DNA structure directs positioning of the mitochondrial genome packaging protein Abf2p

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ABSTRACT

The mitochondrial genome (mtDNA) is assembled into nucleo-protein structures termed nucleoids and maintained differently compared to nuclear DNA, the involved molecular basis remaining poorly understood. In yeast (Saccharomyces cerevisiae), mtDNA is a ∼80 kbp linear molecule and Abf2p, a double HMG-box protein, packages and maintains it. The protein binds DNA in a non-sequence-specific manner, but displays a distinct ‘phased-binding’ at specific DNA sequences containing poly-adenine tracts (A-tracts). We present here two crystal structures of Abf2p in complex with mtDNA-derived fragments bearing A-tracts. Each HMG-box of Abf2p induces a 90° bend in the contacted DNA, causing an overall U-turn. Together with previous data, this suggests that U-turn formation is the universal mechanism underlying mtDNA compaction induced by HMG-box proteins. Combining this structural information with mutational, biophysical and computational analyses, we reveal a unique DNA binding mechanism for Abf2p where a characteristic N-terminal flag and helix are crucial for mtDNA maintenance. Additionally, we provide the molecular basis for A-tract mediated exclusion of Abf2p binding. Due to high prevalence of A-tracts in yeast mtDNA, this has critical relevance for nucleoid architecture. Therefore, an unprecedented A-tract mediated protein positioning mechanism regulates DNA packaging proteins in the mitochondria, and in combination with DNA-bending and U-turn formation, governs mtDNA compaction.

INTRODUCTION

Mitochondria, the organelles to which aerobic eukaryotes owe their existence, possess their own genome, the mitochondrial DNA (mtDNA). This mtDNA codes for components of the electron transport system, which is essential for adenosine triphosphate production (1–3). In addition, it encodes ribosomal proteins, ribosomal RNA and tRNAs (3,4). A species-specific High-Mobility Group-box (HMG-box) protein encoded in the nucleus is the main architectural component that compacts and packages mtDNA into nucleoprotein structures, termed nucleoids (5). In the budding yeast Saccharomyces cerevisiae, y-mtDNA is a predominantly linear molecule (6–8) of 70–85 kilo base pairs (kbps), which is packaged by the ARS (autonomously replicating sequence)-binding factor 2 protein, Abf2p (9–12).

Saccharomyces cerevisiae is a facultative aerobe that grows in both fermentable (e.g. glucose) and non-fermentable (e.g. glycerol) media. In non-fermentable media, the oxidative phosphorylation pathway involves the electron transport system (13), thus requiring intact, transcriptionally active y-mtDNA. Deletion of the Abf2 gene causes loss of mtDNA, resulting in yeast cells that grow in fermentable media but cannot subsequently survive in non-fermentable conditions (9). This suggests that Abf2p holds an essential role in mtDNA maintenance (12,14). Moreover, a 2- and 3-fold increase in Abf2p levels leads to a 2-fold increase in y-mtDNA copy number, while a 8- to 10-fold Abf2p overexpression causes rapid loss of y-mtDNA and generation of mtDNA-depleted (petite) mutant cells (14). The latter has been related to excessive

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y-mtDNA compaction that would impair the access of the replicating machinery to the DNA. Thus, mtDNA transactions and compaction need to be in close balance and are regulated by Abf2p levels.

Abf2p organizes y-mtDNA into nucleoids via sequence-non-specific interactions and DNA bending (15,11). However, the protein shows distinct patterns of DNA ‘phased binding’ at ARS, rich in A-T bases and poly-adenine tracts, such as the ARS1 found in the nucleus, and the 64 bp region downstream of the mitochondrial origin of replication 0rs5/rep2 (9,10) (Figure 1A). This phased binding was attributed to Abf2p exclusion from a 11 bp consensus motif 5′-(A/T)AAA(T/C)ATAAA(A/T)-3′ (ARS-c) found in ARS1, and from its near matches (ARS-m, 9 or 10 out of 11 bp) in the 64 bp sequence (9,10). From a functional point of view, ARS-c promotes DNA replication while ARS-m, though not as efficient, can maintain similar ARS activity when present in sufficient number of copies (16). Such ARS activity in the mitochondria has been related to amplification of residual fragments of wild-type mtDNA that, in the yeast p+ cells (cells that have lost large portions of the wild-type mtDNA), are reiterated as concatamers, restoring the total amount of mtDNA to that found in wild-type mtDNA, are reiterated as concatamers, restoring the total amount of mtDNA to that found in non-auxotrophic BL21(H9262) strain. Cells were grown in LB medium for 2 h at 37°C until the optical density (O.D) at 600 nm reached 0.6. After subsequent induction with 1 mM isopropyl β-d-1-thiogalactopyranoside (IPTG), the culture was grown for 4 h at 37°C and shaking at 225 rpm. Cells were sonicated for 10 min (cycle 2 s on/6 s off) on ice in 50 mM Tris–HCl pH 7.5 and 1M NaCl. The lysate was injected in a Ni-NTA-affinity column (HisTrap HP, GE Healthcare) mounted on an AKTA Purifier (GE Healthcare) system. The proteins were eluted with a linear gradient from buffer A (20 mM imidazole, 50 mM Tris–HCl pH 7.5, 750 mM NaCl) to buffer B (500 mM imidazole, 50 mM Tris–HCl pH 7.5, 750 mM NaCl). Quality of the protein fractions was assessed by sodium dodecyl sulphate-polyacrylamide gel electrophoresis (SDS-PAGE) and the purest fractions pooled and concentrated for gel filtration chromatography using a Superdex 75 10/300 column (GE Healthcare) pre-equilibrated with running buffer (50 mM Tris–HCl pH 7.5, 750 mM NaCl). The purity of the peak fractions was analyzed by SDS-PAGE. Regarding Se-met derivatives, since the wild-type mature Abf2p contains only Met147, Leu52 was mutated to methionine (Abf2p-Semet protein variant) to have sufficient phasing power (23). Non-auxotrophic BL-21 pLys bacterial strain was grown at 37°C to an O.D of 0.6 and methionine synthesis inhibition (23) was carried out (inhibition mix was prepared with amino acids purchased from Sigma) prior to induction with 1mM IPTG (Base medium, amino acid nutrient mix and seleno-methionine were purchased from Molecular Dimensions).

Nucleic acids

The oligonucleotides used for this study are listed in Supplementary Table S1. Duplex DNA fragments for crystallization and isothermal titration calorimetry (ITC) were prepared by annealing complementary oligonucleotides (Sigma) Af222fw and Af222rev, Af2_shift22fw and Af2_shift22rev, GC_22fw and GC_22rev for Af22, Af2_shift22 and GC_22 respectively. The oligonucleotides (100 μM) were resuspended in 20 mM Tris–HCl, pH 7.5 and 100 mM NaCl, and heated at 85°C for 30 min followed by slow cooling to room temperature overnight. The (25 bp × 4) four-way DNA junctions were respectively assembled by equimolar hybridization of the 50-mer oligonucleotides (Seq50.1-4; 40 μM each) in 20 mM Tris–HCl pH 7.8, 50 mM NaCl and 5 mM MgCl2. Oligonucleotides were heated for 5 min at 95°C and cooled down to room temperature overnight. M13mp18 dsDNA plasmids (Bayou Biolabs) were linearized with BsrB I (New England Biolabs) and purified with the illustra GFX PCR DNA purification kit (GE Healthcare).

Crystallization, structure solution and analysis

Crystallization trials were attempted with DNA fragments of different sequences and lengths, both GC and AT rich,
Figure 1. DNA design and Abf2p/DNA complex crystal structures. (A) General organization of the y-mtDNA ori5/rep2 origin of replication, including the location of ARS consensus sequences (ARS-c, 11/11 matches to 5′-(A/T)AAA(T/C)ATAAA(A/T)-3′, in orange) and the downstream 64 bp region (in yellow) in y-mtDNA17. Below, the 64 bp sequence is shown, the near matches (ARS-m; 9/11) to ARS-c are depicted in orange and the position of the 35 bp sequence (see Introduction) is demarcated by vertical bars. Vertical dashed lines indicate the DNA sequence ‘Af22’. (B) The dsDNA sequences used for crystallization are depicted inside arrows that show the head-to-tail arrangement found in the crystals; the DNA regions contacted by Abf2p are framed. Abf2p is represented as two connected L-shaped boxes 1 and 2 (HMG-box 1 and 2, respectively) with arrows representing DNA-inserting residues. The gray HMG-boxes represent the second protein molecule in the asymmetric unit; the dotted lines indicate the domains are connected. From the two ‘Af22’ sequences in tandem, a continuous ‘Af2_shift22’ dsDNA molecule was derived (vertical dashed lines) that yielded the Abf2p/shift22 structure, where the protein again contacts two DNAs. (C) Schematic representation of the Abf2p domains. ‘L’ stands for ‘Linker’, ‘N-flag’ and ‘N-helix’ stand for N-terminal flag and helix, respectively. (D) The crystal structures of Abf2p/22 (left) and Abf2p/shift22 (right) are shown, with secondary structure elements and domains labeled. The intercalating residues Phe51 and Ile124 are highlighted in blue and their insertion sites are shown in insets. The DNA head-to-tail arrangement is indicated with a black arrow. The ARS-m-like regions (see main text) within Af22 and Af2_shift22 are shown in orange, the A-tract sub-region in red. Underneath, the crystallized dsDNA sequences and contacting protein domains are shown; note that the DNA sequences are in the same orientation while the Abf2p domains are in the opposite.
with and without A-tracts. Diffraction quality crystals were obtained only with Abf2p in complex with Af2:22 DNA. The protein/DNA complex for crystallization was prepared by mixing Abf2p and DNA fragment at a molar ratio of 2:1 (protein concentration 0.5 mg/ml) and performing stepwise overnight dialysis to reach a final buffer composition of 50 mM Tris–HCl, pH 7.5, 20 mM NaCl. Crystals were obtained in conditions of 21–25% w/v PEG 4000, 0.1 M tri-sodium citrate pH 4.5 and 0.2 M ammonium acetate at 20°C by sitting drop vapor diffusion method, at a protein concentration of 10–12 mg/ml. Crystals were cryo-protected with a solution of 15% glycerol, 21–25% PEG 4000, 0.1 M tri-sodium citrate pH 4.5, 0.2 M ammonium acetate and vitriﬁed in liquid nitrogen. Data collection of intermediate crystals and optimized ones was performed at synchrotron ALBA (Cerdanyola del Vallès, Spain) and European Synchrotron Radiation Facility, ESRF (Grenoble, France). Specifically, diﬀraction datasets for Abf2p-Semet in complex with Af2:22 were collected at the selenomethionine peak energy of 12.6623 keV, to a nominal resolution of 3 Å and with Friedel pairs collected on the same image. Crystals belonged to the space group C2. Diﬀraction data were processed with XDS (24) and Aimless (25, 26). Single wavelength anomalous diﬀraction phasing was performed with anisotropy corrected data using Sheld C E (27) followed by density modiﬁcation using DM (28). A partial model could be built into the resulting Fourier map through iterative cycles of model building, visual inspection and automated reﬁnement. Subsequently this partial model was used for molecular replacement with native Abf2p/22 crystal data collected at 12.658 eV with a nominal resolution of 2.18 Å, monoclinic space group C2 (see Table 1). Data processing was performed as mentioned above. For conﬁrming sequence register of Abf2p/22, thymine 6 (T6) of Af2:22 was replaced with 5-bromo uracil (5-BrU) (oligonucleotides purchased from Biomers) and crystals were obtained under similar conditions as Abf2p/22, with space group P21 and a nominal resolution of 3.3 Å. The structure was solved by molecular replacement and the position of the Br atoms was located from an anomalous density map (visible peaks above 5 r.m.s.d) obtained using AnoDe (29). Abf2p/shift22 dataset was obtained with a nominal resolution of 2.6 Å, space group P21 (see Table 1). The structure was solved by molecular replacement and omit map was generated to identify the DNA ends.

CCP4i (28) and the tools therein were used for data manipulations such as truncation (Ciruncate) and merging (Aimless). Phenix (30) was used for automated structure reﬁnement. Coot (31) was used for map visualization, model building and validation. PyMOL (DeLano Scientiﬁc) was used for ﬁgure generation. It is to be noted that for both the structures protein chain A and DNA chains B, C, E, F are treated as the reference as other protein chains in the asymmetric unit (a.u.) have regions with poor Fourier map deﬁnition (resulting in a relatively high number of RSRZ outliers), if not otherwise indicated.

Small angle X-ray scattering (SAXS)

Purified Abf2p was dialyzed overnight in 50 mM Tris–HCl, pH 7.5 and 500 mM or 100 mM NaCl. The protein samples were subsequently concentrated. The ﬁltrate was used as a blank to obtain maximal buffer match between the blank and the sample. The Abf2p/22 complex was prepared identically as for crystallization. Small angle X-ray scattering (SAXS) measurements were performed at the ESRF. Data analysis was performed with Primus (32) in ATSAS package. Guinier approximation was used to calculate radius of gyration (Rg). Molecular weight of the protein was estimated from the Porod volume (32). Data from a concentration series (3.8 mg/ml, 5.4 mg/ml, 7.0 mg/ml, 9.5 mg/ml and 13.5 mg/ml) was collected at 50 mM Tris–HCl, pH 7.5, 500 mM NaCl. The estimated molecular masses are 14.3–19.0 kDa, 16.3–21.8 kDa, 15.1–20.1 kDa, 16.1–21.5 kDa and 16.3–21.8 kDa respectively (the lower and upper estimates for each concentration are obtained by dividing the Porod volume by 2 and 1.5 respectively). These values match closely with the 6-histidine-tagged monomer of Abf2p (19.5 kDa). The radii of gyration (Rg), calculated from the Guinier plot are 27.2 +/- 0.1, 27.3 +/- 0.1, 27.4 +/- 0.1, 27.3 +/- 0.1 and 27.8 +/- 0.1 Å, respectively. Additional data was collected at 50 mM Tris–HCl, pH 7.5, 100 mM NaCl (0.36 mg/ml; no reliable data could be obtained at other concentrations) for which the calculated molecular mass and Rg are 12.6–16.8 kDa and 28.4 (+/- 0.6) Å respectively. The curve corresponding to the highest concentration (13.5 mg/ml) was used for all other subsequent analyses as it showed lesser noise at higher angles. Pair distribution (p(r)) plot and maximum particle dimension (Dmax) were calculated from scattering data using GNOM (32). Ensemble analysis was performed using EOM 2.0 (33, 34) with standard parameters for the genetic algorithm and for Crysol (32). For EOM, we generated a collection of 10,000 models starting from the crystal structure coordinates. The N-flag+N-helix+HMG-box 1 assembly and HMG-box 2 were treated as two rigid domains while allowing ﬂexibility to the linker (113-Lys-Leu-Pro-115, 113 added to include the unwinding of HMG-box 1 helix C-terminus as observed from MD simulations), the N-terminal His-tag and the last two C-terminal residues. Fitting the calculated scattering proﬁle to the experimental SAXS data, yielded χ² = 1.379. As an alternative, we used a frame from the MD simulation (extracted from simulation run after the r.m.s.d. had converged) as a starting model for EOM (Figure 4B). Using the same assignment of ﬂexible regions, EOM generated a subset of five conformations which, collectively, were in better agreement (χ² = 0.957) with the experimental data.

Molecular dynamics simulations

Molecular dynamics (MD) simulations for the protein, DNA and protein–DNA complexes were performed starting from the X-ray crystal structures using Amber 14 package. All simulations were carried out using state-of-the-art simulation conditions, the amber14SB with parmbsc1 reﬁnement (35) force ﬁelds and explicit TIP3P water model. Models were prepared for simulation using tleap in Amber Tools 15. We added sodium and chloride ions to neutralize the systems and give a realistic ionic environment (150 mM NaCl). All simulations were performed using a truncated octahedron periodic box, periodic boundary conditions and the particle-mesh-ewald for long range electro-
Table 1. Data collection and refinement statistics

|                  | Abf2p/22         | Abf2p/shift22     |
|------------------|------------------|-------------------|
| Wavelength       | 0.97950          | 0.97625           |
| Resolution range | 43.48 - 2.18     | 48.05 - 2.6       |
| Space group      | C 1 2 1          | P 1 2 1           |
| Unit cell        | 88.49 113.41 67.71 90 103.803 90 | 71.989 131.991 71.994 90 103.211 90 |
| Total reflections| 126763 (6246)    | 88589 (9083)      |
| Unique reflections| 33305 (3075) | 38396 (3889)      |
| Multiplicity     | 3.8 (2.0)        | 2.3 (2.3)         |
| Completeness (%) | 0.98 (0.92)      | 0.95 (0.97)       |
| Mean l/σ(l)      | 12.01 (1.99)     | 6.09 (1.05)       |
| Wilson B-factor  | 49.68            | 55.15             |
| R-merge          | 0.06649 (0.5319) | 0.1022 (0.8523)   |
| R-meas           | 0.07359 (0.6935) | 0.1296 (1.085)    |
| CC1/2            | 0.998 (0.632)    | 0.995 (0.207)     |
| CC*              | 1.0 (0.88)       | 0.999 (0.585)     |
| Reflections used in refinement | 33305 (3074) | 38396 (3885)      |
| Reflections used for R-free | 1881 (164)  | 1828 (166)        |
| R-work           | 0.2090 (0.3061)  | 0.2267 (0.3656)   |
| R-free           | 0.2364 (0.3213)  | 0.2601 (0.3623)   |
| CC(work)         | 0.956 (0.754)    | 0.948 (0.466)     |
| CC(free)         | 0.938 (0.772)    | 0.907 (0.339)     |
| Number of non-hydrogen atoms | 4520           | 8996              |
| Macromolecule atoms | 4445           | 8772              |
| Protein residues | 313              | 618               |
| RMS(bonds)       | 0.008            | 0.008             |
| RMS(angles)      | 0.95             | 1.00              |
| Ramachandran favored (%) | 99                  | 98               |
| Ramachandran allowed (%) | 0.65            | 1.6              |
| Ramachandran outliers (%) | 0                   | 0                |
| Rotamer outliers (%) | 0.7               | 1.3              |
| Clashscore       | 8.45             | 12.79             |
| Average B-factor | 58.93            | 61.38             |
| B-factor macromolecules | 59.04            | 61.53             |
| B-factor solvent | 52.14            | 50.10             |

* Statistics for the highest-resolution shell are shown in parentheses.

DNA deformation energy and DNA stiffness calculations
DNA structures can be described in terms of base-pair and base-step parameters that consist of three translations (shift, slide and rise) and rotations (tilt, roll and twist) and the DNA deformability along these six directions can be described by the associated stiffness constant matrix (40). From the ensemble of MD simulations, the covariance matrix describing the deformability of the helical parameters for a given DNA fragment (e.g. a dinucleotide step) is computed and is inverted to generate the 6×6 stiffness matrix for each fragment. Pure stiffness constants corresponding to the six parameters mentioned above (kshift, kslide, krise, ktilt, kroll and ktwist) are extracted from the diagonal of the matrix while the total stiffness (Ktot) is obtained as a product of these six constants and provides a rough estimate of the flexibility of each base pair step. The normalized Boltzmann-like probability distribution, exp(-ΔEdef/kBT), was derived from the deformation energy (ΔEdef) and was used to determine the relative probability of positioning the HMG-boxes on a given DNA segment. The deformation energy was calculated using a mesoscopic energy model (40,41), which is based on a harmonic approximation to describe deformability along DNA helical parameters. The equilibrium geometry and stiffness force constants were extracted from a dataset built from long all-atoms MD simulations of short oligonucleotides in water using the parmbsc1 force field (35).

Abf2p in vivo assays
Diploid ABF2 (YMR072W) hemizygous strain Y26205 (BY4743; MATa/MATa; ura3Δ0/ura3Δ0; leu2Δ0/leu2Δ0; his3Δ1/his3Δ1; met15Δ0/MET15; LYS2/lys2Δ0; YMR072/w/YMR072/w:KANMX4) was obtained from the European S. cerevisiae Archive for Functional Analysis (Euroscarf). A plasmid-borne ABF2 was made by cloning a PCR fragment including Abf2 CDS plus 500 nt on both sides into pRS316 (42). This construct was introduced in Y26205 diploid cells, which were grown on uracil deficient synthetic dextrose (SDC) medium and subsequently sporulated. Kan" (ABF2 disrupted) Ura" (pRS316 transfected) cells were isolated and the genotype was verified by PCR (43). ABF2 truncations were made by...
PCR while preserving the 500 nt flanks, and were cloned into pRS315 (a plasmid providing leucine prototrophy (42)). To test ABF2 truncations, we followed a plasmid-shuffling strategy using 5-fluororotic acid (5-FOA), toxic for Ura+ cells (44,45). The mitochondrial DNA integrity of the corresponding Kan+ Ura+ Leu+ cells was tested by growth in glycerol (respiration-only conditions). For this purpose, a spot assay was performed with 1/5 serial dilutions, starting from cultures grown in leucine deficient SDC medium to an O.D of 0.5. All cultures and plates were grown at 30°C.

Isothermal titration calorimetry (ITC)

Samples for ITC were prepared by simultaneously dialyzing the protein and DNA in the same buffer (25 mM Hepes pH 7.5, 150 mM NaCl) to obtain maximal buffer match between samples. Titrations were performed with a VP ITC instrument with 1400 μl cell volume and 300 μl syringe volume, at a temperature of 25°C. The general setup was designed with protein in the cell and DNA in the syringe. The protein was used at a concentration of 8–16 μM and the DNA at a concentration of 100–150 μM. DNA into buffer titrations were performed as control and since the profile was flat, subtraction of the control was not performed. The resulting data were analyzed using Origin (Origin Lab, Northampton, MA, USA). The dsDNA fragment GC_22 (complementary strands 5′ GAAGATATCCGGGTCCCA ATAA 3′ and 5′ TTATTGGGACCCGGGATATCC 3′) was used as a control.

Electrophoretic mobility shift assays (EMSA)

Two-fold serial dilutions of Abf2p and deletion mutants were performed on ice in 20 mM Tris–HCl pH 7.5 and 750 mM NaCl. Reactions were initiated by addition of proteins as appropriate and incubated for 30 min at 25°C. dsDNA binding experiments contained 2ng-μl−1 linearized M13mp18 and were performed in a binding buffer with a final composition of 20 mM Tris–HCl pH 7.5 and 100 mM NaCl. Subsequently, samples were resolved on 25 cm long, 1.2% (w/v) agarose gels (SeaKem LE Agarose, Lonza) in 0.5× TBE (Sigma). Gels were run in a Sub-Cell GT cuvette (BioRad) for 17–18 h at room temperature at 3V/cm. The reactions with the four-way junctions contained 100 nM of the (25 bp × 4) DNA and the buffer was supplemented with 1mM MgCl2. After addition of a loading dye, the binding reactions were loaded on a 7% polyacrylamide gel (Sigma) in 0.5 × TBE. Gels were pre-run for 15 min and run at 11V/cm on a HoefferMiniVE cuvette placed on ice. Agarose and polyacrylamide gels were stained with SybrSafe (Molecular Probes) and scanned for fluorescence using a Typhoon 8600 scanner (GE Healthcare).

Fluorescence confocal microscopy

Kan+ Ura+ Leu+ cells harboring different plasmid borne Abf2p truncations were grown in leucine deficient SDC media to O.D. of 0.5–1. Positive and negative controls were grown in glycerol and YPD respectively. Positive control consisted of wild-type (ABF2 intact) haploids and negative control consisted of Kan+ (ABF2 disrupted) haploids. Cells were fixed with 4.5% formaldehyde for 1 h, treated with zymolyase Z100T (Ecogen) for 20 min at 30°C, deposited on poly-lysine coated slides and stained with 300 mM solution of 4′, 6-Diamidine-2-phenylindole dihydrochloride (DAPI) for 5 min prior to mounting. Fluorescence confocal microscopy images were acquired on a Zeiss LSM780, with 64X oil immersion objectives, numerical aperture 1.4. Excitation wavelength was 405 nm (Diode laser) and emission range was 419.47–535.16 nm. Pixel size for acquired images was 0.05 μm. For each sample, maximum intensity projection of the stacks was performed with Fiji (46). Counting of cytoplasmic DAPI foci (proxy for mtDNA content) was performed manually for 50 cells per sample type and an average was obtained.

Circular dichroism

The protein samples were dialyzed overnight in 10 mM potassium phosphate pH 7.5, 100 mM ammonium sulfate. A quartz sample cell of 1 mm path length was used for data collection on a JascoJ-815CD spectrophotometer, with a scan rate of 50 nm/min and data pitch of 0.5 nm resulting in 150 data points between 260–185 nm. Protein concentration was kept constant at 0.1mg/ml for all samples.

**RESULTS**

Well-diffracting crystals were obtained for Abf2p (mature form, without the mitochondrial targeting sequence) bound to an A-T rich double-stranded (ds) DNA, A1f2,22 (5′-AATAAATAATTATATATAA-3′, Figure 1B), spanning 22 of the 35 bp sequence (19477 to 19511bp in ρ− mtDNA (4), Figure 1A) that is present in the wild-type *S. cerevisiae* ρ− mtDNA and additionally amplified as concatemers in ρ− yeast (*S. cerevisiae*). This sequence is expected to be recognized by Abf2p in vivo. The crystal structure (hereafter termed Abf2p/22; PDB ID 5JH0; Table 1; Supplementary Figure S1) reveals an asymmetric unit (a.u) composed of two Abf2p molecules (polypeptide chains Abf2p-A and -D) and two double-stranded (ds) DNAs (chains-BC and -EF, 22 bp each) (Figure 1B and D). Abf2p consists of two HMG-boxes in tandem, HMG-box 1 and 2. At the N-terminus of the protein, three residues in extended conformation (27-Lys-Ala-Ser-29, henceforth referred to as ‘N-flag’) and a 12 amino-acid (aa) N-terminal helix (N-helix) (Figures 1C, D and 2A) are found. Notably, such a helix is not present in any structure containing HMG-boxes reported so far. The N-helix leads to HMG-box 1 (42–113) (Figures 1C and 2A), which displays a typical HMG-box ‘L-shape’ consisting of an extended segment (42–48), helix1 (49–64, with a kink at position 57), a four-residue turn, helix2 (70–83, which with helix1 forms the short L-arm), a short two-residue connecting loop and finally, helix3 (86–113, antiparallel to the extended segment and forming the long L-arm of the HMG-box L-shape). A Linker of two residues (Leu-Pro) follows, which is much shorter than the predicted length of 10aa (9). Similar to HMG-box 1, HMG-box 2 (116–179) consists of an extended segment (116–121), helix1 (122–137), a five-residue turn, helix2 (143–156, with a kink at position 130) and a two-residue connecting loop

*6 Nucleic Acids Research, 2016*
leading to helix3 (159–179, Figures 1C, D and 2A). Finally, the last four residues 180-Tyr-Pro-Leu-Asn-183 form a short extended C-terminal segment.

**Abf2p binds to two separate DNAs via its two domains**

In the a.u. of the Abf2p/22 crystal, two curved dsDNA molecules of 22 bp are arranged head-to-tail by two protein molecules, forming a pseudo-continuous DNA circle exhibiting stacking interactions between respective 3'-5' DNA ends (Figure 1D). Thus, each protein holds together and bends two different DNA molecules via its HMG-boxes, much like a staple (hereafter referred to as dual-binding) (Figure 2A). Specifically, HMG-box 1 from Abf2p-A contacts dsDNA EF chains (from T13 to A19 of chain-E), while HMG-box 2 contacts the other DNA molecule BC (from A2-A5 chain-B), with the DNA ends participating in perfect stacking interactions between HMG-box binding sites (Figures 1B, D and 2A). A similar scheme applies for the second protein molecule in the a.u. where HMG-box 1 and HMG-box 2 contact dsDNA chains BC and EF respectively. Each HMG-box engages in mostly electrostatic interactions with the DNA minor groove (Supplementary Figure S2), opens it up to 12 Å (ideal B-DNA standard minor groove width 5.9 Å, Figure 2B) and concomitantly bends the DNA by ~90°. Additionally, each HMG-box inserts hydrophobic residues between base steps, thus causing considerable alterations of DNA parameters (Figure 2B). Specifically, HMG-box 1 Phe51 inserts at step A8/T9/A14/T15 (chains F/E), the phenyl ring stacking with the six-membered ring of A8 (Figure 1B and D). This base-step shows a high positive roll and the involved base pairs show a prominent buckle (Figures 1D, and 2B). In comparison, Ile124 from HMG-box 2 inserts partially at T3/A4/T19/A20 (B/C), and induces a positive roll (Figures 1D and 2B). Superposition of both HMG-boxes indicates high structural similarity (r.m.s.d. of 1.1 Å for 66 Cα), both Phe51 and Ile124 being at equivalent positions in helix1. Thus, the two DNAs that are contacted by a single protein together form a U-turn (or a half DNA circle, Figure 2A). Importantly, the Abf2p mutant K44A, R45A, K117A and K118A causes 80% of *S. cerevisiae* cells to lose mtDNA (14,47) (and thus only survive in fermentable conditions). These amino acid residues correspond to a conserved Pro-B-B-Pro (B, basic aa) motif found in the N-terminal extended segment of either HMG-box domains, 43-Pro-Lys-Arg-Pro-46 in HMG-box 1 and 116-Pro-Lys-Arg-Pro-119 in HMG-box 2. The structure shows the NZ at the tip of these lysine side-chains being involved
Figure 3. Abf2p truncations and their effect on DNA binding. (A) Schematic representation of the Abf2p deletion mutants. The domains are designated as in the text except for the linker, indicated with an ‘L’. Dashed lines indicate deleted regions. (B) Titration of Abf2p and Mut1 to Mut6 constructs binding on linearized M13mp18 dsDNA (400 pM) at 25°C by EMSA. The protein/bp ratios are indicated and correspond respectively to 0.37, 0.75, 1.5, 3, 6 and 12 pmoles of Abf2p. (C) Titration of Abf2p and Mut1 to Mut6 binding to four-way junction DNA (100 nM) by EMSA. (D) Effect of Abf2p truncations in vivo. Spot assay of cells with 1/5 serial dilutions on plate containing glycerol as carbon-source (‘YPG plate’) shows the effect of different Abf2p truncations on the ability to maintain y-mtDNA and thus on capability of yeast (Saccharomyces cerevisiae) cells to use non-fermentable carbon source. Full-length Abf2p and Mut6 (Abf2p without HMG-box 2) are capable of protecting mtDNA whereas Mut2 (Abf2p without N-flag and N-helix), Mut3 (N-helix+HMG-box 1) and Mut5 (HMG-box 2 alone) are not functional. (E) Visualization of mitochondrial nucleoids in cells harboring different Abf2p truncations by DAPI staining of DNA and fluorescence confocal microscopy. Abf2p and Mut6 maintain mtDNA whereas Mut2, Mut3 and Mut5 cannot. C(+) corresponds to wild-type haploids, and C(-) to Abf2Δcells. The scale bars correspond to 2.5 μm.

[Text continues with detailed analysis of the figure and the figures themselves]
Figure 4. Abf2p flexibility and dynamics. (A) The Abf2p domains in the Abf2p/22 crystal structure are shown (same color-code as in Figure 1, the DNA molecule is not displayed for clarity) together with the amino-acid side-chains (depicted in gray ball-and-sticks and labeled) that participate in the N-hydrophobic core that stabilizes the contact between the N-helix, the C-terminal region of HMG-box 1 helix3 and the linker. (B) A frame from the 500 ns MD simulation of Abf2p that shows the hydrophobic residues from the N-helix exposed to the solvent, the changes in the orientation of N-helix and HMG-box 2 (HMG2) (relative to HMG-box 1 in the crystal structure above) indicated by arrows. (C) Root mean square fluctuation (r.m.s.f, in Å) of individual residues during MD simulation (500 ns) of Abf2p alone (red dots) and bound to DNA (orange dots). The different regions of the protein are schematically represented along the sequence. Note that the Abf2p flexibility reduces in the protein/DNA complex.

We reasoned that if the dual binding mode was caused by a specific preference of the HMG-boxes for the contacted DNA sequence patches, then, joining both patches into a single 22 bp DNA molecule should result in a complex of one protein bound to a single, continuous-DNA. Accordingly, we designed the sequence ‘Af2 shift22’ (5′-TTATATAATATAAAATAATAAA-3′) (Figure 1B). To our surprise, the resulting structure (hereafter referred to as Abf2p/shift22; PDB ID 5JGH; Table 1) again showed dual binding in an arrangement highly similar to the previous Abf2p/22 complex (Figure 1B and D). To explain this, we performed a thorough analysis of both DNA sequences, and noticed the presence of an ARS-m-like sequence containing poly-adenine tracts (A-tracts) in both DNAs (chainB (A7)A8A9T10T11 in Af22, and chainC A12A13A14A15 in Af2 shift22, Figure 1D). Noticeably, the A-tracts are not contacted by the protein in either crystal. Moreover, crystallization trials with additional DNA fragments devoid of A-tracts (see ‘Materials and Methods’ section), yielded poorly diffracting crystals, suggesting high disorder. Thus, the A-tracts are not contacted by the protein in either crystal. To discern the role of the different Abf2p regions (N-flag, N-helix, HMG-box 1 and HMG-box 2) in binding DNA we produced deletion mutants and analyzed their assembly on linearized M13mp18 dsDNA plasmids (5435bp) by electrophoretic mobility shift assays (EMSA). We constructed mutants Mut1 (no N-flag), Mut2 (HMG-box 1+HMG-box 2), Mut3 (N-helix+HMG-box 1), Mut4 (HMG-box 1), Mut5 (HMG-box 2) and Mut6 (N-flag+N-helix+HMG-box 1) (Figure 3A). Circular-dichroism experiments confirmed that all mutants were properly folded (Supplementary Figure S3A). The EMSA analyses (Figure 3B) showed that, under our experimental conditions, increasing concentrations of full length protein, in presence of a constant concentration of DNA, progressively up-shifted the band with a clear shift at a ratio of 1Abf2p:10 bp. Removal of the N-flag in Mut1 did not modify DNA binding, although DNA migration showed some smearing effect, suggesting some destabilization of the nucleoprotein complex. A similar effect was observed when the entire N-terminal segment was removed (N-flag+N-helix in Mut2). In contrast, neither HMG-box 1 (Mut4) nor HMG-box 2 (Mut5) showed any appreciable binding in this assay. However, HMG-box 1, in presence of N-flag+N-helix (Mut6), surprisingly showed a DNA binding efficiency similar to the full-length protein. Dissecting the latter, Mut3 (N-helix+HMG-box 1) behaved identically to Mut6, apparently suggesting that the N-flag does not have an additional, predominant functional role in DNA binding. Thus, the N-helix and HMG-box 2 individually endow DNA binding ability to HMG-box 1 in vitro. On the other hand, Abf2p levels have been additionally shown to be positively correlated to the level of recombination intermediates in mtDNA (14,47) and Abf2p has...
been shown to possess high affinity for four-way junctions in vitro (50). Thus, we additionally performed EMSA assays with four-way junction templates (Figure 3C). We observed that full length Abf2p binds a four-way junction efficiently (Figure 3C). All of the DNA was bound at a protein:DNA ratio of 4:1, which possibly involves oligomerization of the protein on DNA. However, binding efficiency deteriorates slightly for Mut1 and significantly for Mut2, indicating that the absence of the N-flag or the entire N-module affects binding. The HMG-boxes alone i.e. Mut4 and Mut5 are inefficient binders. In contrast, HMG-box 1 bound to the N-helix (Mut3) slightly improves binding while presence of N-flag+N-helix (Mut6) restores binding markedly. Summarizing, while all mutants except the HMG boxes alone have a similar behavior in binding dsDNA, they present differences when binding to four-way junctions, suggesting different binding modes.

In order to observe effects of the deletions in vivo, we subsequently generated haploid yeast Abf2Δ cells, bearing plasmids encoding the full length or mutant Abf2 constructs described above. Cells expressing full length, wild-type Abf2p grew normally on non-fermentable carbon source (e.g. glycerol), indicating a functional respiratory chain (Figure 3D). Strikingly, cells with Abf2p lacking the N-flag+N-helix (Mut2) were able to ferment glucose but failed to grow on glycerol. This indicates that the two HMG-box domains, coordinated by the Leu-Pro linker enable Mut2 to bind DNA (see above) but this is not sufficient for y-mtDNA stability and maintenance, which underlines the functional importance of the N-terminal module. In contrast, and as previously shown (51), cells carrying the N-flag+N-helix+HMG-box 1 construct (Mut6) grew normally on glycerol indicating that this Abf2p mutant is sufficient for maintaining y-mtDNA. Removal of the N-flag (Mut3) from Mut6 though results in loss of viability on glycerol. In light of the crystal structure, this is attributable to a role of the N-flag in establishing additional contacts with the DNA. Previous work with an Abf2p construct spanning amino acids 54–183, thus lacking the N-flag+N-helix module and an N-terminal segment of HMG-box 1, did not shift DNA in vitro but did impair cell growth in YPG media containing traces of galactose (51). The different length of this and our constructs together with the different experimental conditions tested (use of an inducible (51) or endogenous (this work) promoter; different strain background and temperature and media of growth) may explain the different results. To get further insight into the effect of our mutants on mtDNA, we visualized mitochondrial nucleoids by fluorescence confocal microscopy (Figure 3E, Supplementary Figure S3B). This revealed that Abf2Δ cells expressing plasmid-borne full length Abf2p or Mut6 have an average count of cytoplasmic y-mtDNA foci (15.5 and 10.9 respectively, see ‘Materials and Methods’ section) comparable to wild-type haploids (C(+), 22.2 foci). In contrast, Abf2Δ cells expressing Mut2, Mut3 or Mut5 lose almost all of their mtDNA (average foci count of 0.42, 0.48 and 0.9 respectively), similar to Abf2Δ cells (C(-), average count of 0.4). Thus, combining our in vitro and in vivo results, we conclude that an intact N-flag+N-helix module is essential for mtDNA maintenance.

Abf2p dynamics provides clues to DNA binding mechanism

Our EMSA and in vivo assays with different deletion mutants pointed to the relevance of the N-flag and helix in y-mtDNA binding and metabolism. On the other hand, the crystal structure shows a detailed conformation of the final DNA bound state but does not provide mechanistic information about the involved binding process. To obtain this information, we performed MD simulations on the isolated protein and the complex in aqueous solution. A 500 ns MD simulation of the free protein shows that, in absence of bound DNA, Abf2p adopts a rather extended conformation where the two HMG-boxes separate from each other and the contacts between the N- and C-termini are lost (Figure 4B). This movement is allowed by the flexibility of the short two-residues Linker (114-Leu-Pro-115) and can be aided by occasional unwinding of the C-terminal end of HMG-box 1 helix3 (specifically Lys-113), which undergoes helix-turn-coil transitions (Supplementary Figure S4A). In addition, the N-helix shows a great variability in its orientation (Supplementary Figure S4B) and turns along its main axis resulting in Leu34, Leu38 and Ile39 side chains being exposed to the solvent (Figure 4B) and a consequent disruption of the N-hydrophobic core. The calculated root mean square fluctuation (r.m.s.f) along the simulation demonstrates that the N-helix is highly flexible (Figure 4C). In contrast, a 500ns all-atom MD simulation of Abf2p/DNA complex (1protein:2DNA) shows that the protein remains tightly bound to the DNA (Figure 4C and Supplementary Figure S5A), with overall lesser fluctuation. In this bound state, the stacking between the DNA ends is preserved (Supplementary Figure S5A) and the Phe51 and Ile124 insertions are maintained. The N-flag, N-helix and the N-hydrophobic core are also stably maintained, as in the crystal structures (Figure 4C). Therefore, the orientation of the N-flag and helix are fixed during complex formation with simultaneous consolidation of the N-hydrophobic core.

Abf2p compacts upon DNA binding in solution

The inter-domain flexibility observed from MD simulations was analyzed experimentally by SAXS, a technique that allows extraction of information about the structure and dynamics of macromolecules in solution (52). The data indicated that Abf2p is a monomer in solution (estimated 16.3–21.8 kDa, which match closely with a Abf2p his-tagged monomer of 19.5 kDa, see ‘Materials and Methods’ section). The calculated radius of gyration (Rg = 27.8 Å) is larger than that expected for a ~20 kDa protein (17Å according to Flory equation Rg = 3 x Ne0.33 (53)) and agrees well with that calculated from MD simulation of the free protein (Rg = 29.9Å).These results indicate an extended state of the protein. Accordingly, the pair-wise distribution function (P(r)), which reflects the distribution of intraparticle distances (52), yielded a large maximal particle dimension Dmax of 130Å (Supplementary Figure S6A). Additional information regarding the conformational variability of macromolecules can be obtained from the Kratky representation (52). In this case, the Kratky plot showed a flat profile that is typical of a flexible particle (Supplementary Figure S6B). To further describe Abf2p flexibility...
and the expanse of its conformational variability, we applied the ensemble optimization method (33,34) (see ‘Materials and Methods’ section) which, from a large pool of static structures (thousands, generated from an initial starting model), selects a sub-ensemble of conformations that best describes the SAXS data. In the present case, the sub-ensemble selected from this calculation describes the experimental data with significant accuracy ($\chi^2 = 0.957$, Figure 5A), and shows an important variability in the relative orientations of the HMG-boxes (Figure 5B). The $R_g$ estimates for the sub-ensemble (Figure 5C) show a bimodal distribution suggesting two alternative predominant conformations of the free protein in solution, one being more compact than the other. The Abf2p/Af2_22 complex, in contrast to the free protein, showed characteristics of a more compact species, as reflected in the respective radii of gyration and maximal particle dimensions ($R_g = 13.8\text{Å}$ versus $27.8\text{Å}$; $D_{\text{max}} = 65\text{Å}$ versus $130\text{Å}$, Supplementary Figure S6A). This is supported by the corresponding Kratky plot (Supplementary Figure S6B), which shows a bell-shaped profile, characteristic of a compact particle (52). Therefore, free Abf2p is highly flexible and extended in solution while DNA binding reduces its conformational freedom leading to a more compact overall arrangement.

Poly-adenine tracts direct Abf2p binding on DNA

In order to unveil the features of the DNA molecules responsible for the dual-binding mode observed in the crystal structures (one protein binding to two DNA molecules), we analyzed the structural properties of the Af2_22 and Af2_shift22 DNAs by all-atoms MD simulations, starting from ideal B-DNA models. Averages of the intra- and interbase pair parameters and groove parameters from the trajectories revealed that the A-tracts showed conformational features distinct from standard B-DNA (40,54). In general terms, there are two types of A-tracts. Asymmetric A-tracts are a contiguous stretch of adenine bases ($A_n$ where $n \geq 4$), which show a progressive narrowing of the DNA minor groove going from the $5'$ to the $3'$ end, while symmetric A-tracts are of the form $A_{n_1}T_{n_2}$ (where $n_2 \geq 2$) and exhibit narrowest minor groove at the central AT junction (54). Both A-tract types are represented in our crystal structures. The Af2_22 sequence has the symmetric tract ($A_9T_9$) and Af2_shift22 DNA possesses the asymmetric tract $A_{12}A_{13}A_{14}A_{15}$ (chainC). The MD simulations showed that the symmetric A-tract at Af2_shift22 narrows from both ends to a minimum width of $3.75\text{Å}$ (compared to the $5.91\text{Å}$ in an ideal B-form), precisely at the $A_9T_{10}$ junction (Figure 2B). For $A_{12}A_{13}A_{14}A_{15}$, the minor groove width shows a prominent decrease from $5'$ to $3'$, reaching a minimum of $4.0\text{Å}$. Notably, a MD simulation of the Abf2p/22 complex (1protein:2DNA) showed that the A-tract maintains the structural characteristics of the unbound form (Supplementary Figure S5B) and is not affected by the DNA distortions due to protein binding at adjacent non-A-tract regions. Furthermore, the crystallographic structures show remarkably narrow minor grooves precisely at the A-tracts ($3.08\text{Å}$ and $2.8\text{Å}$ for Abf2p/22 and /shift22, respectively; Figure 2B). Thus, it can be deduced that Abf2p, a protein that binds to and opens up DNA minor grooves, finds A-tracts inhospitable. Values for the DNA stiffness descriptor (40) ($K_{\text{tot}}$, see ‘Materials and Methods’ section, Supplementary Figure S5C), revealed that the above A-tract regions possess high intrinsic rigidity (Supplementary Figure S5C and ‘Materials and Methods’ section) that make them difficult to be distorted. All these observations suggest that the unusually narrow minor groove of A-tracts together with their inherent rigidity impede Abf2p binding, diverting the protein towards more permissive DNA sequences. This presumably causes the unexpected dual-binding observed in both the crystals. Specifically, in the crystal structures, HMG-box 2 binds inside the ARS-m-like sequence, but avoids the A-tracts (Figure 1B and D), further indicating that the A-tracts are the key elements responsible for positioning Abf2p on DNA.

In order to explore how the DNA conformational properties affect Abf2p binding, we used our Elastic Deformation Energy model (40,41,55) to calculate the energy required to contour segments of the natural y-mtDNA 64bp sequence into the conformations induced by HMG-box 1 or HMG-box 2 (see ‘Materials and Methods’ section). Based on the rationale that binding propensity will be higher for those segments that require less energy to be distorted, we derived a sequence-dependent probability of binding (see ‘Materials and Methods’ section, Figure 6). The results show exclusion of the HMG-boxes from the A-tracts (Figure 6). Additionally, HMG-box 1 binding probability (Figure 6A) shows a more restrained pattern compared to HMG-box 2 (Figure 6B), indicating that the former has more stringent sequence requirements and a higher tendency to avoid A-tracts. The binding probability of full-length Abf2p can be expected to be a combination of that observed for the individual boxes. In summary, the narrow minor groove and rigidity of the A-tracts make them unsuitable as binding sites for Abf2p, while the adjacent sequence stretches are well suited for insertion and are favored for positioning the HMG boxes. This points to a mechanism of protein positioning mediated by DNA structure, which somehow resembles that suggested for nucleosome placements in nuclear chromatin (56–59). Interestingly, we calculated that $33.3\%$ of the y-mtDNA is composed of A-tracts (3) (symmetric and asymmetric; Supplementary Table S2 and Supplementary ‘A-tracts in y-mtDNA’), consistent with previous estimates (60). According to the above results, Abf2p DNA binding would follow a similar trend at other y-mtDNA regions possessing A-tracts.

The thermodynamics of Abf2p DNA binding is altered by A-tracts

Analysis of the thermodynamics of Abf2p DNA recognition by ITC showed that the presence of A-tracts modifies the mode of binding. Titration of Abf2p with Af2_22 DNA (protein in cell, DNA in syringe), produced a thermogram with an exothermic phase until a DNA:protein molar ratio of 1:1 was reached. Notably, a second endothermic phase followed (Figure 7A, top left). The isotherm was fitted to a model for two independent sets of sites (Figure 7A, bottom left), which would correspond to two binding sites on the protein. For site1, a stoichiometry of 0.86 was obtained corresponding to a $\sim 1:1$ DNA:site1 ratio. For
site2, a binding stoichiometry of 0.49 was obtained, corresponding to a 1:2 (DNA:site2) ratio. In addition, the derived affinity for site1 ($K_d = 52.1 \text{ nM}$) was higher than for site2 ($K_d = 166.4 \text{ nM}$). Our EMSA assays demonstrate that the N-flag+N-helix+HMG-box 1 construct has a much higher DNA binding efficiency than HMG-box 2. Additionally, the crystal structures and the predicted probability of binding to the 64bp sequence indicate that the N-flag+N-helix+HMG-box 1 module is constrained to be positioned outside the A-tracts and has a single preferred binding site on the $Af2_{22}$ sequence (in grey in Figure 7B, top, Figure 6). In contrast, HMG-box 2 possesses two putative binding sites on $Af2_{22}$. Therefore, we rationalized that the first exothermic phase is mostly attributable to the binding of N-helix+HMG-box 1, until a 1:1 (DNA:site1) molar ratio is reached. After the N-flag+N-helix+HMG-box 1 module is saturated, the HMG-box 2 domain has access to two sites on each new injected DNA molecule, yielding a 1:2 (DNA:site2) stoichiometry. Thus, the two binding events together lead to an overall molar ratio of 3:2 (DNA:protein).

Additional ITC experiments were performed with a 22 bp DNA sequence that did not contain any A-tract (see ‘Materials and Methods’ section and Figure 7A, right). These showed an exothermic phase until a molar ratio close to 1:1 was reached, followed by a slightly endothermic phase (Figure 7A, top right). Fitting the data to the two-sites model yielded stoichiometry values of 0.55 and 0.50 for sites 1 and 2 respectively (Figure 7A, bottom right). Thus, both sites correspond to a 1:2 (DNA:site) molar ratio. In this case we propose that the absence of A-tracts on GC$_{22}$ allows the N-flag+N-helix+HMGbox 1 module (site 1) to bind freely, giving a 1:2 (DNA:site1) complex (Figure 7B, bottom). On near-saturation of the N-flag+N-helix+HMGbox 1 fragment, HMG-box 2 (site 2) binding becomes predominant, giving a 1:2 ratio for site 2. Therefore, for the GC$_{22}$ complex the overall molar ratio (DNA:protein) is 1:1, whereas for the $Af2_{22}$ it is 3:2. Thus, the ITC experiments provide further experimental evidence that the A-tracts dictate the mode of DNA binding by Abf2p.

**DISCUSSION**

Abf2p functions like a staple on DNA with each of the two HMG-boxes bending the DNA by 90°, thus generating a U turn. In this arrangement, the N- and C-terminus of Abf2p are close to each other. In the free protein the two ends are far apart, suggesting that the protein conformation observed in the crystal is induced by DNA binding. We also show that the HMG-boxes alone have very low DNA binding efficiency, and the presence of the N-flag+N-helix module is crucial for mtDNA maintenance, as in vivo and microscopy results clearly demonstrate. Our findings suggest a sequential DNA binding model where the N-flag+N-helix+HMG-box 1 module initiates DNA binding. In this event, the N-helix functions as a pin-lock that consolidates...
Figure 6. Binding probability of HMG-boxes on the 64 bp sequence. (A) Normalized probability for HMG-box 1 to bind and distort a sequence stretch (see ‘Materials and Methods’ section). Each peak corresponds to the probability of HMG-box 1 binding to the 7 bp stretch starting from the peak position. The said sequence stretches are indicated in boxes above each relevant peak and additionally highlighted in green on the 64 bp sequence. HMG-box 1 is represented by a green L shape (1) and the site of insertion is indicated by an arrow in each case. The location of the AF2_22 sequence within the 64 bp sequence is demarcated. The A-tracts within the 64 bp sequence are shown in red. (B) Normalized probability for HMG-box 2. Features are represented similar to (A). The 4 bp HMG-box 2 binding sites are shown for representative peaks.

Abf2p and its human counterpart TFAM are the best characterized mtDNA packaging proteins. Both show universal features of mtDNA bending and compaction through their HMG-box domains, but present very low sequence similarity. We don’t know if this reflects different roles played by the two proteins in nucleoid organization and segregation or possibly, differences in mtDNA base composition and topology. Indeed, striking differences are found between respective mtDNAs: human mtDNA is a small G-C rich circular molecule of 16.5 kbp, whilst S. cerevisiae mtDNA is an A-T rich linear molecule of ~80 kbp. Both proteins generate a U-turn on the DNA \((61,19)\). However, TFAM, by virtue of its long and flexible 30aa linker, intertwines and bends the DNA, positioning the HMG-boxes at opposite faces on the DNA. In contrast, Abf2p possesses a short linker and thus the two vicinal HMG-boxes interact with the DNA from one side. As a result, the N- and C-termini are distant in space in TFAM, but come close in...
Figure 7. Thermodynamics of interaction of Abf2p with DNA sequences with and without A-tracts (A) Isothermal titration calorimetry thermograms (above) and fitting of the binding isotherms (below) to a model with two independent binding sites for Af22 (left) and GC22 (right) DNA. The corresponding DNA sequences are shown at the top of each thermogram. (B) Schemes of the binding model with two independent binding sites for Af22 (top) and GC22 DNA (bottom). Abf2p is depicted schematically with the HMG-boxes 1 and 2 represented in green (1) and orange (2) respectively. The non-A-tract part of the ARSm-like sequence in Af22 is represented in orange whereas the A-tract is depicted in red. At the very bottom, the 1:1 stoichiometry is represented as a plausible circle in order to show its compatibility with the crystal structure (the two dsDNA molecules in the circle are depicted in different gray tones).

Abf2p. Therefore, the two homologs utilize different binding strategies to achieve a similar DNA bending. This suggests that U-turns constitute the basic structural unit of mitochondrial nucleoid organization in both systems.

Our EMSA assays (Figure 3B) reveal that Abf2p shifts dsDNA at a ratio of 1Abf2p:10-20 bps, which agrees with the number of DNA bps (~19 bps) contacted by the protein in the crystal structures. Additionally, previous atomic force microscopy studies showed that Abf2p induces compaction at a 1Abf2p:20 bps ratio (15), being more prominent at 1:10. At these ratios TFAM showed higher compaction, which correlated with a reduction in in vitro DNA replication and transcription (62). A similar effect was observed for Abf2p, where Abf2p overexpression resulted in loss of y-mtDNA, suggesting excessive DNA compaction and consequent reduced access for the replication machinery (14).

The dual-binding found in the crystal structures and the relative conformational freedom of the two HMG-boxes observed from SAXS-EOM analyses demonstrates the capability of the protein to bind two DNA molecules simultaneously. This ability would be compatible with ‘cross-strand binding’ or DNA looping by Abf2p, as also described for TFAM (63,64). Thus, both DNA bending and cross-strand binding might be underlying the nucleoid packaging mechanism of the two proteins.

We have also demonstrated that the A-tracts direct positioning of Abf2p on DNA. Since Abf2p has been shown to bind GC rich sequences with relatively higher preference (11) additional crystallization trials were made with DNA fragments of different lengths that are either GC-rich or devoid of A-tracts. Both trials however yielded poorly diffracting crystals, suggesting high disorder. Presumably, by specifying positioning of the protein, the A-tracts induced a complex that formed the ordered crystals reported here. Accordingly, previous competition experiments showed that binding of Abf2p to an ARS1 probe could not be competed by a poly(dA)-poly(dT) double-stranded homo-polymer (10,11), indicating the incapability of this protein to bind to poly-adenine DNA. A-tract guided nucleoprotein interactions have also been proposed in the nucleus, where biochemical studies have shown that A-tracts help in positioning nucleosomes (56–59). This has been further corroborated by computational approaches (41,55). Additionally, other biochemical experiments have demonstrated that the A-tracts are involved in DNA synthesis termination in the human immunodeficiency virus (65,66) and in regulation of transcription in pathogenic bacteria (67–69). Thus, A-tract mediated control of protein/nucleic acid interactions appear to be a general strategy across species. We and others detected that y-mtDNA has a high content of A-tracts (3,60). Thus, location of these tracts might determine strategic positioning of Abf2p, suggesting a DNA-mediated control of nucleoid architecture and of mtDNA accessibility.

Y-mtDNA undergoes frequent recombination ((7) and references therein). Several studies support that recombination is involved in y-mtDNA replication and maintenance, and a mechanism of recombination-dependent replication has been proposed (70,71). In mitochondria, Abf2p levels correlate with the level of recombination intermediates (14,47) and in vitro studies showed that Abf2p and its orthologs from different yeast species show preference for
such DNA structures (50). This suggested that Abf2p could be involved in maintenance of y-mtDNA copy number via interaction with recombination intermediates (14,47). Our studies confirm the binding of Abf2p to four-way junctions. Our mutants additionally show that such binding involves cooperativity between different protein regions and indicate an important role of the N-flag+N-helix module. Mut2 and 3 bind weakly to four-way junction and cannot support growth on glycerol plates, whereas growth is observed for Mut6 which shows better binding. This is consistent with the hypothesis that Abf2p influences the levels of recombination intermediates in vivo, thus putatively playing a role in recombination-dependent replication and DNA maintenance. The preference of Abf2p for four-way junctions over dsDNA (50) suggests a second level of regulation in which binding to replication intermediates is favored over DNA compaction that, in turn, is regulated by A-tracts. Binding to four-way junctions, in conjunction with DNA packaging, might provide a protective effect against mutagens through involvement in DNA repair. We can speculate that such a mechanism might be responsible for similar effects observed for HMG-box proteins in general (72,73). In connection to this, a novel and unexpected feature observed in the crystals is DNA end-joining by Abf2p. This could be relevant for cellular processes where DNA ends are available, such as in DNA breaks or recombination. However, any potential biological implication of Abf2p DNA end-joining in recombination or DNA repair remains to be analyzed by further studies.

To conclude, the binding of Abf2p to DNA is modulated by the positioning of poly-adenine tracts, posing a molecular strategy for y-mtDNA regulation and nucleoid organization. Our results demonstrate how the intrinsic structural properties of DNA can play a key role in orchestrating localization of non-specific DNA binding proteins that are essential for mitochondrial maintenance, and thus for cell life.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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