

The ancillary N-terminal region of the yeast AP-1 transcription factor Yap8 contributes to its DNA binding specificity

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ABSTRACT

Activator protein 1 (AP-1) is one of the largest families of basic leucine zipper (bZIP) transcription factors in eukaryotic cells. How AP-1 proteins achieve target DNA binding specificity remains elusive. In *Saccharomyces cerevisiae*, the AP-1-like protein (Yap) family comprises eight members (Yap1 to Yap8) that display distinct genomic target sites despite high sequence homology of their DNA binding bZIP domains. In contrast to the other members of the Yap family, which preferentially bind to short (7–8 bp) DNA motifs, Yap8 binds to an unusually long DNA motif (13 bp). It has been unclear what determines this unique specificity of Yap8. In this work, we use molecular and biochemical analyses combined with computer-based structural design and molecular dynamics simulations of Yap8–DNA interactions to better understand the structural basis of DNA binding specificity determinants. We identify specific residues in the N-terminal tail preceding the basic region, which define stable association of Yap8 with its target promoter. We propose that the N-terminal tail directly interacts with DNA and stabilizes Yap8 binding to the 13 bp motif. Thus, beside the core basic region, the adjacent N-terminal region contributes to alternative DNA binding selectivity within the AP-1 family.

INTRODUCTION

Yap8 protein is one of eight members of the yeast AP-1 (Yap) family (1) that belongs to the fungal specific Pap1 subfamily of basic leucine zipper (bZIP) transcription factors (2) (Figure 1). The bZIP proteins regulate transcription by binding as dimers to specific DNA motifs. Yap1 preferen-

tially binds to a 7 bp pseudo-palindromic sequence TTACTAA called the Yap response element (YRE) (1). However, Yap1 can also recognize TGAATA (3,4), TGAGTAA (5) and TGACAAA (5) motifs. Other members of the Pap1 subfamily, like *Schizosaccharomyces pombe* Pap1, and *Saccharomyces cerevisiae* Yap4 and Yap6, have preferences for an 8 bp palindromic version of the YRE (TTACGTAA) (2,6). DNA binding of bZIP transcription factors involves amino acids in the conserved basic region that precedes the leucine zipper region involved in dimerization. The crystal structure of the Pap1 bZIP domain bound to the 8 bp YRE revealed that five amino acid residues of the basic region make direct contacts with a TTAC half-site, and these residues constitute the signature DNA recognition NxxAQxxFR sequence (2). This motif is highly conserved among members of the Pap1 subfamily suggesting that Yap proteins share a common mechanism of DNA binding (Figure 1 and Supplementary Figure S1). Despite the high similarity of DNA binding regions and corresponding recognition elements of 7–8 bp, little is known how individual Yap proteins achieve their specificity of transcriptional regulation.

The transcription factors Yap1 and Yap8 are key components of the cellular response to arsenite [As(III)], arsenate [As(V)] and antimonite [Sb(III)] stress. Yap1 and Yap8 sense the presence of these agents and coordinate activation of gene expression required for alleviation of metalloid toxicity (7–10). Yap1 stimulates transcription of a large set of genes encoding proteins that are involved in adaptation to arsenic-induced oxidative stress and metalloid detoxification (7,9,11,12). In contrast, Yap8 is highly specific and seems to activate transcription of only two genes (13); *ACR2* that encodes an arsenate reductase (14) and *ACR3* that encodes an As(III)/Sb(III) efflux transporter (15,16).

Yap8 is the only member of the Yap family that recognizes a long 13 bp TGATTAATAATCA sequence, called the Yap8 response element (Y8RE), that consists of a 7 bp

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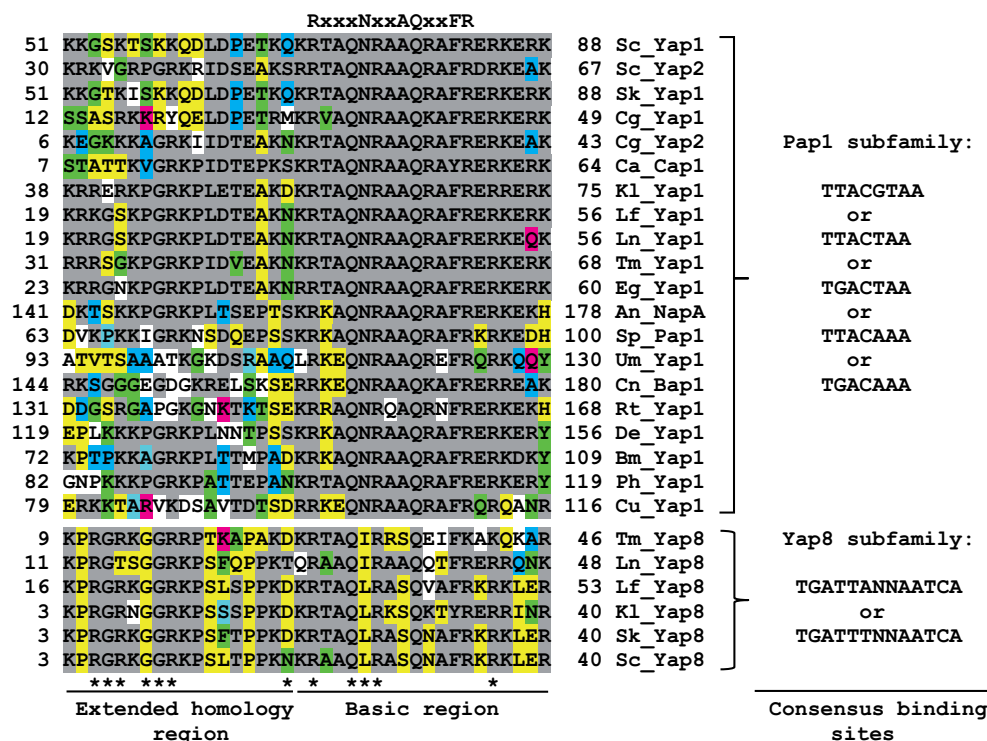


Figure 1. Comparison of basic regions, the N-terminal adjacent sequences and consensus DNA binding motifs of Yap1/2 and Yap8 orthologues. The following proteins are from the Saccharomycotina (Ascomycota) species: Sc_Yap1 (NCBI accession no. NP_013707), Sc_Yap2 (NP_010711) and Sc_Yap8 (NP_015525) proteins are from *S. cerevisiae*, Sk_Yap1 (EJT43841) and Sk_Yap8 (EJT44313) are from *S. kudriavzevii*, Cg_Yap1 (XP_446996) and Cg_Yap2 (XP_446103) are from *Candida glabrata*, Ca_Cap1 (EEQ44283) is from *C. albicans*, Kl_Yap1 (CAH02665) and Kl_Yap8 (CAG99045) are from *Kluyveromyces lactis*, Lf_Yap1 (SCW02819) and Lf_Yap8 (SCW01455) are from *Lachancea fermentati*, Ln_Yap1 (SCU88970) and Ln_Yap8 (SCV05062) are from *L. nothofagi*, Tm_Yap1 (TOMI0S02e09538g1, Genome Resources for Yeast Chromosomes, <http://gryc.inra.fr>) and Tm_Yap8 (TOMI0S05e00210g1) are from *Torulopsis microellipsoides*, e.g._Yap1 (NP_984291) is from *Eremothecium gossipii*, An_NapA (XP_680782) is from *Aspergillus nidulans* (Pezizomycotina, Ascomycota), Sp_Pap1 (NP_593662) is from *Schizosaccharomyces pombe* (Taphrinomycotina, Ascomycota), Cn_Bap1 (XP_012046219) is from *Cryptococcus neoformans* (Agaricomycotina, Basidiomycota), Um_Yap1 (KIS70678) is from *Ustilago maydis* (Ustilaginomycotina, Basidiomycota), Rt_Yap1 (CEE11106) is from *Rhodotorula toruloides* (Pucciniomycotina, Basidiomycota), De_Yap1 (RHZ80237) is from *Diversispora epigaea* (Mucormycota), Br_Yap1 (ORY02218) *Basidiobolus meristosporus* (Zoopagomycota), Ph_Yap1 (TPX58997) is from *Powellomyces hirtus* (Chytridiomycota), Cu_Yap1 (ORZ35932) is from *Catenaria anguillulae* (Blastocladiomycota). Conserved amino acid residues involved in direct binding to DNA bases as determined for the *S. pombe* Pap1 protein (2) are indicated at the top of sequence alignment. Known residues that are important for Yap8 function are marked with asterisks (18, this work). Identical or similar amino acid residues are highlighted accordingly. Experimentally confirmed consensus DNA binding motifs for each subfamily are indicated on the right panel.

core similar to the canonical YRE flanked by TGA bases (7,13). We recently showed that the Yap8 ortholog from *Kluyveromyces lactis* binds to multiple variants of Y8RE with different 7 bp core sequences flanked by conserved TGA bases (17). That study together with mutational analysis of the Y8RE sequence in *S. cerevisiae*, highlighted the importance of flanking TGA bases for Yap8–DNA interactions (Figure 1) (13,17). This distinct DNA binding property of Yap8 is reflected in its basic region in which invariant Asn and Ala residues of the NxxAQxxFR consensus sequence are replaced with Leu and Ser (LxxSQxxFR) (Figure 1). Indeed, Leu26 is essential for Yap8 binding to Y8RE and has, together with Asn31 and Leu26, been proposed to contribute to the DNA binding specificity of Yap8 (18).

In this study, we determined that a Yap8 variant with a core basic region identical to that of Yap1 still binds to Y8RE and fully activates transcription of *ACR3*. Such Yap8 variants also acquire capacity to bind to some, but not all, 7 bp motifs recognized by Yap1. Mutational analysis of the N-terminal tail adjacent to the basic region revealed spe-

cific residues that are required for stable association of Yap8 with the Y8RE-containing *ACR3* promoter and its activation. Based on a Yap8–DNA interaction model and *in vitro* DNA binding assays, we suggest that the N-terminal tails of Yap8 homodimer directly interact with the A/T-rich regions flanking the core Y8RE and stabilize Yap8 binding to the central 13 bp motif. We propose that the N-terminal tail of Yap8 constitutes an ancillary region that contributes to a unique DNA binding activity of Yap8 toward the 13 bp-long Y8RE motif. We hypothesize that the N-terminal region preceding the core basic region may influence the DNA binding specificity of other AP-1 proteins.

MATERIALS AND METHODS

Strains, plasmids and growth conditions

The *S. cerevisiae* strains used in this study were wild type W303-1A (*MATa ade2-1 can1-100 ura3-1 his3-11,15 leu2-3,112 trp1-1*), RW104 (*acr3Δ::kanMX*), RW117 (*yap8Δ::loxP*), RW120 (*yap8Δ::loxP*)

yap1Δ::loxP::kanMX::loxP), and RW124 (*yap1Δ::loxP*) (7). Plasmids used in this study are described in Supplementary Table S1. Standard yeast methods and growth conditions were used. Growth assays in the presence of sodium arsenite (Sigma-Aldrich) were carried out as previously described (19).

Mutagenesis

Site-directed mutagenesis of *YAP8* was performed using pYX122-YAP8 (20) and pGEX4T-1-GST-YAP8 (13) plasmids as templates, the oligonucleotides listed in Supplemental Table S2 and QuikChange Lightning Site-Directed Mutagenesis Kit (Agilent Technologies) according to the protocol provided by the manufacturer. All mutations were confirmed by commercial DNA sequencing.

β-Galactosidase assay

Yeast cells expressing various versions of *ACR3-lacZ* gene fusions were grown in selective minimal medium in the presence of 0.1 mM As(III) for 6 h or left untreated. The β-galactosidase activity was measured at least three times in triplicates on permeabilized cells as described previously (21).

RNA extraction and quantitative real-time PCR (qRT-PCR)

Total RNA was isolated from exponentially growing cells that were either untreated or exposed to 0.1 mM As(III) and collected at the indicated time points using RNeasyMini Kit (Qiagen). Reverse transcription was performed with 1.5 μg of purified RNA using High-Capacity cDNA Reverse Transcription Kit (Applied Biosystems) according to the manufacturer's instruction. Quantitative real-time PCRs were performed in the LightCycler 480 Instrument (Roche), using RealTime 2xPCR Master Mix SYBR (A&A Biotechnology) and *ACR3-fw/rv* primers listed in Supplemental Table S2 as described previously (22). *IPP1* was used as a reference gene. All assays were performed at least three times (biological replicas) in triplicates (technical replicas).

Protein extraction and western blot analysis

Cell extracts were prepared by TCA precipitation and proteins were separated by 10% SDS-PAGE followed by immunoblotting with anti-HA antibody (Sigma-Aldrich, ref: H6908, lot: 015M4868V, 1:2500 dilution) and anti-PGK1 antibodies (Abcam, ref: ab11368, lot: GR254438-1; 1:5000 dilution).

Immunofluorescence microscopy

Immunofluorescent labeling of yeast cells was performed as described earlier (23). Cells were fixed in 3.7% formaldehyde for 2 h, washed and digested with Zymolyase (BioShop) for 30 min. The efficiency of spheroplasting was monitored by phase microscopy. Spheroplasts were washed twice and suspended in PBS buffer supplemented with 0.1% BSA. Yeast cells were stained with primary antibody (anti-HA, Sigma-Aldrich, ref: H6908, lot: 015M4868V, 1:1000 dilution) for 12 h at 4°C. The samples were washed with PBS

containing 0.1% BSA after exposed secondary antibody Alexa Fluor® 488 goat anti-rabbit IgG (H+L, Life Technologies, ref: A11008, lot: 1470706, 1:200 dilution) at room temperature for two hours. After triple washing with PBS, cells were labeled with DAPI (Life Technologies, 1:5000 dilution) to visualize nuclei and examined with a fluorescence microscope (Axio Imager M2, Carl Zeiss) equipped with a 100× oil immersion objective, differential interference contrast and appropriate filters. Images were collected using Zeiss AxioCam MRm digital camera and processed with Zeiss Zen 2012 software.

Expression and purification of GST-Yap8 variants

Expression of wild type and mutant versions of GST-Yap8 was induced by incubating *Escherichia coli* BL21(DE3)pLysS cells with 1 mM IPTG (isopropyl β-D-thiogalactoside) for four hours at 30°C in LB medium (1% tryptone, 0.5% yeast extract, 1% NaCl) in the presence of 100 μg/ml ampicillin and 34 μg/ml chloramphenicol. Cells were harvested and disrupted by sonication in cold PBS buffer containing protease inhibitor cocktail (Roche), 10 mM β-mercaptoethanol, 1% Triton X-100 and 10% glycerol. All GST-tagged proteins were purified using glutathione beads (GE Healthcare) according to the protocol supplied by the manufacturer.

Electrophoretic mobility-shift assay (EMSA)

The 5' end biotinylated complementary oligonucleotide pairs (Sigma-Aldrich) were annealed to make double-stranded and biotin-labeled probes by mixing in a buffer (10 mM Tris-HCl, pH 8.0, 1 mM EDTA), boiling for 5 min and cooling slowly to room temperature. Unlabeled complementary oligonucleotide pairs were also annealed to make double-stranded competitor probes. EMSA reaction solutions were prepared by adding the following components according to the manufacturer's protocol (LightShift Chemiluminescent EMSA kit; Thermo Fisher Scientific): 1× binding buffer, 50 ng poly (dI-dC), 2.5% glycerol, 0.05% Nonidet P-40, 5 mM MgCl₂, 10 ng of purified recombinant GST-tagged protein, competitor (4 pmol) and biotin-labeled probes (20 fmol). Reaction solutions were incubated for twenty minutes at room temperature. The protein-probe mixture was separated in a 6% polyacrylamide native gel in a standard 0.5× TBE buffer. Electrophoresis was performed on ice (100 V, 1 h). The DNA was transferred (100 V, 30 min) to a positive nylon membrane (Amersham Hybond-N+, GE Healthcare) and UV crosslinked (1200 uJ/cm², UVP TL-2000 Ultraviolet Translinker). Migration of biotin-labeled probes was detected in the ChemiDoc MP Imager (BioRad) using streptavidin-horseradish peroxidase conjugates that bind to biotin and chemiluminescent substrate according to the manufacturer's protocol. The sequences of the oligonucleotides used are listed in Supplementary Table S2.

Alternatively, the oligonucleotide probes were 5' end labeled with [γ-³²P]ATP using polynucleotide kinase (Thermo Scientific), purified through Sephadex G-50 chromatography, annealed with complementary oligonucleotides in the presence of 100 mM NaCl at 75°C for 10 min

and gradually cooled to room temperature. Purified recombinant GST-tagged proteins (at indicated concentrations) were incubated with ^{32}P -labeled oligonucleotide probes (40 000 cpm) in a 20 μl reaction containing EMSA buffer (10 mM Tris-HCl, pH 8.0, 50 mM NaCl, 1 mM DTT, 0.05% NP-40, 100 ng poly(dI-dC) and 6% glycerol) for 30 min at 4°C. The reaction mixtures were subjected to electrophoresis on 5% non-denaturing polyacrylamide gels in a standard 0.5 \times TBE buffer. Electrophoresis was performed on ice (100 V, 1 h). The gels were dried and analyzed using a phosphorimager (Molecular Imager FX, Bio-Rad).

Fluorescence anisotropy assay

The fluorescence anisotropy of FAM-labeled *ACR3* oligonucleotides (labeled on the 5' end with 6-carboxyfluorescein) was measured on two-four independent repetitions with different protein to DNA ratios and one reference solution without protein in buffer A (10 mM Tris-HCl pH 8, 50 mM NaCl, 0.5% NP-40, 1 mM DTT and 5% glycerol). The total volume of the working solutions was 100 μl , and the added protein elution buffer amount was kept constant to 50 μl by adding buffer A when necessary. Measurements were performed on a spectrofluorometer FS5 (Edinburgh Instruments) in a temperature-controlled microcuvette at 25°C. Fluorescence emission intensity was recorded at 515 nm, with excitation at 490 nm, and emission and excitation slits set to 2 nm. All titrations were performed using 1 nM of DNA, and after each addition the sample was equilibrated for 6 min. Stoichiometric binding curves were fit to the equation: $\Delta A = \Delta A_T / 2D_T \{ (E_T + D_T + K_d) - [(E_T + D_T + K_d)^2 - 4E_T D_T]^{1/2} \}$, where ΔA is the change in anisotropy, ΔA_T is the total anisotropy change, E_T is the total protein concentration, D_T is the total DNA concentration, and K_d is the dissociation constant.

Chromatin immunoprecipitation (ChIP)

ChIP was performed as described previously (24). Sheared chromatin was immunoprecipitated using anti-HA antibody (Sigma-Aldrich, ref: H6908, lot: 015M4868V, 1:2500 dilution) overnight followed by incubation with sepharose protein G beads (Dynabeads Protein G, Life Technologies). Precipitates and input DNA were analyzed by qRT-PCR using PRACR3-fw/rv oligonucleotides listed in Supplementary Table S2, for the *ACR3* promoter region from -251 to -100 relative to the ATG translation initiation codon. qPCR data are presented as percentages of input DNA normalized to the *IPPI* gene region as a control. Results are representative of at least two independent biological replicas and four PCR reactions and error bars indicate \pm standard deviations.

Molecular modeling

The model structure of the basic-leucine-zipper domain (residues 7–89) of the Yap8 protein homodimer was created using the homology building functionality of Yasara program (25). The DNA sequence of 25 bp (TTTGTT-TG

Table 1. Binding affinity (K_d) of the Yap8 protein and its mutant variants to DNA fragments corresponding to the Y8RE-containing *ACR3* promoter region

Protein	K_d (nM)	DNA probe
Yap8 (WT)	9.9 \pm 1.4 ^a	ACR3-WT
Yap8-7aa	9.3 \pm 1.7 ^a	ACR3-WT
Yap8-8aa	15.1 \pm 2.1 ^a	ACR3-WT
Yap8-N20A	25.5 \pm 4.3 ^a	ACR3-WT
Yap8(WT)	10.8 \pm 1.7 ^b	ACR3-WT
Yap8-R7A	13.4 \pm 1.7 ^b	ACR3-WT
Yap8-R11A	18.9 \pm 2.9 ^b	ACR3-WT
Yap8 (WT)	10.8 \pm 3.5 ^c	ACR3-WT
Yap8 (WT)	19.4 \pm 4.1 ^c	ACR3-M3

The K_d values of dissociation constant were derived after curve fitting to anisotropy binding data shown in Figure 3C^a, Figure 5E^b and Figure 8C^c, respectively.

ATTAATAATCA-**ACTTTA**) contains Yap8-response element, Y8RE, shown in bold. The structure of the Yap8–DNA complex was modeled using HADDOCK molecular multi-body docking server (26,27). The residues: Asn20, Arg22, Gln25, Leu26 and Phe33 were indicated as ‘active’, as their alanine-mutants show sufficient reduction in the proteins activity and/or ability to bind DNA (Table 1). Residues Arg27 and Arg36 were defined as ‘passive’ as their alanine mutants show only partial resistance to As(III). The Y8RE-DNA residues were identified as ‘passive’. Out of the 29 structure clusters provided by the HADDOCK server, one of the clusters had a significantly higher score, which was selected for further analysis. Lastly, the N-terminal fragments (residues 7–18) of the protein were added manually in a random coil configuration using program USCF Chimera (28). The random coil configuration of the N-terminal tails was justified by protein secondary structure prediction servers Jpred4 (29), PredictProtein (30) and PSIPRED (31). Additionally, the complex structure containing Asn20Ala Yap8 mutants was created using the ‘Rotamers’ functionality of USCF Chimera program.

The two complex structures were subjected to subsequent studies by molecular dynamics simulations (MD), using GROMACS MD software package, version 5.1 (32). Simulations were carried out using a combination of the latest AMBER all-atom nucleic acid Parmbsc1 (33) and ff14SB (34) force fields in implicit solvent using of SCP/E water molecules (35) and 150 mM KCl. MD simulations were carried out at constant pressure and temperature (1 atm, 300 K). Further details of the simulation protocols can be found in Supplemental Information. Each productive MD run was 500 ns long. MD trajectories were analyzed using CPPTRAJ program (36), focusing on the analysis of the protein–DNA interactions, including hydrogen bonds, salt bridges, and hydrophobic (apolar) interactions. Dynamic contacts maps were created by summing up the hydrogen bonds and the salt bridge interactions for each pair of Yap8–DNA interacting residues, which resulted in a contact strength value. We also performed conformational clusters analysis following the protocol described by Lavery *et al.* (37,38) for the basic-regions (residues 17–40) of the protein dimer. For the random-coil N-terminal regions (residues 7–16) conformational clusters were identified with the cluster feature of

CPTRAJ program (36), using DBSCAN (density-based) clustering algorithm (39). RMSD of heavy atoms of DNA outside YRE region and excluding two terminal base pairs on both ends and the protein residues 7–16 was used as a distance metric. The Yap8–DNA complex structure that represents the biggest conformational cluster was selected to represent the model structure. Molecular graphics were created with USCF Chimera.

RESULTS

Construction of Yap8 variants containing the Yap1-like basic region

The DNA binding basic region is highly conserved in the fungal AP-1 family (Figure 1 and Supplementary Figure S1). Mutations of several such conserved residues in the Yap8 basic region, including Arg22, Gln25, Arg27 and Arg36, were previously reported to impair the transcriptional activity of Yap8 toward a Y8RE-containing promoter (18). Also in our hands, Yap8-Q25A was not able to induce expression of the *ACR3-lacZ* reporter gene (Figure 2B) or rescue As(III) sensitivity of cells lacking the *YAP8* gene (Figure 2C). Likewise, Yap8-R36A appeared partially defective as we observed weak activation of *ACR3-lacZ* expression (Figure 2B) and partial complementation of *yap8*Δ (Figure 2C). Importantly, the corresponding residues in the Yap1–DNA complex, Gln85 and Arg96, were shown to interact with DNA phosphate backbone (2) and appears in all members of the fungal AP-1 family, with the exception of *K. lactis* Yap3 (Gln→Leu substitution) and *Torulaspora microellipsoides* Yap8 (Arg→Lys substitution) (Figure 1 and Supplementary Figure S1). This suggests that Yap8 shares similar DNA binding properties with other members of the fungal AP-1 family. However, Yap8 contains several amino acid substitutions within its basic region at positions conserved in other members of Yap family (Figure 1); these amino acid residues may contribute to the specificity of Yap8 toward the extended Y8RE motif as well as its inability to bind to short YRE motifs.

To investigate this, we stepwise replaced amino acid residues in the basic region of Yap8 into the corresponding residues present in Yap1 and functionally characterized the resulting Yap8 variants. First, we constructed a quadruple A23T L26N S29A N31R mutant (or Yap8-4aa) to make the core of the basic region identical with the Yap1 basic region, including the NxxAQxxFR consensus sequence (Figure 2A). The Yap8-4aa mutant was able to fully activate expression of the *ACR3-lacZ* reporter gene (Figure 2B) and to complement the As(III) sensitivity of the *yap8*Δ mutant (Figure 2C). Next, we introduced three additional mutations (K35E, L37E and E39R) to make the C-terminal region adjacent to the core basic region identical to the Yap1 sequence (Figure 2A). The septuple Yap8-7aa mutant also behaved like wild type Yap8 in terms of *ACR3* expression and *yap8*Δ complementation (Figure 2B and C). Finally, we additionally replaced Asn20, located adjacent to the core of the basic region with Gln (corresponding amino acid in Yap1) in Yap8-7aa (Figure 2A). The octuple Yap8-8aa mutant failed to *trans*-activate the *ACR3-lacZ* reporter gene and complement As(III) sensitivity of *yap8*Δ (Figure 2B and C). In this regard, Yap8-8aa behaved

like Yap1, which is not able to activate *ACR3* expression (Figure 2B). However, if the central adenine residue in the Y8RE element is replaced with cytosine, Yap1 can weakly induce *ACR3* expression (13) (Figure 2D). Thus, we analyzed activity of the *MUT3-ACR3-lacZ* promoter with the TGATAACTAATCA sequence containing both Y8RE and YRE (underlined) motifs in a single element (Figure 2D). Wild type Yap8, Yap8-4aa and Yap8-7aa variants strongly induced expression of the *MUT3-ACR3-lacZ* reporter gene whereas Yap8-8aa behaved like Yap1 and weakly activated the *MUT3-ACR3* promoter (Figure 2D). In sum, these results suggest that Asn20 contributes to Yap8 binding to the *ACR3* promoter.

Interestingly, Asn20 is often conserved in Yap1/2 and Yap8 orthologues or substituted for either Asp/Glu or Thr/Ser (Figure 1) suggesting a functional importance of this site in some Yap proteins. To get a better insight into the role of Asn20, we replaced this residue with glutamine (Yap8-N20Q) or aspartate (Yap8-N20D), which are present in the corresponding positions in *Saccharomyces cerevisiae* Yap1 and *K. lactis* Yap8, respectively (Figure 1). Asn20 was additionally replaced with alanine (Yap8-N20A). Yap8-N20A and Yap8-N20Q failed to induce expression of both *ACR3-lacZ* and *MUT3-ACR3-lacZ* upon As(III) stress (Figure 2B and D) and poorly complemented As(III) sensitivity of the *yap8*Δ mutant (Figure 2C). In contrast, Yap8-N20D showed wild type activity in both assays (Figure 2B and C). We confirmed that all Yap8 variants tested were present at the same amounts as the wild type Yap8 protein (Supplementary Figure S2) and that all tested variant proteins were correctly localized to the nucleus (Supplementary Figure S3). Thus, the observed effects are likely directly related to Yap8 function/activity. We conclude that both Asn20/Asp20 preceding the basic region as well as highly conserved residues (Gln25, Arg36) within the basic region are important for Yap8 function.

DNA binding properties of the Yap8 basic region and N20A mutants

To characterize the DNA binding properties of the Yap8 variants, we performed EMSAs using purified GST-Yap8 proteins (Supplementary Figure S4) and biotin-labeled oligonucleotides corresponding to the *ACR3* promoter sequence with the TGATTAATAATCA motif (Figure 3A). It is important to point out that we have previously shown that the GST-Yap8 fusion protein is fully functional *in vivo* (13). In agreement with published data (18) and our *in vivo* assays (Figure 2), Yap8-Q25A exhibited markedly reduced ability to bind to the *ACR3* oligo (Figure 3A). In line with the inability to induce *ACR3* expression (Figure 2C), Yap8-8aa and Yap8-N20A variants also showed highly reduced capacity to bind to the *ACR3* oligo (Figure 3A). Accordingly, *in vivo* ChIP experiment revealed that Yap8-8aa and Yap8-N20A do not stably associate with the *ACR3* promoter in living cells, neither in the absence nor presence of As(III) (Figure 3B). Importantly, the Yap8-7aa variant with Yap1-like core basic region retained the wild type activity both *in vitro* (Figure 3A) and *in vivo* (Figure 3B).

To more accurately measure the affinity of Yap8–DNA interactions, we performed a fluorescence anisotropy bind-

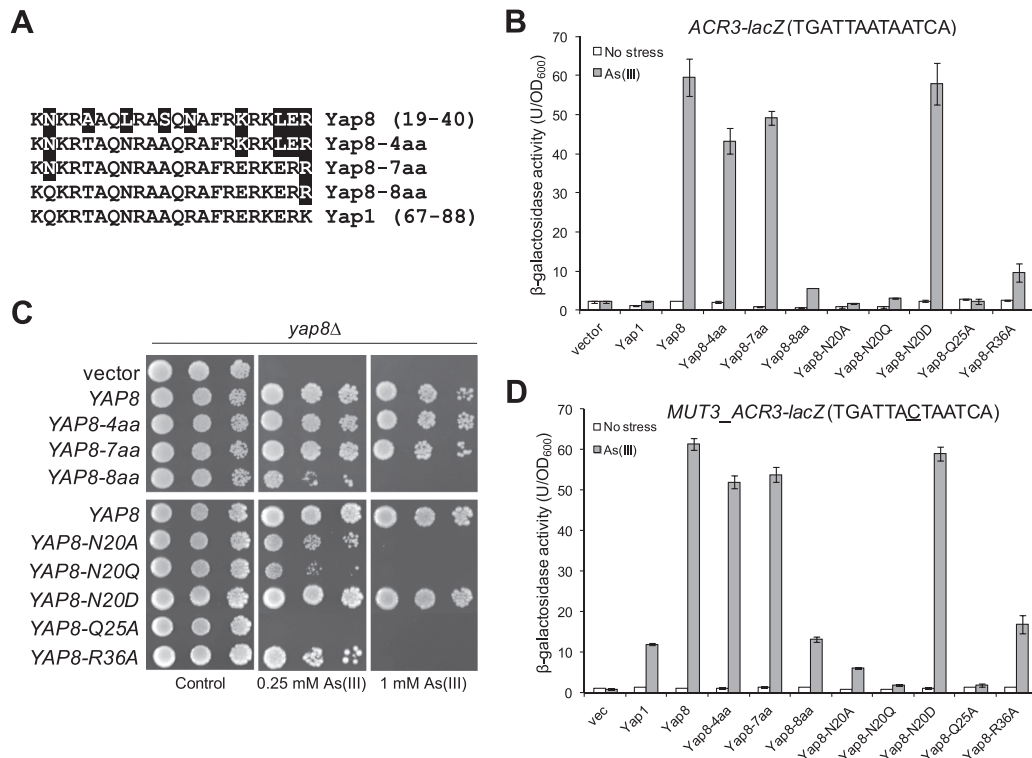


Figure 2. Analysis of Yap8 basic region variants. (A) Mutagenesis strategy to stepwise turn the Yap8 basic region into Yap1-like sequences. (B) β -galactosidase activity driven by the *ACR3-lacZ* promoter was measured in the *yap1* Δ *yap8* Δ mutant expressing either Yap1, Yap8 or Yap8 mutant proteins. Cells were exposed to 0.1 mM As(III) for 6 h or left untreated for the control. The values are the means of three biological replicas performed in triplicate \pm S.D. (C) Complementation of As(III) sensitivity of *yap8* Δ by Yap8 variants. The *yap8* Δ mutant was transformed with empty vector (pYX122) or plasmids expressing indicated Yap8 variants. The resulting transformants were spotted on minimal selective plates containing various concentrations of As(III) and incubated 3 days at 28°C. (D) β -Galactosidase activity driven by the *MUT3-ACR3-lacZ* promoter was measured in the *yap1* Δ *yap8* Δ mutant expressing either Yap1, Yap8 or Yap8 mutant proteins. Cells were exposed to 0.1 mM As(III) for 6 h or left untreated for the control. The values are the means of three biological replicas performed in triplicate \pm S.D.

ing assay. Binding titrations were performed as a function of increasing concentration of Yap8 and its mutated forms at fixed DNA concentration corresponding to the *ACR3* promoter sequence. From fluorescence anisotropy measurements it is clear that the Yap8-7aa mutant showed virtually identical binding affinity as the wild-type protein (Figure 3C, Table 1). In contrast, Yap8-8aa and Yap8-N20A variants showed significantly weaker binding to the Y8RE-containing DNA fragment (Figure 3C, Table 1). The results obtained by this solution-based, true-equilibrium method are consistent with *in vivo* (complementation tests, lacZ assay, ChIP) and *in vitro* (EMSA) data shown above (Figures 2B–D, 3A and B). Together, our data strongly suggest that the N-terminal Asn20 residue is important for high affinity binding of Yap8 to the 13 bp long Y8RE motif.

Yap8 variants with the Yap1-like basic region cannot bind to all YREs

We next investigated whether the Yap8 variants with Yap1-like basic regions had acquired ability to bind to 7 bp YRE motifs. For this, we performed EMSAs using oligos corresponding to *GSH1* (contains one YRE with sequence TTAGTCA) and *TRX2* (contains two YREs with sequence TTACTAA) promoters. None of these YREs contain TGA flanks. As expected, wild type Yap8 did not bind to the

GSH1 oligo (Figure 4A). However, Yap8-4aa, Yap8-7aa and Yap8-8aa bound weakly to the *GSH1* oligo at higher protein concentrations (stable binding required 100 ng protein for the *GSH1* oligo compared to 10 ng for the *ACR3* oligo), suggesting low-affinity binding of these Yap8 variants to the YRE TTAGTCA (Figure 4). Neither Yap8-N20A nor Yap8-N20Q bound to the *GSH1* oligo. None of the Yap8 variants bound stably to the *TRX2* promoter fragment (Figure 4). Thus, replacing up to eight amino acid residues to make the Yap8 basic region more Yap1-like was not sufficient to enable binding of the modified Yap8 to the YRE motif (TTACTAA), present in *TRX2*. This suggests that the amino acid residues outside the basic region may contribute to YRE recognition and/or stable DNA binding.

The N-terminal tail adjacent to the basic region contributes to Yap8 DNA binding activity

The N-terminus of the bZIP domain of Yap1 (Asp63-Pro64) and Yap8 (Thr16-Pro17-Pro18) (Figure 1) include the N-capping motifs containing Asn, Asp, Ser, Thr, or Gly followed by single or double Pro residues. The N-capping motif is believed to stabilize the helical structure of the basic region upon DNA binding, without a direct interaction with DNA (40). To investigate the role of the putative N-

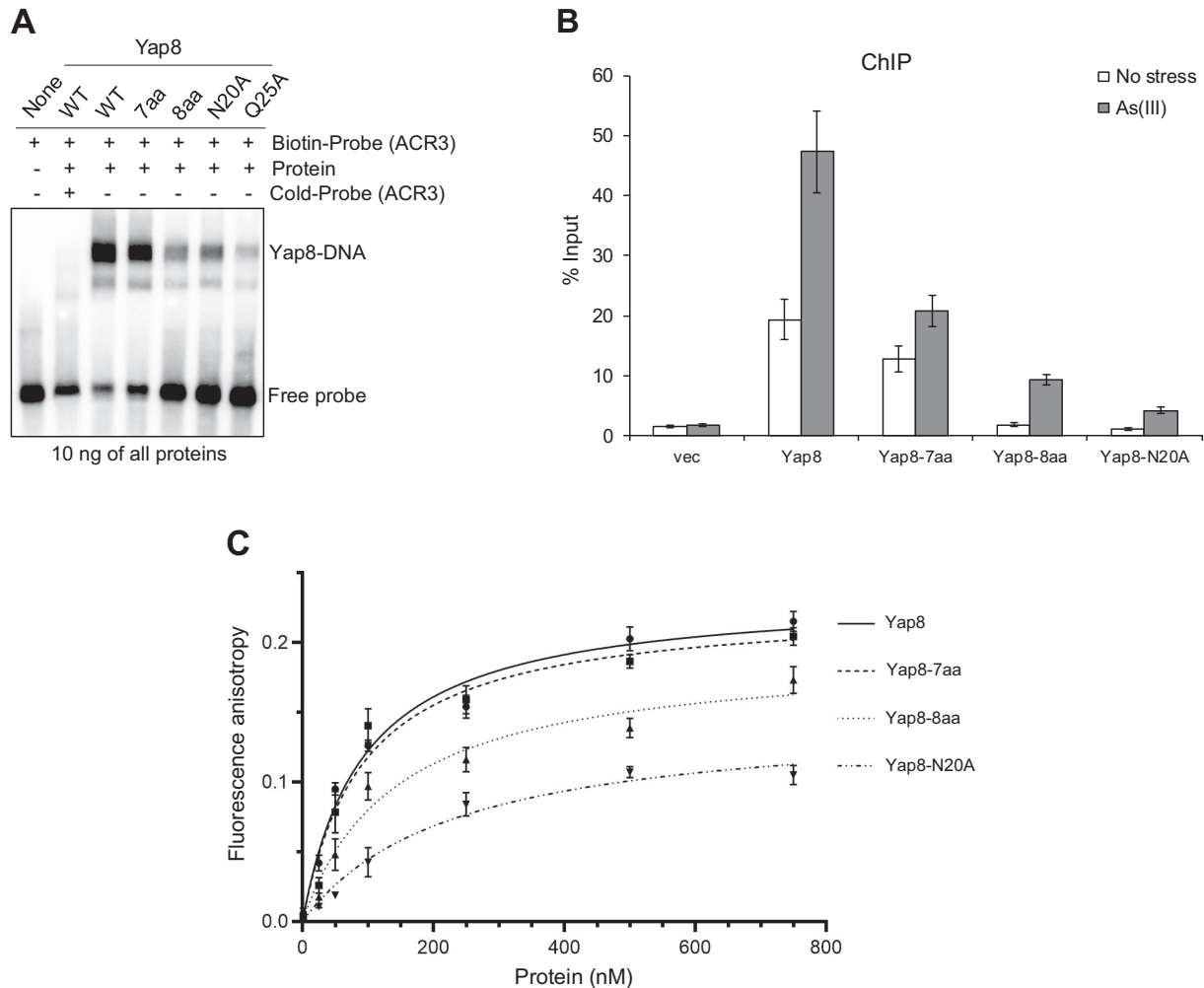


Figure 3. DNA binding activity of Yap8 basic region mutants. (A) Binding of Yap8 variants to the *ACR3* promoter as determined by EMSA. Purified GST-Yap8 variants at indicated concentration were incubated with biotin-labeled oligonucleotides corresponding to Y8RE-containing promoter fragments of *ACR3* gene followed by electrophoresis. (B) Binding of Yap8 variants to the *ACR3* promoter as determined by ChIP. *yap8* Δ cells bearing plasmids expressing the indicated Yap8-HA fusion proteins or the control vector were exposed to 0.5 mM As(III) for 30 min or left untreated. qRT-PCR was performed with chromatin fragments immunoprecipitated with anti-HA antibodies and primers amplifying the Y8RE-containing *ACR3* promoter region. Error bars indicate standard error of the mean from at least two independent biological replicas and four PCR reactions. (C) Fluorescence anisotropy assays performed with indicated variants of purified GST-Yap8 and the FAM-labeled *ACR3* promoter fragment as described in Materials and Methods.

cap of Yap8, we constructed and functionally characterized Yap8-T16A, Yap8-P17A and Yap8-P18A mutants. We found that all tested mutants fully complemented the arsenic sensitivity of *yap8* Δ suggesting that the Thr-Pro-Pro motif does not affect the Yap8 function (Supplementary Figure S5). In addition, we tested the significance of adjacent Ser14, Leu15, Lys19 and Lys21 residues for Yap8 function by alanine replacement and found that the resulting mutants showed wild type phenotype (Table 2, Supplementary Figure S5). To summarize, it seems that in the Yap8 region of Ser14-Lys21 only Asn20 residue is important for Yap8 function.

Members of the mammalian Maf subfamily of bZIP superfamily that recognize a 13–14 bp consensus element (TGCTGAC(G)TCAGCA) called the Maf recognition element (MARE) (41) require the N-terminal extended homology region (EHR) preceding the basic region for high-affinity binding to DNA (42,43). Interestingly, the N-

terminal regions of Yap8-like proteins as well as many Yap1/2 orthologues found in most phyla of fungi, with the exception of Blastocladiomycota and Basidiomycota, exhibit evolutionary conservation (Figure 1). It is important to note that this is not the case for *S. cerevisiae* and *C. glabrata* Yap1 proteins, nor for Yap3, Yap5/7 (with the exception of HapX-like proteins) and Yap4/6 orthologues (Figure 1 and Supplementary Figure S1). We hypothesized that the Pro4-Pro13 region of Yap8, which is rich in basic residues and shows the highest level of conservation among Yap1 and Yap8-like proteins may contribute to high-affinity binding to DNA. To test this, we constructed a Yap8 variant lacking residues from Arg5 to Pro13; the resulting Yap8- Δ 5–13 mutant failed to complement arsenic sensitivity of *yap8* Δ strain (Figure 5A) and to induce expression of the *ACR3-lacZ* reporter gene (Figure 5B). Importantly, the Yap8- Δ 5–13 mutant was expressed at the wild type level (Supplementary Figure S2) and showed nuclear localiza-

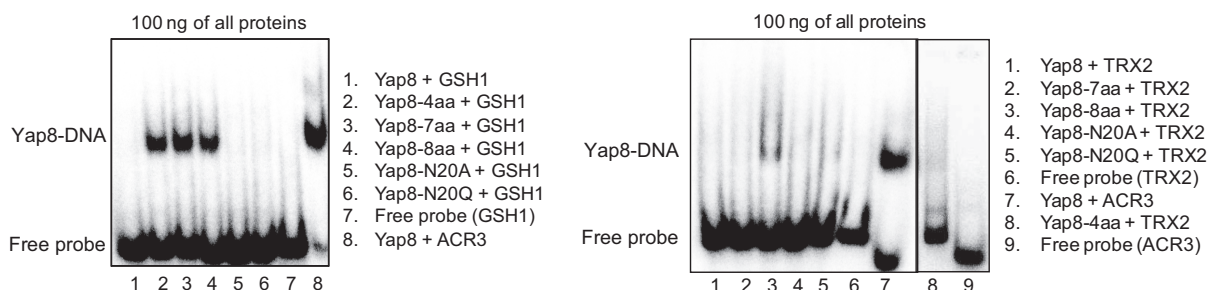


Figure 4. DNA binding of Yap8 protein variants with the Yap1-like basic region determined by EMSA. Purified GST-Yap8 variants at indicated concentrations were incubated with 32 P-labeled oligonucleotides corresponding to Y8RE/YRE-containing promoter fragments of *ACR3*, *GSH1* and *TRX2* genes followed by electrophoresis.

tion (Supplementary Figure S3). This suggests that the N-terminal tail is critical for Yap8 function.

To identify residues in the N-terminal tail that are important for Yap8 binding to DNA, we generated ten single alanine-replacement mutations covering residues from Pro4 to Pro13 and tested the functionality of the resulting Yap8 mutants. Of these: Yap8-R5A, Yap8-G6A, Yap8-R7A and Yap8-G9A partially complemented arsenic sensitivity of *yap8* Δ strain (Figure 5A) and showed residual ability to induce expression of the *ACR3-lacZ* reporter gene (Figure 5B). Yap8-G10A and Yap8-R11A exhibited the strongest phenotype with no ability to confer resistance to As(III) (Figure 5A) or to activate the *ACR3* promoter (Figure 5B). Except for Yap8-G9A, which showed reduced protein level, all tested Yap8 mutants showed protein accumulation at the wild type level in response to As(III) treatment (Supplementary Figure S2) and nuclear localization both in the absence and presence of As(III) (Supplementary Figure S3). Based on these results, we conclude that the arginine and glycine-rich N-terminal region is important for Yap8 ability to activate the *ACR3* promoter.

We hypothesized that arginine residues of the Yap8 N-terminal tail may be involved in DNA binding, whereas glycine residues may contribute to plasticity of this region allowing tighter contact with DNA. To test this, we investigated the ability of purified Yap8-R7A, Yap8-G10A and Yap8-R11A protein variants tagged with GST (Supplementary Figure S4) to bind the *ACR3* promoter *in vitro* by EMSA (Figure 5C) and *in vivo* by ChIP (Figure 5D). Yap8-G10A showed no binding to DNA fragment containing the Y8RE motif, whereas Yap8-R7A and Yap8-R11A exhibited reduced binding to DNA compared to the wild type Yap8 (Figure 5C). Likewise, little (Yap8-R7A) or no association (Yap8-G10A and Yap8-R11A) of Yap8 variants to the *ACR3* promoter was observed by ChIP (Figure 5D). Next, we performed a fluorescence anisotropy binding assay to measure binding affinity of these Yap8 mutants to a DNA fragment comprising the *ACR3* promoter. The Yap8-R7A mutant protein showed moderately decreased binding affinity to the Y8RE-containing DNA fragment, whereas the K_d value determined for the Yap8-R11A version is twice as high as K_d of the wild type Yap8 (Figure 5E, Table 1). However, the affinity of Yap8-R7A and Yap8-R11A variants for the Y8RE-containing DNA fragment was higher than the affinity observed for Yap8-N20A (Table 1). We were not able to perform this experiment for Yap8-G10A

because of the high tendency of the purified protein to precipitate. Basic residues of Yap8 N-tails (R7 and R11) may be engaged in either non-specific electrostatic interactions with negatively charged DNA backbone, and thus stabilizing Yap8 binding to DNA, or specific interactions with bases of the Y8RE-containing region. To test these hypotheses, we generated single Yap8-R7K and Yap8-R11K variants and double Yap8-R7K,R11K mutant and analyzed their ability to bind the *ACR3* promoter fragment by EMSA (Figure 5F). Despite the presence of positively charged lysine residue, Yap8-R7K showed poor ability to bind to the *ACR3* promoter fragment, whereas both Yap8-R11K and Yap8-R7K,R11K variants showed no DNA binding activity. This strongly suggests that Arg7 and Arg11 residues of the Yap8 N-tail are engaged in specific interactions with the *ACR3* promoter sequence and contribute to recognition of the Y8RE-containing region by Yap8. In sum, we conclude that the N-terminal tail is required for stable binding of Yap8 to DNA and may contribute to a unique specificity of Yap8 toward the long (13 bp) Y8RE motif.

Molecular modeling and structural analysis of Yap8–DNA complex

To get an atom-level understanding of the structural basis of Yap8–DNA recognition we created an all-atom model of Yap8–DNA complex (Figure 6). The model consists of Yap8 homodimer and 25 bp DNA segment containing the Y8RE motif. Each Yap8 monomer contains an α -helical basic-leucine-zipper (residues 17–89) domain and an unstructured N-terminal region (residues 7–16). Details of the model building are provided in the methods section. According to the model, the protein inserts its α -helical basic-leucine-zipper domain in the DNA major groove of the Y8RE sequence, with the N-terminal regions making contacts with the DNA minor groove of the Y8RE flanks. To verify the model we performed 500 ns all-atom molecular dynamics (MD) simulations. The MD simulation allowed construction of the dynamic interactions maps (Figure 7A), which describe the details of the specific Yap8–DNA interactions and the dynamics of the intermolecular interface. In addition, the protein–DNA interactions were characterized by the occupancy (percentage present) during the MD simulation and the average lifetime (Supplementary Table S3). The MD simulations show that the interactions patterns dif-

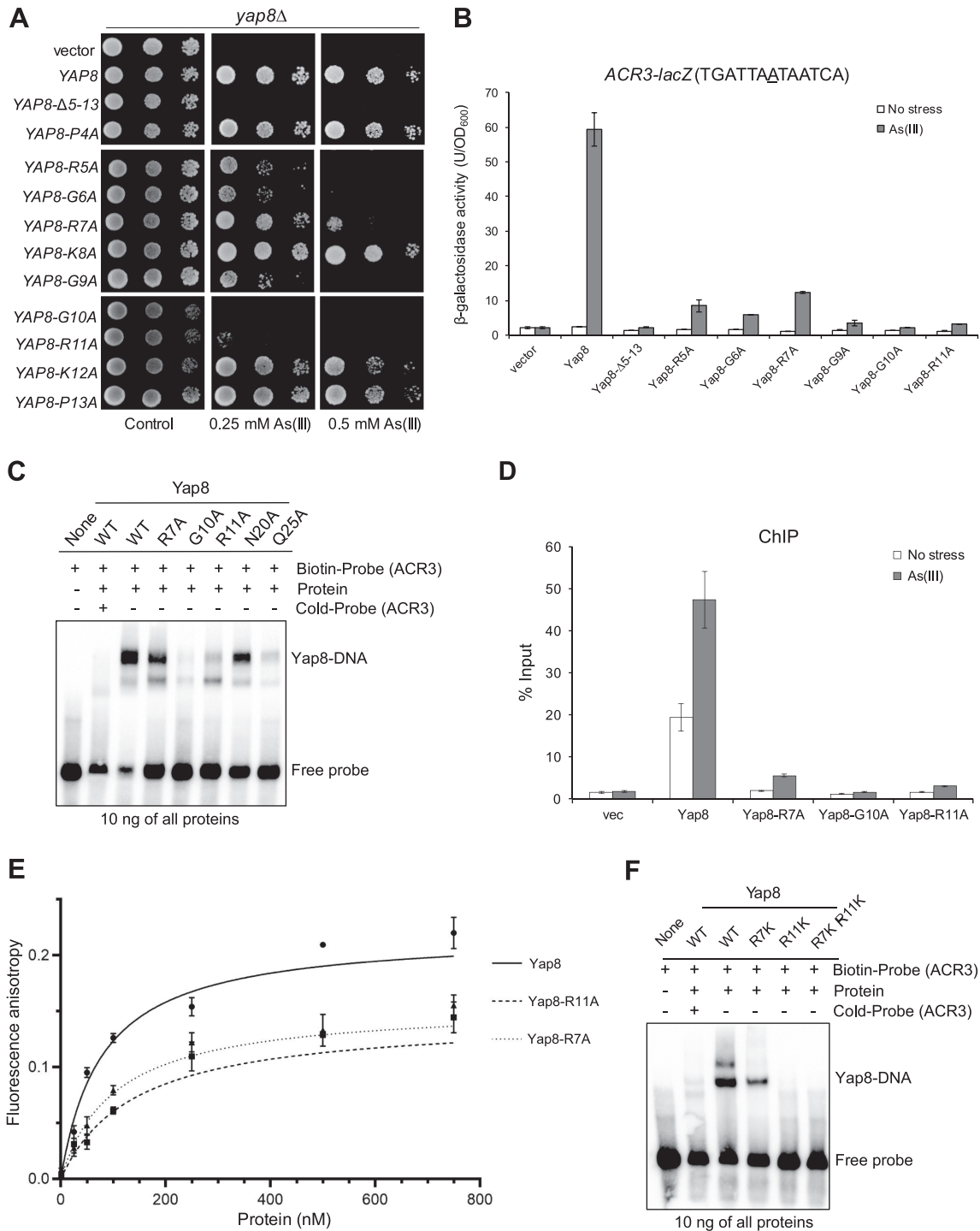


Figure 5. Functional analysis of Yap8 N-terminal tail mutants. **(A)** Complementation of As(III) sensitivity of *yap8Δ* by indicated Yap8 variants. The *yap8Δ* mutant was transformed with empty vector (pYX122) or plasmids expressing Yap8 variants. The resulting transformants were spotted on minimal selective plates containing various concentrations of As(III) and incubated 3 days at 28°C. **(B)** β-galactosidase activity driven by the *ACR3-lacZ* promoter was measured in the *yap1Δ yap8Δ* mutant expressing indicated Yap8 mutant proteins. Cells were exposed to 0.1 mM As(III) for 6 h or left untreated for the control. The values are the means of three biological replicas performed in triplicate ± S.D. **(C)** Binding of Yap8 variants to the *ACR3* promoter as determined by EMSA. Purified GST-Yap8 variants at indicated concentration were incubated with biotin-labeled oligonucleotides corresponding to Y8RE-containing promoter fragments of *ACR3* gene followed by electrophoresis. **(D)** Binding of Yap8 variants to the *ACR3* promoter as determined by ChIP. *yap8Δ* cells bearing plasmids expressing Yap8-HA variant proteins or the control vector were exposed to 0.5 mM As(III) for 30 min or left untreated. qRT-PCR was performed with chromatin fragments immunoprecipitated with anti-HA antibodies and primers amplifying the *ACR3* promoter region containing Y8RE1 and Y8RE2 motifs. Error bars indicate standard error of the mean from at least two independent biological replicas and four PCR reactions. **(E)** Fluorescence anisotropy assays performed with indicated variants of purified GST-Yap8 and the FAM-labeled *ACR3* promoter fragment as described in Materials and Methods. **(F)** R7K and R11K variants of Yap8 are defective in binding to the *ACR3* promoter as determined by EMSA.

Table 2. Summary of functional analysis of Yap8 mutant proteins

Mutant	Mutated	As(III)	Mutant	Model
Name	Region	Resistance	Class	Prediction
P4A	N-term	+++	F	Not included in the model
R5A	N-term	+	PF	Not included in the model
G6A	N-term	+	PF	Not included in the model
R7A	N-term	+	PF	interacts with DNA bases and backbone
K8A	N-term	+++	F	Residue's backbone interacts with DNA backbone
G9A	N-term	+	PF	Plasticity of N-term region, tighter contact with DNA
G10A	N-term	–	NF	Plasticity of N-term region, tighter contact with DNA, interacts with DNA backbone
R11A	N-term	+	PF	Interacts with DNA backbone
K12A	N-term	+++	F	Interacts with DNA backbone
P13A	N-term	+++	F	Model shows no contact with DNA
S14A	N-term	+++	F	Interacts with DNA backbone
L15A	N-term	+++	F	Model shows no contact with DNA
T16A	N-term	+++	F	Model shows no contact with DNA
P17A	N-term	+++	F	Model shows no contact with DNA, caps the basic region
P18A	N-term	+++	F	Model shows no contact with DNA, caps the basic region
K19A	N-term	+++	F	Some contact with DNA backbone
N20A	N-term	+	PF	No contact with DNA, defines the conformational space of N-terminal region to enable tighter protein-DNA contacts
K21A	N-term	+++	F	Interacts with DNA backbone, seen only for monomer 2
R22A*	Basic	–	NF	Direct contact with TGA bases of Y8RE
A23	Basic	ND	ND	Model shows no contact with DNA
A24	Basic	ND	ND	Model shows no contact with DNA
Q25A	Basic	–	NF	Interacts with DNA backbone
L26A*	Basic	–	NF	Model shows no contact with DNA, but this observation is sensitive to definition of a hydrophobic interaction
R27A*	Basic	++	PF	Interacts with DNA backbone
A28	Basic	ND	ND	Model shows no contact with DNA
S29A*	Basic	+++	F	Interacts with DNA backbone
Q30	Basic	ND	ND	Forms several H-bonds with bases from the central region of Y8RE
N31A*	Basic	+++	F	Some contact with DNA backbone
A32	Basic	ND	ND	Model shows no contact with DNA
F33	Basic	–	NF	Hydrophobically interacts with TGATT
R34	Basic	ND	ND	Interacts with DNA backbone and central A base of Y8RE
K35A	Basic	+++	F	Model shows no contact with DNA
R36A	Basic	++	PF	Direct contact with DNA backbone
K37A	Basic	+++	F	Model shows no contact with DNA
L38A	Basic	+++	F	Model shows no contact with DNA
E39A	Basic	+++	F	Model shows no contact with DNA
R40A	Basic	++	PF	Model shows no contact with DNA

Mutated region: N-term refers to N-terminal region preceding the basic region. As(III) resistance: –, none; + or ++, partial; +++, full. Mutant class: F – functional, PF – partially functional, NF – non-functional. ND – not determined. *Determined by Amaral *et al.* (18).

fer between the monomers (Figure 7A, Supplementary Table S3).

In the unstructured N-terminal regions (residues 7–16), we observe that Arg7 of both monomers form strong and stable hydrogen bonds with the T3_W and T22_W bases of the flanking sequences (subscripts ‘W’ and ‘C’ indicate correspondingly the 5′-3′ and the 3′-5′ DNA strands); Arg11 residues form salt bridges with the DNA backbone. But the occupancies and the lifetimes of the contacts vary between the monomers, presumably reflecting the different nucleotide composition of the Y8RE flanking sequences. There is also a number of hydrogen bonds formed between the backbones of the protein and DNA, which stabilize the N-terminals–DNA interactions, involving Arg7, Lys8, Gly10, Arg11 and Lys12. In the basic region (residues 17–40), we observe Arg22 and Asn25 residues interacting with the T7_W and G8_W bases of the TGA-triplet of the Y8RE-sequence. The model structure suggests that Leu26 of monomer 1 could participate in hydrophobic interactions with the methyl groups of T10_W, and Leu26 of monomer 2—with either T14_W or T15_C DNA bases. However, we

do not observe these interactions during the MD simulations, although this observation is sensitive to the definition of a hydrophobic interaction. Here, we employed the 6 Å distance between centers of masses of the corresponding residues. Gln30 of monomer 1 forms hydrogen bonds with the T10_W and A11_C bases, while Gln30 of monomer 2 forms hydrogen bonds with the T14_W and T15_C bases. Arg34 residue of monomer 2 forms a hydrogen bond with the A13_W base. Arg34 of monomer 1 does not exhibit a symmetric interaction and only participates in a number of salt bridge contacts with the DNA backbone. Salt bridge contacts with the DNA backbone are also observed for Arg22, Asn25, Arg27, and Arg34 residues of monomer 1; and for Lys21, Asn25, Arg27, Asn31, Arg34 and Arg36 residues of monomer 2, though again the occupancies and the average lifetimes of the interactions differ between the monomers (Supplementary Table S3). Overall, monomer 2 appears to have a tighter interaction interface with the lower half of the YRE sequence (Figure 7A, Supplementary Table S3).

To investigate the role of Asn20 for the protein-DNA complexation, we repeated the 500 ns MD simulations for

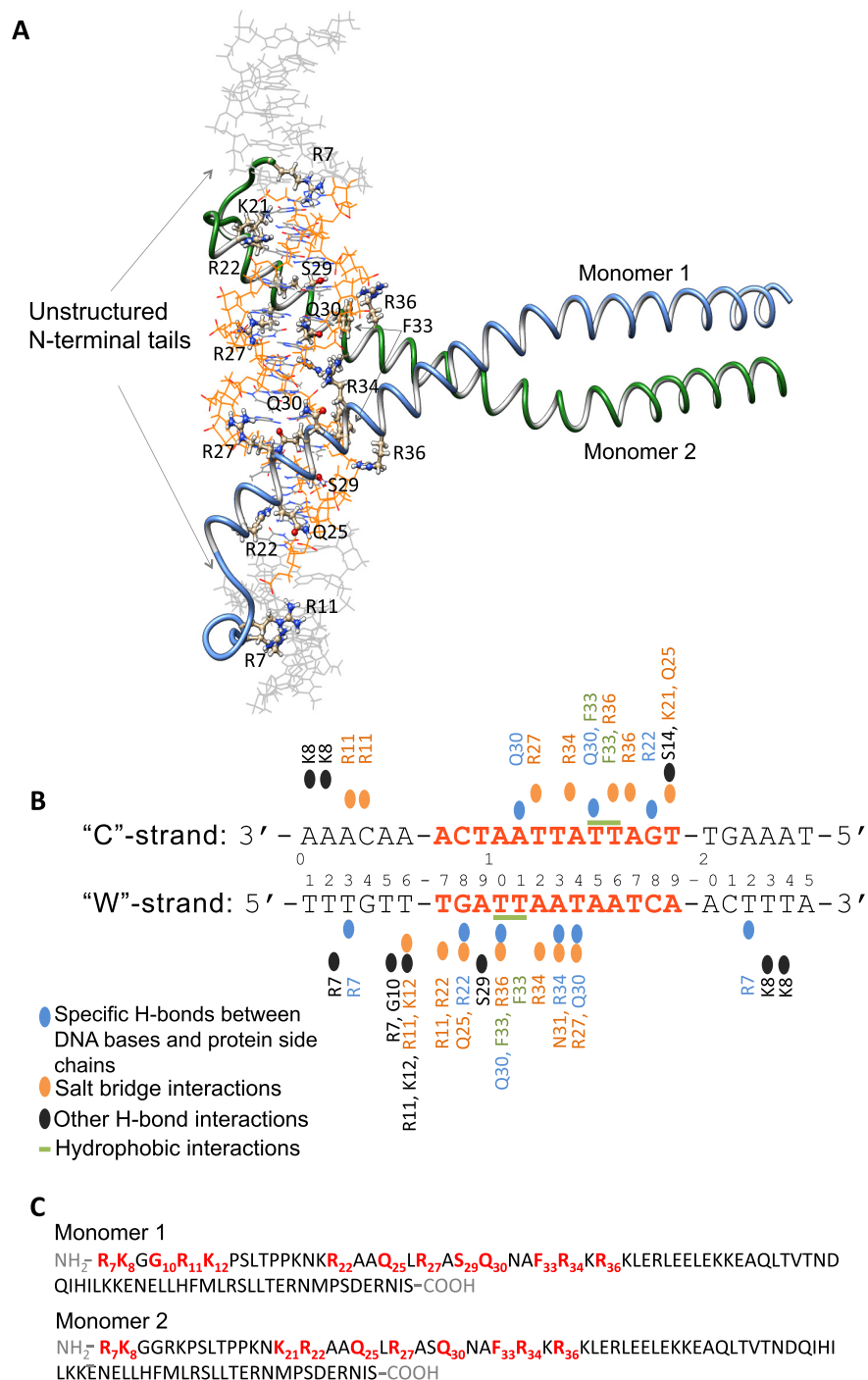


Figure 6. Model structure and interactions network of the Yap8–DNA complex. (A) Model structure of Yap8 protein homodimer in complex with the 25 bp long DNA segment containing Yap8 response element (Y8RE), in orange. Each Yap8 monomer, shown with labeled major DNA-interacting residues, includes an unstructured N-terminal region (residues 7–16) and basic leucine zipper domain (residues 17–89). (B) Schematic overview of the protein–DNA interactions. The DNA sequence, used in the model, is numbered 1–25 with the ‘Watson’ (‘W’) strand representing the 5’-3’ direction and the ‘Crick’ (‘C’) strand – the 3’-5’ direction. Only the interactions that occur at least 25% of the time of the 0.5 μ s MD simulation are depicted. (C) Amino-acid sequences, residues 7–89, of Yap8 monomers included in the model. In bold-red are the residues that show the stable interactions with DNA.

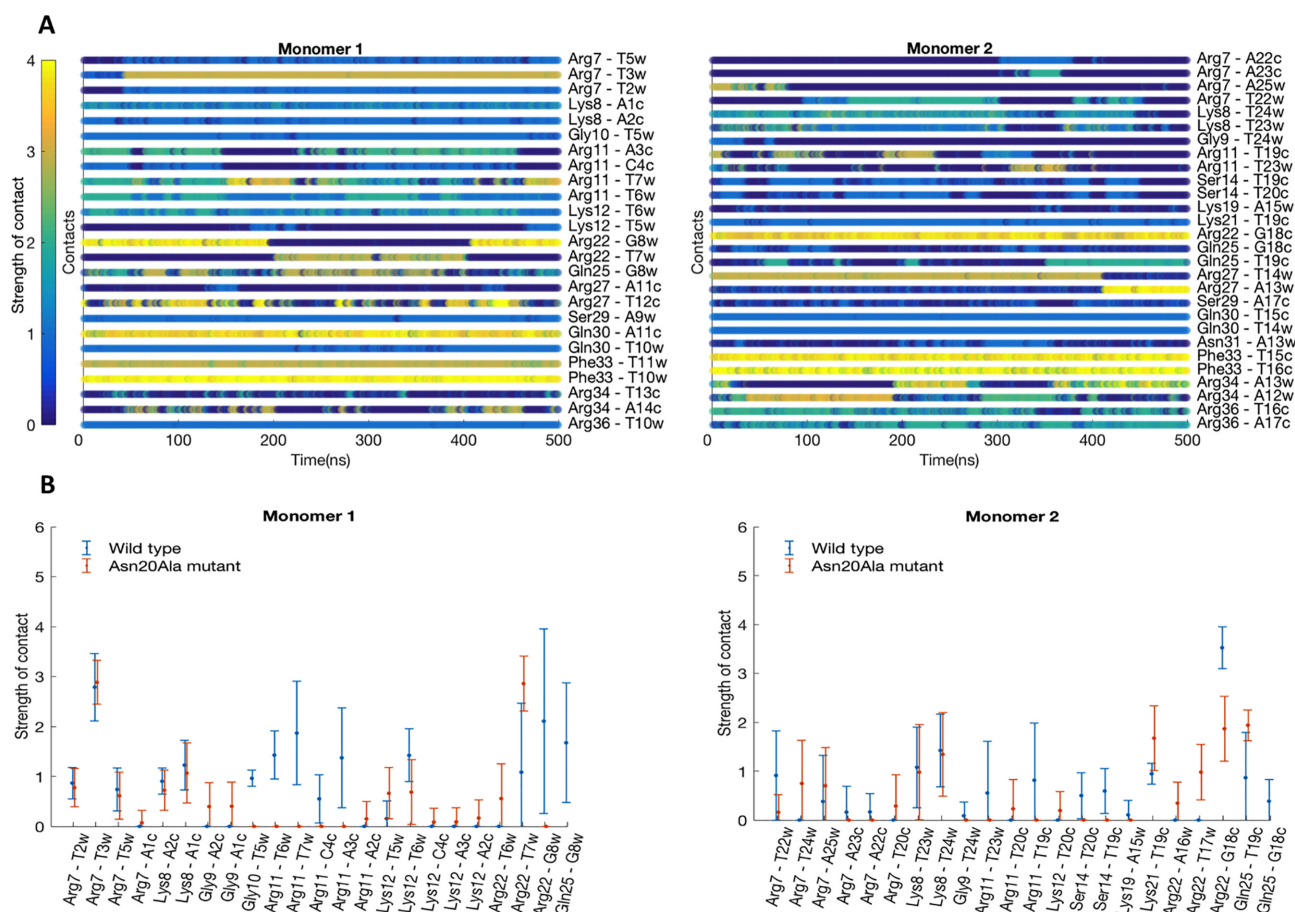


Figure 7. Characterization of Yap8–DNA interactions derived from the MD simulations. **(A)** Dynamic interactions maps illustrating the intermolecular wild-type Yap8–DNA interface. The interactions between pairs of the protein–DNA residues are characterized by a contact strength and its occurrence during the 0.5 μ s MD simulation. **(B)** Comparison of the interaction patterns between pairs of residues of Yap8 wild-type or N20A-mutant proteins and DNA observed during the 0.5 μ s MD simulations. Each specific contact is characterized by mean value of the contact strength and its standard deviation.

the N20A Yap8 mutant. Except for the N to A mutation in Yap8 N20A, the starting structures of the wild-type and the mutant complexes were identical. The intermolecular interface was again characterized by the dynamic interactions maps (Supplementary Figure S6), the contacts occupancies and average lifetimes (Supplementary Table S4). When comparing the wild-type Yap8–DNA and the N20A mutant–DNA interaction patterns, we observe a number of deviations in the N-terminal regions right before the start of the basic domain, residues 9–16. Interestingly the interactions with DNA exhibited by the residues further toward the N-terminus, Arg7 and Lys8, of both proteins are nearly identical in their strength, occupancy, and average lifetimes (Figure 7B). This observation suggests that Asn20 residue, even though not directly interacting with DNA might influence the conformational space of the N-terminal tails enabling a tighter protein–DNA contacts network.

Regions flanking the core Y8RE contribute to stable binding of Yap8 to the *ACR3* promoter

Our model of the Yap8–DNA complex predicts that the unstructured N-terminal tails of Yap8 interact with A/T-rich

regions flanking Y8RE and thus stabilizing binding of the Yap8 basic region to the 13 bp (TGATAAATAATCA) motif (Figure 6). We compared sequences of known and putative promoter regions targeted by Yap8 orthologues and found that the Y8RE flanks are indeed enriched for A/T bp (Figure 8A). Importantly, no such enrichments were found in regions flanking YREs in *S. cerevisiae* genome (data not shown). To test the importance of A/T-rich flanks of Y8RE for Yap8 binding to the *ACR3* promoter, we designed three variants of Y8RE-containing DNA fragments that are enriched in A/T base pairs only in 5' (*ACR3*-M1) or 3' (*ACR3*-M2) flank of the core Y8RE or show no A/T enrichment in both flanking regions (*ACR3*-M3) (Figure 8B). Next, we analyzed Yap8 binding to these variants of the *ACR3* promoter by EMSA. Yap8 showed moderately reduced binding ability to *ACR3*-M1 and *ACR3*-M2 DNA variants, whereas Yap8 binding to the *ACR3*-M3 fragment enriched in G/C bases in both flanks of Y8RE was significantly compromised (Figure 8B). Moreover, binding affinity of Yap8 to the G/C-rich version of the Y8RE-containing region was reduced by half compared to the wild type sequence (Figure 8C; Table 1). In sum, our data suggest that Yap8 binding to an unusually long Y8RE of 13 bp via the basic region is stabilized by specific interactions of un-

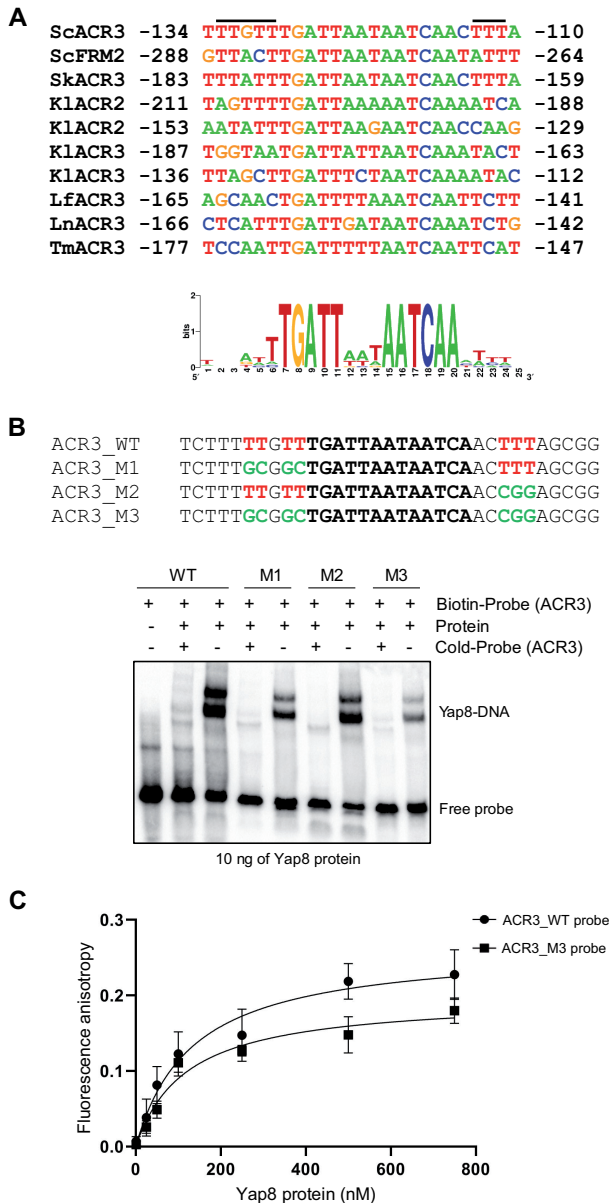


Figure 8. The importance of A/T-rich regions flanking Y8RE for Yap8 binding to the *ACR3* promoter. (A) Alignment of confirmed (ScACR3, ScFRM2, KlACR2, KlACR3) and putative (SkACr3, LfACR3, LnACR3, TmACR3) Y8RE promoter regions targeted by Yap8 orthologues from *S. cerevisiae* (Sc), *S. kudriavzevii* (Sk), *K. lactis* (Kl), *L. fermentati* (Lf), *L. nothofagi* (Ln) and *T. microellipsoides* (Tm). Putative interaction sites of unstructured N-tails of Yap8 are indicated with black bars. The sequence conservation logo of 13-bp Y8REs together with 6-bp flanking regions was determined by the WebLogo application (<https://weblogo.berkeley.edu>). (B) Binding of Yap8 protein to the *ACR3* promoter variants as determined by EMSA. Purified GST-Yap8 protein was incubated with indicated biotin-labeled oligonucleotides corresponding to Y8RE-containing promoter fragments of *ACR3* gene followed by electrophoresis. (C) Fluorescence anisotropy assays performed with purified GST-Yap8 and the FAM-labeled wild type and G/C-rich *ACR3* promoter (ACR3-M3) fragments.

structured N-terminal tails with A/T-rich regions flanking Y8RE.

DISCUSSION

How does Yap8 achieve binding specificity toward its 13 bp recognition element? The characteristic feature of the Pap1 subfamily of bZIP proteins, including Yap1 to Yap7, is the presence of the conserved RxxxNxxAQxxFR motif in the DNA binding basic region (Figure 1). It has been shown for the Pap1-DNA complex that the signature residues Asn86, Ala89, Gln90, Phe93 and Arg94 are involved in direct interactions with DNA bases of the TTAC half-site of the 8 bp YRE whereas Arg82 binds to the guanine flanking the TTAC sequence (2). Four additional conserved Pap1 residues (Gln85, Arg87, Arg91 and Arg96) interact with the phosphate backbone (2). In Yap8, the conserved Asn and Ala residues in the DNA recognition sequence (NxxAQxxFR) are replaced with Leu26 and Ser29, and Arg91 involved in interaction with phosphate in Pap1 is replaced with Asn31 (Figure 1). Recently, alanine replacement analysis within the Yap8 basic region revealed the importance of the highly conserved residues Arg22, Gln25, Arg27, and Arg36 (corresponding to Arg82, Gln85, Arg87, and Arg96 of Pap1) for Yap8-DNA binding (Table 1) (18). In the case of the Yap8-specific residues Leu26, Ser29 and Asn31 (corresponding to Asn86, Ala89 and Arg91 of Pap1), only the L26A mutation impaired the DNA binding activity of Yap8. Interestingly, concomitant replacement of Leu26 and Asn31 with Asn and Arg (present in the corresponding positions in Pap1 and Yap1) extended the binding ability of Yap8 to the YRE motif (TTACTAA) as shown by *in vitro* EMSA assay (18). At the same time, the double L26N N31R Yap8 variant retained its ability to bind to Y8RE and complemented the arsenic sensitivity of the *yap8*Δ mutant (18). Consistent with these findings, we showed that Gln25 and Arg36 are important for Yap8 activity. Moreover, the quadruple A23T L26N S29A N31R mutant (or Yap8-4aa), having the core of the basic region identical with that of Yap1, retained full ability to induce *ACR3* expression and to bind Y8RE *in vitro* (Figures 2 and 3). Yap8-4aa showed low-affinity binding to the *GSH1* oligo containing the TTAGTCA motif (Figure 4) but was unable to bind to the *TRX2* promoter with two YREs (TTACTAA) (Figure 4). This suggests that structural elements outside the basic region may contribute to the DNA binding specificity of Yap proteins.

It has been previously shown that amino acid residues flanking the basic region are important for DNA-binding activity and DNA-target specificity of bZIP proteins (40,41,44). For example, transcription factors belonging to the mammalian Maf subfamily, bind to a 13–14 bp MARE consensus element (TGCTGAC(G)TCAGCA) (41,42). MARE consists of TGC and GCA flanks and the core motif TGACTCA (12-*o*-tetradecanoylphorbol 13-acetate (TPA)-responsive element, TRE) or TGACGTCA (cyclic AMP-responsive element, CRE). The CRE motif is also recognized by mammalian AP-1 (Jun-Fos heterodimer) and CRE binding protein (CREB/ATF), respectively. It was proposed that the N-terminal extended homology re-

gion (EHR) preceding the basic region (42) together with the substitution of the basic region Ala—a highly conserved residue, critical for DNA recognition in other AP-1 proteins (12)—with Tyr (RxxxNxxYAxCR) (45) determines the atypical binding specificity of Maf proteins. Unexpectedly, the X-ray crystal structure of the MafG-DNA complex revealed that the MafG-specific Tyr64 and EHR are not involved in MARE recognition (43). Instead, the invariant Arg57 and Asn61 residues (RxxxNxxYAxCR), corresponding to Arg82 and Asn86 of Pap1, or Arg22 and Leu26 of Yap8 (Figure 1) directly contact the GC bases of the flanks instead of MARE. Binding of the basic region helix is stabilized by a network of hydrogen bonds formed by the residues of the basic region, including MafG-specific Tyr64, and several N-terminal residues either adjacent to the basic region or those forming short α -helices of EHR (43). Interestingly, the yeast bZIP transcription factor Hac1 involved in the unfolded protein response exhibits dual DNA binding specificity, and recognizes either short (6–7 bp) or extended (11–13 bp) motifs within target gene promoters (46). Importantly, the N-terminal region of Hac1 is required for the dual site recognition: the individual basic residues within this region contribute to the alternative specificities (46). To summarize, these observations suggest that N-terminal regions preceding the bZIP domains facilitate DNA binding and contribute to target gene specificity.

Here, we show that the N-terminal region adjacent to the basic region is critical for high affinity binding of Yap8 to the long (13 bp) Y8RE motif (Figures 3 and 5) and for induction of *ACR3* expression *in vivo* (Figures 2 and 5; Table 1). Mutational analysis of the Yap8 EHR revealed several arginines (Arg5, Arg7, Arg11) and glycines (Gly6, Gly9, Gly10), which are important for Yap8-dependent *ACR3* activation (Figure 5B). Moreover, we confirmed that Arg7, Gly10 and Arg11 facilitate Yap8 high affinity binding to the Y8RE motif (Figure 5C–F). The model of Yap8–DNA complex (Figures 6 and 7A) generally supports the experimental data (Figures 3, 5 and 8), except for the predicted salt-bridge nature of Arg11–DNA contacts since the R11K variant shows the *ACR3* promoter binding deficiency suggesting specific interactions with DNA (Figure 5F). The model suggests that the N-terminal tail is engaged in a tight network of contacts between the protein and the Y8RE-DNA flanking sequences enriched in A/T bases, which enables stable positioning of the α -helical basic region in the major grooves of the Y8RE motif. The binding pose of the α -helical basic region allows contacts between Arg22, Asn25, Arg27, Ser29, Gln30, Phe33 and Arg36 residues and the extended TGATT half-site, while the central adenine base is recognized by Arg34 of Yap8 monomer 2. Interestingly, the contacts occupancies and average lifetimes, observed in 0.5 μ s MD simulation, differ between the two monomers (Figure 7B). This observation could result from the asymmetry of bZIP coil-coil protein dimerization, or the varied interactions patterns of the Y8RE–DNA flanks and the N-terminal regions.

Our data show that Asn20 adjacent to the basic region is critical for high affinity binding of Yap8 to the 13 bp long Y8RE motif (Figures 2 and 3). The model suggests that Asn20 is not in direct contact with DNA, but influences the conformational space of the N-terminal tails (re-

gion 9–19). The MD simulations of N20A Yap8 mutant-DNA complex showed that the mutant exhibits less stable contacts between the N-terminal and DNA (Figure 7B and Supplementary Figure S6), which could influence the overall stability of Yap8–DNA complexation.

Alignment of fungal AP-1 protein sequences revealed that residues corresponding to Gly10 and Arg11 in Yap8 are conserved in several member-proteins, including Pap1 (Figure 1). Of these, *Kluyveromyces lactis* Yap1 shows the closest similarity to the N-terminal tail of Yap8 and contains five of seven residues found to be important for Yap8 binding to the 13 bp Y8RE motif (Figure 1). We have previously shown that K1Yap1 contributes to activation of *ACR2* and *ACR3* genes in *K. lactis* suggesting that it exhibits broader DNA binding specificity (17). Importantly, *K. lactis* Yap1, which contains the N-terminal region similar to Yap8, is able to partially complement lack of Yap8 in *S. cerevisiae* (our unpublished data). We propose that the composition of the N-terminal region preceding the basic region influences the repertoire of DNA motifs recognized by AP-1 proteins and dictate target gene specificities. Interestingly, it has been recently shown that the high-affinity binding of the bZIP HapX-CBC (CCAAT-binding complex) transcription factor from *Aspergillus* species to the target promoters requires the conserved KPGRK motif (corresponding to 8-KGGRK-12 of Yap8) that binds to a A/T-rich region located upstream of YRE half-sites (47). It is important to emphasize that the MafG-DNA complex is the only crystal structure of bZIP domain dimer bound to DNA obtained with the protein fragment containing the N-terminal region (43). Investigating the significance of the N-terminal region of other bZIP proteins for DNA binding specificity will unveil the mechanisms employed by bZIP transcription factors for recognition of target gene sites.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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