Structural Basis for the Interaction between Yeast Spt-Ada-Gcn5 Acetyltransferase (SAGA) Complex Components Sgf11 and Sus1

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Sus1 is a central component of the yeast gene gating machinery, the process by which actively transcribing genes such as GAL1 become associated with nuclear pore complexes. Sus1 is a component of both the SAGA transcriptional co-activator complex and the TREX-2 complex that binds to nuclear pore complexes. TREX-2 contains two Sus1 chains that have an articulated helical hairpin fold, enabling them to wrap around an extended α-helix in Sac3, following a helical hydrophobic stripe. In SAGA, Sus1 binds to Sgf11 and has been proposed to provide a link between SAGA and TREX-2. We present here the crystal structure of the complex between Sus1 and the N-terminal region of Sgf11 that forms an extended α-helix around which Sus1 wraps in a manner that shares some similarities with the Sus1-Sac3 interface in TREX-2. However, the Sus1-binding site on Sgf11 is somewhat shorter than on Sac3 and is based on a narrower hydrophobic stripe. Engineered mutants that disrupt the Sgf11-Sus1 interaction in vitro confirm the importance of the hydrophobic helical stripe in molecular recognition. Helix α1 of the Sus1-articulated hairpin does not bind directly to Sgf11 and adopts a wide range of conformations within and between crystal forms, consistent with the presence of a flexible hinge and also with results from previous extensive mutagenesis studies (Klöckner, C., Schneider, M., Lutz, S., Jani, D., Kressler, D., Stewart, M., Hurt, E., and Köhler, A. (2009) J. Biol. Chem. 284, 12049–12056). A single Sus1 molecule cannot bind Sgf11 and Sac3 simultaneously and this, combined with the structure of the Sus1-Sgf11 complex, indicates that Sus1 forms separate subcomplexes within SAGA and TREX-2.

There is an emerging consensus that the different steps of the gene expression pathway are tightly coupled and show a high degree of interdependence (2–5). Integration of the steps that lead from transcription to mRNA nuclear export (including splicing, 5’ cap addition, and polyadenylation) relies on a complex network of interactions between activated genes, processing factors and the nuclear pore complex (NPC).3 For a subset of genes, this integration is achieved by tethering actively transcribed genes to the NPC, a process known as “gene gating” (6), in which the small nuclear protein, Sus1, is a central component (7–10). Sus1 is part of both the 2-MDa Spt-Ada-Gcn5 acetyltransferase (SAGA) complex, that is a co-activator for transcription by RNA polymerase II, and the TREX-2 (transcription and export-2) complex that tethers SAGA to NPCs (reviewed in Ref. 11). The SAGA complex possesses histone acetyltransferase and deubiquitination activities and is bound to chromatin during active transcription (12). The TREX-2 complex contains Sac3, Thp1, Sem1, and Cdc31 in addition to Sus1 and has roles in both nuclear mRNA export and transcription elongation (9, 11, 13–16). The TREX-2 complex binds the mRNA export factors Mex67 and Mtr2 as well as NPCs (13) and, by interacting with both SAGA and NPCs, facilitates gene gating (9). It has been proposed that gated genes may benefit from enhanced or optimal rates of messenger synthesis compared with genes residing in an intranuclear position (17), and this may be of particular importance in producing a rapid cellular response to stress, consistent with the involvement of a significant proportion of SAGA regulated genes in stress response (18).

Yeast SAGA is a large 2-MDa complex that functions as a transcriptional co-activator for a number of RNA polymerase II dependent genes (12, 19). The Gcn5 histone-acetyltransferase forms a submodule with multiple Ada proteins that regulate gene expression by acetylation of histones H3 and H2B (20–22). The SAGA complex also contains a second enzymatic activity through the deubiquitinating (DUB) module formed by the ubiquitin-specific protease, Ubp8, together with SAGA subunits Sgf73, Sgf11, and Sus1 (23, 24). The complex deubiquitinates Lys123 on histone H2B that has been shown to have downstream effects on the methylation state of histone H3 and a regulatory role in gene activation (25, 26). Within the yeast SAGA complex, Sgf11 binds Sus1 directly, forming a stable dimer (23). However, both the Sgf11/Sus1 heterodimer and Ubp8 are required for all three proteins to bind to Sgf73 (23, 24). Analogous interactions are found between Sgf11, Sus1, and Ubp8 homologues in both Drosophila and human cells (27–29).

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3 The abbreviations used are: NPC, nuclear pore complex; SAGA, Spt-Ada-Gcn5 acetyltransferase; DUB, deubiquitinating; GST, glutathione S-transferase.
Sgf11 is also associated with mRNA biogenesis during transcription elongation, being observed at coding regions in a SAGA-dependent manner (30). This is consistent with the presence of SAGA subunits within both open reading frames and at promoter regions (31). Histone deubiquitination and Ubp8 catalytic activity appear to be required for the association of actively transcribing genes with NPCs during gene gating (24). Although the precise details of the arrangement of the individual proteins within the SAGA DUB module remains to be established, the interaction between Sgf11 and Sus1 is necessary for its integrity (11, 23, 29).

Within the TREX-2 complex, residues 723–805 of Sac3 (the "CID" domain) bind two Sus1 chains and one chain of the calmodulin-like centrin, Cdc31. The crystal structure of this complex (10) shows that the Sac3 CID region adopts an extended α-helical conformation about which Sus1 and Cdc31 wrap. Both Sus1 chains have an articulated helical hairpin fold that is based on five α-helical segments linked by putative hinges. This conformation has been proposed to enable them to wrap around the Sac3 helix, like fingers gripping a thin rod (10). The binding interface contains few hydrogen bonds or polar interactions and is based on a hydrophobic stripe that winds around the Sac3 helix. Both Sus1 molecules (designated Sus1A and Sus1B) bind to stripes within the Sac3 CID region. In each case, the hydrophobic stripe is generated by a four-residue sequence repeat in which the first two residues are Phe, Tyr, Ile, Leu, or Met or have a side chain containing a considerable hydrophobic portion, such as Arg or Glu. In both Sus1-binding sites on Sac3, the binding motif is ~25-residues-long. Both Sus1 chains and Cdc31 are required for optimal NPC association of TREX-2 (10, 24).

Because Sus1 is found in both SAGA and TREX-2, it was initially proposed that Sus1 may physically bridge the two complexes (7). More recently, a more dynamic role for Sus1 in linking the two complexes has been proposed, whereby the competitive exchange of Sus1 molecules between SAGA and TREX-2 would serve to both physically link the complexes and modulate their function (1). Alternatively, the presence of Sus1 in both complexes may be coincidental, and this protein may have mechanistically separate roles in SAGA and TREX-2. Other proteins may also be important in linking the two complexes and, for example, deletion of the SAGA component Sgf73 alters the association of Sus1 with the TREX-2 complex (24, 30). Sgf73 also plays a role in recruiting Sac3 and Thp1 to SAGA, and it has been suggested that Sgf73 may alter the TREX-2 component Sac3 allowing for efficient TREX-2 assembly (24). Distinguishing between these different putative functions of Sgf11 and Sus1 has been difficult because it was not known how Sus1 binds to Sgf11 (11, 32). Extensive mutagenesis studies of Sus1 showed that, although it was possible to generate mutants in which Sac3 binding was lost while Sgf11 binding was retained, it was not possible to generate mutants in which Sgf11 binding was lost while Sac3 binding was retained (1). These mutagenesis results indicated that Sus1 binding to these two partners was in some way different but did not indicate what these differences were.

Here, we present the crystal structure of the N-terminal region of Sgf11 bound to Sus1, its direct binding partner in the SAGA complex. Sgf11 forms an extended α-helix around which Sus1 wraps in a manner that has some similarities to the way in which it binds to Sac3 in the Sac3-Cdc31-Sus1 complex. However, the Sus1-binding site on Sgf11 is somewhat shorter than on Sac3 and is based on a narrower hydrophobic stripe. As a consequence, helix α-1 of the Sus1-articulated hairpin fold does not bind directly to Sgf11 and, consistent with the presence of a flexible hinge between it and the next helix (α2), takes up a remarkably wide range of conformations within and between crystal forms. We have engineered mutants based on the structure of the complex that disrupt the Sgf11-Sus1 interaction and have used these to confirm the importance of the hydrophobic stripe in molecular recognition.

**EXPERIMENTAL PROCEDURES**

**Protein Expression and Purification**—Full-length and truncated Sgf11 variants were generated by PCR using yeast genomic DNA (Novagen; Beston, UK) and cloned into pGEX-TEV (33), a modified version of pGEX-4T-1 (GE Healthcare) in which the thrombin site has been replaced with a TEV protease site. Sus1 cDNA was cloned into an untagged expression vector, pNMTK, a modified pOPT vector (34) with kanamycin resistance. Sgf11 Ala insertion mutations were introduced using the QuickChange site-directed mutagenesis kit (Stratagene). The cloning of Cdc31 and Sac3 variants has been described elsewhere (10).

For crystallography, GST-Sgf11 constructs (amino acids 1–33 and 7–33) were coexpressed with untagged Sus1 in BL21 (DE3) CodonPlus RIL cells in ZYM-5052 autoinducing medium (35) at 20 °C. Cells were lysed by high pressure cavitation (10–15k pounds per square inch) in 50 mM Tris-HCl (pH 8.0), 25% w/v sucrose, 1 mM EDTA, and 1 mM phenylmethylsulphonyl fluoride. Complete EDTA-free protease inhibitor mixture (Roche) was added to the lysed cells. Cells were clarified by centrifugation, filtered through a 0.45-μm membrane, and bound to glutathione-Sepharose 4B resin (GE Healthcare) for 1 h at 4 °C. The resin was washed with 500 ml of 50 mM Tris-HCl (pH 8.0), 150 mM NaCl, 1 mM EDTA, and 1 mM dithiothreitol to remove nonspecifically bound proteins, and the Sgf11-Sus1 complex released from the GST tag by overnight incubation with 100 μg of His-TEV protease (S219V mutant; (36)). The complex was further purified on a HiLoad Superdex 75 26/60 prep-grade column (GE Healthcare) in 20 mM Tris-HCl (pH 8.0), 50 mM NaCl, 1 mM EDTA, and 1 mM dithiothreitol.

**Crystallization and Data Collection**—Crystals were grown at 19 °C by hanging drop vapor diffusion in either 3.2 M sodium formate (Sgf11-(7–33)-Sus1) or 1.1 M Na₂HPO₄ and 20 mM K₂HPO₄ (Sgf11-(1–33)-Sus1). Crystals of Sgf11-(7–33)-Sus1 required no cryoprotectant; however, crystals of Sgf11-(1–33)-Sus1 were exposed to mother liquor containing 30% (w/v) sucrose before flash cooling in liquid nitrogen prior to data collection. Crystals of Sgf11-(1–33)-Sus1 were also dehydrated by vapor diffusion, by adding increasing amounts of sucrose into the mother liquor for up to a week before flash freezing. Crystallographic data were collected either in-house using a Rigaku FR-E+ SuperBright generator equipped with Osmic...
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Table 1

| Crystal data                | 7–33 | 1–33 |
|-----------------------------|------|------|
| Sgf11 fragment              |      |      |
| Space group                 |      |      |
| Unit cell dimensions (Å)    |      |      |
| a                           | 75.15| 68.67|
| b                           | 75.15| 68.67|
| c                           | 197.37| 232.86|

Data collection
- Wavelength (Å): 1.54059
- Resolution range (Å): 25.2–1.54
- Total observations\(^a\): 503,521
- Unique observations\(^a\): 36,176
- Completeness (%): 98.3 (95.3)
- Multiplicity: 13.9 (7.2)
- \(R_{	ext{merge}}\) (%): 1.5 (29.2)
- \(R_{	ext{o}}\) (%): 5.9 (75.7)
- Mean \(I/|\sigma(I)|\): 26.2 (2.4)

Reconstruction
- \(R_{	ext{cryst}}/R_{	ext{free}}\): 19.0/23.7
- Bond length r.m.s.d. (Å): 0.6 (0.62)
- MolProbity score/percentile: 1.41/99

Ramachandran plot (%)
- Favourable: 99.5
- Allowed: 0.5
- Forbidden: 0

PDB accession code\(^c\): 3KIK, 3KJL

\(^a\) Parentheses refer to final resolution shell.
\(^b\) r.m.s.d., root mean square deviation.
\(^c\) PDB, Protein Data Bank.

RESULTS AND DISCUSSION

Sus1 Wraps around a Helix Formed by Residues 8–30 of Sgf11—In contrast to Sac3, each Sus1 chain binds only a single Sgf11 chain (10). Inspection of the Sgf11 sequence (24, 29) indicated the presence of a putative CCCH zinc finger domain at its C terminus (residues \(\sim 65–99\)). Deletion mutants indicated that Sus1 binding involved primarily the N terminus of Sgf11 (residues 1–33; Fig. 1). Crystals suitable for high resolution structure determination were obtained with Sgf11 fragments comprising residues 1–33 and 7–33. The crystals containing Sgf11 residues 1–33 showed anisotropic diffraction, whereas those with 20 mM Tris-HCl (pH 8.0), 200 mM NaCl, 1 mM EDTA, and 1 mM dithiothreitol. Whereas resin from the full-length Sgf11 variants was washed with 20 mM Tris-HCl (pH 8.0), 200 mM NaCl, 10 \(\mu\)M ZnCl\(_2\), and 5 \(\mu\)M \(\beta\)-mercaptoethanol. Samples were analyzed by SDS-PAGE.

Binding studies demonstrating the mutually exclusive binding of Sus1 to Sac3 or Sgf11 involved immobilizing coexpressed GST-Sac3-Cdc31 complexes onto glutathione 4B resin from clarified bacterial lysates. After extensive washing, bound material was analyzed by SDS-PAGE.

**FIGURE 1.** An N-terminal region of Sgf11 comprising residues 7–33 is sufficient for binding to Sus1. GST fusions of Sgf11 and a series of fragments were coexpressed with Sus1 in *Escherichia coli*, and complexes were immobilized on glutathione-Sepharose 4B resin from clarified lysates. After extensive washing, bound material was analyzed by SDS-PAGE.
containing residues 7–33 diffracted to 2.1 Å resolution and were solved first using molecular replacement based on a model of residues 21–90 of Sus1 from the Cdc31-Sac3-Sus1 complex (10). The structure obtained in this way was then used to obtain a structural model for the complex between Sus1 and Sgf11 (10). The structure obtained in this way was then used to obtain a structural model for the complex between Sus1 and Sgf11. Both crystal forms had perfect merohedral twinning and the asymmetric unit of each contained four copies of the Sgf11-Sus1 complex. In each case, this helix had been rotated by almost 180° between different copies. Moreover, the structural integrity of this region of Sus1 was reduced in the Sgf11 complex, and often, the density for the flexibility of the hinge between helices 2–5, with little contact with helix α1 and was thus less extensive than in Sac3 (Figs. 2 and 3). However, in the DUB complex, Sus1 helix α1 may be involved with interactions with other components such as Sgf73 and/or Ubp8. In contrast to the interaction with Sac3, Sus1 does not encircle the Sgf11 helix completely and instead leaves one side exposed.

There was a striking variation in the position of Sus1 helix α1 between Sgf11-Sus1 complexes within and between crystals (Fig. 4 and supplemental Movie S3) so that in the most extreme case, this helix had been rotated by almost 180° between different copies. Moreover, the structural integrity of this region of Sus1 was reduced in the Sgf11 complex, and often, the density in this region was less clear than in other regions of the structure. This remarkable structural plasticity provided direct evidence for the flexibility of the hinge between helices α1 and -2.

The structure of the Sgf11-Sus1 interface and comparison with Sus1 bound to Sac3 is consistent with the results obtained from a mutagenesis study of Sus1 (1), in which it was possible to obtain mutants that retained affinity for Sgf11, whereas affinity for Sac3 was lost. The predominantly hydrophobic nature of the Sgf11-Sus1 interface was consistent with the failure of mutagenesis of charged residues in Sus1 (mutants sus1-1 to sus1-9) to impair the interaction with either Sac3 or Sgf11 (1). The lack of contact between Sgf11 and the N and C termini of Sus1 seen in the crystal structures of the complex was consistent with the failure of deletions of Sus1 residues 1–10 or 91–96 to alter its binding to Sgf11. Mutations of Gly37 and Trp38 to Ala (sus1-11) impaired binding to both Sac3 and Sgf11. Trp38 is

which was slightly lower than the 1,324 – 1,546 Å² observed for the two Sus1-Sac3 interfaces in the Sac3 CID complex (10). Molecular recognition in the Sgf11-Sus1 complex was based on the hydrophobic helical stripe on the Sgf11 helix derived from an ~four-residue-repeating sequence motif (Fig. 3). A similar pattern of hydrophobic residues was seen in the sequences of Sgf11 from Kluyveromyces lactis and Candida albicans (Fig. 3C). Although the helical stripe motif on Sgf11 was similar in some respects to that observed for the Sac3-Sus1 interfaces, it was narrower and shorter than in Sac3. The interaction interface involved residues 8 – 32 of Sgf11 and was primarily hydrophobic with few H-bonds or salt bridges (Figs. 2D and 3B). The Sgf11 interaction interface on Sus1 was based primarily on the inner surface of helices α2–5, with little contact with helix α1 and was thus less extensive than in Sac3 (Figs. 2 and 3). However, in the DUB complex, Sus1 helix α1 may be involved with interactions with other components such as Sgf73 and/or Ubp8. In contrast to the interaction with Sac3, Sus1 does not encircle the Sgf11 helix completely and instead leaves one side exposed.

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strongly conserved in Sus1 (10), and Gly37 forms the hinge between helices α2 and α3, and so these mutations would be expected to cause major disruption to the molecular architecture of the Sus1 molecule, consistent with this mutant binding neither Sgf11 nor Sac3. Mutations within the hinge between Sus1 helices α1 and α2 (sus1-10) had little effect on the interaction with Sgf11 but did impair binding to Sac3, consistent with helix α1 making considerable contact in the Sus1–Sac3 complexes. Finally, mutation in the hinge between Sus1 helices α4 and α5 (sus1–12 V73A/D75A) caused a less severe phenotype for Sus1 binding but did not alter the Sgf11 interaction, probably because these residues are not intimately involved in the Sgf11–Sus1 interface (Fig. 2, D and E). As illustrated in Figs. 2 and 3, the Sus1-binding interface on Sac3 was more extensive than the Sgf11-binding interface and overlapped it completely, accounting for the inability to engineer Sus1 mutants in which Sac3 binding was retained, whereas Sgf11 binding was impaired.

Mutations That Interfere with the Sus1–Sgf11 Interaction—Previous studies (10, 39) highlighted the difficulty of interfering with interactions of the type observed between Sgf11 and Sus1 using simple point mutations. The interaction interface was primarily hydrophobic, and so mutations based on charge repulsion could not be employed, whereas dramatic mutations, involving substituting large hydrophobic residues or substituting charged for hydrophobic residues, would increase the likelihood of introducing major conformational changes. In analogous studies on the binding of calmodulin to helical peptides, Ala substitutions sometimes even enhanced affinity (39). We therefore used an alternative strategy to disrupt the interaction, based on inserting an Ala residue in the Sgf11 helix, so that the hydrophobic stripe to which Sus1 binds was disrupted. As illustrated in Fig. 5, A and B, insertion of a single residue in an α-helix results in a rotation of ~100°. Insertion of a single Ala after Sgf11 residues 18 or 19 (lanes 19A and 20A in Fig. 5C) resulted in a dramatic decrease in the affinity of Sgf11 for Sus1 in vitro pulldown assays (Fig. 5C). The effectiveness of this mutagenic approach is highlighted by comparison with a point mutation in the hydrophobic stripe at Ile15 (I15A) that had a less dramatic effect on Sus1 binding. In summary, the results obtained with these mutants confirm the importance of the hydrophobic stripe on Sgf11 in its interaction with Sus1.

Sus1 Binding to Sac3 and Sgf11 Is Mutually Exclusive—The observation that Sus1 wrapped around an α-helix in a similar way when complexed to either Sgf11 or Sac3, indicated that its binding to these two partners should be mutually exclusive. We tested this hypothesis by challenging two Sac3-Cdc31 complexes with the Sgf11–Sus1 complex (Fig. 6). The two Sac3 complexes, containing the Sus1B and Sus1A/B-binding sites, were unable to bind the Sgf11–Sus1 complex. These results indicated that a Sus1 chain binds to Sac3 and Sgf11 in a mutually exclusive manner because Sgf11 is not observed in the bound material, whereas Sus1 is evident. Inter-
Interestingly, in vitro Sac3 can extract Sus1 from a Sus1/Sgf11 heterodimer (Fig. 6) and this may reflect Sus1 equilibrating between the two complexes.

It was initially suggested that Sus1 could function to physically bridge the SAGA and TREX-2 complexes directly (7). However, the current study and previous work tend to support a more complex model for SAGA and TREX-2 interaction (1, 24). The similarity of the Sus1-Sgf11 and Sus1-Sac3 interaction interfaces (Figs. 2 and 3), combined with the indication that the binding of these two partners to Sus1 was mutually exclusive (Fig. 6), would make it unlikely that a single Sus1 molecule could link the SAGA and TREX-2 complexes. In principle, the Sus1 chains in each complex could interact with one another or with other components of the complexes to form an interface. We have not observed any indication of dimerization of either the Sgf11-Sus1 or Sac3-Sus1-Cdc31 complexes that would be consistent with a Sus1-Sus1 interaction. However, our present data cannot exclude the possibility that Sus1 may bridge the two complexes with a low affinity interaction that could, for example, be between the Sus1 chains of each complex (perhaps mediated by an additional factor) or the C-terminal zinc finger of Sgf11 interacting with the Sac3 CID region of the TREX-2 complex. Alternatively, it has been proposed that Sus1 could link the SAGA and TREX-2 complexes functionally by a dynamic exchange mechanism in which the competitive capture of Sus1 from TREX-2 by the DUB complex could dissociate TREX-2 from NPCs and up-regulate histone deubiquitination (1). This regulation would be reversed upon the reciprocal exchange of Sus1 from the DUB to TREX-2 and thereby could synchronize the NPC binding of TREX-2 to transcriptional events (1). It has also been suggested that post-translational modifications of Sus1 or other components of TREX-2 or SAGA could play a role in regulating gene gating (1). For example, Lys9 of Sus1 forms a salt bridge in the Sus1A-Sac3 and Sus1B-Sac3 interface but does not participate in the Sgf11-Sus1 interface (Figs. 2 and 3), and so post-translational modification of this residue could potentially contribute to switching the
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binding preference of Sus1 from Sac3 (TREX-2) to Sgf11 (SAGA).

In summary, we have determined the structure of the Sgf11-Sus1 complex and shown that Sus1 functions in each complex that is based on stabilizing extended single α-helices that would normally be anticipated to have very low stability when isolated in solution. However, the hydrophobic stripe that formed the Sus1-binding interface on Sgf11 was less extensive than that observed on Sac3 and provided a structural explanation for how it was possible to produce Sus1 mutants, in which binding to Sgf11 was retained while binding to Sac3 was lost (1), but how it was not possible to impair Sac3 binding without also losing Sgf11 binding. Overall, these structural results also favor a model in which Sus1 forms a component of separate subcomplexes in SAGA and TREX-2 rather than a single Sus1 chain linking the two complexes. Although further work will be required to establish precisely how Sus1 and Sgf11 participate in the linkage of SAGA to TREX-2, the structural and biochemical data obtained here identify the stabilizing of long α-helices in both complexes as an important function that contributes to their acting as interaction platforms that promote integration of transcriptional and export events in the gene expression machinery.

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