SAGA histone acetyltransferase module facilitates chromatin accessibility to SMC5/6

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Abstract

Structural Maintenance of Chromosome (SMC) complexes are molecular machines driving chromatin organization at higher levels. In eukaryotes, three SMC complexes (cohesin, condensin, and SMC5/6) play key roles in cohesion, condensation, replication, transcription and DNA repair. Here, we performed a fission yeast genetic screen to identify novel factors required for the SMC5/6 complex with compromised binding to DNA. We identified 79 genes of which the histone acetyltransferases (HATs) were the most represented. Genetic and phenotypic analyses suggested a particularly strong functional relationship between SMC5/6 and SAGA complexes. Furthermore, several SMC5/6 subunits physically interacted with SAGA HAT module components Gcn5 and Ada2. As Gcn5-dependent acetylation facilitates accessibility of chromatin to DNA repair proteins, we first analyzed the formation of DNA damage-induced SMC5/6 foci in the Δgcn5 mutant. The SMC5/6 foci formed normally in Δgcn5, suggesting SAGA-independent SMC5/6 localization to DNA damaged sites. In unchallenged cells, we used Nse4-FLAG chromatin-immunoprecipitation (ChIP-seq) analysis to assess SMC5/6 distribution. A significant portion of SMC5/6 accumulated within gene regions in WT cells, and it was reduced in Δgcn5 and Δada2 mutants. The drop in SMC5/6 levels was also observed in gcn5-E191Q acetyltransferase-dead mutant, suggesting that the SAGA HAT module may facilitate chromatin accessibility to SMC5/6.

Keywords: genetic interactions; SMC5/6 complex; Nse3 KITE; SAGA histone acetyltransferase module; Gcn5; Ada2; histone H3K9ac acetylation; chromatin accessibility; DNA repair; rDNA; gene regions
Introduction

Chromatin is composed of DNA and protein complexes structured at multiple levels to ensure its spatial and functional organization [1]. At the basic level, histone proteins pack DNA into nucleosomes and their arrays. At the higher levels, Structural Maintenance of Chromosome (SMC) complexes (cohesin, condensin and SMC5/6) assist in the formation of higher-order structures like topologically associated domains or condensed mitotic chromosomes [2]. Chromatin compaction at each level affects DNA accessibility. At the nucleosomal level, histone chaperons, modifiers, and remodelers can loosen, move or remodel chromatin fibers to enable essential processes like transcription or DNA repair [3, 4]. For example, the histone-modifying SAGA complex acetylates H3 histones at the transcription start sites, inducing chromatin structure opening, which facilitates the accessibility of chromatin to transcription initiation complexes or other complexes requiring direct contact with DNA [5, 6].

The SMC complexes play roles in all key chromatin processes, like cohesion, condensation, replication, transcription, and DNA repair. Their cores comprise long-armed SMC, kleisin, and kleisin-associated (KITE or HAWK) subunits [7, 8]. Uniquely, SMC5/6 complexes contain highly conserved Nse1 and Nse2, ubiquitin- and SUMO-ligases, respectively [9, 10]. The Smc subunits are primarily built of head ATPase domains, long anti-parallel coiled-coil arms, and hinges [11-13]. Two Smc molecules form stable dimers via their hinge domains, and without ATP, their arms align to rod-like structures [14, 15]. The binding of ATP molecules to the ATPase head domains promotes the formation of large annular structures [16, 17]. The ATP binding-hydrolysis cycle drives ring-to-rod dynamic changes and promotes DNA translocation or loop extrusion [18-21].

SMC complexes were believed to interact only topologically with chromatin fibers via their large annular structures, which can embrace and traverse large chromatin complexes
(including nucleosomes; [13]). However, growing evidence suggests their requirement for open chromatin and direct physical binding to DNA [22-24]. Moreover, Piazza et al. [23] described the preferential binding of condensin kleisin-associated HAWK subunits to free DNA over nucleosomal DNA. Recent cryoEM analyses showed the formation of K-compartments within all SMC complexes, consisting of ATP-bound Smc heads, kleisin, and kleisin-associated subunits, which can accommodate only free DNA [17, 25-27]. In line with these findings, the SAGA complex assists in loading condensin at open chromatin regions of highly transcribed genes [24]. Similarly, the RSC chromatin remodelling complex recruits the Scc2-Scc4 factor to nucleosome-free regions, assisting in cohesin loading at these sites [28-30].

Recently, it was shown that the SMC5/6 K-compartment, composed of Smc5-Smc6 heads, Nse4 kleisin, and Nse1-Nse3 kleisin-associated KITE subunits, binds free DNA [17, 22]. In our previous study, we described the essential role of Nse3-DNA interaction for SMC5/6 loading or accumulation [22]. Here, we performed a genetic screen with the nse3-R254E fission yeast mutant with reduced DNA-binding affinity to identify new factors required for its viability. We found strong genetic interactions with histone acetyltransferase (HAT) complexes, SAGA and NuA4. Using chromatin-immunoprecipitation (ChIP-seq) analysis, we observed a significant portion of SMC5/6 accumulated within gene regions, which was reduced in SAGA HAT deletion (Δada2 and Δgcn5) mutants. The magnitude of decrease in SMC5/6 occupancy correlated with the SAGA-modified H3K9ac levels around the transcription start sites. The SMC5/6 reduced levels were also observed in gcn5-E191Q acetyltransferase-dead mutant, suggesting that the SAGA HAT module may facilitate the accessibility of chromatin for SMC5/6 loading.
Results

Genetic screen with a DNA-binding defective allele of the SMC5/6 complex

To identify factors that facilitate the loading or accumulation of the SMC5/6 complex on chromatin, we performed a genetic search for genes affecting the survival of cells with compromised DNA-binding ability of the SMC5/6 complex [22]. First, we created a query fission yeast strain with the DNA-binding defective nse3-R254E mutation in the PEM2 background (Suppl. Fig. S1A; [31]) and crossed it against the whole gene deletion yeast collection from BIONEER [32]. Using the yeast colony size phenotypic readout [33], we identified 79 deletion strains that exhibited a negative genetic interaction with nse3-R254E (Suppl. Table ST1).

To validate our results, we randomly selected 19 of the 79 strains and crossed them with the original nse3-R254E strain [22]. Tetrad analysis confirmed that these mutations are synthetically sick or lethal with the nse3-R254E mutation (Suppl Fig. S1B). Analysis of all 79 interacting genes for the Biological process category using Gene ontologies (GO; [34]) showed the highest scores for DNA repair, chromatin organization, meiosis, and replication processes (Fig. 1A and Suppl Table ST2), in line with previous studies (reviewed in [35, 36]). Reassuringly, several nse3-R254E genetic interactions overlapped with the genetic interactions of other smc5/6 mutants [22, 37-42], supporting the validity of our screen results.

Genetic interactions between SMC5/6 and histone acetyltransferase complexes

The analysis of the Cellular components GO of the 79 interacting genes showed the highest scores for histone acetyltransferase (HAT) complexes (SAGA and NuA4; Fig. 1B and Suppl Table ST3). Three (Ada2, Ada3/Ngg1, Gcn5) out of four HAT module subunits and one (Ubp8) out of four DUB module subunits of the SAGA complex were identified as hits in our screen [43]. We verified their genetic interactions using tetrad analysis (Figs. 1C and
S1B) and also tested the other non-essential SAGA subunits not detected in our screen. Although the meiotic defects of the \( \Delta ada1, \Delta spt7, \Delta spt8, \) and \( \Delta spt20 \) deletion mutants hampered the double mutant preparation [44], the tetrad analysis of the other non-essential SAGA subunits (Suppl. Fig. S1C) showed negative genetic interactions with the \( nse3-R254E \) mutation (Suppl. Fig. S1D), suggesting a strong functional relationship between the SMC5/6 and SAGA complexes.

Interestingly, the temperature-sensitive (ts) phenotypes of the SAGA HAT module \( \Delta ada2, \Delta ada3, \) and \( \Delta gcn5 \) mutants were enhanced by \( nse3-R254E \) (Fig. 1D; [44]). The ts phenotypes were also enhanced by the \( smc6-74 \) and \( smc6-X \) hypomorphic mutations (Suppl. Fig. S2A; [38, 45]). We also used \( nse1-R188E \) and \( nse2-SA \) mutations, which specifically abrogate DNA-repair function (ubiquitin- and SUMO- ligase activity, respectively), but not the SMC5/6 essential function [9, 46]. However, these mutations did not affect the ts phenotype of \( \Delta gcn5 \) (Suppl. Fig. S2A; not shown). These data suggest that the SMC5/6 essential function supports cell survival at higher temperatures in the absence of the HAT module and vice versa; the viability of the \( nse3-R254E \) mutant is compromised in the absence of a functional SAGA HAT module.

**SMC5/6 and SAGA physically interact**

The strong genetic relationship between the SAGA HAT module and the SMC5/6 complex prompted us to test their mutual physical interactions. First, we performed co-immunoprecipitation of Gcn5-myc and Nse4-FLAG kleisin subunit to show the association between the SAGA and SMC5/6 complexes *in vivo* (Fig. 2A). The fission yeast cells carrying Nse4-FLAG (with or without Gcn5-myc) were lysed, and proteins were precipitated with an anti-myc antibody. A small amount of Nse4-FLAG was specifically recovered in the Gcn5-myc precipitates but not in the control experiment without myc-tagged Gcn5, suggesting a weak or transient association between SAGA and SMC5/6 in the yeast cells.
Next, we used our panel of yeast two-hybrid (Y2H) SMC5/6 subunits to test their interactions with the HAT module (Ada2, Ada3, Gcn5, and Sgf29) subunits. Two of the HAT module subunits, Ada2 and Gcn5, bound the Nse3 subunit (Figs. 2B and S2B). The fragment analysis showed that Gcn5 bound the N-terminal part of Nse3(aa1-190), while Ada2 interacted with its C-terminal WHB domain (Nse3(aa200-307); Figs. 2B and S2C). In addition, Ada2 bound the N-terminal region of Nse2 and the coiled-coil arm of SMC5 next to the SMC5-Nse2 binding interface [47]. Altogether, our data suggest a functional relationship between SAGA and SMC5/6 assisted by a network of physical interactions.

**SMC5/6 and SAGA are required for efficient DNA repair**

The SAGA complex plays an important role in transcription regulation and DNA repair by facilitating the accessibility of chromatin to transcription factors and repair proteins [44, 48-51]. Deletion mutants of most SAGA genes were sensitive to hydroxyurea (HU; [44]) and SAGA mutations increased the sensitivity of the smc5/6 hypomorphic mutants to HU and other DNA-damaging agents (Figs. 3A, S2A, and S3A). These data suggest either a direct role of the SAGA complex in facilitating SMC5/6 access to chromatin at sites of DNA damage or its indirect additive role.

Upon DNA damage, SMC5/6 accumulates in foci in a Brc1-dependent way [52, 53]. To assess whether SMC5/6 chromatin accessibility at these DNA damage sites is directly facilitated by SAGA, we analyzed the Nse4-GFP foci in the Δgcn5 and Δada2 mutants (Fig. 3B). We observed no difference between the frequency of cells with foci in the wild-type (WT), Δgcn5, and Δada2 after treatment with either MMS (methyl methane sulfonate) or HU. In contrast, the number of foci in the Δbrc1 mutant was strongly reduced (Fig. 3C). These results indicate no direct involvement of SAGA in SMC5/6 localization to sites of DNA damage. Instead, they suggest additive effects of the SAGA and SMC5/6 complexes on DNA repair.
SAGA facilitates chromatin accessibility to SMC5/6 in gene regions

In addition to the accessibility of chromatin to DNA repair or transcription factors, SAGA facilitates chromatin accessibility to condensin complexes [24]. Therefore, we determined the SMC5/6 localization in the unchallenged WT, Δgcn5, Δada2, and Δubp8 cells using chromatin immunoprecipitation of Nse4-FLAG followed by deep sequencing (ChIP-seq). In the WT cells, most SMC5/6 localized to the repetitive regions (like rDNA, centromeres, or tDNA copies), with the highest occupancy at the rDNA repeats (Fig. 4A) consistent with previous reports [54-59]. Interestingly, a significant portion of Nse4-FLAG was localized within the intergenic regions and, most notably, genes. We identified 331 Nse4-FLAG peaks (representing ¼ of the total Nse4 occupancy; Fig. 4A) localized to the gene regions.

In Δgcn5 and Δada2 mutants, SMC5/6 distribution was altered, while SMC5/6 occupancy in the Δubp8 mutant was similar to the WT cells. A heatmap clustering analysis showed that the Nse4-FLAG peaks were either enhanced, unchanged, or reduced in the Δgcn5 and Δada2 mutants (Fig. 4B). The Nse4-FLAG signals were mainly enhanced at the repetitive sequences (Fig. 4C), while most peaks in the intergenic regions were not changed (Fig. 4D). Strikingly, most peaks within the gene regions showed reduced Nse4 occupancy (252 out of 331; Figs. 4C and D), suggesting that the SAGA HAT module may facilitate chromatin accessibility to SMC5/6 at gene loci.

The SAGA HAT module acetylates histone H3 at its lysine K9 and K14 residues [60, 61]. To determine the H3K9ac distribution in our fission yeast cells, we performed ChIP-seq using a specific anti-H3K9ac antibody. A heatmap clustering analysis showed that the magnitude of decrease in SMC5/6 occupancy in gene bodies in the Δgcn5 mutant (Δgcn5-WT plot; Figs. 5A and B) correlated with the level of H3K9ac around the transcription start site.
and with the transcript levels [5], suggesting that the SAGA-dependent H3K9 acetylation may facilitate the accessibility of chromatin to SMC5/6.

To assess the role of the SAGA-dependent acetylation further, we used the *gcn5-E191Q* acetyltransferase-dead mutant [51]. First, we found that the *gcn5-E191Q* ts phenotype was exacerbated by *nse3-R254E*, similar to Δ*gcn5*. Second, the *gcn5-E191Q nse3-R254E* double mutant was as sensitive to DNA damaging agents as the Δ*gcn5 nse3-R254E* double mutant (Suppl. Fig. S3D), suggesting that the Gcn5 acetyltransferase activity plays a crucial role in the SAGA-SMC5/6 genetic interactions. Finally, using ChIP-qPCR analysis, we observed reduced Nse4-FLAG occupancy in the *gcn5-E191Q* mutant at selected gene regions similar to Δ*gcn5* (Fig. 5C), although some loci exhibited only a modest drop in SMC5/6 levels. Altogether, our results suggest an important role for the SAGA HAT module in facilitating the accessibility of chromatin to SMC5/6 at gene loci.

**Discussion**

Here, we discovered the SAGA HAT module as a new chromatin factor assisting in the localization of SMC5/6 to specific genomic loci. In line with SAGA's best-known function in transcription [43], Δ*gcn5* deletion reduced SMC5/6 occupancy specifically at the gene loci (Fig. 4). In addition, the magnitude of decrease in SMC5/6 occupancy in gene bodies correlated with the level of H3K9 acetylation and transcript levels (Fig. 5), suggesting that the SAGA-dependent H3K9 acetylation may facilitate the accessibility of chromatin to SMC5/6. Consistent with this notion, SMC5/6 occupancy was also reduced in the *gcn5-E191Q* acetyltransferase-dead mutant. The fact that occupancy reduction was less pronounced in the *gcn5-E191Q* mutant could be explained by the physical interactions between SAGA and SMC5/6. Such interactions may target SMC5/6 to SAGA-rich regions, while H3K9 acetylation opening chromatin would enable DNA entrapment. In the *gcn5-E191Q* mutant, SMC5/6 could still be targeted to gene regions, but unmodified, inaccessible chromatin would hamper DNA entrapment. Presumably, the DNA-unbound SMC5/6 in the *gcn5-E191Q*
mutant would cause a similar defect as in the Δgcn5 mutant and result in the same phenotypes of gcn5-E191Q nse3-R254E and Δgcn5 nse3-R254E double mutants (Suppl. Fig. S3D and E) as the binding to free DNA is essential for the SMC5/6 function [17, 22].

The additive phenotypes of the double mutants (Fig. 3) probably stem from non-redundant functions of SMC5/6 and SAGA complexes. For example, SMC5/6 is essential for replication processes (like replication restart; [35]), while SAGA is necessary for the timely transcription of S-phase specific factors [60]. Defects in these functions probably combine in double mutants into high HU sensitivity (Figs. 3, S3A, and D). Similarly, the reduced DNA-binding affinity of the nse3-R254E mutant is combined with the reduced binding to gene regions upon gcn5 deletion, resulting in reduced growth phenotypes of double mutants. Interestingly, enhanced SMC5/6 accumulation at repetitive regions (Fig. 4) could not rescue these growth defects suggesting an important role of SMC5/6 binding to the gene regions.

It was recently shown that plant ADA2b binds SMC5 and assists in the localization of SMC5/6 to DNA-damage sites [62]. Later studies suggested ADA2b-diRNA specific pathway targeting SMC5/6 to DNA-damage foci in plants [63]. However, it is unclear whether this pathway involves other SAGA components. In fission yeast, it was shown that the SMC5/6 targeting to DNA-damage sites depends on the BRCT domain-containing Brc1 protein but not on Ada2 or Gcn5 (Fig. 3; [54]). It suggests a specific function of the ADA2b-SMC5 interaction in DNA-damage response in plants. Future studies may show if SAGA likewise facilitates the accessibility of chromatin to SMC5/6 at gene loci in plants. Interestingly, we found an interaction between human hADA2b and hNSE4a/b subunits (Suppl. Fig. S2D), suggesting diverse interaction networking between SAGA and SMC5/6 complexes across different organisms. It will be interesting to learn about the functional relationships between human SAGA and SMC5/6 complexes, particularly in facilitating SMC5/6 targeting to gene regions, in the future.
Consistent with the above findings, rDNA loci and other repetitive regions, which are prone to DNA damage even without any genotoxic treatment, were not reduced for SMC5/6 in \( \Delta gc n 5 \) and \( \Delta ada 2 \) deletion mutants (Fig. 4). Instead, repetitive loci were enriched for SMC5/6 upon SAGA HAT deletion. We speculate that reduced acetylation levels promoted an increase in methylation levels, indirectly stimulating the SMC5/6 localization to repetitive heterochromatic regions [54, 64]. Altogether, our findings support previous views that different factors target SMC5/6 to different genomic regions [35, 53, 54] and show a new role of SAGA in SMC5/6 targeting to gene regions.

**Materials and methods**

**Yeast techniques**

Standard fission yeast genetic techniques were used [65]. Yeast strains were crossed and sporulated either at 25°C (ts mutants) or 28°C (non-ts mutants). Tetrad analysis was carried out on Singer MSM300 (Singer, UK). The deletion integrations were verified on both ends by PCR with specific primers to the G418 cassette and genomic sequence of a deleted gene (approximately 600-800bp from start or end). The PCR products were sequenced.

* S. pombe cultures were grown to the mid-log phase, and serial 10-fold dilutions were spotted onto rich media with the indicated dose of DNA damaging agent (hydroxyurea or methyl methane sulfonate). Subsequently, plates were incubated at the indicated temperatures (25, 28, or 37°C) for 3–4 days. Selective media were supplemented with Nourseothricin (cloNAT, 100 µg/ml; Jena Bioscience), G418 (100 µg/ml; Applichem), and/or cycloheximide (100 µg/ml; Sigma).

**Yeast genetic screens**

The pAW8-Nse3 integration construct [22] was modified for use in the PEM2 strain as follows. The SpHl site was mutated to XhoI using a site-directed mutagenesis kit (Agilent Technologies; primers: LJ48 and LJ49; Suppl. Table ST4). The cloNAT cassette was
amplified (LJ42 and LJ43) and inserted into the XhoI site in front of the Nse3 gene using the In-fusion cloning kit (Takara). A 650bp-long genomic sequence (upstream of Nse3; LJ44 and LJ45) was inserted in front of the cloNAT cassette (using XhoI) to ensure its proper integration into the S. pombe genome. The mutant cloNAT-nse3-R254E construct was created using site-directed mutagenesis (R254E_F and R254E_R primers; [22]). For the yeast transformation, the WT and mutant cloNAT constructs were cleaved by SpeI, and the 3246bp long fragment was purified from agarose gel by Gel Extraction Kit (Qiagen). Approximately 1 μg of purified DNA was transformed into the PEM2 strain [31] by standard LiAc protocol. The proper integration of the cloNAT-Nse3 cassette and rlp42 mutation was checked by PCR and sequencing. The nse3-R254E PEM2 strain phenotypes were compared with the original nse3-R254E strain (Suppl. Fig. S1A; [22]).

The WT and nse3-R254E mutant PEM2 strains (YLJ222 and YLJ228; Suppl. Table ST5) were crossed with the S. pombe haploid deletion library (BIONEER, version 5, us.bioneer.com) according to the published protocol [31]. The screen was repeated twice using the Rotor HDA robot (Singer, UK). The plate images were taken by a Canon EOS Rebel T3i camera, and the individual colony size was measured. The viability of single deletion mutants (control WT plates) against double mutants (test nse3-R254E plates) was compared using SGAtools online platform [33]. Genes with a score less than -0.25 were chosen as potential negative interactors (Suppl. Tab. ST1).

The resulting group of 79 genes was analyzed by the Gene ontology tool BiNGo [66], which is a plugin of the Cytoscape online platform [34]. The genes were classified according to the Pombase GO database into Biological processes and Cellular component categories, respectively [67]. The default parameters with a 0.05 significance level were applied for both categories.

**Y2H analysis**
The Gal4-based Y2H system was used to analyze SMC5/6-SAGA interactions [68]. S. pombe ada2, ada3, gcn5, and sgd29 genes were PCR amplified from genomic DNA (primers used for ada2 and gcn5 cloning are listed in Suppl. Table ST4). All inserts were cloned into respective sites of the pGBK7T or pGADT7 vectors using the In-Fusion cloning system. pGBK7T-Nse2(aa2-178) was described in [9]. pGADT7-Nse3(aa1-190) was prepared by mutagenesis of 191st aa to STOP codon in pGADT7-Nse3(aa1-328) [69]. The Nse3(aa200-307) fragment was cut out from pTriEx4-Nse3(aa200-307) [70] by Ncol-XhoI enzymes and cloned into pGADT7. The Smc5(aa170-225+837-910) fragment was amplified from the Smc5(aa2-225+837-1065) construct [71] and inserted into the NcoI-NotI sites of pGBK7T.

The pairs of pGBK7T and pGADT7 constructs were co-transformed into the Saccharomyces cerevisiae PJ69–4a strain by standard LiAc transformation protocol and selected on SD -Leu, -Trp plates. Drop tests were carried out on SD -Leu, -Trp, -His (with 0.3, 0.5, 1, 3, 5, or 10 mM 3-aminotriazole) plates at 28°C. Each combination of partners was co-transformed and tested at least twice.

Co-immunoprecipitation of S. pombe proteins

Logarithmically growing YLJ507 and MMP21 cells (Suppl. Table ST5; Suppl. Fig. S3B) were cultivated in a rich medium at 28°C (OD955 = 0.4 – 0.7). 5x10^8 cells were harvested by centrifugation (3 min, 4°C, 5000 rpm) and washed with 10 ml of ice-cold PBS. Pellets were stored in the 2 ml screw cup tubes at -80°C. The crude yeast extracts were prepared in 400 μl CHIP lysis buffer (50 mM HEPES, pH 7.5, 140 mM NaCl, 1 mM EDTA, 1% Triton X-100, Complete EDTA-free protease inhibitor cocktail tablets, Roche) with half volume of glass beads (Sigma) in 2 ml low binding tubes using FASTprep-24 (MP Biomedicals; 5 times, 30 sec, 6.5 speed). The suspension was recovered by piercing the bottom of the tube with a needle, placing it into a new 2 ml tube, and centrifugation (3 min, 4°C, 5000 rpm). The beads
were washed with 200 μl of CHIP lysis buffer (3 min, 4°C, 5000 rpm). The collected suspensions were clarified by centrifugation (15 min, 4°C, 15000 rpm), and the supernatant was transferred to new low-binding tubes. 40 μl of cell extract was taken for input control. Immunoprecipitation was carried out by adding 2 μl mouse anti-myc antibody (2276S, Cell Signalling) to the cell extracts and incubation for 2 hours at 4°C. 20 μl of protein G coated Dynabeads (Invitrogen) were washed twice with 1 ml CHIP lysis buffer and resuspended in 60 μl of CHIP lysis buffer, then added to the extract with anti-myc antibody and incubated overnight at 4°C. The beads were pelleted using a magnetic rack, and the unbound fraction (40 μl) was taken. Beads were washed four times with 1 ml CHIP lysis buffer, and proteins were eluted by 40 μl of 1xSDS loading buffer. After 15 min incubation at room temperature, the supernatant (bound fraction) was recovered. All fractions were analyzed by western blotting using mouse anti-myc-HRP (R951-25, Thermo Fisher) and mouse anti-FLAG-HRP (F1804-1MG, Sigma) antibodies, respectively.

Chromatin immunoprecipitation analysis

Nse4-FLAG ChIP-seq

All strains (Suppl. Table ST5) were cultivated into the mid-log phase (OD = 0.4 – 0.6) and incubated with 1 % formaldehyde for 15 min at room temperature to cross-link DNA–protein complexes. Glycine was added to a final concentration of 125 mM, and the incubation continued for 5 min. 5x10^8 cells were harvested and washed with 10 ml of ice-cold PBS. The yeast cell wall breakage was performed in 400 μl CHIP lysis buffer with half the volume of glass beads in 2 ml low binding tubes using FASTprep-24. The suspension was washed two times with CHIP lysis buffer (15 min, 4°C, 15000 rpm), and 300 μl of the extract was sonicated with Bioruptor (Diagenode, 30 sec ON/30 sec OFF, High Power, 25 times) and clarified by centrifugation (15 min, 4°C, 15000 rpm), resulting in an average DNA fragment
size of 300-500 bp. 5 μl of the sonicated precleared extract was taken as an input control sample.

Monoclonal anti-FLAG M2 antibody (F1804, Sigma) was diluted 1:150, incubated with precleared cell extract in 1.5 ml low binding tube for 2 h on ice, and precipitated overnight with Dynabeads protein G (Invitrogen). Precipitates were washed with 1 ml of CHIP lysis buffer, 1 ml of High Salt buffer (CHIP lysis buffer with 500 mM NaCl), 1 ml of Wash buffer (10 mM Tris-HCl at pH 8.0, 0.25 M LiCl, 0.5% NP-40, 1 mM EDTA), and 1 ml of TE buffer (20 mM Tris-HCl at pH 8.0, 1 mM EDTA). After elution (50 mM Tris at pH 8, 0.1% SDS, 10 mM EDTA) and de-crosslinking overnight at 65°C, the DNA was purified using QIAquick PCR Purification Kit (Qiagen).

For ChIP-seq analysis, the input DNA samples were tested for DNA fragmentation and determination of DNA concentration by the Fragment analyzer (Agilent). Input and immunoprecipitation (IP) samples with the best fragmentation and high concentration were used for the creation of NGS libraries (NEBNEXT ULTRA II DNA Library Prep kit, NEB) and sequencing (Illumina Next seq 500, Illumina).

**H3K9ac/H3 ChIP-seq**

Two independent replicates were performed. Cells were grown to exponential phase (OD_{600} = 0.5) in the complex YES medium and fixed by adding formaldehyde to the final concentration of 1 %. After 30 min incubation, the remaining formaldehyde was quenched by 125 mM glycine. Cells were washed with PBS and broken with glass beads. Extracted chromatin was sheared with the Bioruptor sonicator (Diagenode) using 15 or 30 cycles (for biological replicate 1 and 2, respectively) of 30 sec ON/30 sec OFF at high power settings. For all immunoprecipitations (IP) within a biological replicate, the same amount of chromatin extract was used (2.5 or 3.7 mg of total protein); 1/10 of the total chromatin extract amount was kept for input DNA control. For each IP, 5 μg of antibody (H3: Ab1791, H3K9ac: Ab4441, all
Abcam) were incubated with the chromatin extract for 1 hour at 4°C with rotation. Then, 50 μl of BSA-blocked Protein A-coated magnetic beads (10002D, ThermoFisherScientific) were added to the chromatin extract-antibody suspension and incubated for additional 4 hours at 4°C with rotation. The precipitated material and input chromatin extract were de-crosslinked and treated with RNase A and proteinase K. DNA was purified using phenol-chloroform extraction and sodium acetate/ethanol precipitation. In biological replicate 2, DNA purification on AMPure XP beads (AC63880, Beckman Coulter) was performed after the phenol-chloroform extraction to remove low-molecular fragments and RNA. DNA concentration was measured using the Quantus fluorometer (Promega), and fragment size distribution was checked on Agilent Bioanalyzer using the High Sensitivity DNA Assay. Library construction and sequencing (50 nt SE) were performed by BGI Tech Solutions (Hong Kong) using the BGISEQ-500 sequencing system.

**NGS data analysis**

The reference fission yeast *S. pombe* genome (2018-09-04) and annotation (2019-11-15) were downloaded from PomBase ([https://www.pombase.org/](https://www.pombase.org/); [67, 72]). Read quality was checked using FastQC version 0.11.8 ([https://www.bioinformatics.babraham.ac.uk/projects/fastqc/](https://www.bioinformatics.babraham.ac.uk/projects/fastqc/)), and reads were aligned to the *S. pombe* genome using HISAT2 2.1.0 [73] and SAMtools 1.9 [74, 75]. Read coverage tracks (i.e., target protein occupancy) were then computed and normalized to the respective mapped library sizes using deepTools 3.5.1 [76]. The raw ChIP-seq data are available from the ArrayExpress ([https://www.ebi.ac.uk/](https://www.ebi.ac.uk/)) database under the accession number E-MTAB-11081.

WT fission yeast RNA-seq data were obtained from the NCBI Sequence Read Archive ([https://www.ncbi.nlm.nih.gov/sra](https://www.ncbi.nlm.nih.gov/sra); datasets SRR8742773-SRR8742775; [5]). Reads were processed and analyzed with the same tools as above. All relevant scripts for (ChIP-seq and
RNA-seq) data processing and analysis are available from https://github.com/mprevorovsky/Palecek-Nse-SAGA.

**ChIP-qPCR**

The Nse4-FLAG strains (crossed with the Δgcn5 or *gcn5-E191Q* strain; Suppl. Table ST5) were used. The untagged wild-type strain was used as a negative control. All cells were cultivated into the mid-log phase. Cells were then incubated with 1% formaldehyde for 10 min at room temperature to cross-link DNA–protein complexes. Chromatin immunoprecipitation was performed using a protocol described above for H3K9ac ChIP with the following modifications. Monoclonal anti-FLAG M2 antibody (F1804; Sigma) was diluted at 1:350 (5 µg/sample), incubated with 2 mg of total cell extract for 2 h at 4°C with rotation, and precipitated with Dynabeads protein G (Invitrogen). After overnight incubation, several washes, elution, and de-crosslinking, the DNA was purified using Phenol/Chloroform method.

The relative amount of PCR product was quantified by qPCR using SensiFAST™ SYBR® Hi-ROX Kit (Bioline). The sequences of primers used for the quantitative detection of the chromosomal loci are listed in Suppl Table ST6. Input DNA recovery was calculated as $2^{\Delta [CT(immunoprecipitate)−CT(input)]}$ and normalized to a negative locus *slx9*. Melt curve analysis was performed for each sample after PCR amplification to ensure that a single product was obtained.

**Microscopy**

For the Nse4-GFP foci number determination, cells were grown in YES medium overnight, diluted to OD = 0.4 in the morning, and treated with 0.03% MMS or 20 mM HU for 5 h at 30°C. 2.5 µl of cell culture was mounted on the slides, and GFP fluorescence was observed. Pictures were taken on the Axio Imager Z1 microscope, using a Plan-Apochromat 63x oil objective, the Axiocam CCD camera, and processed with the AxioVision software (all by
Zeiss). A minimum of 500 cells were counted in three independent experiments. For statistical
evaluation, p-values were calculated using the \( \chi^2 \) test.

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Figure legends

Figure 1. Strong genetic relation between the SMC5/6 and histone acetyltransferases

(A) Summary of significantly enriched GO categories for Biological processes of genes with genetic interactions with nse3-R254E (P < 0.005). (B) Negative log10 (P-values) evaluating the significance of the main GO Cellular component terms identified in the set of the 79 genes. Only the molecular complexes are shown. (C) Tetrad analysis of the heterozygous diploid fission yeast strains. The colony size of the nse3-R254E, Δgcn5 and nse3-R254E, Δada2 double mutant is significantly reduced (triangle). Single and double mutant alleles are indicated. (D) Ten-fold serial dilutions of the indicated strains were plated onto YES media and grown at 25°C (control) or 37°C. The nse3-R254E (nse3/RE) mutation enhanced the sensitivity of the Δada2, Δada3, and Δgcn5 mutants to the higher temperature.

Figure 2. Interactions between the SMC5/6 and SAGA complexes

(A) Extracts from fission yeast strains MMP21 (Nse4-FLAG) and YLJ507 (Nse4-FLAG and Gcn5-myc) were immunoprecipitated using the anti-myc antibody. The input (I), unbound (U) and bound (B) fractions were separated by 12% SDS-PAGE. The Nse4-FLAG and Gcn5-myc proteins were analyzed on a western blot using anti-FLAG-HRP and anti-myc-HRP, respectively. (B) The yeast two-hybrid (Y2H) system was used to determine individual protein-protein interactions between SMC5/6 and SAGA HAT module subunits. The Gal4AD or Gal4BD domains fused to the full-length Ada2 or Gcn5 subunits were co-transformed together with the fragments of SMC5/6 subunits into the PJ69 cells and grown on the plates without Leu, Trp (-L, W; control plates). The protein-protein interactions between SMC5/6 and SAGA were scored by the growth of the yeast PJ69 transformants on the plates without Leu, Trp, and His, containing 3-Amino-1,2,4-triazole (0.5 mM AT or 10 mM AT plates).
fragments were as follows: Gal4BD-Nse2 (aa2-178), Gal4AD-Nse3 (aa1-190), Gal4AD-Nse3 (aa200-307), and Gal4BD-SMC5 CC arm (aa170-225 + 837-910). In control experiments, respective empty pGADT7 (AD) or pGBK7 (BD) vector was co-transformed with either SAGA or SMC5/6 construct. Note that the Gal4BD-SMC5 CC arm construct self-activated (SMC5-vector combination) and was therefore grown on 10 mM AT plates to assess its binding to Ada2 (SMC5-Ada2 combination).

**Figure 3. SMC5/6 and SAGA are required for efficient DNA repair**

(A) Sensitivity of the SMC5/6 and SAGA mutants to genotoxins. Ten-fold serial dilutions of the yeast strains were plated onto YES media containing indicated concentrations of the hydroxyurea (HU) or methyl methane sulfonate (MMS). The double mutants were more sensitive than their respective single mutant counterparts, suggesting the non-redundant functions of SMC5/6 and SAGA in DNA repair. (B) Live-cell microscopy of endogenous Nse4-GFP upon HU and MMS treatment, respectively. The Nse4-GFP foci were present in the WT and Δgcn5 cells but largely absent in Δbrc1 cells. (C) Quantification of the data in panel B suggests that the localization of SMC5/6 to the DNA-damage foci is independent of SAGA.

**Figure 4. SMC5/6 distribution is dependent on the SAGA HAT module**

(A) The pie chart shows the distribution of the Nse4-FLAG peak areas in the different genome regions in the WT fission yeast cells. Most SMC5/6 is localized to the repetitive regions (like rDNA and centromeres), with the highest occupancy of the rDNA repeats. A significant portion of the Nse4-FLAG is localized within the intergenic regions or genes. (B) The heatmap diagrams compare the occupancy of Nse4-FLAG peaks in the WT, Δgcn5, Δada2, and Δubp8 mutant cells (as identified in the WT). The top part shows peaks enhanced in Δgcn5 and Δada2 (enhanced), while the bottom part clusters peaks reduced in the SAGA HAT module deficient cells (reduced). Peaks in the rDNA repeats are shown separately as
these chromosome regions are not fully assembled and annotated in the *S. pombe* reference genome, and exert a different range of coverage values (rDNA). (C) The pie charts show the distribution of the enhanced (top) and reduced (bottom) Nse4-FLAG peak areas in Δ*gcn5*. The SMC5/6 accumulation is mainly enhanced at the repetitive loci. The SMC5/6 localization is primarily reduced in gene regions. (D) The box plot graph compares the Nse4-FLAG occupancy in WT and Δ*gcn5* cells.

**Figure 5. The SMC5/6 accumulation correlates with the H3K9 acetylation status**

(A) Heatmap statistical analysis of the loci with reduced SMC5/6 occupancy upon the *gcn5* deletion. The Nse4-FLAG signals from WT and Δ*gcn5* are compared, and their differential plot is shown in the middle panel (Δ*gcn5*-WT). Results from ChIP-seq analysis of H3 and H3K9ac are shown (H3K9ac/H3). In addition, transcriptomic (RNA-seq) data are included (mRNA). (B) Scatter plot analysis shows a strong correlation between the drop of SMC5/6 accumulation upon *gcn5* deletion (X-axis) and H3K9-acetylation status (Y-axis) of gene regions (and their transcription levels; colour scale). (C) Results of chromatin immunoprecipitation followed by quantitative PCR (ChIP–qPCR) at selected gene loci are shown. Strains 503 (neg. control), MMP21 (WT, containing Nse4-FLAG), Nse4-FLAG Δ*gcn5* (Δ*gcn5*), and Nse4-FLAG *gcn5-E191Q* (*gcn5-E191Q*) were analyzed. The fold enrichment was calculated against the negative *slx9* locus (mean ± standard deviation, n ≥3 biological replicates).
**A. Biological Processes**

| p-value     | Description                                                                 |
|-------------|----------------------------------------------------------------------------|
| 3.27E-13    | DNA metabolic process                                                      |
| 1.69E-12    | response to DNA damage stimulus                                            |
| 4.47E-08    | DNA repair                                                                 |
| 1.79E-07    | cell cycle checkpoint                                                      |
| 1.66E-06    | post-translational protein modification                                    |
| 7.32E-06    | chromosome organization                                                    |
| 2.50E-05    | histone acetylation                                                        |
| 6.27E-05    | meiosis                                                                    |
| 1.15E-04    | DNA replication                                                            |
| 1.20E-04    | chromatin remodeling                                                       |
| 2.61E-04    | base-excision repair                                                       |
| 7.50E-04    | nucleotide-excision repair                                                 |
| 1.33E-03    | telomere organization                                                      |
| 4.50E-03    | positive regulation of ubiquitin-protein ligase activity                   |

**B. Molecular Complexes**

- histone acetyltransferase complex
- SAGA complex
- ribonuclease H2 complex
- NuA4 histone acetyltransferase complex
- H4/H2A histone acetyltransferase complex
- nucleotide-excision repair complex
- SMC5/6 complex

![Graph showing molecular complexes with -log(p) values](image)

**C. Δgcn5 x nse3/RE**

- △ WT
- □ nse3/R254E mutant
- ○ single deletion mutant
- △ double mutant

**D.**

|       | Ctrl (25°C) | 37°C |
|-------|-------------|------|
| WT    |             |      |
| Δgcn5 |             |      |
| Δada2 |             |      |
| Δada3 |             |      |
| Δsgf29|             |      |
| nse3/RE|            |      |
| Δgcn5, nse3/RE|    |      |
| Δada2, nse3/RE| |      |
| Δada3, nse3/RE|   |      |
| Δsgf29, nse3/RE| |      |
A.

|        | control | Gcn5-myc |
|--------|---------|----------|
| I      | U       | B        |

Gcn5-myc

Nse4-FLAG

B.

- LW

0.5mM AT

10mM AT

ADA vector

BD vector

Nse2 (aa2-178)

Nse3 (aa1-190)

Nse3 (aa200-307)

Smc5 CC arm

Ada2

Gcn5

vector
A. Nse4 distribution

- rDNA repeats
- centromeric regions
- intergenic regions
- gene regions

B. Diagram showing the effects of WT, Δgcn5, Δada2, and Δubp8 on rDNA, enhanced, unchanged, and reduced regions.

C. Enhanced and reduced regions with gene regions, repeats, and intergenic regions highlighted.

D. Box plot showing Nse4 occupancy in WT and Δgcn5 for rDNA, centromeric regions, intergenic regions, and gene regions.
A. Nse4-FLAG

Gene regions with reduced Nse4 occupancy in Δgcn5

WT | Δgcn5 | Δgcn5-WT | H3K9ac/H3 | mRNA

0.5 gene 1.0Kb | 0.5 gene 1.0Kb | 0.5 gene 1.0Kb | 0.5 gene 1.0Kb

B. n=234, Pearson’s r=-0.296, p=3.97e-6

H3K9ac/H3 vs. mRNA

C. ChIP-qPCR

Fold enrichment

- vht1
- rps502
- fip1
- ser3
- tef3

neg. control
WT
Δgcn5
gcn5-E191Q