain (Papai et al. 2020; Wang et al. 2020), which agrees with earlier work that demonstrated that Ada3 links the HAT module to the SAGA complex [Han et al. 2014]. It is interesting that the Taf6 HEAT repeat domain is important for anchoring the HAT module. As eluded to earlier, this domain is also important for the modulation of TAF9–TAF6 dimerization by TAF5 [Scheer et al. 2012; Antonova et al. 2018]. Deletion of the Taf6 HEAT domain does not affect the stability of the yeast SAGA complex, even though it does disrupt the H3/H4-like Taf6/9 dimer pair [Dahiya and Natarajan 2018], which agrees with the notion that other interactions stabilize Taf6 and Taf9. Indeed, in the human TAF5/TAF6/TAF9 subcomplex, the TAF9 C-terminal domain wraps around the TAF5 WD40 repeat domain like a clamp with three major anchor points [Antonova et al. 2018]. In TAF9 mutants where the TAF9/TAF5 interaction surface is completely disrupted, SAGA fails to form completely. The only interaction that this mutant TAF9 can maintain is that with TAF6. TFIIID is less stable, and interference with one of the three major anchor points is already sufficient to affect complex stability [Antonova et al. 2018]. This indicates that TAF5/TAF6 have different stabilizing protein contacts in each complex also (for review, see Timmers 2020). In TFIIID, Taf5 docks to the octamer-like fold, where it is stabilized by Taf4 (Wang et al. 2020). In SAGA, which lacks Taf4, Ada1 assumes the position of Taf4 and stabilizes Taf5, and the Taf5 N-terminal tail occupies a rather distant position where it is further stabilized by Spt20 and contacts Taf6 [Wang et al. 2020]. Thus, in the SAGA complex, Taf5, Spt20, and Ada1 are important stabilizers of Taf6 and Taf9, which is in agreement with all the genetic and biochemical studies that have uncovered since the very early discovery of the SAGA complex that Spt20 and Ada1 are critical core components for structural integrity and SAGA function.

The composition of the HAT module facilitates its function

The SAGA HAT module contains four Ada subunits that are critical for the regulation of the enzymatic lysine acetyltransferase activity as well as specific lysine residue recognition [Piña et al. 1993; Horiuchi et al. 1995; Syntichaki and Theoreos 1998; Grant et al. 1999; Balasubramanian et al. 2002; Ringel et al. 2015]. Recombinant Gcn5 acetylates free histones [Kuo et al. 1996], but the Gcn5-associated proteins enhance the ability of Gcn5 to acetylate nucleosomal histones and confer upon Gcn5 the ability to acetylate an expanded set of lysine residues [Grant et al. 1997, 1999] on both histone and nonhistone substrates [Kim et al. 2010].

The particular requirement for Ada2 became evident when Gcn5 with an active HAT domain but missing Ada2-interacting portion could not sustain yeast growth [Candau et al. 1997]. Recent structural work reveals the molecular explanation. Ada2 contains the SANT histone-binding domain [found in Swi3, ADA2, N-Cor, and TFIIIB, hence named SANT] [Sterner et al. 2002], a ZZ-type zinc finger (ZZ) domain involved in histone tail binding [Zhang et al. 2019] and a Swi3p, Rsc8p, and Moitra (SWIRM) domain, a eukaryotic domain found in proteins implicated in chromatin remodeling and gene expression [Boyer et al. 2002; Sterner et al. 2002; Qian et al. 2005; Da et al. 2006]. The SANT domain does not appear to act as a histone tail-binding module, because it is positioned away from the Gcn5 peptide-binding pocket. Instead, the structure of Gcn5 and Ada2 associated with acetyl-CoA substrate suggests that the SANT domain enhances binding of the acetyl-CoA substrate [Sun et al. 2018]. Gcn5, Ada2, and Ada3 form a heterotrimeric complex when expressed and combined in vitro. They form an oblong complex where Ada2 as the linchpin holds Ada3 and Gcn5 together [Horiuchi et al. 1995; Balasubramanian et al. 2002]. Ada3 links the HAT module to SAGA, but deleting the most C-terminal portion of Ada3 does not compromise SAGA stability, nor does it lead to dissociation of Gcn5 from the SAGA complex [Han et al. 2014]. However, it still causes the classical Ada+ growth phenotype, and to an extent equivalent to complete deletion of the ADA3 gene [Han et al. 2014]. Therefore, the C-terminal domain of Ada3 is required for full Gcn5 activity in vivo.

In vitro, the trimeric complex of Ada3, Ada2, and Gcn5 is sufficient for nucleosomal H3K9 and H3K14 acetylation [Balasubramanian et al. 2002]. In vivo, Sgf29 is likely required for promoter recruitment and H3K9 acetylation at these regions is impaired in Sgf29 mutants [Bian et al. 2011]. Sgf29 contains a tandem tudor domain that binds histone H3 methylated on lysine 4 [Bian et al. 2011]. H3K4me3 recognition allows SAGA to dock to pre-existing trimethyl marks at promoter regions and promotes subsequent processive acetylation [Ringel et al. 2015]. The C-terminal Ada3 mutant likely cannot bind Sgf29 and processive acetylation is disturbed such that the Ada− growth phenotype presents itself.

Distinct requirements of different modules during gene regulation

Although SAGA holistically integrates multiple coactivator functions, there is a distinct genetic requirement for each module during gene regulation. For example, even though Tra1 is important for recruitment of SAGA to promoter regions, S. pombe SAGA complexes lacking Tra1 display only very limited defects in gene regulation and show no global decrease in gene expression [Helmlinger et al. 2011]. In addition, gene expression profiles of spt3Δ or spt8Δ mutants are distinct from those of ubp8Δ or sgf11Δ, indicating that loss of TBP-binding ability affects gene expression to a different degree than loss of DUB activity [Ingvarsdottir et al. 2005]. Deletion strains that interfered with HAT function (gcn5Δ, ada2Δ, or ada3Δ mutants) behaved comparably, but affected only a moderate number of genes [178], and only a subset of these were affected by loss of Sgf29 [Helmlinger et al. 2011]. Together, these data suggest that SAGA’s coactivator function depends on its structural integrity, ability to engage TBP,
anchor the DUB module, and HAT activity. However, not all genes require all of these activities.

One striking observation is that loss of Spt20 causes severe defects in gene regulation in S. pombe and S. cerevisiae (Lee et al. 2000; Helmlinger et al. 2011; Bonnet et al. 2014; Warfield et al. 2017), which suggests that its deletion disrupts multiple coactivating functions simultaneously, which is in keeping with its role as a key structural component. However, Spt20, Ada1, and Spt7 mutants do not behave identically, suggesting that even though these are all critical for the structure of the SAGA complex, they do not govern the same coactivating functions equally (Helmlinger et al. 2011). Spt20 and Spt8 mutants grossly affect the same genes. This indicates that a portion of the genes that require Spt20 rely on TBP interactions. However, the number of genes deregulated in Spt20 mutants exceeded that of Ada1, Spt7, and also exceeded the number of genes that are misexpressed in the Sgf73 and Ubp8 mutants. Therefore, the additional defects in Spt20 mutants compared with Ada1 and Spt7 can be explained by the inability to anchor the complete DUB module to SAGA, as has been observed in multiple studies (Lee et al. 2011; Elías-Villalobos et al. 2019). A dis-associated submodule of the four DUB subunits in yeast is likely still catalytically active (Köhler et al. 2010; Samara et al. 2010; Lim et al. 2013). This suggests that not associating the active DUB module to SAGA leads to more severe consequences than a catalytically dead module and raises intriguing questions about how DUB function is regulated.

Different studies have addressed how DUB activity is regulated at the promoter and gene body. In S. cerevisiae, global ubH2B levels increase upon deletion of Ubp8 or the anchoring subunit Sgf73, indicating that the DUB is inactive when not anchored to SAGA (Bonnet et al. 2014). However, in humans, loss of the anchoring subunit leads to a decrease in global ubH2B levels and increased ubH2B levels in gene bodies of expressed genes (Bonnet et al. 2014). Also, in Drosophila, a global decrease in ubH2B is observed upon deletion of the anchoring subunit (Mohan et al. 2014). In Drosophila, DUB module activity is regulated slightly differently than in yeast, in that Sgf11, e(y)2, and Nonstop without Sgf73 form a catalytically active complex (Mohan et al. 2014; Cloud et al. 2019). In fact, even in wild-type conditions, the DUB module (Sgf11 and Nonstop) associates with ~600 genomic regions that are not targeted by the canonical SAGA complex. Approximately half of these map near TSS and ~10% require ubiquitin protease activity, indicating that the Nonstop regulates gene expression independent of canonical SAGA (Li et al. 2017).

Other studies in S. cerevisiae suggest that Spt20 is also important for integration of the SAGA HAT module. spt20a mutants fail to integrate the HAT module, yet the HAT module exists as stable subcomplex (Lee et al. 2011). Spt20 and Ada2 assemble cotranslationally (Kassem et al. 2017), which raises the possibility that the SAGA core module already engages the HAT module in the cytoplasm. In humans, recombinant expression of all four HAT module subunits leads to the formation of a stable HAT subcomplex (Nguyen-Huynh et al. 2015). The same components form the Drosophila ADA complex (Soffers et al. 2019), which is similar to the yeast ADA complex, which preferentially acetylates H3K14, H3K18, and H2B (Eberharter et al. 1999; Grant et al. 1999) but lacks the yeast-specific Ada HAT components Ahc1 and Ahc2. The Drosophila nuclear complex does not associate with Spt20; thus, it is not the imported form of a Spt20–Ada2b cotranslational assembly complex (Soffers et al. 2019). The ADA complex localizes to sites that canonical SAGA does not occupy, and it has HAT activity not only on recombinant histones and core histones, but also on nucleosome substrates, suggesting that it can also acetylate chromatin in a cellular context. Loss of Drosophila Ada2b influences the expression of ~600 genes in larvae and ~900 genes in pupa (Zsindely et al. 2009; Pankotai et al. 2013). These observations raise questions about which genes depend on the metazoan ADA complex and which are regulated by SAGA, and also what advantage such redundant complexes may confer.

Conclusion

Combined biochemical, genetic, and structural studies have painted a picture of the versatile and diverse functions of the SAGA complex during gene regulation, starting from its role in transcription initiation to functions in elongation and RNA processing and export. Our understanding of its molecular mode of action keeps evolving with the identification of new subunits that account for additional functions, substrates, and knowledge about the roles of individual modules. Current questions are how the conformational flexibility directs function in vivo, and we are eagerly awaiting further structures of nucleosome-bound SAGA. These will help understand whether SAGA clamps one nucleosome at a time or can modify histone tails of adjacent nucleosomes. From here, further questions arise. Are pre-existing methylation marks directing levels of acetylation and ubiquitination at the same nucleosome and vice versa? Do multiple SAGA copies bind at different regions of a gene and exert different activities, and does this explain why some SAGA-associated marks are prevalent upstream of the transcription start site, yet others in the gene body? Can we account for these differences by enzymatic activities of isolated SAGA modules? Indeed, 23 yr after its discovery, the SAGA histone acetyltransferase complex keeps generating new paradigms for how the components of chromatin-modifying complexes function together and independently.

Acknowledgments

We thank the Workman laboratory members, Dr. Randal Halfmann, Dr. Ryan D. Mohan, Professor Patrick Cramer, and their group members for critical reading of the manuscript, discussion, and suggestions. These studies were supported by funds from the Stowers Institute and the National Institutes of General Medical Sciences grant R35GM118068 [to J.L.W.].
Boija A, Klein IA, Sabari BR, Dall’Agnese A, Coffey EL, Zamudio AV, Li CH, Shrinivas K, Manteiga JC, Hannett NM, et al. 2018. Transcription factors activate genes through the phase-separation capacity of their activation domains. Cell 175: 1842–1855.e16. doi:10.1016/j.cell.2018.10.042

Bolognesi B, Lorenzo Gotor N, Dhar R, Cirillo D, Baldrighi M, Tartaglia GG, Lehner B. 2016. A concentration-dependent liquid phase separation can cause toxicity upon increased protein expression. Cell Rep 16: 222–231. doi:10.1016/j.celrep.2016.05.076

Bonnet J, Wang CY, Baptista T, Vincent SD, Hsiao WC, Stierle M, Kao CF, Tora L, Devys D. 2014. The SAGA coactivator complex acts on the whole transcribed genome and is required for RNA polymerase II transcription. Genes Dev 28: 1999–2012. doi:10.1101/gad.250225.114

Boyer LA, Langer MR, Crowley KA, Tan S, Denu JM, Peterson CL. 2002. Essential role for the SANT domain in the functioning of multiple chromatin remodeling enzymes. Mol Cell 10: 935–942. doi:10.1016/S1097-2765(02)00664-2

Brand M, Leurent C, Mallouh V, Tora L, Schultz P. 1999a. Three-dimensional structures of the TAFII-containing complexes TFIIH and TFIID. Science 286: 2151–2153. doi:10.1126/science.286.5447.2151

Brand M, Yamamoto K, Staub A, Tora L. 1999b. Identification of TATA-binding protein–free TAF2-containing complex subunits suggests a role in nucleosome acetylation and signal transduction. J Biol Chem 274: 18285–18289. doi:10.1074/jbc.274.26.18285

Brown CE, Howe L, Sousa K, Alley SC, Carrozza MJ, Tan S, Workman JL. 2001. Recruitment of HAT complexes by direct activator interactions with the ATM-related Tra1 subunit. Science 292: 2333–2337. doi:10.1126/science.1060214

Brownell JE, Zhou J, Ranalli T, Kobayashi R, Edmondson DG, Roth SY, Allis CD. 1996. Tetrahydromena histone acetyltransferase A: a homolog to yeast Gcn5p linking histone acetylation and signal transduction. Cell 84: 843–851. doi:10.1016/S0092-8674(00)81063-6

Burley SK, Roeder RG. 1996. Biochemistry and structural biology of transcription factor IID [TFIID]. Annu Rev Biochem 65: 769–799. doi:10.1146/annurev.bi.65.070196.004005

Candau R, Berger SL. 1996. Structural and functional analysis of yeast putative adaptors. Evidence for an adaptor complex in vivo. J Biol Chem 271: 5237–5245. doi:10.1074/jbc.271.9.5237

Candau R, Zhou JX, Allis CD, Berger SL. 1997. Histone acetyltransferase activity and interaction with ADA2 are critical for Gcn5 function in vivo. EMBO J 16: 555-565. doi:10.1093/emboj/16.3.555

Carpentier AE, Memedula S, Plutz MJ, Belmont AS. 2005. Common effects of acidic activators on large-scale chromatin structure and transcription. Mol Cell 25: 958–968. doi:10.1016/j.mcb.2005.08.007

Cavusoglu N, Brand M, Tora L, Van Dorselaer A. 2003. Novel subunits of the TATA binding protein free TAFII-containing transcription complex identified by matrix-assisted laser desorption/ionization-time of flight mass spectrometry following one-dimensional gel electrophoresis. Proteomics 3: 217–223. doi:10.1002/pmc.200390030

Cheon Y, Kim H, Park K, Kim M, Lee D. 2020. Dynamic modules of the coactivator SAGA in eukaryotic transcription. Exp Mol Med 52: 991–1003. doi:10.1038/s41227-020-0463-4

Cheung ACM, Diaz-Santin LM. 2019. Share and share alike: the role of Tra1 from the SAGA and NuA4 coactivator complexes. Transcription 10: 37–43. doi:10.1080/21541264.2018.1530936

---

**References**

Antonova SV, Haffke M, Corradini E, Mikuciunas M, Low TY, Sigur L, van Es RM, Gupta K, Scheer E, Vos HR, et al. 2018. Chaperonin CCT checkpoint function in basal transcription factor TFIIH assembly. Nat Struct Mol Biol 25: 1119–1127. doi:10.1038/s41594-018-0156-z

Arents G, Burlingame RW, Wang BC, Love WE, Moudrianakis EN. 1991. The nucleosomal core histone octamer at 3.1 A resolution: a tripartite protein assembly and a left-handed superhelix. Proc Natl Acad Sci 88: 10148–10152. doi:10.1073/pnas.88.22.10148

Atanassov BS, Mohan RD, Lan X, Kuang X, Lu Y, Lin K, McIvor E, Li W, Zhang Y, Flores L, et al. 2016. ATXN7L3 and ENY2 coordinate activity of multiple H2B dequibutinases important for cellular proliferation and tumor growth. Mol Cell 62: 558–571. doi:10.1016/j.molcel.2016.03.030

Balasubramanian R, Pray-Grant MG, Selleck W, Grant PA, Tan S, Baptista T, Grunberg S, Minoungou N, Koster MJE, Timmers H, Hahn S, Devys D, Tora L, Karagiannis J, Brandl CJ. 2018. The pseuderogon L, van Es RM, Gupta K, Scheer E, Vos HR, et al. 2018. Chaperonin CCT checkpoint function in basal transcription factor TFIIH assembly. Nat Struct Mol Biol 25: 1119–1127. doi:10.1038/s41594-018-0156-z

Arents G, Burlingame RW, Wang BC, Love WE, Moudrianakis EN. 1991. The nucleosomal core histone octamer at 3.1 A resolution: a tripartite protein assembly and a left-handed superhelix. Proc Natl Acad Sci 88: 10148–10152. doi:10.1073/pnas.88.22.10148

Bian C, Xu C, Ruan J, Lee KK, Burke TL, Tempel W, Barsyte D, Li J, Wu M, Zhou BO, et al. 2011. Sgf29 binds histone H3K4me2/3 and is required for SAGA complex recruitment and histone H3 acetylation. EMBO J 30: 2829–2842. doi:10.1038/emboj.2011.193

Bieniossek C, Papai G, Schaffitzel C, Garzoni F, Chaillet M, Scheer E, Papadopoulos P, Tora L, Schultz P, Berger I. 2013. The architecture of human general transcription factor TFIID core complex. Nature 493: 699–702. doi:10.1038/nature11791

Birck C, Poch O, Romier C, Ruif M, Mengus G, Lavigne AC, Davidson I, Moras D. 1998. Human TAFII28 and TAFII18 interact through a histone fold encoded by atypical evolutionary conserved motifs also found in the SPT3 family. Cell 94: 239–249. doi:10.1016/S0092-8674(00)81423-3

Biswas D, Imbalzano AN, Eriksson P, Yu Y, Stillman DJ. 2004. Role for Nhp6p, Gcn5p, and the Swi/Snf complex in stimulating formation of the TATA-binding protein–TFIIB–DNA complex. Mol Cell Biol 24: 8312–8321. doi:10.1128/MCB.24.18.8312-8321.2004

---

**GENES & DEVELOPMENT**

1297
Claro, V., Thapa, A., Morales-Sosa, P., Miller, T.M., Miller, S.A., Holsaple, D., Gerhart, P.M., Mottaher, E., Jack, J.L., Leiva, E., et al. 2019. Ataxin-7 and non-stop coordinate SCAR protein levels, subcellular localization, and actin cytoskeleton organization. *Elife* 8: e49677. doi:10.7554/eLife.49677

Da, C., Lenkart, J., Zhao, K., Shiekhattar, R., Cairns, B.R., Marmoerstein, R. 2006. Structure and function of the SWIRM domain, a conserved protein module found in chromatin regulatory complexes. *Proc Natl Acad Sci* 103: 2057–2062. doi:10.1073/pnas.0510949103

Dahiri, R., Natarajan, K. 2018. Mutational analysis of TAF6 revealed the essential requirement of the histone-fold domain and the HEAT repeat domain for transcriptional activation. *FEBS* 285: 1491–1510. doi:10.1111/febs.14423

Daniel JA, Torok MS, Sun ZW, Schietz D, Allis CD, Yates JR III, Grant PA. 2004. Deubiquitination of histone H2B by a yeast acetyltransferase complex regulates transcription. *J Biol Chem* 279: 1867–1871. doi:10.1074/jbc.C300494200

Diaz-Santin LM, Lukoyanova N, Aciyan E, Cheung AC. 2017. Durso RJ, Fisher AK, Albright-Frey TJ, Reese JC. 2001. Analysis of Elías-Villalobos A, Fort P, Helmlinger D. 2019. New insights into Elías-Villalobos A, Toullec D, Faux C, Séveno M, Helmlinger D. 2019. Unit Tra1 at 3.7 angstrom resolution. *Genes Dev* 13: 2940–2945. doi:10.1101/gad.13.22.2940

Durand A, Bonnet J, Fournier M, Chavant V, Schultz P. 2014. Mapping the deubiquitination module within the SAGA complex. *Structure* 22: 1553–1559. doi:10.1016/j.str.2014.07.017

Durso RJ, Fisher AK, Albright-Frey TJ, Reese JC. 2001. Analysis of TAF90 mutants displaying allele-specific and broad defects in TAFII40 interacting with both a VP16 activation domain and the basal transcription factor TFIIB. *Cell* 75: 1837–1848. doi:10.1016/S0092-8674(00)50105-3

Eberharter A, Sterner DE, Schietz D, Hassan A, Yates JR III, Berger SL, Workman JL. 1999. The ADA complex is a distinct histone acetyltransferase complex in *Saccharomyces cerevisiae*. *Mol Cell Biol* 19: 6621–6631. doi:10.1128/MCB.19.10.6621

Eisenmann DM, Arndt KM, Ricupero SL, Rooney JW, Winston F. 1992. SPT3 interacts with TFIIH to allow normal transcription in *Saccharomyces cerevisiae*. *Genes Dev* 6: 1319–1331. doi:10.1101/gad.6.7.1319

Eisenmann DM, Chapon C, Roberts SM, Dollard C, Winston F. 1994. The *Saccharomyces cerevisiae* SPT8 gene encodes a very acidic protein that is functionally related to SPT3 and TATA-binding protein. *Genetics* 137: 647–657.

Elías-Villalobos A, Fort P, Helmlinger D. 2019. New insights into the evolutionary conservation of the sole PIKK pseudokinase Tra1/18RAP. *Biochem Soc Trans* 47: 1597–1608. doi:10.1042/BST20180496

Elías-Villalobos A, Toullec D, Faux C, Séveno M, Helmlinger D. 2019. Chaperone-mediated ordered assembly of the SAGA and NuA4 transcription co-activator complexes in yeast. *Nat Commun* 10: 5237. doi:10.1038/s41467-019-13243-w

Ellisdon AM, Jani D, Köhler A, Hurt E, Stewart M. 2010. Structural basis for the interaction between yeast Spt-Ada-Gcn5 acetyltransferase (SAGA) complex components Sgf11 and Sus1. *J Biol Chem* 285: 3850–3856. doi:10.1074/jbc.M109.070839

Evangelista FM, Maglott-Roth A, Sierle M, Brino L, Soutoglou E, Tora L. 2018. Transcription and mRNA export machineries SAGA and TREX-2 maintain monoubiquitinated H2B balance required for DNA repair. *Cell Biol* 217: 3382–3397. doi:10.1083/jcb.201803074

Fassler JS, Winston F. 1988. Isolation and analysis of a novel class of suppressor of Ty insertion mutations in *Saccharomyces cerevisiae*. *Genetics* 118: 203–212.

Ferreira ME, Hermann S, Prochasson P, Workman JL, Berndt KD, Wright AP. 2005. Mechanism of transcription factor recruitment by acidic activators. *J Biol Chem* 280: 21779–21784. doi:10.1074/jbc.M502627200

Fishburn J, Mohibullah N, Hahn S. 2005. Function of a eukaryotic transcription activator during the transcription cycle. *Mol Cell* 18: 369–378. doi:10.1016/j.molcel.2005.03.029

Fleming AB, Kao CF, Hillyer C, Pikaart M, Osley MA. 2008. H2B ubiquitylation plays a role in nucleosome dynamics during transcription elongation. *Mol Cell* 31: 57–66. doi:10.1016/j.molcel.2008.04.025

Gamer AM, Kim J, Roeder RG. 2009. The STAGA subunit ADA2b is an important regulator of human GCN5 catalysis. *Mol Cell Biol* 29: 266–280. doi:10.1128/MCB.00315-08

Gangloff YG, Romier C, Thuault S, Werten S, Davidson L 2001a. The histone fold is a key structural motif of transcription factor TFIIH. *Trends Biochem Sci* 26: 250–257. doi:10.1016/S0968-0004(00)01741-2

Gangloff YG, Sanders SL, Romier C, Kirschner D, Weil PA, Tora L, Davidson L. 2001b. Histone folds mediate selective heterodimerization of yeast TAFII55 with TFIIID components yTAFII7 and yTAFII65 and with SAGA component ySPT7. *Mol Cell Biol* 21: 1841–1853. doi:10.1128/MCB.21.5.1841-1853.2001

Ganshcroff LJ, Dollard C, Tan P, Winston F. 1995. The *Saccharomyces cerevisiae* SPT7 gene encodes a very acidic protein important for transcription in vivo. *Genetics* 139: 523–536.

Garcia-Oliver E, Garcia-Moliner O, Rodriguez-Navarro S. 2012. mRNA export and gene expression: the SAGA–TREX-2 connection. *Biochim Biophys Acta* 1819: 555–565. doi:10.1016/j.bbadis.2011.11.011

Georgakopoulos T, Thireos G. 1992. Two distinct yeast transcriptional activators require the function of the GCN5 protein to promote normal levels of transcription. *EMBO J* 11: 4145–4152. doi:10.1002/j.1607-1603.1992.tb05507.x

Georgieva S, Kirschner DB, Jagla T, Nabirockina E, Hanke S, Schenkel H, de Lorenzo C, Sinha P, Jagla K, Mechler B, et al. 2000. Two novel *Drosophila* TAFII56s have homology with human TAFII30 and are differentially regulated during development. *Mol Cell Biol* 20: 1639–1648. doi:10.1128/MCB.21.5.1639-1648.2000

Gödde JS, Nakatani Y, Wolffe AP. 1995. The amino-terminal tails of the core histones and the translational position of the TATA box determine TBP/TFIID association with nucleosomal DNA. *Nucleic Acids Res* 23: 4557–4564. doi:10.1093/nar/23.22.4557

Goodrich JA, Hoey T, Thut CJ, Admon A, Tjian R. 1993. *Drosophila* TAFI40 interacts with both a VP16 activation domain and the basal transcription factor TFIIB. *Cell* 75: 519–530. doi:10.1016/0092-8674(93)90386-5

Grant PA, Duggan L, Cote J, Roberts SM, Brownell JE, Candau R, Ohba R, Owen-Hughes T, Allis CD, Winston F, et al. 1997. Yeast Gcn5 functions in two multisubunit complexes to acetylate nucleosomal histones: characterization of an Ada complex and the SAGA (Spt/Ada) complex. *Genes Dev* 11: 1640–1650. doi:10.1101/gad.11.13.1640

Grant PA, Schietz D, Pry-Grant MG, Steger DJ, Reese JC, Yates JR III, Workman JL. 1998a. A subset of TAFII56s are integral components of the SAGA complex required for nucleosome acetylation and transcriptional stimulation. *Cell* 94: 45–53. doi:10.1016/S0092-8674(00)81220-9
Grant PA, Schieltz D, Pray-Grant MG, Yates JR III, Workman JL. 1998b. The ATM-related cofactor Tra1 is a component of the purified SAGA complex. Mol Cell 2: 863–867. doi:10.1016/ S1097-2765(00)80300-7

Grant PA, Sterner DE, Duggan LJ, Workman JL, Berger SL. 1998c. Expanded lysine acetylation specificity of Gcn5 in native complexes. J Biol Chem 273: 5900–5906. doi:10.1074/jbc.273.9.5895

Guarente L. 1995. Transcriptional activators in yeast and beyond. Trends Biochem Sci 20: 517–521. doi:10.1016/S0968-0004(00)89120-3

Guelman S, Sugarman T, Flores L, Swanson SK, Kiesecker CL, Kusch T, Anderson S, Yates JR III, Washburn MP, Ahmayer SM, et al. 2006a. Host cell factor and an uncharacterized SANT domain protein are stable components of ATAC, a novel dAda2A/dGcn5-containing histone acetyltransferase complex in Drosophila. Mol Cell Biol 26: 871–882. doi:10.1128/MCB.26.3.871-882.2006

Guelman S, Sugarman T, Flores L, Weake V, Swanson SK, Washburn MP, Ahmayer SM, Workman JL. 2006b. The essential gene wda encodes a WD40 repeat subunit of Drosophila SAGA required for histone H4 acetylation. Mol Cell Biol 26: 7178–7189. doi:10.1128/MCB.00130-06

Hall DB, Struhl K. 2002. The VP16 activation domain interacts with multiple transcriptional components as determined by protein-protein cross-linking in vivo. J Biol Chem 277: 46043–46050. doi:10.1074/jbc.M208911200

Han Y, Luo J, Ranish J, Hahn S. 2014. Architecture of the Saccharomyces cerevisiae SAGA transcription coactivator complex. EMBO J 33: 2534–2546. doi:10.15252/embj.201488638

Helmlinger D, Tora L. 2017. Sharing the SAGA. Trends Biochem Sci 42: 850–861. doi:10.1016/j.tibs.2017.09.001

Helmlinger D, Marguerat S, Villén J, Swaney DL, Gygi SP, Bähler J, Winston F. 2011. Tra1 has specific regulatory roles, rather than global functions, within the SAGA co-activator complex. EMBO J 30: 2834–2852. doi:10.1038/emboj.2011.181

Henry KW, Wyce A, Lo WS, Duggan LJ, Emre NC, Kao CF, Pillus LG, Shilatifard A, Osley MA, Berger SL. 2003. Transcriptional activation via sequential histone H2B ubiquitylation and deubiquitylation, mediated by SAGA-associated Ubp8. Genes Dev 17: 2648–2663. doi:10.1101/gad.114403

Hisatke K, Ohta T, Takada R, Guermah M, Horikoshi M, Nakatani Y, Roeder RG. 1995. Evolutionary conservation of human TATA-binding-polypeptide-associated factors TAFIIs1 and TAFII80 and interactions of TAFII80 with other TAFs and with general transcription factors. Proc Natl Acad Sci 92: 8195–8199. doi:10.1073/pnas.92.18.8195

Hoffmann A, Chiang CM, Oelgeschläger T, Xie X, Burley SK, Nakatani Y, Roeder RG. 1996. A histone octamer-like structure within TFIID. Nature 380: 356–359. doi:10.1038/380356a0

Horiiuchi J, Silverman N, Marcus GA, Guarente L. 1995. ADA3, a putative transcriptional adaptor, consists of two separable domains and interacts with ADA2 and GCN5 in a trimeric complex. Mol Cell Biol 15: 1203–1209. doi:10.1128/MCB.15.3.1203

Horiiuchi J, Silverman N, Piña B, Marcus GA, Guarente L. 1997. ADA1, a novel component of the ADA/GCN5 complex, has broader effects than GCN5, ADA2, or ADA3. Mol Cell Biol 17: 3220–3228. doi:10.1128/MCB.17.6.3220

Ingvarsdottr K, Krogan NJ, Emre NC, Wyce A, Thompson NJ, Emili A, Hughes TR, Greenblatt JF, Berger SL. 2005. H2B ubiquitin protease Ubp8 and Sgf11 constitute a discrete functional module within the Saccharomyces cerevisiae SAGA complex. Mol Cell Biol 25: 1162–1172. doi:10.1128/MCB.25.3.1162-1172.2005

Kassem S, Villanyi Z, Collart MA. 2017. Nost5-dependent co-translational assembly of Ada2 and Spt20 is essential for functional integrity of SAGA. Nucleic Acids Res 45: 1186–1199. doi:10.1093/nar/gkw1059

Kim JH, Saraf A, Flores L, Washburn M, Workman JL. 2010. Gen5 regulates the dissociation of SWI/SNF from chromatin by acetylation of Swi2/Snf2. Genes Dev 24: 2766–2771. doi:10.1101/gad.1979710

Klein J, Nolden M, Sanders SL, Kirchner J, Weil PA, Melcher K. 2003. Use of a genetically introduced cross-linker to identify interaction sites of acidic activators within native transcription factor IID and SAGA. J Biol Chem 278: 6779–6786. doi:10.1074/jbc.M212514200

Knutson BA, Hahn S. 2011. Domains of Tra1 important for activator recruitment and transcription coactivator functions of SAGA and NuA4 complexes. Mol Cell Biol 31: 818–831. doi:10.1128/MCB.00687-10

Köhler A, Pascual-Garcia P, Llopis A, Zapater M, Posas F, Hurt E, Rodriguez-Navarro S. 2006. The mRNA export factor Sus1 is involved in Spt/Ada/Gcn5’s acetyltransferase-mediated H2B deubiquitinylation through its interaction with Ubp8 and Sgf11. Mol Cell Biol 17: 4228–4236. doi:10.1091/mbc.e06-02-0098

Köhler A, Schneider M, Cabal GG, Nehrbass U, Hurt E. 2008. Yeast Ataxin-7 links histone deubiquitination with gene gating and mRNA export. Nat Cell Biol 10: 707–715. doi:10.1038/ncb1733

Köhler A, Zimmerman E, Schneider M, Hurt E, Zheng N. 2010. Structural basis for assembly and activation of the heterotetrameric SAGA histone H2B deubiquitinase module. Cell 141: 606–617. doi:10.1016/j.cell.2010.04.026

Kokubo T, Gong DW, Wootton JC, Horikoshi M, Roeder RG, Nakatani Y. 1994. Molecular cloning of Drosophila TFIID subunits. Nature 367: 484–487. doi:10.1038/367484a0

Kolesnikova O, Ben-Shem A, Luo J, Ranish J, Schultz P, Papai G. 2016. Molecular structure of promoter-bound yeast TFIID. Nat Commun 9: 4666. doi:10.1038/ncomms14167

Kotani T, Zhang X, Schiltz RL, Ogryzko VV, Howard T, Swanson MJ, Vassilev A, Zhang H, Yamaiuchi J, Howard BH, et al. 1998. TBP-associated factors in the PCAF histone acetylase complex. Cold Spring Harb Symp Quant Biol 63: 493–500. doi:10.1101/sq.98.196.493

Kuo MH, Brownell JE, Sobel RE, Ranalli TA, Cook RG, Edmondson DG, Roth SY, Allis CD. 1996. Transcription-linked acetylation by Gcn5 of histones H3 and H4 at specific lysines. Nature 383: 269–272. doi:10.1038/383269a0

Kusch T, Guelman S, Abmayr SM, Workman JL. 2003. Two Drosophila Ada2 homologues function in different multiprotein complexes. Mol Cell Biol 23: 3305–3319. doi:10.1128/MCB.23.9.3305-3319.2003

Lang G, Bonnet J, Umlauf D, Karmodyi K, Koffler J, Steierle M, Devys D, Tora L. 2011. The tightly controlled deubiquitination activity of the human SAGA complex differentially

The SAGA chromatin-modifying complex

GENES & DEVELOPMENT 1299
modifies distinct gene regulatory elements. *Mol Cell Biol* **31**: 3734–3744. doi:10.1128/MCB.05231-11

Laprade L, Rose D, Winston F. 2007. Characterization of new Spt3 and TATA-binding protein mutants of *Saccharomyces cerevisiae*: Spt3–TBP allele-specific interactions and bypass of Spt8. *Genetics* **177**: 2007–2017. doi:10.1534/genetics.107.081976

Larschan E, Winston F. 2001. The *S. cerevisiae* SAGA complex functions in vivo as a coactivator for transcriptional activation by Gal4. *Genes Dev* **15**: 1946–1956. doi:10.1101/gad.911501

Lee TJ, Causton HC, Holstege FC, Shen WC, Hannett N, Jennings EG, Winston F, Green MR, Young RA. 2000. Redundant roles for the TFIIID and SAGA complexes in global transcription. *Nature* **405**: 701–704. doi:10.1038/35015104

Lee KK, Flores L, Swanson SK, Washburn MP, Workman JL. 2005. The deubiquitylation activity of Ubp8 is dependent upon Sgf11 and its association with the SAGA complex. *Mol Cell Biol* **25**: 1173–1182. doi:10.1128/MCB.25.3.1173-1182.2005

Lee KK, Swanson SK, Flores L, Washburn MP, Workman JL. 2009. Yeast Sgf73/Ataxin-7 serves to anchor the deubiquitination module into both SAGA and Slik(SALSA) HAT complexes. *Epigenetics Chromatin* **2**: 2. doi:10.1186/1756-8935-2-2

Leurent C, Sanders S, Ruhlmann C, Mallouh V, Weil PA, Kirschner DB, Tora L, Schultz P. 2002. Mapping histone fold TAFs within yeast TFIID. *EMBO J* **21**: 4324–4334. doi:10.1093/emboj/cdf342

Li H, Cuenin C, Murr R, Wang ZQ, Herczy J. 2004. HAT cofactor Trtap regulates the mitotic checkpoint by modulation of Mad1 and Mad2 expression. *EMBO J* **23**: 4824–4834. doi:10.1038/sj.emboj.7600479

Li B, Carey M, Workman JL. 2007. The role of chromatin during transcription. *Cell* **128**: 707–719. doi:10.1016/j.cell.2007.01.015

Li J, Xue X, Ruan J, Wu M, Zhu Z, Zang J. 2010. Cloning, purification, crystallization and preliminary crystallographic analysis of the tandem tudor domain of Sgf29 from *Saccharomyces cerevisiae*. *Acta Crystallogr Sect F Struct Biol Cryst Commun* **66**: 902–904. doi:10.1107/S1744309110016726

Li X, Seidel CW, Szerszen LT, Lange JF, Workman JL, A bmayr SM. 2017. Enzymatic modules of the SAGA chromatin-modifying complex play distinct roles in *Drosophila* gene expression and development. *Genes Dev* **31**: 1588–1600. doi:10.1101/gad.300988.117

Lim S, Kwak J, Kim M, Lee D. 2013. Separation of a functional deubiquitylation module from the SAGA complex by the proteasome regulatory particle. *Nat Commun* **4**: 2641. doi:10.1038/ncomms3641

Lin L, Chamberlain L, Zhu LJ, Green MR. 2012. Analysis of Gal4-directed transcription activation using Tra1 mutants selectively defective for interaction with Gal4. *Proc Natl Acad Sci U S A* **109**: 1997–2002. doi:10.1073/pnas.1116340109

Liu X, Tesfai J, Evrard YA, Dent SY, Martinez E. 2003. e-Myc transformation domain recruits the human SAGA complex and requires TRRAP and GCN5 acetylase activity for transformation activation. *J Biol Chem* **278**: 20405–20412. doi:10.1074/jbc.M211795200

Luger K, Mäder AW, Richmond RK, Sargent DF, Richmond TJ. 1997. Crystal structure of the nucleosome core particle at 2.8 Å resolution. *Nature* **389**: 251–260. doi:10.1038/38444

Marcus GA, Silverman N, Berger SL, Horiiuchi J, Guarente L. 1994. Functional similarity and physical association between GCN5 and ADA2: putative transcriptional adaptors. *EMBO J* **13**: 4807–4815. doi:10.1002/j.1460-2075.1994.tb06806.x

Marcus GA, Horiiuchi J, Silverman N, Guarente L. 1996. ADA5/SPT20 links the ADA and SPT genes, which are involved in yeast transcription. *Mol Cell Biol* **16**: 3197–3205. doi:10.1128/MCB.16.6.3197

Martinez E, Kundu TK, Fu J, Roeder RG. 1998. A human SPT3–TAF15–GCN5-L acetylase complex distinct from transcription factor IID. *J Biol Chem* **273**: 23781–23785. doi:10.1074/jbc.273.37.23781

Martinez E, Pallan VB, Tjernberg A, Lymar ES, Camper AM, Kundu TK, Chait BT, Roeder RG. 2001. Human STAGA complex is a chromatin-acetylating transcription coactivator that interacts with pre-mRNA splicing and DNA damage-binding factors in vivo. *Mol Cell Biol* **21**: 6782–6795. doi:10.1128/MCB.21.20.6782-6795.2001

McMahon SB, Van Buskirk HA, Dugan KA, Copeland TD, Cole MD. 1998. The novel ATM-related protein TRRAP is an essential cofactor for the e-Myc and E2F oncoproteins. *Cell* **94**: 363–374. doi:10.1016/S0092-8674(00)81479-8

McMahon SJ, Pray-Grant MG, Schielitz D, Yates JR III, Grant PA. 2005. Polyglutamine-expanded spinocerebellar ataxia-7 protein disrupts normal SAGA and SLIK histone acetyltransferase activity. *Proc Natl Acad Sci U S A* **102**: 8478–8482. doi:10.1073/pnas.0503493102

Mengus G, May M, Iacq X, Staub A, Tora L, Chambon P, Davidson L. 1995. Cloning and characterization of hTAFII18, hTAFII20 and hTAFII28: three subunits of the human transcription factor TFIIH. *EMBO J* **14**: 1520–1531. doi:10.1002/j.1460-2075.1995.tb07138.x

Mohan RD, Dialynas G, Weake VM, Liu J, Martin-Brown S, Flores L, Washburn MP, Workman JL, A bmayr SM. 2014. Loss of *Drosophila* Ataxin-7, a SAGA subunit, reduces H2B ubiquitination and leads to neural and retinal degeneration. *Genes Dev* **28**: 259–272. doi:10.1101/gad.225151.113

Morgan MT, Haj-Yahya M, Ringel AE, Bandi P, Brik A, Wolberger C. 2016. Structural basis for histone H2B deubiquitination by the SAGA DUB module. *Science* **351**: 725–728. doi:10.1126/science.aac5681

Muratoglu S, Georgieva S, Papai G, Scheer E, En ünül ü I, Komonyi C, Cserpán I, Lebedeva L, Nabirochkina E, Udvardy A, et al. 2003. Two different *Drosophila* ADA2 homologues are present in distinct GCN5 histone acetyltransferase-containing complexes. *Mol Cell Biol* **23**: 306–321. doi:10.1128/MCB.23.1.306-321.2003

Nakatani Y, Bagby S, Ikura M. 1996. The histone folds in transcription factor TFIIH. *J Biol Chem* **271**: 6575–6578. doi:10.1074/jbc.271.12.6575

Natarajan K, Jackson BM, Rhein Hebbeus AG. 1998. γ-TAFI61 has a general role in RNA polymerase II transcription and is required by Gcn4p to recruit the SAGA coactivator complex. *Mol Cell* **2**: 683–692. doi:10.1016/S1079-7595(00)80166-5

Nguyen-Huynh NT, Sharov G, Potel C, Fichter P, Trowitzsch S, Leitz Wagn N. 2011. Chemical cross-linking and mass spectrometry to determine the subunit interaction network in a recombinant human SAGA HAT subcomplex. *Protein Sci* **20**: 1232–1246. doi:10.1002/pro.2676

Ogryzko VV, Kotani T, Zhang X, Schiltz RL, Howard T, Yang XJ, Howard BH, Qin J, Nakatani Y. 1998. Histone-like TAFs within the PCAF histone acetylase complex. *Cell* **94**: 35–44. doi:10.1186/S0002-8674(00)81219-2
Smith ER, Belote JM, Schiltz RL, Yang XJ, Moore PA, Berger SL, Nakatani Y, Allis CD. 1998. Cloning of Drosophila GCN5: conserved features among metazoan GCN5 family members. *Nucleic Acids Res* **26**:2948–2954. doi:10.1093/nar/26.12.2948

Soffers JHM, Li X, Saraf A, Seidel CW, Florens L, Washburn MP, Abmayr SM, Workman JL. 2019. Characterization of a metazoan ADA acetyltransferase complex. *Nucleic Acids Res* **47**:3383–3394. doi:10.1093/nar/gkz042

Stegeaman R, Spearecker PJ, Swanson SK, Stephenson R, Florens L, Washburn MP, Weake VM. 2016. The splicingosomal protein SF3B5 is a novel component of Drosophila SAGA that functions in gene expression independent of splicing. *J Mol Biol* **428**:3632–3649. doi:10.1016/j.jmb.2016.05.009

Sterner DE, Grant PA, Roberts SM, Duggan LJ, Belotserkovskaya R, Pacella LA, Winston F, Workman JL, Berger SL. 1999. Functional organization of the yeast SAGA complex: distinct components involved in structural integrity, nucleosome acetylation, and TATA-binding protein interaction. *Mol Cell Biol* **19**:86–98. doi:10.1128/MCB.19.1.86

Sterner DE, Wang X, Bloom MH, Simon GM, Berger SL. 2002. The SANT domain of Ada2 is required for normal acetylation of histones by the yeast SAGA complex. *J Biol Chem* **277**:8179–8186. doi:10.1074/jbc.M108601200

Suganuma T, Gutierrez JL, Li B, Florens L, Swanson SK, Washburn MP, Abmayr SM, Workman JL. 2008. ATAC is a double histone acetyltransferase complex that stimulates nucleosome sliding. *Nat Struct Mol Biol* **15**:364–372. doi:10.1038/nsmb.1397

Sun J, Paduch M, Kim SA, Kramer RM, Barrios AF, Lu V, Luke J, Usatyuk S, Kossiakoff AA, Tan S. 2018. Structural basis for activation of SAGA histone acetyltransferase Gcn5 by partner subunit Ada2. *Proc Natl Acad Sci USA* **115**:10010–10015. doi:10.1073/pnas.1805343115

Syntichaki P, Theiroes G. 1998. The Gcn5-Ada complex potentiates the histone acetyltransferase activity of Gcn5. *J Biol Chem* **273**:24414–24419. doi:10.1074/jbc.273.38.24414

Thomas MC, Chiang CM. 2006. The general transcription machinery and general cofactors. *Crit Rev Biochem Mol Biol* **41**:105–178. doi:10.1080/10409230600648736

Timmers HTM. 2020. SAGA and TFIDD: friends of TBP drifting apart. *Biochim Biophys Acta Genet Regul Mech* doi:10.1016/j.bbrgm.2020.194604

Torres-Zelada EF, Stephenson RE, Alpsoy A, Anderson BD, Swanson SK, Florens L, Dykhuizen EC, Washburn MP, Weake VM. 2019. The *Drosophila* Dbf4 ortholog Chiffon forms a complex with Gcn5 that is necessary for histone acetylation and viability. *J Cell Sci* **132**:jes214072. doi:10.1242/jcs.214072

Tran HG, Steger DJ, Iyer VR, Johnson AD. 2000. The chromo domain protein chdlp from budding yeast is an ATP-dependent chromatin-modifying factor. *EMBO J* **19**:2323–2331. doi:10.1093/emboj/19.10.2323

Tuttle LM, Pacheco D, Warfield L, Luo J, Ranish J, Hahn S, Klevit RE. 2018. Gcn4-mediator specificity is mediated by a large and dynamic fuzzy protein–protein complex. *Cell Rep* **22**:3251–3264. doi:10.1016/j.celrep.2018.02.097

Uesugi M, Nyangule O, Lu H, Levine AJ, Verdine GL. 1997. Induced a helix in the VP16 activation domain upon binding to a human TAF. *Science* **277**:1310–1313. doi:10.1126/science.77530.1310

Vermeulen M, Eberl HC, Matarese F, Marks H, Denissov S, Butter F, Lee KK, Olsen JV, Hyman AA, Stunnenberg HG, et al. 2010. Quantitative interaction proteomics and genome-wide profiling of epigenetic histone marks and their readers. *Cell* **142**:967–980. doi:10.1016/j.cell.2010.08.020

Wang L, Mizzen C, Ying C, Candau R, Barlev N, Brownell J, Allis CD, Berger SL. 1997. Histone acetyltransferase activity is conserved between yeast and human GCN5 and is required for complementation of growth and transcriptional activation. *Mol Cell Biol* **17**:519–527. doi:10.1128/MCB.17.1.519

Wang H, Dienemann C, Stutzer A, Urlaub H, Cheung ACM, Cramer P. 2020. Structure of the transcription coactivator SAGA. *Nature* **577**:717–720. doi:10.1038/s41586-020-1933-5

Warfield L, Ranish JA, Hahn S. 2004. Positive and negative functions of the SAGA complex mediated through interaction of Spt8 with TBP and the N-terminal domain of TFIIA. *Genes Dev* **18**:1022–1034. doi:10.1101/gad.1192204

Weake VM, Swanson SK, Mushegian A, Florens L, Washburn MP, Abmayr SM, Workman JL. 2009. A novel histone fold domain-containing protein that replaces TAF6 in Drosophila SAGA is required for SAGA-dependent gene expression. *Genes Dev* **23**:2818–2823. doi:10.1101/gad.1846409

Weinzierl RO, Ruppert S, Dylacht BD, Tanese N, Tjian R. 1993. Cloning and expression of *Drosophila* TAFII60 and human TAFI70 reveal conserved interactions with other subunits of TFIIID. *EMBO J* **12**:5303–5309. doi:10.1002/j.1460-2075.1993.tb06226.x

Wieczorek E, Brand M, Jacq X, Tora L. 1998. Function of TAFII-containing complex without TBP in transcription by RNA polymerase II. *Nature* **393**:187–191. doi:10.1038/30283

Winston F, Carlson M. 1992. Yeast SNF/SWI transcriptional activators and the SPT/SIN chromatin connection. *Trends Genet* **8**:387–391. doi:10.1016/0168-9525(92)90300-S

Winston F, Dollard C, Malone EA, Clare J, Kapakos IG, Farabaugh P, Minehart PL. 1987. Three genes are required for trans-activation of Ty transcription in yeast. *Genetics* **115**:649–656.

Wu PY, Winston F. 2002. Analysis of Spt7 function in the *Saccharomyces cerevisiae* SAGA coactivator complex. *Mol Cell Biol* **22**:5367–5379. doi:10.1128/MCB.22.15.5367-5379.2002

Wu PY, Ruhlmann C, Winston F, Schultz P. 2004. Molecular architecture of the *S. cerevisiae* SAGA complex. *Mol Cell* **15**:199–208. doi:10.1016/j.molcel.2004.06.005

Xie X, Kobuku T, Cohen SL, Mirza UA, Hoffmann A, Chait BT, Xu C, Min J. 2011. Structure and function of WD40 domain containing protein that replaces TAF6 in *Drosophila* SAGA. *EMBO J* **30**:391–398. doi:10.1038/emboj.2010.375

Xu YZ, Quercy CC. 2007. Competition between the ATPase Prp5 and branch region-U2 snRNA pairing modulates the fidelity of spliceosome assembly. *Mol Cell* **28**:838–849. doi:10.1016/j.molcel.2007.09.022

Xu C, Min J. 2011. Structure and function of WD40 domain proteins. *Protein Cell* **2**:202–214. doi:10.1016/s1323-0111-2018-1

Xu W, Edmondson DG, Roth SY. 1998. Mammalian GCN5 and P/CAF acetyltransferases have homologous amino-terminal domains important for recognition of nucleosomal substrates. *Mol Cell Biol* **18**:5659–5669. doi:10.1128/MCB.18.10.5659

Yang XJ, Ogryzko VV, Nishikawa J, Howard BH, Nakatani Y, Brownell J. 1996. A p300/CBP-associated factor that competes with the adenoviral oncprotein E1A. *Nature* **382**:319–324. doi:10.1038/382319a0

Zhang N, Ichikawa W, Faiola F, Lo SY, Liu X, Martinez E. 2014. MYC interacts with the human STAGA coactivator complex via multivalent contacts with the GCN5 and TRRAP
Molecular architecture of the human 17S U2 snRNP. *Nature* **583**: 310–313. doi:10.1038/s41586-020-2344-3

Zsindely N, Pankotai T, Ujfaludi Z, Lakatos D, Komonyi O, Bodai L, Tora L, Boros IM. 2009. The loss of histone H3 lysine 9 acetylation due to dSAGA-specific dAda2b mutation influences the expression of only a small subset of genes. *Nucleic Acids Res* **37**: 6665–6680. doi:10.1093/nar/gkp722