

Structural characterization of the Asf1–Rtt109 interaction and its role in histone acetylation

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ABSTRACT

Acetylation of histone H3 at lysine-56 by the histone acetyltransferase Rtt109 in lower eukaryotes is important for maintaining genomic integrity and is required for *C. albicans* pathogenicity. Rtt109 is activated by association with two different histone chaperones, Vps75 and Asf1, through an unknown mechanism. Here, we reveal that the Rtt109 C-terminus interacts directly with Asf1 and elucidate the structural basis of this interaction. In addition, we find that the H3 N-terminus can interact via the same interface on Asf1, leading to a competition between the two interaction partners. This, together with the recruitment and position of the substrate, provides an explanation of the role of the Rtt109 C-terminus in Asf1-dependent Rtt109 activation.

INTRODUCTION

Eukaryotic DNA is packaged into a protein–DNA complex called chromatin. Nucleosomes, consisting of two copies each of histones H2A, H2B, H3 and H4, are the basic packing unit of chromatin, binding ~145 bp of DNA (1). Histones can undergo post-translational modifications (PTMs), which play an important role in a multitude of processes including transcriptional regulation and DNA repair (2). Incorporation of PTMs begins immediately after synthesis (3), when histones are bound to both general and histone-specific chaperones (3,4).

Asf1 is a histone-specific chaperone necessary for both histone processing and deposition (5–8), as well as transcriptional regulation (9) and DNA repair (5,10). It binds the H3:H4 hetero-dimer, thereby preventing formation of the (H3:H4)₂ tetramer (11,12).

In lower eukaryotes, newly synthesized histone H3 is acetylated on lysine-56 (K56) by Rtt109 and on lysine-9 (K9) by Rtt109 and Gcn5 (8,13–14). The role of K9 acetylation is largely unknown, while K56 acetylation has been shown to be important for genomic stability (15) and is

required for recognition of the (H3:H4)₂ tetramer by the downstream chaperone Rtt106 (16).

Rtt109 is a fungal histone acetyltransferase with only remote homology to human p300 and is required for *Candida albicans* pathogenesis (17). The lack of close homologues in higher eukaryotes makes it an attractive target for antifungal therapy, although recent high throughput screens have failed to identify suitable lead candidates (18–20). Alone, Rtt109 has very low acetyltransferase activity, but is strongly activated by the two histone chaperones Asf1 and Vps75 (13–14,21–22). Depending on chaperone association and substrate, different levels of activation and specificity have been observed *in vitro* (23,24). In addition, Rtt109 has been reported to regulate transcription together with Asf1 via an acetylation-dependent pathway that does not target H3:H4 (25).

Previously, it was reported that the Rtt109 C-terminus is required for Asf1-dependent activation of Rtt109-mediated acetylation of K56 *in vitro* (26). *In vivo*, the C-terminal tail of Rtt109 is necessary for K9 acetylation, but K56 acetylation is impaired only by the concurrent absence of both the Rtt109 C-terminus and Vps75 (26). This data prompted us to study the functional role of the Rtt109 C-terminus in the activation of acetylation. While Rtt109 forms a tight complex with Vps75 (27,28), its interaction with Asf1 has remained elusive. Here, we show that in *Saccharomyces cerevisiae* the Rtt109 C-terminus directly interacts with Asf1 via a surface shared by multiple Asf1 interaction partners (29,30). Interestingly, the same surface can also interact with the H3 tail in the Asf1–H3:H4 complex, indicating a role for Asf1 in chaperoning the H3 tail. Using NMR methods designed for the study of high-molecular-weight complexes, we show that the interaction between the Rtt109 C-terminus and Asf1 is retained when Rtt109 is bound to the Vps75 homodimer (Vps75)₂ and when Asf1 forms a heterotrimer with H3:H4. These findings suggest that the interaction of Asf1 with the Rtt109 C-terminus contributes to Rtt109 activation both *in vitro* and *in vivo*, where Rtt109 is in complex with Vps75.

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MATERIALS AND METHODS

Protein expression and purification

Plasmids for *Xenopus laevis* histone H3/H4 (pET3d/a), *Saccharomyces cerevisiae* Vps75 (pET28a) and *Saccharomyces cerevisiae* Asf1 (2–169, pGEX6P-2) were a kind gift from K. Luger (Colorado State).

Asf1^{2–169} was sub-cloned into pETM11, containing an N-terminal hexa-histidine tag, followed by a TEV protease cleavage site. The plasmid was transformed into *Escherichia coli* BL21(DE3) and expression was induced by addition of 0.5 mM IPTG at OD₆₀₀ = 0.6–0.8 and continued overnight at 16°C. Cells were lysed by sonication in lysis buffer (1× phosphate-buffered saline (PBS) + 500 mM NaCl, 10 mM imidazole, 5 mM BME (β-mercaptoethanol), 1× protease inhibitor (Roche; 11697498001)). The protein was purified by immobilized-metal-ion affinity-chromatography (IMAC, GE Healthcare HisTrap HP 5 ml). The column was equilibrated and washed with wash buffer (25 mM Tris–HCl pH 7.5, 500 mM NaCl, 5 mM BME, 10 mM imidazole) and eluted with a gradient to 75% elution buffer (25 mM Tris–HCl pH 7.5, 500 mM NaCl, 5 mM BME, 500 mM imidazole) over 15 CV (column volumes). The fractions containing the tagged protein were pooled and the his-tag removed by overnight incubation with TEV protease. To remove the tag and the TEV protein, the solution was passed over a HisTrap HP column equilibrated in wash buffer and the flow-through collected. Cleaved Asf1 was further purified by ion-exchange (GE Healthcare HisTrap Q, 25 mM Tris–HCl pH 7.5, 100–1000 mM NaCl, 5 mM BME) and size-exclusion chromatography (Superdex S200 16/60; 25 mM Tris–HCl pH 7.5, 500 mM NaCl, 5 mM BME). Full-length human Asf1b was purified identically to *S. cerevisiae* Asf1. Vps75 was purified identically to Asf1 except that different buffers were used during the first HisTrap step (wash buffer: 25 mM Tris–HCl pH 7.5, 500 mM NaCl, 5 mM BME, 20 mM imidazole; elution buffer: 25 mM Tris–HCl pH 7.5, 500 mM NaCl, 5 mM BME, 400 mM imidazole).

¹⁵N- and ¹⁵N,¹³C-labeled proteins were expressed in M9 minimal medium containing ¹⁵NH₄Cl or ¹⁵NH₄Cl and ¹³C glucose and purified as described above. Perdeuterated and ¹H,¹³C methyl isoleucine-, leucine- and valine-labeled Asf1 (Val-[¹³CH₃]⁷, Leu-[¹³CH₃]⁸, Ile-[¹³CH₃]^{δ1}) was obtained by expression in *E. coli* BL21(DE3) grown in M9 minimal medium in 100% D₂O. α-ketobutyric acid sodium salt (60 mg/l, Sigma, 589276) and α-ketoisovaleric acid sodium salt (120 mg/ml, Sigma, 691887) were added 45 min before induction at OD₆₀₀ = 0.6–0.8.

Codon-optimized full-length *Saccharomyces cerevisiae* Rtt109 was cloned in a pETM-11 vector containing a N-terminal hexa-histidine tag, followed by a TEV protease cleavage site. Rtt109 was expressed in *E. coli* BL21(DE3) and purified as previously described (31). *Xenopus laevis* histones were expressed and purified as reported by Luger *et al.* (32). Labeled histones were expressed in M9 minimal medium containing ¹⁵NH₄Cl and purified identically to the unlabeled proteins.

Peptides were synthesized by the Peptide Synthesis Unit of the Helmholtz Center for Infection Research (Braunschweig) under the supervision of Dr Werner Tegge.

Microscale thermophoresis measurement

Asf1 was labeled with NT-647 via an NHS coupling using a kit from Nanotemper (MO-L0001) following the manufacturer's instructions. Thermophoresis temperature-jump titration series were performed in standard capillaries (Nanotemper, MO-K002) using 100 nM labeled Asf1 and concentrations of 0.058–475 μM of Rtt109 (2× serial dilutions) and 0.228–590 μM of Rtt109–Vps75₂ (2× serial dilutions). Experiments were performed at 25°C, in 50 mM citrate buffer at pH 6.5, 150 mM NaCl, 5 mM BME, with an excitation power of 20% and an MST power of 10–20%. Measurements were performed in triplicates, K_D values were calculated independently and used to obtain the average and error values. Higher MST powers resulted in poor-quality curves at higher Rtt109 concentrations, probably due to Rtt109 aggregation. This effect was not observed for the Rtt109–Vps75₂ complex. Data was analyzed using MO.Affinity Analysis (Nanotemper, v2.1.2333).

Isothermal titration calorimetry (ITC)

ITC measurements were performed on a NANO ITC (TA Instruments) in PBS pH 7.4 with 5 mM BME at 20°C using an equilibration time of 3000 s and a stir-rate of 200 rpm. The cell concentration was 185 μM Asf1 (in a volume of 220–230 μl) and the peptide concentration in the syringe was 2.78 mM for Rtt109^{419–436} and 2.76 mM for Rtt109^{419–428}. For the measurement of the Rtt109^{419–433} peptides, the cell concentration was 160 μM Asf1 (in a volume of 240–250 μl) and the peptide concentration in the syringe was 1.95 mM for Rtt109^{419–433}, 2.75 mM for Rtt109^{419–433} RK427DD and 1.45 mM for Rtt109^{419–433} KK428DD. The first injection of 0.48 μl was not considered for the analysis. Twenty to twenty-two injections of 1.98 μl of peptide followed, with 300 s between injections. Data was analyzed using NanoAnalyze (TA Instruments, v3.6.0). Data points that suffered from baseline distortions were not included in the analysis.

Nuclear magnetic resonance spectroscopy

NMR data were acquired on Bruker Avance III HD 850 MHz and 600 MHz spectrometers equipped with He-cooled (TCI) and N₂-cooled (CPP) cryogenic inverse HCN probe-heads, respectively. All experiments were performed at 298 K.

Asf1 data collection and assignment

2D ¹H–¹⁵N and ¹H–¹³C (ILV-methyl) correlation spectra were acquired using ¹⁵N-HSQC (33–35) and ¹³C-HMQC (methyl-TROSY) (36–39) pulse sequences. All assignment experiments were performed in PBS pH 7.4 with 5 mM BME for free Asf1 and PBS pH 6.2 with 5 mM BME for peptide-bound Asf1. Free and peptide-bound Asf1 backbone resonances were assigned using standard TROSY-HNCA, TROSY-HNCACB, TROSY-HN(CO)CACB and

TROSY-HNCO experiments (40,41) with a deuterated and ^{13}C , ^{15}N -labeled Asf1 sample at 500 μM protein and 3 mM Rtt109 peptide. All other experiments for structure determination were recorded using ^{13}C , ^{15}N -labeled Asf1. Side-chain resonances for the peptide-bound Asf1 were assigned using an HC(C)H-TOCSY experiment (42,43) and assignment of the aromatic resonances was aided by an (HB)CB(CGCD)HD spectrum (44). ILV methyl resonances in the Asf1–H3:H4 complex (Asf1 perdeuterated and ^1H , ^{13}C methyl-ILV-labeled; H3:H4 unlabeled) were assigned using a 4D ^{13}C -HMQC–NOESY– ^{13}C -HMQC experiment (45,46) recorded with non-uniform-sampling. The sampling schedule was generated using the ist@HMS online schedule generator (<http://gwagner.med.harvard.edu/intranet/istHMSv2/index.html>). The 4D spectrum was reconstructed with iterative-soft-thresholding as implemented in the istHMS software (47–49). Distances for structure calculation of the Asf1–Rtt109^{419–433} complex were derived from 3D NOESY– ^{15}N -HSQC and NOESY– ^{13}C -HSQC spectra (120 ms NOESY mixing time) (50–53). Intermolecular Asf1–Rtt109^{419–433} NOEs in the NOESY– ^{13}C -HSQC spectrum were identified with the assistance of a 3D ^{13}C , ^{15}N -double-filtered-NOESY– ^{13}C -HSQC experiment (54). Intramolecular Rtt109^{419–433} NOEs were measured from a 2D ^1H -NOESY (150 ms NOESY mixing time) spectrum (55–57) recorded on a sample containing 150 μM Rtt109^{419–433} and 500 μM perdeuterated and ^{13}C , ^{15}N -labeled Asf1. Because the Rtt109 peptide has only moderate affinity for Asf1 and is in fast exchange, the intermolecular Asf1–Rtt109^{419–433} resonances observed in the 3D ^{13}C , ^{15}N -double-filtered-NOESY– ^{13}C -HSQC spectrum of Asf1 in the presence of excess Rtt109 peptide correspond closely to the free form of the peptide. To assign these resonances, we recorded 2D ^1H -NOESY (150 ms NOESY mixing time) and ^1H -TOCSY (70 ms DIPSI-2 mixing) experiments (57–59) with 3 mM Rtt109^{419–433} peptide in the presence of 500 μM perdeuterated and ^{13}C , ^{15}N -labeled Asf1.

Non-exchangeable protons were identified by rapidly buffer-exchanging ^{15}N -labeled Asf1 into 100% D_2O buffer (15 mM NaH_2PO_4 pH 6.2, 150 mM NaCl, 5 mM BME) using a HiTrap desalting column (GE Healthcare). Backbone amide ^1H – ^{15}N RDCs were determined by measuring the ^1H – ^{15}N splitting in isotropic and anisotropic samples from the difference in the ^{15}N frequencies of the amide peaks in the ^{15}N -HSQC spectrum and a ^{15}N -TROSY-HSQC spectrum recorded with $\text{H}^\alpha/\text{H}^\beta$ band-selective decoupling (60,61). The anisotropic sample was prepared by addition of Pfl bacteriophage (ASLA biotech) (62) to a concentration of 8.7 mg/ml (giving a residual ^2H quadrupolar splitting of the HOD signal of 7.3 Hz).

Titration experiments were performed with 50–100 μM Asf1 with the indicated equivalents of interaction partners.

Data processing and analysis

NMR data was processed with Topspin (Bruker), istHMS and nmrPipe (63) and analysed using CcpNmr Analysis (64). The magnitude (D_a) and rhombicity (R) of the alignment tensor were derived by fitting the experimental RDCs to PDB IROC using singular-value decomposition as im-

plemented in PALES (65). CSPs were calculated as $d = \sqrt{\frac{1}{2}(\Delta\delta_H^2 + (0.15 \cdot \Delta\delta_N)^2)}$ or $d = \sqrt{\frac{1}{2}(\Delta\delta_H^2 + (0.09 \cdot \Delta\delta_C)^2)}$.

Structure calculation and analysis

Structure calculations were performed using ARIA v2.3/CNS v1.21 (66,67) via the standard ARIA v2 iterative-assignment protocol. Distance restraints were generated from NOESY peak heights with application of spin-diffusion correction for the calibration factors. For the ^{15}N -edited NOESY spectrum, peak heights were scaled according to the intensities of the corresponding amide peaks in the ^{15}N HSQC spectrum prior to generation of distance restraints. Amide-to-carbonyl hydrogen-bond restraints were applied for non-exchangeable amide protons. RDC restraints for backbone amide bonds of residues in canonical secondary structure elements were applied using the SANI potential of CNS ($D_a = 7.2$ Hz; $R = 0.52$). Dihedral-angle restraints for backbone ϕ and ψ angles were generated from backbone chemical shifts using TALOS-N (68). Restraints were applied for residues where the TALOS-N classifications was ‘strong’ or ‘generous’, using respective error ranges of twice and three times the uncertainty reported by TALOS-N. Initial structure calculations indicated a conflict between the hydrogen-bond and dihedral restraints for residue F106. A calculation where both types of restraints were omitted for this residue revealed a well-defined conformation in good agreement with the predicted hydrogen bond (and differing from the TALOS-predicted dihedral restraints). Therefore the dihedral restraints for F106 were removed for the final structure calculation. The numbers of steps for the hot, cool1, cool2 and refine stages of the molecular-dynamics simulated-annealing protocol were 40 000, 20 000, 16 000 and 16 000, respectively. All distance restraints were used in every iteration. One hundred structures were calculated in the final iteration, of which the 10 lowest-energy structures were refined in explicit water to generate the final structural ensemble.

Structural quality reports and statistics were generated using the PSVS web-server (69). Buried surface area was determined using the Intersurf tool (prune distance: 6.0, molecule bias: 1.0) and the measure volume and area tools in Chimera (70). Figures were prepared using gnuplot, CcpNmr Analysis (71) and Pymol (The PyMOL Molecular Graphics System, Version 1.8 Schrödinger, LLC).

Acetylation assays

Asf1–H3:H4, Vps75₂–Rtt109 and Vps75₂–Rtt109^{1–424} complexes were reconstituted by size-exclusion chromatography (Superdex S200 10/300; 10 mM HEPES pH 8.0, 100 mM NaCl, 5 mM BME for complexes with Rtt109, 10 mM HEPES pH 8.0, 500 mM NaCl, 5 mM BME for Asf1–H3:H4). Acetylation reactions were performed using 0.3 μM Rtt109–Vps75₂ or Rtt109, 0.3 μM Asf1–H3:H4 and 3 μM Ac-CoA in 10 mM HEPES pH 8.0, 100 mM NaCl, 5 mM BME. Reactions were stopped with 0.5 M sodium acetate, pH 5.5; 1.6 μl of the reaction mixtures were spotted on a nitrocellulose membrane. Membranes

were air-dried for ca. 30 min, blocked for 1 h in 5% BSA–TBST buffer (20 mM Tris–HCl pH 7.5, 150 mM NaCl, 0.05% Tween-20), and subsequently incubated for 1 h with a 1:2000 dilution of rabbit anti-histone H3 (acetyl K9) primary antibody (ab4441 Abcam, lot no. GR304381–1) in TBST with 0.1% BSA. After washing three times with TBST buffer, the membranes were incubated for 1 h with the secondary antibody, an anti-rabbit IgG (whole molecule)–alkaline phosphatase (A3687 Sigma, lot no. SLBK3154V) solution diluted 1:10 000 with 0.1% BSA and TBST. After three washes with TBST buffer, the blots were developed with BCIP®/NBT Liquid Substrate System (B1911 Sigma, lot no. SLBQ3317V).

The density of the dots was quantified using the ImageJ software (71). After background subtraction, the measured intensities were normalized to the negative control (Asf1–H3:H4 alone). The results are the average of three experiments with the standard deviation representing the error.

RESULTS

Asf1 interacts with the C-terminus of Rtt109

We characterized the interaction of Rtt109 and Asf1 by NMR spectroscopy (Figures 1 and 2, Supplementary Figures S1–S3). Rtt109 is prone to aggregation and precipitation in commonly used buffers. In previous work, we used a thermal-shift assay to optimize Rtt109 stability and found optimal conditions in a buffer containing citric acid at low pH (5.5–6.5) (31). Using this buffer, we titrated ¹⁵N-labeled Asf1 with increasing amounts of unlabeled full-length Rtt109 and observed significant changes in the Asf1 ¹⁵N-HSQC spectrum (33–35) (Figure 1A and Supplementary Figure S2A). Mapping these chemical shift perturbations (CSPs) on the structure of Asf1 revealed that they cluster on the surface opposite to the histone interaction surface (Figure 1B), suggesting that the interaction of Asf1 with Rtt109 is compatible with histone binding. Next we performed an identical titration with a truncated Rtt109 construct (1–424) lacking the C-terminal 12 amino acids (26). In contrast to full-length Rtt109, only small changes in the Asf1 spectrum were observed upon addition of Rtt109^{1–424} (Figure 1A), indicating that the interaction between Asf1 and Rtt109 is dependent on the C-terminus of Rtt109. Similarly, when we added a peptide corresponding to the C-terminal 18 residues of Rtt109 (Rtt109^{419–436}) to labeled Asf1 (Figure 1C and Supplementary Figure S2B), we observed large CSPs, which mapped onto the same Asf1 surface as those from adding full-length Rtt109 (Figure 1D).

To better define the residues of Rtt109 responsible for the interaction with Asf1, we performed NMR titrations with successively shorter peptides derived from the Rtt109 C-terminus. Rtt109^{419–433} interacted with Asf1 in an identical manner to Rtt109^{419–436} (Figure 2A and C). Rtt109^{419–428} showed a somewhat changed pattern of CSPs (Figure 2A and C), while deletion of the 10 C-terminal amino acids (Rtt109^{419–426}) completely abolished the interaction with Asf1 (Figure 2A and C).

Next, we used isothermal titration calorimetry (ITC) (72) and microscale thermophoresis (MST) (73) to quantify the interactions of Asf1 with the Rtt109 C-terminal peptides

and full-length Rtt109 (Figure 2D, Supplementary Figures S4 and S5). The ITC titrations revealed comparable K_D values for Rtt109^{419–436} and Rtt109^{419–433} ($6.91 \pm 1.74 \mu\text{M}$ and $7.78 \pm 1.23 \mu\text{M}$, respectively), which increased to $44.1 \pm 6.93 \mu\text{M}$ for the shorter peptide Rtt109^{419–428}. MST titrations with full-length Rtt109 gave a K_D of $7.63 \pm 1.31 \mu\text{M}$, in agreement with the hypothesis that the C-terminal residues of Rtt109 are solely responsible for the interaction with Asf1. In the presence of Vps75₂, the affinity of full-length Rtt109 for Asf1 was slightly weaker ($K_D = 42.7 \pm 1.27 \mu\text{M}$).

Structural characterization of the Asf1–Rtt109 interaction

We used NMR to solve the structure of Asf1 in complex with the Rtt109 C-terminus (⁴¹⁹LAITMLKPRKKAKAL⁴³³) (Figure 3 and Table 1). The Rtt109 C-terminal peptide structure was defined to a precision of 0.69 Å (backbone root-mean-square deviation, RMSD, calculated over residues 421–429 for the 10 lowest-energy structures) by 231 (199 unambiguous) intrapeptide NOEs (derived from a 2D ¹H NOESY spectrum (55–57) of the peptide in complex with perdeuterated Asf1) and 159 (67 unambiguous) intermolecular Asf1–peptide NOEs (assigned using a 3D ¹³C,¹⁵N-filtered-NOESY–¹³C-HSQC spectrum (54) of the peptide in complex with ¹³C,¹⁵N-labeled Asf1). A further 6732 (4117 unambiguous) Asf1 intramolecular NOEs derived from standard 3D NOESY–¹⁵N-HSQC and NOESY–¹³C-HSQC spectra (50–53) defined the complex to a precision of 0.47 Å (backbone RMSD calculated over residues 2–47 and 54–156 of Asf1 and residues 421–429 of Rtt109^{419–433} for the 10 lowest-energy structures). Comparison with a crystallographic structure of Asf1 (PDB ID: 1ROC) demonstrates that the Asf1 conformation is not significantly perturbed by Rtt109 binding (backbone RMSD of 0.80 Å for residues 1–155).

The peptide interacts with a hydrophobic patch composed of residues 60–72 (⁶⁰ILVGPVPGVGNKF⁷²) on the side of Asf1 opposite to the histone interaction surface, resulting in a buried surface area of 739 Å². This interaction interface is identical to the one identified by CSP analysis of Asf1 titrated with either full-length Rtt109 or Rtt109^{419–436} (Figures 1 and 2). Residues T422, L424 and P426 of the Rtt109 peptide are largely responsible for the hydrophobic interactions (Figure 3C). In addition, the basic residues R427, K428 and K429 participate in ionic interactions with the negatively charged surface adjacent to the hydrophobic patch, contacting residues D37, D58 and E39, respectively. The interaction is further stabilized by a hydrogen bond between the carbonyl group of peptide residue R427 and the amide group of Asf1 residue L61 (Figure 3C).

To validate this structure we generated two Rtt109^{419–433} mutants with two aspartate substitutions at either R427 and K428 or K428 and K429, referred to as Rtt109^{419–433} RK427DD and Rtt109^{419–433} KK428DD respectively. ITC measurements showed a complete loss of binding affinity for both mutant peptides, confirming that the ionic interactions mediated by these residues are necessary for Asf1 recognition (Supplementary Figure S4).

Next, we compared the Asf1–Rtt109^{419–433} structure with those available for Asf1 in complex with other interaction

Table 1. Assignment and restraint statistics and structural quality parameters for the structure of the Asf1²⁻¹⁶⁹–Rtt109⁴¹⁹⁻⁴³³ complex. For the structural quality parameters, the first column (headed (SA)) contains average values for the ensemble (standard deviations in parentheses), while the second column (headed (SA)_c) contains the corresponding values for the structure closest to the mean. Structural RMSDs were calculated over the well-ordered residues, defined as residues 2–47 and 54–156 of Asf1²⁻¹⁶⁹ and residues 421–429 of Rtt109⁴¹⁹⁻⁴³³

Resonance assignments		
All protons	98.9%	
Asf1 ²⁻¹⁶⁹	98.8%	
Rtt ⁴¹⁹⁻⁴³³	100.0%	
Asf1 ²⁻¹⁶⁹ backbone atoms (N, C ^α , C', H ^N , H ^α , C ^β)	94.9%	
NMR restraints		
NOE distance restraints	7121	
Unambiguous	4383	
Classified by sequence-separation:		
Intra-residue	1584	
Sequential	1051	
Short-range	410	
Medium-range	73	
Long-range (incl. inter-molecular)	1265	
Classified by intra-/inter-molecular:		
Inter-molecular	67	
Intra-Asf1 ²⁻¹⁶⁹	4117	
Intra-Rtt109 ⁴¹⁹⁻⁴³³	199	
Ambiguous	2738	
Classified by sequence-separation:		
Intra-residue	375.7	
Sequential	562.9	
Short-range	368.5	
Medium-range	98.3	
Long-range (incl. inter-molecular)	1332.6	
Classified by intra-/inter-molecular:		
Inter-molecular	91.9	
Intra-Asf1 ²⁻¹⁶⁹	2614.6	
Intra-Rtt109 ⁴¹⁹⁻⁴³³	31.5	
Hydrogen-bond restraints	81	
Dihedral-angle restraints	270	
φ angle	135	
ψ angle	135	
RDC restraints (N–H)	61	
Structural quality	(SA)	(SA) _c
RMSDs from experimental restraints		
NOE distances (Å)	0.0216 (0.0005)	0.0217
H-bond distances (Å)	0.0134 (0.0029)	0.0130
Dihedral angles (°)	0.817 (0.056)	0.825
Violations		
NOE distances > 0.5 Å	0 (0)	0
NOE distances > 0.3 Å	2.3 (0.9)	3
Dihedral angles > 10°	0 (0)	0
Dihedral angles > 5°	0.8 (0.9)	0
RMSDs from ideal geometry		
Bond lengths (Å)	0.00420 (0.00011)	0.0044
Bond angles (°)	0.5472 (0.0087)	0.5421
Improper angles (°)	1.592 (0.078)	1.613
Final energy (kcal mol ⁻¹)	–6914 (82)	–7006
Structural RMSDs to mean structure		
Backbone atoms (Å)	0.47 (0.07)	0.38
Heavy atoms (Å)	0.80 (0.11)	0.70
Ramachandran plot summary (PROCHECK)		
Most favoured regions	88.2%	
Additionally allowed regions	9.8%	
Generously allowed regions	0.5%	
Disallowed regions	1.6%	
Global quality scores (–Raw/Z-score)		
Verify3D	0.35/–1.77	
ProsaII	0.30/–1.45	
Procheck	–0.54/–3.19	
Molprobrity Clashscore	34.34/–4.37	

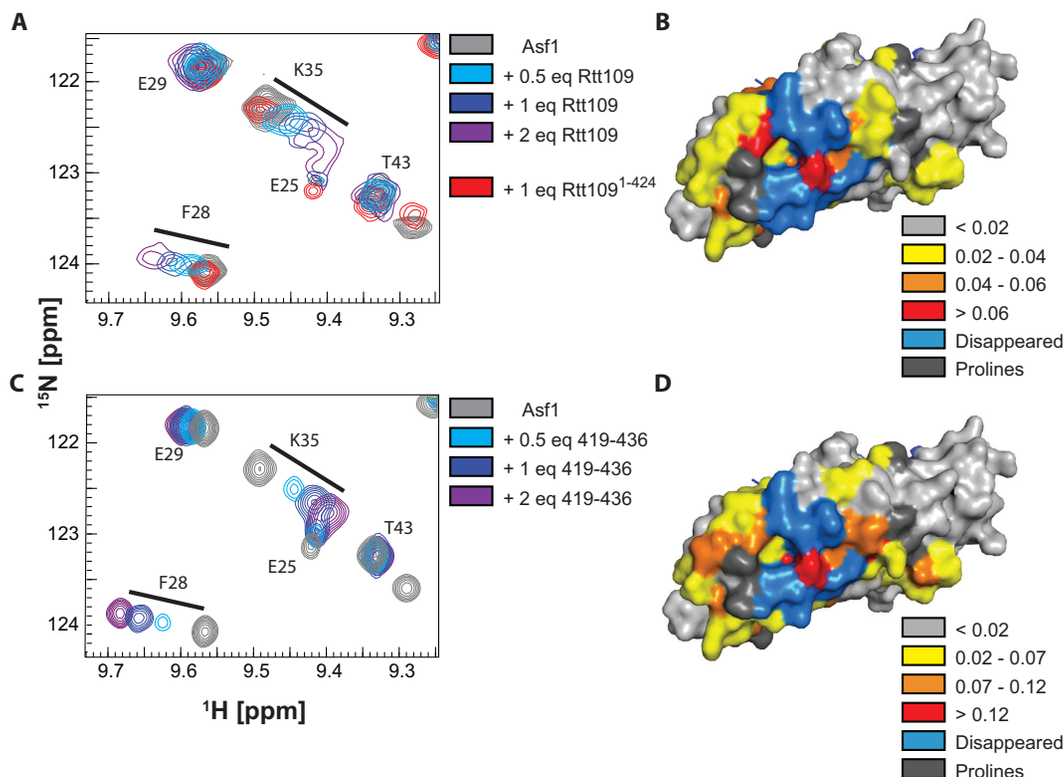


Figure 1. Rtt109 interacts with Asf1 via its C-terminus. (A) Overlay of ^{15}N -HSQC spectra of Asf1 (100 μM) in the presence of Rtt109 and Rtt109 $^{1-424}$. The target concentrations of full-length Rtt109 corresponded to 0.5, 1 and 2 equivalents of Rtt109, but the actual achieved concentrations (estimated from 1D ^1H spectra) were lower due to partial precipitation of Rtt109 in the NMR tube. The spectra were acquired in 50 mM sodium citrate pH 6.5, 150 mM NaCl, 5 mM BME at 600 MHz and 298 K. (B) Mapping of the CSPs observed upon addition of 1 equivalent of full-length Rtt109 onto the Asf1 structure in complex with the H3:H4 dimer (PDB ID 2HUE). (C) Overlay of ^{15}N -HSQC spectra of Asf1 (100 μM) in the presence of Rtt109 $^{419-436}$. The spectra were acquired in 50 mM sodium citrate pH 6.5, 150 mM NaCl, 5 mM BME at 850 MHz and 298 K. (D) Mapping of the CSPs observed upon addition of 1 equivalent of Rtt109 $^{419-436}$ onto the Asf1 structure as in (B). The chemical shifts perturbations observed upon addition of Rtt109 $^{419-436}$ were larger than those observed upon addition of full-length Rtt109, which can be recapitulated by the on-going precipitation of full-length Rtt109 during the NMR experiment. Asf1 is in surface representation.

partners that engage the same surface (HIRA, Cac2, Rad53 and Hip1). We identified a common binding motif consisting of $\Phi\text{X}\Phi\text{R}+$, in which Φ and $+$ represent a small hydrophobic amino acid and a positively charged residue, respectively (Figure 3D). Interestingly, the Rtt109 peptide is reversed with respect to the other Asf1 interaction partners (Supplementary Figure S6A). The salt-bridge observed between D37 and R427 in our structure is conserved in all other Asf1-peptide complexes (Supplementary Figure S6B).

The Asf1–Rtt109 interaction in the presence of histone H3:H4 dimer

Rtt109 probably interacts with Asf1 while it is in complex with the Rtt109 substrate, the histone H3:H4 dimer. On the other hand, Rtt109 also forms a tight complex with Vps75 $_2$. To determine whether the interaction between the Rtt109 C-terminus and Asf1 is retained in higher-order complexes, we produced Asf1 with ^1H , ^{13}C -methyl-labeled Ile, Leu and Val residues in a perdeuterated background (74). This labeling scheme yields high-quality methyl group spectra for high-molecular-weight complexes that would otherwise be inaccessible by solution NMR methods (36,75). The Asf1–

H3:H4 complex is prone to aggregation at high concentration in low-salt conditions, but by using a citrate buffer (pH 6.5), which also increases Rtt109 stability, we obtained a soluble complex at physiological ionic strength. 60% of the Asf1 methyl resonances in the Asf1–H3:H4 complex could be assigned by analysis of a 4D ^{13}C -HMQC–NOESY– ^{13}C -HMQC spectrum (45,46) in conjunction with the Asf1 structure.

To investigate the Rtt109–Asf1 interaction in the presence of both Vps75 and histones, we first titrated the Asf1–H3:H4 complex with unlabeled Vps75 $_2$. Surprisingly, we observed noticeable CSPs for several residues, which clustered on the same surface engaged by the Rtt109 C-terminal peptide (Figure 4A and Supplementary Figure S7). In contrast, residues belonging to the Asf1 surface in contact with the core of the H3–H4 dimer did not display any CSPs upon addition of Vps75 $_2$ (e.g. L6 in Figure 4A). Interestingly, adding unlabeled Vps75 $_2$ to Asf1 alone did not give any CSPs, suggesting that Vps75 does not interact directly with Asf1 (Supplementary Figure S8). To explain these observations, we reasoned that Vps75 modulates the interaction between histones and Asf1, leading to CSPs of Asf1 resonances in the Asf1–H3:H4 complex upon addition of Vps75 $_2$. Because the region of Asf1 showing these CSPs is

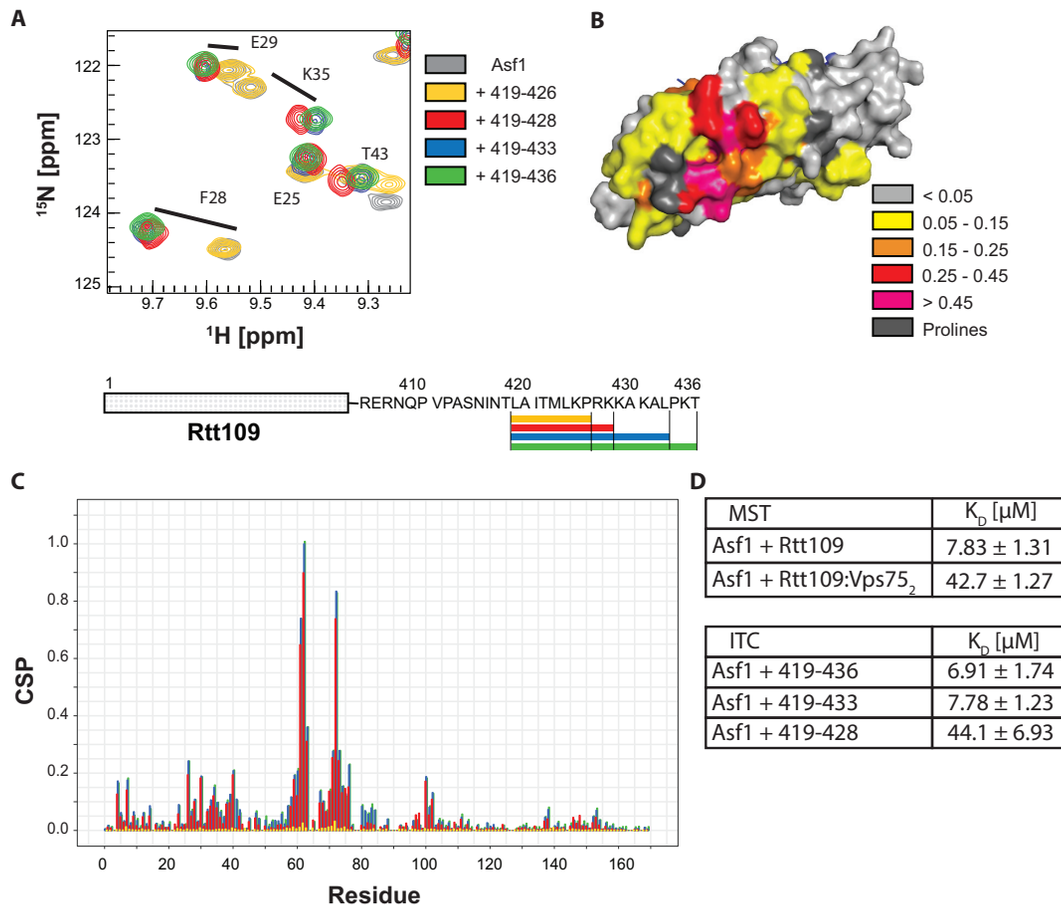


Figure 2. Rtt109 interacts with Asf1 via its C-terminus. (A) Overlay of ^{15}N -HSQC spectra of Asf1 (100 μM) in the presence of Rtt109-derived peptides (all at 3 mM) of various lengths. The spectra were acquired in PBS pH 6.2 with 5mM BME at 850 MHz and 298 K. (B) Mapping of the CSPs observed upon addition of 3 mM Rtt109⁴¹⁹⁻⁴³⁶ onto the Asf1 structure in complex with the H3:H4 dimer (from PDB ID 2HUE). (C) CSP analysis of the spectra in (A). (D) Results of ITC and MST titrations of Asf1 with different peptides and proteins. Experiments were performed in triplicate.

located far from the canonical histone interaction interface, we hypothesized that the long unstructured histone H3 tail folds back onto Asf1 in the Asf1–H3:H4 complex. To test this hypothesis, we reconstituted the H3:H4 complex using truncated H3 that lacks the N-terminal 28 aa. The chemical shift perturbations observed for the Asf1–H3:H4–Vps75₂ and Asf1–H3(Δ 28):H4 complexes relative to Asf1–H3:H4 were very similar (Figure 4A), providing strong evidence that Vps75 binding results in release of the H3 tail from Asf1. In addition, we titrated the H3:H4 complex containing ^{15}N -labeled H3 with unlabeled Asf1. As expected, we observed CSPs (Figure 4B) for H3 residues 19–26, confirming the interaction between the H3 tail and Asf1. When Rtt109 is added to this mixture, the CSPs of the H3 tail relative to free H3:H4 are reduced, showing that Rtt109 can out-compete the H3 tail for the same interaction surface on Asf1 (Figure 4B). Having discovered a previously unknown H3–Asf1 interaction, we wondered whether the same interaction mode is also preserved in higher eukaryotes. To address this question, we recorded spectra of the H3:H4 complex containing ^{15}N -labeled H3 in the presence of human Asf1b. Interestingly, similar chemical shifts were observed for H3 in both the Asf1 and hAsf1b complexes (Supplemen-

tary Figure S9), suggesting conservation across species of the interaction of Asf1 with the H3 tail.

The Asf1–Rtt109 interaction is retained in a complex containing Vps75 and histone H3:H4

Next, we probed the binding of the Asf1–H3:H4 substrate to the Rtt109–Vps75₂ complex by monitoring CSPs of the Ile, Leu and Val methyl resonances of Asf1. In the presence of Vps75, we observe large Asf1 CSPs upon addition of Rtt109 (Figure 5A and Supplementary Figure S10). When comparing the Asf1 CSPs from the Rtt109⁴¹⁹⁻⁴³³-to-Asf1 and Rtt109-to-Asf1–H3:H4–Vps75₂ titrations, we observe similar shifts on the same Asf1 interface described before, confirming that the interaction between Asf1 and the Rtt109 C-terminus is retained in the Asf1–H3:H4–Rtt109–Vps75₂ complex. Differences in CSPs were observed only for residues 6–9, which we presume is due to this region of Asf1 experiencing a different chemical environment in the Asf1–H3:H4–Rtt109–Vps75₂ complex compared to the Asf1–Rtt109 complex.

dependent K9 acetylation (in the absence of Vps75) was not detected by western blotting in Fillingham *et al.* (26), but has been previously observed by mass spectrometry, albeit to a lower extent than K56 acetylation (76). We observe that the absence of Vps75 slows but does not abolish K9 acetylation and that the Rtt109 C-terminus is required for K9 acetylation only in the absence of Vps75.

DISCUSSION

We have shown that the Rtt109 C-terminus directly interacts with Asf1 via both hydrophobic and charge-mediated contacts. The interaction mode is similar to other reported Asf1–peptide complexes, with the exception that the Rtt109 peptide is inversely oriented relative to the peptide in all other complexes. Using NMR techniques designed for studying high-molecular-weight complexes, we showed that the interaction between Asf1 and the C-terminus of Rtt109 is retained in a complex containing histone H3:H4 dimer and Vps75₂. Interestingly, the histone H3 tail interacts with the same surface of Asf1 as the Rtt109 C-terminus in the Asf1–H3:H4 complex; the Rtt109 C-terminus can out-compete this interaction (Figure 5B), releasing the H3 tail from Asf1. Similarly, Vps75 binding also causes Asf1 to release the H3 tail (Figure 5B).

In acetylation assays, we observe that *in vitro* deletion of the Rtt109 C-terminus impedes Asf1-dependent H3-K9Ac, but does not affect acetylation in the presence of both Asf1 and Vps75. Similar results have been reported in the literature for H3-K56Ac both *in vitro* and *in vivo* (26). Our structural findings provide a rationale for this data. The Rtt109–Asf1 interaction can out-compete binding of the H3 tail to Asf1, releasing it for modification. In the absence of both the Rtt109 C-terminus and Vps75, the H3 tail remains bound to Asf1 and protected from the enzyme. In agreement with this scenario, the Rtt109 C-terminus has been demonstrated to be redundant for Vps75-dependent H3-K9Ac *in vitro* in the absence of Asf1 (77). In addition, the Asf1–Rtt109 interaction may help in recruiting the Asf1-bound histone substrate to Rtt109, possibly limiting the orientations of H3:H4 with respect to the catalytic site and thereby promoting productive binding of the substrate. In agreement with the observation that Vps75 can rescue the effect of Rtt109 C-terminus deletion *in vitro*, we find that both Vps75₂ and the Rtt109 C-terminus modulate the Asf1–H3 tail interaction (Figure 5B). *In vivo*, however, both Vps75 and the Rtt109 C-terminus seem to be required for K9 (but not for K56) acetylation (26). This discrepancy suggests that there are additional factors contributing to or competing with H3-K9Ac by Rtt109 *in vivo*.

The chaperoning of the H3 tail by Asf1 might have implications beyond the Rtt109 enzymatic activation in the other Asf1-dependent regulatory pathways. The interaction of the H3 tail with Asf1 is conserved in humans (Supplementary Figure S9), suggesting a general role of Asf1 in modulating the conformation and accessibility of the H3 tail. Binding to Asf1 may protect the H3 tail from either interaction or modification and represent a mechanism to regulate access to the tail with respect to other cellular factors. In support of this hypothesis, the presence of Asf1 can inhibit H3 and H4 acetylation by the SAS complex (78).

AVAILABILITY

The atomic coordinates reported in this paper have been deposited in the Protein Data Bank, www.pdb.org (PDB ID: 6F0Y); the NMR data has been deposited in the Biological Magnetic Resonance Bank (BMRB ID: 34201).

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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