Architecture of *Saccharomyces cerevisiae* SAGA complex

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Dear Editor,

Eukaryotic gene transcription by RNA polymerase II (Pol II) is highly regulated and fine-tuned by diverse epigenetic complexes, which often function as co-activators or co-repressors to modulate transcription in concert with sequence-specific transcription factors (TFs). In *Saccharomyces cerevisiae*, the 1.8-MDa coactivator SAGA (Spt–Ada–Gcn5–acetyltransferase) complex1 was reported to be required for genome-wide transcription by RNA Pol II, and proposed to function as a general cofactor2. Yeast SAGA is composed of 19 subunits, which are categorized into four modules, namely HAT (Gcn5, Ada2, Ada3, and Sgf29), DUB (Ub8, Sgf11, Sgf73, and Sus1), TAF (Taf5, Taf6, Taf9, Taf10, and Taf12), and SPT (Tra1, Ada1, Spt3, Spt7, Spt8, and Spt20). Two catalytic subunits, Gcn5 within HAT1 and Ub8 within DUB4, permit histone H3 acetylation and ub-H2B deubiquitylation activities of the SAGA complex. Remarkably, acetylation and deubiquitylation carried out by SAGA are crucial for transcriptional initiation and elongation; in yeast, inactivation of SAGA leads to a global decrease of RNA Pol II-mediated transcription.

The largest SAGA component Tra1 (3744 residues) in the SPT module was shown to directly interact with the acidic activation domain of TFs, such as VP16, GCN4, Gal4, and HAP44, which help recruit SAGA to promoter regions of target genes. Tra1 is also a subunit of the yeast NuA4 acetyltransferase complex5. Roles of Tra1 in SAGA and NuA4 assembly, as well as TF binding await further investigation in molecular detail. SAGA was first reported as a histone H3 acetyltransferase in 19976, and the biological function of SAGA has been widely studied ever since then. Despite recent cryo-electron microscopy (cryo-EM) studies of Tra1 from *S. cerevisiae* (3.7 Å)8 and of the SAGA complex from *Pichia pastoris* (11.7 Å)9, the structure of an intact SAGA complex at higher resolution is needed to further elucidate its assembly and regulation.

We successfully purified an endogenous SAGA complex that contains all 19 subunits from fermented *S. cerevisiae* cells after optimization of growth conditions (Fig. 1a; Supplementary Fig. S1b). To avoid sample disassembly, the purified native SAGA complex was subject to cross-linking prior to negative stain and cryo-EM specimen preparation (Supplementary Fig. S1a, c). The structure of SAGA was determined by single-particle cryo-EM analysis at a resolution up to 6.9 Å (Fig. 1b; Supplementary Fig. S2 and Table S1). The overall shape of SAGA resembles an open padlock, and can be divided into lobes A and B (Fig. 1b, c; Supplementary Fig. S2c). To further improve the map, lobe A and lobe B were individually filtered out for focused refinement. Finally, the cryo-EM map of lobe A was reconstructed at 4.6 Å at FSC = 0.143 (Supplementary Fig. S2c, d, g). Consistent with a previous report6, the 3744-residue Tra1 was assigned to lobe A. After docking of the 3.7 Å free Tra1 model (PDB ID: 5OJ5)9, the structure of Tra1 in the context of SAGA was further modeled and refined to fit into the 4.6 Å cryo-EM map. By contrast, lobe B is less well resolved, and an overall resolution of 9.3 Å was calculated at FSC = 0.5 (Supplementary Fig. S2d, f). The two enzymatic modules, HAT and DUB, are located within lobe B. The blurry density of lobe B likely reflects conformational dynamics of the catalytic modules in the absence of the nucleosomal substrate.

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Fig. 1 (See legend on next page.)
Tra1 is composed of HEAT (Finger, N-clasp, Ring, and C-clasp), FAT, FRB, PI3K/Pi4K, and FATC domains from the N- to the C-terminus (Fig. 1d). We eventually generated an alanine-substituted structure of Tra1 consisting of alanine, glycine, and proline residues (Fig. 1e). Tra1 is characteristic of a helical solenoid fold and arranged into a “Θ”-like structure, in which a Ring motif (1140–2442) within the HEAT domain forms the ring (Fig. 1f). Tra1 functions as a base to organize other subunits in lobe B of SAGA (Fig. 1c). Structural comparison of free Tra1 with its complex state revealed only regional conformational adjustments upon SAGA assembly, which reflects the structural rigidity of the solenoid fold of Tra1 (Fig. 1f; Supplementary Movie S1).

A comparison of the cryo-EM map of S. cerevisiae SAGA with the reported one from P. pastoris revealed two better-defined regions in lobe B, referred to as R-right and R-left regions (Fig. 1b, c; Supplementary Fig. S3). The R-right region is located at the middle of the SAGA next to Tra1. Under lower-threshold levels, the density of the R-right region extends from the C-terminal region of Ring (Ring-C) (~2130–2266) toward the middle region of Finger (~579–691) of the HEAT domain, thus spanning over Tra1 like an arch (Fig. 1f; g; Supplementary Fig. S3a, b). In addition, the elongated ridge surface formed by FAT, PI3K/Pi4K, C-Clasp, and Ring-C of Tra1 constitutes the major contact interface with lobe B (Fig. 1f). Previous negative staining studies of Spt8-GFP SAGA suggested positioning of Spt8 next to Tra1 (Supplementary Fig. S4a). Direct comparison of our 2D class average images (both negative staining and cryo-EM) with the reported one of Spt8-GFP suggests that Spt8 is likely located to the R-right region (Fig. 1g; Supplementary Fig. S4b).

To further dissect the subunit organization of lobe B, we next performed chemical cross-linking of proteins coupled with mass spectrometry (CXMS) analyses. In total, 139 unique inter- and 184 unique intra-subunit cross-linking pairs were identified (Fig. 1h; Supplementary Table S2), which reflects an extensive interconnectivity among SAGA subunits. Direct cross-linking of Tra1 with Taf12, Ada1, Spt20, Spt3, and Taf9 indicated that these subunits in lobe B are located next to Tra1 (Fig. 1h).

Remarkably, high-confidence cross-linking peaks suggest that Taf12/Ada1/Spt20/Spt3 are clustered to contact with Tra1 through a surface formed by Ring-C, C-Clasp, and FAT (Fig. 1h; Supplementary Figs. 5–7). Aided by GFP tracing, the HAT (Gcn5, Ada2, Ada3, and Sgf29) and DUB (Ubp8, Sgf11, Sgf73, and Sus1) modules have been previously shown to locate at the Tra1-distal end of lobe B. Consistently, few cross-linking pairs were detected between Tra1 and DUB/HAT (Fig. 1h).

The second largest subunit, Spt7, plays a central role in lobe B organization, as reflected by its broad interactions with most components from SPT (Spt3 and Spt8), TAF (Taf5, Taf6, and Taf12), and HAT (Ada2 and Ada3) modules (Fig. 1h). Meanwhile, few cross-linking pairs were detected between Tra1 and Spt7/Taf5/Taf6/Taf10, suggesting that these subunits are clustered in the middle part of lobe B with Spt7 positioned at the core. Unlike Taf5, Taf9, Taf10, and Taf12 that are cross-linked to multiple subunits, Taf6 is rather isolated and mainly cross-linked to Spt7 (similar to Spt8, ~66.2 kDa; Fig. 1h). Combining with our EM analysis, Taf6 (~58 kDa) is probably located to the relatively exposed R-left region. This is consistent with previous reports in which the “R-left” region was assigned as part of TAF. Collectively, we proposed an assembly model of SAGA (Fig. 1; Supplementary Figs. S4b, S8a), which lays out a structural framework for functional analysis of the SAGA holo enzyme.

SAGA and NuA4 are two major acetyltransferases in yeast and share the largest subunit Tra1. To compare the role of Tra1 in organizing SAGA and NuA4, we superimposed the two complexes by aligning Tra1 together (Fig. 1j). Distinct surfaces of Tra1 were involved in the assembly of SAGA and NuA4, and the rest parts of the two enzymatic machineries are built up in perpendicular directions (Fig. 1k). Notably, NuA4 extends from the lateral surface of the Tra1 ring, while the SAGA grows up
from the central surface of Tra1. We noticed that the Eaf5 subunit of NuA4 clashes with lobe B of SAGA (Fig. 1j)\(^9\), which suggests that SAGA and NuA4 unlikely function as one super-complex, unless an event of subunit switch occurs.

SAGA serves as a general cofactor in yeast, genome-wide linking epigenetic modifications to transcriptional activation in concert with sequence-specific TFs. We next performed surface charge analysis of the reported 3.7 Å Tra1 cryo-EM structure (PDB ID: 5OJS)\(^6\) in the context of SAGA (Fig. 1k; Supplementary Fig. S8). Intriguingly, the exposed Tra1 surface upon complex formation displayed bipolarized electrostatic potential distribution patterns, in which the front-view surface is mainly negatively charged and the back-view surface is largely positively charged (Fig. 1k). The positively charged back-view surface of Tra1 is likely involved in binding with DNA or the acidic activation domain of TFs, hence orchestrating physical TF recruitment, cooperative DNA binding, and histone modifications, in particular nuclear processes regulated by SAGA or NuA4 (Fig. 1l). In support, the Vp16/Gal4 activator interaction domain (AID) of Tra1 has been mapped to segment 2233–2836 that spans Ring-C to FAT in a previous report (Fig. 1e, k)\(^4\).

Histone H3 acetylation and ub-H2B deubiquitylation are two tightly coupled activities of SAGA, which raises the question of how the HAT and DUB modules are coordinated during transcription. Hence, we docked ub-nucleosome-DUB structure into the SAGA (PDB ID: 4ZUX, Supplementary Fig. S9)\(^10\), which might constitute one of the possible intermediate states for nucleosome recognition by SAGA. Intriguingly, a ubiquitylated nucleosome was captured by the DUB module and swung away from the HAT module. It appears that the HAT and DUB modules could not engage in the catalysis of one nucleosome substrate concurrently. Our structural analysis suggests that the DUB and HAT modules of SAGA likely act sequentially, in which histone H3 acetylation by HAT is preceded by the ubiquitin removal and ub-nucleosome recognition by DUB. Such a sequential catalysis model awaits further complex structural studies.

**Accession codes**

The electron density maps for *S. cerevisiae* full SAGA complex and subunit Tra1 have been deposited in EMDB under accession codes EMD-9663 and EMD-9664, respectively. The atomic coordinates for a pseudo-structure of Tra1 have been deposited in the Protein Data Bank under accession code 6IG9.

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**Authors’ contributions**

H.L. supervised this study; X.Z., G.L., H.G., and H.L. designed the experiments; H. G., X.Z., H.Q., J.K., and X.R. purified the proteins; G.L. prepared the cryo-EM specimen, collected the data, and reconstructed the map; X.Z. built the model; Y.C. and M.D. performed the mass spectrometry analysis; X.L. and J.L. provided cryo-EM support and critical input; X.Z., G.L., and H.L. wrote the paper.

**Conflict of interest**

The authors declare that they have no conflict of interest.

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