

Analysis of the XPA and ssDNA-binding surfaces on the central domain of human ERCC1 reveals evidence for subfunctionalization

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ABSTRACT

Human ERCC1/XPF is a structure-specific endonuclease involved in multiple DNA repair pathways. We present the solution structure of the non-catalytic ERCC1 central domain. Although this domain shows structural homology with the catalytically active XPF nuclease domain, functional investigation reveals a completely distinct function for the ERCC1 central domain by performing interactions with both XPA and single-stranded DNA. These interactions are non-competitive and can occur simultaneously through distinct interaction surfaces. Interestingly, the XPA binding by ERCC1 and the catalytic function of XPF are dependent on a structurally homologous region of the two proteins. Although these regions are strictly conserved in each protein family, amino acid composition and surface characteristics are distinct. We discuss the possibility that after XPF gene duplication, the redundant ERCC1 central domain acquired novel functions, thereby increasing the fidelity of eukaryotic DNA repair.

INTRODUCTION

During DNA replication, recombination and repair, double-stranded DNA inevitably forms three- or four-way junctions, bubbles, flaps or broken ends with single-stranded extensions. These irregular structures must be processed correctly in order to successfully complete DNA metabolism and thereby maintain genome integrity. This task is accomplished by structure-specific endonucleases specialized in pruning downstream of branch, flap or bubble structures by incision at junctions between double- and single-stranded DNA (1). Inactivation or malfunctioning of these enzymes causes genetic defects or cancer, underlying their importance in genome stability.

A remarkable class of structure-specific endonucleases in humans is XPF. The protein family is characterized by the presence of the ERCC4 domain and consists of seven members (XPF, MUS81, ERCC1, EME1, EME2, FANCM and FAAP24) (2). Only XPF and MUS81 have nuclease activity, which is mediated by the conserved core nuclease motif (ERKX₃D) (3–5). Their catalytic function depends on heterodimer formation with the non-catalytic family members. XPF forms an obligate complex with ERCC1 and functions primarily in nucleotide excision repair (NER), a versatile pathway able to detect and remove a variety of DNA lesions induced by UV light and environmental carcinogens. The ERCC1/XPF heterodimer has additional roles in DNA interstrand cross-link (ICL) repair (6) and telomere maintenance (7). The symptoms of the first patient with inherited ERCC1 deficiency (8) and of a patient with a novel XPF mutation (9) are distinct from the classical NER phenotype, and underscore the pleiotropic function of ERCC1/XPF.

In contrast to eukaryotes, archaea have a single homolog of the XPF endonuclease that forms homodimers. The archaeal XPF minimally consists of the catalytic nuclease domain followed by a DNA-binding domain containing two consecutive helix-hairpin-helix motifs (HhH₂ domain). Dimerization occurs between both the nuclease and HhH₂ domains of each subunit (Figure 1A) (3). The strong preference of the archaeal endonucleases for 3' flap DNA substrates is explained by the function of their individual domains. Structural data suggest a model for DNA binding where one HhH₂ domain of the dimer binds to an upstream and the other to a downstream DNA duplex, sharply bending the DNA substrate and thereby allowing one active site of the dimeric nuclease domain to cleave the 3' protruding single strand (10). Combination of mutations in nuclease and HhH₂ domains that do not disrupt the dimeric interfaces but impair the individual functions (DNA cleavage or DNA binding), provide evidence for this model (11).

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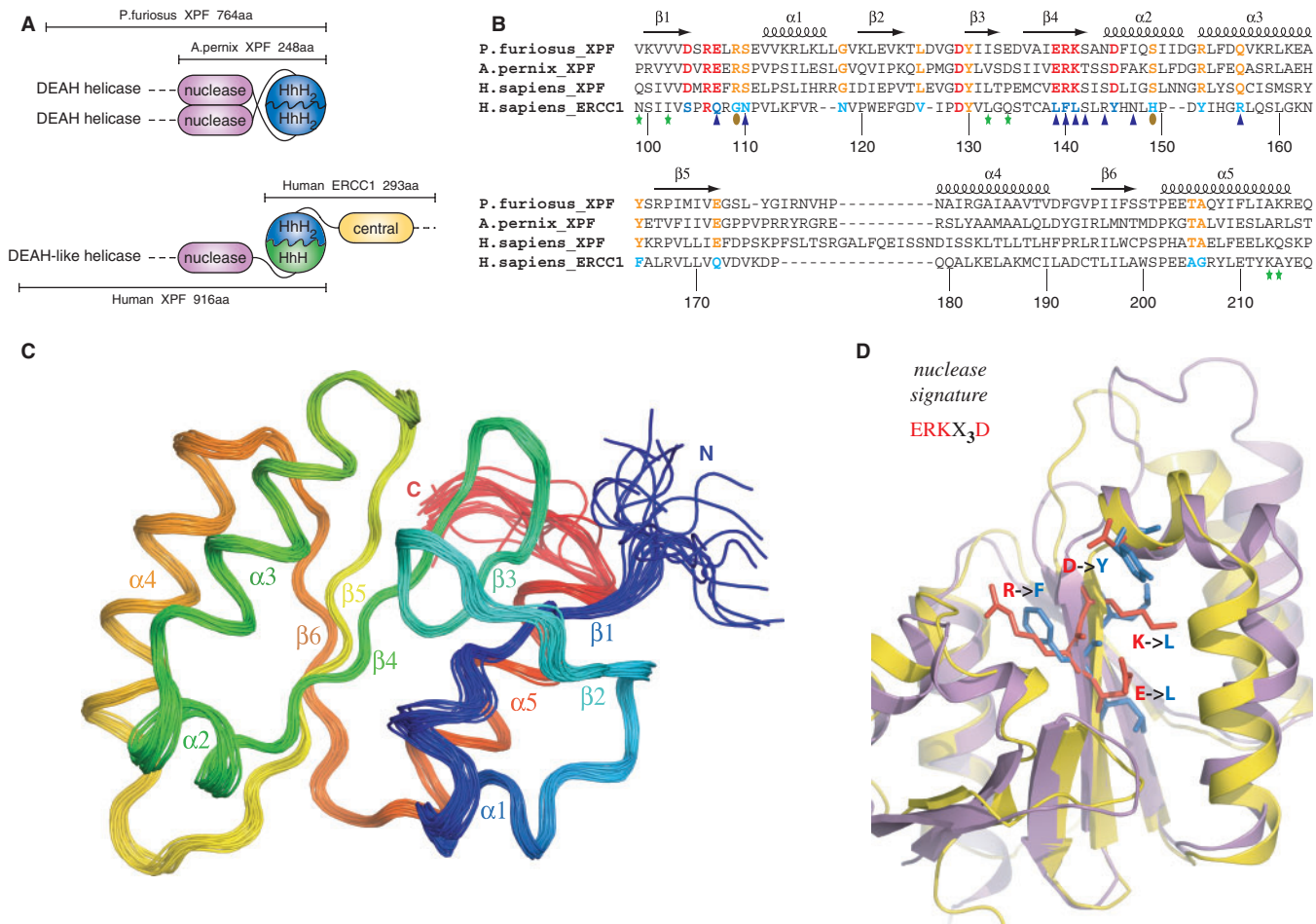


Figure 1. (A) Domain organization of the archaeal XPF homodimeric members and the human ERCC1/XPF heterodimer. (B) Structure-based sequence alignment of the nuclease XPF domains from archaea to human and the corresponding central domain of human ERCC1. Secondary structure elements of the prototype XPF nuclease fold are indicated at the top of the sequences. Catalytic residues in the XPF nucleases are colored red and their corresponding substitutions in ERCC1 blue. Other invariant residues in XPF domains are depicted in orange, while the ERCC1 equivalents are depicted in cyan. Residues of cERCC1 perturbed largely upon XPA titration are indicated by blue triangles and those appear only in the final complex by brown ellipses. Green asterisks indicate cERCC1 residues perturbed by DNA titration. cERCC1 sequence is numbered at the bottom. (C) Ensemble of the final 20 structural conformers of cERCC1 as determined by solution NMR. Secondary structure elements and N- and C-termini are labeled. (D) Superposition of the crystal XPF nuclease structure (2bgw) from *A. pernix* (purple) and the solution NMR structure (2jpd) of human cERCC1 (yellow). Emphasis is given to the nuclease signature and the corresponding substitutions in cERCC1.

The DNA substrate specificity is different for the human ERCC1/XPF heterodimer. The human enzyme is known to specifically incise hairpin, bubble or splayed-arm DNA substrates (12,13), which consist of only one duplex (upstream). Previous data indicated the role of the C-terminal ERCC1 'canonical' HhH₂ domain in engaging the upstream DNA duplex of a hairpin substrate (14). This function requires heterodimerization with the C-terminal HhH domain of the human XPF partner to form a complex analogous to the archaeal HhH₂ dimer interface (12,14). Previous structural data (12) and the data presented here illustrate the structural similarity between the human ERCC1 central domain and the archaeal XPF nuclease domain. Unlike the archaeal nuclease dimer interface, there is no evidence that the corresponding domains of human ERCC1 and XPF interact with each other in solution (Figure 1A) (12).

The catalytic function of the ERCC1/XPF endonuclease is crucial for NER. NER operates through a 'cut and patch' mechanism by excising and removing a short stretch of DNA containing the lesion, and subsequently restoring the genetic information by repair synthesis using the undamaged strand as the template. The incision step involves the sequential and coordinated assembly of the DNA damage sensor XPC-HR23B, the transcription/repair factor TFIIH, the architectural protein XPA, the ubiquitous ssDNA-binding protein RPA, and the two structure-specific endonucleases ERCC1/XPF and XPG, responsible for the incisions 5' and 3' to the damaged site, respectively (15). A main role in the progress of the reaction is attributed to XPA, which in complex with RPA probes the DNA helix conformations (16) and participates in multiple interactions with the other NER factors (15). XPA is required for the recruitment of ERCC1/XPF in the

NER pre-incision complex (17,18). Consistent with this regulatory function, *in vitro* studies with recombinant proteins and *in vivo* studies using a yeast two-hybrid system have demonstrated interactions between XPA and ERCC1 (19).

Here, we report the solution structure of the ERCC1 central domain (cERCC1) and investigate its interactions with XPA and DNA at the molecular level. Using biochemical studies and NMR titration experiments, we have identified two distinct interaction surfaces of cERCC1 that mediate XPA and DNA binding. Interestingly, the two interactions can take place simultaneously to yield a cERCC1/DNA/XPA ternary complex, which in turn explains the important role of ERCC1 in targeting its catalytic XPF partner to the NER pre-incision complex.

MATERIALS AND METHODS

Cloning, protein expression and protein purification

ERCC1 central domain (cERCC1, residues 96 to 219) was PCR-amplified from a vector containing the full-length ERCC1 gene and subcloned into a pET28b (Novagen) expression vector between BamHI and XhoI sites. Downstream of the XhoI site a linker together with a his-tag has been engineered. XPA constructs were derived from a full-length XPA vector purchased from RZPD (clone ID IRAUp969B1273D6) and subcloned into a pLICHIS vector using the enzyme free cloning (EFC) strategy (20). The same procedure was followed for the GST fusion proteins, either XPA or cERCC1 (pLICHISGST vector). As a result all the EFC constructs contain an N-terminal his-tag, while the cERCC1 construct bears a C-terminal his-tag. The nucleotide sequences of the cloned DNAs were confirmed by sequencing. The cERCC1-HIS, HIS-XPA, HIS-GST-XPA and HIS-GST-cERCC1 proteins were expressed in BL21(DE3) Rosetta cells (Novagen) and were subject to two-step purification as has been described (21). When appropriate, HIS-GST-cERCC1 was cleaved with thrombin in the presence of 1 mM Ca^{2+} , 50 mM Tris (pH 8.0) and 100 mM NaCl in order to obtain the untagged protein. Finally, the protein samples were exchanged either to 50 mM sodium phosphate (pH 5.5), 100 mM NaCl for the cERCC1 structure determination, or to 50 mM Tris (pH 8.0), 100 mM NaCl for the binding studies.

NMR spectroscopy

Multidimensional NMR experiments were carried out at 290 K on Bruker AVANCE 600 and 900 NMR spectrometers equipped with TXI triple-resonance probes in 50 mM sodium phosphate (pH 5.5), 100 mM NaCl by using the cERCC1-HIS protein. Spectra were processed using the NMRPipe software package (22) and analyzed with Sparky (23). The ^1H , ^{15}N and ^{13}C resonance assignments are 97% complete and were made using standard triple resonance techniques, 3D NOESY- $(^{15}\text{N}, ^1\text{H})$ -HSQC spectra, and 3D NOESY- $(^{13}\text{C}, ^1\text{H})$ -HSQC spectra (both with a mixing time of 60 ms) (24).

The chemical shifts have been deposited at the BMRB with accession number 15240.

Structure calculations

Automatic NOE assignment and structure calculations were performed using the program CYANA version 2.0 (25). Hydrogen bond restraints were defined when they were consistent with the secondary chemical shift data and expected NOE contacts and only for the helical parts of the protein. The set of 2669 NOE restraints determined by CYANA, together with restraints for 31 hydrogen bonds and ϕ and ψ torsion angle restraints (174) derived from TALOS (26) were used in a water refinement run according to the standard RECOORD protocol (27) utilizing CNS (28). Molecular images were generated with PyMol (29). Coordinates have been deposited at the PDB with accession code 2jpd and the structural restraints at the BMRB with entry number 15240.

EMSAs

All the DNA substrates were 5' ^{32}P -labeled with T4 polynucleotide kinase and purified on a polyacrylamide gel under native conditions. The nucleotide sequences for ss20, b10 and ds30 have been described before (30). Per reaction, 100 fmol of substrate was incubated with the appropriate amount of ERCC1 protein in binding buffer (50 mM Tris, pH 8.0, 100 mM NaCl, 10% glycerol and 1 mg/ml BSA). After incubation for 30 min at 4°C, samples were loaded onto 7.5% native polyacrylamide gels containing 0.5× TBE and run at 4°C. Alternatively Electrophoretic mobility shift assay (EMSA) reactions were loaded on a 3.5% agarose gel containing 0.5% TBE, run at 4 or 20°C yielding essentially identical results. Gels were visualized and quantified using a phosphor imager (BioRad) as described before (30). For depletion experiments, 20 μl of MagneHis beads (Promega) or GST agarose beads (Sigma), were extensively washed with binding buffer, and after buffer removal the EMSA reaction was added to the beads. After 15 min incubation with regular mixing, the beads were removed from the EMSA reaction using a magnet or centrifugation and the reaction mixture was loaded on acrylamide gel or, when appropriate, the substrate was added to the depleted reaction mixture.

GST pull-down assay

GST pull-down assay was performed and quantified as described before (31) using the indicated cERCC1 concentrations and 3–5 μg of GST proteins in 150 μl of 50 mM Tris, 100 mM NaCl, 10% glycerol, 1 mM DTT, 0.2 mM PMSF and 20 mg/ml of BSA (pH 8.0). The semi-quantitative experiments were performed in 50, 150, 500, 1500 and 5000 μl buffer with a constant amount of cERCC1 and GST-XPA.

NMR titrations

cERCC1 titrations with XPA and DNA were performed on a Bruker AVANCE 700 NMR spectrometer and were monitored with 2D ^1H - ^{15}N HSQC experiments. In all

Table 1. Structural statistics of the structure ensemble of human cERCC1 (residues 96–219)

Rmsd (Å) with respect to mean ^a	
Backbone/Heavy atoms	0.40 ± 0.04/0.89 ± 0.08
Number of experimental restraints	
Intraresidue NOEs	632
Sequential NOEs ($ i-j = 1$)	678
Medium range NOEs ($1 < i-j < 4$)	479
Long-range NOEs ($ i-j > 4$)	880
Total NOEs	2669
Dihedral angle restrains	174
Hydrogen bonds	31
Restraint violations	
NOE distances with violations >0.3 Å	0.00 ± 0.00
Dihedrals with violations >3°	0.00 ± 0.00
Rmsd for experimental restraints	
All distance restraints (2669) (Å)	0.013 ± 0.005
Torsion angles (174) (°)	0.579 ± 0.077
CNS energies after water refinement	
E_{vdw} (kcal/mol)	-614 ± 17
E_{elec} (kcal/mol)	-4914 ± 61
Rmsd from idealized covalent geometry	
Bonds (Å)	0.01 ± 0.00
Angles (°)	1.15 ± 0.03
Impropers (°)	1.34 ± 0.06
Ramachandran analysis	
Residues in the favored regions (%)	90.68 ± 2.04
Residues in additional allowed regions (%)	8.16 ± 1.71
Residues in generously allowed regions (%)	0.81 ± 0.65
Residues in disallowed regions (%)	0.36 ± 0.68

^aResidues 101–215 of cERCC1.

cases, the concentration of cERCC1 was 0.2 mM in 50 mM Tris (pH 8.0), 100 mM NaCl. XPA and DNA were dissolved in the same buffer and therefore salt and pH did not vary throughout the experiments. DNA binding was feasible only with the cERCC1 construct lacking the C-terminal his-tag, while XPA binding was identical, regardless of the his-tag presence. Normalized chemical shift changes were calculated by using the equation: $\delta = ([\delta_{\text{HN}}]^2 + [\delta_{\text{N}}/6]^2)^{0.5}$.

RESULTS AND DISCUSSION

Structure analysis

We have determined the solution structure of the ERCC1 central domain (cERCC1) to elucidate the molecular details of its proposed functions. We have collected a large number of distance and dihedral angle restraints that yielded an ensemble of 20 conformers with very good convergence (Figure 1C). The quality of the structure can be judged by the summary of the structural and restraint statistics given in Table 1. All the secondary structure elements are well defined including the short loops that join them. cERCC1 comprises a compact architecture and folds as a six-stranded β -sheet flanked by five α -helices on both sides. As described before (12), this fold is reminiscent of the type II restriction endonucleases, to which the catalytic domain of XPF also belongs.

Overall, the solution NMR structure of cERCC1 is in very good agreement with the crystal structure of the same domain (2a1i) (rmsd 1.1 Å for 108 C $_{\alpha}$ atoms). The only noticeable difference relates to the last few residues of

helix α_3 and the loop connecting this structure element with β_5 . The presence of a mercury atom (linked to C137) in the crystal structure may account for the substantial side-chain rearrangements within this loop. When compared to the crystal structures of the archaeal nucleases, helices α_1 , α_2 , and strands β_1 , β_2 appear to be shorter in the solution structure of cERCC1. Solution cERCC1 and the crystal structures of archaeal XPF nucleases (2bgw and 1j23) contain the same number of secondary structure elements arranged in a very similar manner (rmsd 1.8 and 2.0 Å for *Aeropyrum pernix* and *Pyrococcus furiosus* nucleases, respectively, for 108 C $_{\alpha}$ atoms), although in both cases the primary sequence homology is very low (Figure 1B and D).

Remarkably, conserved residues of XPF nuclease scattered in the primary sequence superimpose structurally with residues conserved in the ERCC1 sequence family (Figures 1 and 3). Whereas the conservation in XPF is directly related to the catalytic function, ERCC1 has preserved the same fold but lacks the essential residues for catalysis. The fold similarities, coupled with the obligate nature of the heterodimerization (32), are in full agreement with the common origin of the two proteins (33).

Minimal domains for ERCC1–XPA interaction

Truncation studies of ERCC1 have mapped the XPA interaction site to a region between residues 91 and 118 (34). From the cERCC1 structure and its compact fold, we predict that such truncations will have a devastating effect on the structural integrity of the central domain (96–219). For XPA, the ERCC1 interaction region seems to be located in a small stretch containing two highly conserved motifs rich in glycine and glutamic acid residues (72-GGGFILEEEEEE-84, conserved residues are underlined) (19). We have prepared a HIS-GST-XPA construct (59–99) containing the conserved stretch, and examined its ability to interact with cERCC1 in a GST pull-down assay. Indeed, the GST XPA fusion protein was able to specifically bind cERCC1 (Figure 2A), confirming that this domain of ERCC1 is sufficient for interactions with XPA. No binding of cERCC1 to GST bound agarose beads or uncharged agarose beads was observed (Figure 2A, data not shown). This XPA peptide is unstructured (35) and appeared very sensitive to proteolysis both during overexpression in *Escherichia coli* and as pure protein after extensive purification (Figure 2A and B). We next performed a semi-quantitative GST pull-down assay (31) and estimated an apparent K_d of 1 μ M. Since we reached only 20% binding saturation at the two highest protein concentrations (3 and 10 μ M of cERCC1) under these experimental conditions, we could not determine the binding constant more accurately (Figure 2B).

Functional disparity of the XPF nuclease and ERCC1 central domains

The cERCC1–XPA interaction specificity has been explored using NMR titration experiments. Titration with the HIS-XPA peptide causes extensive specific

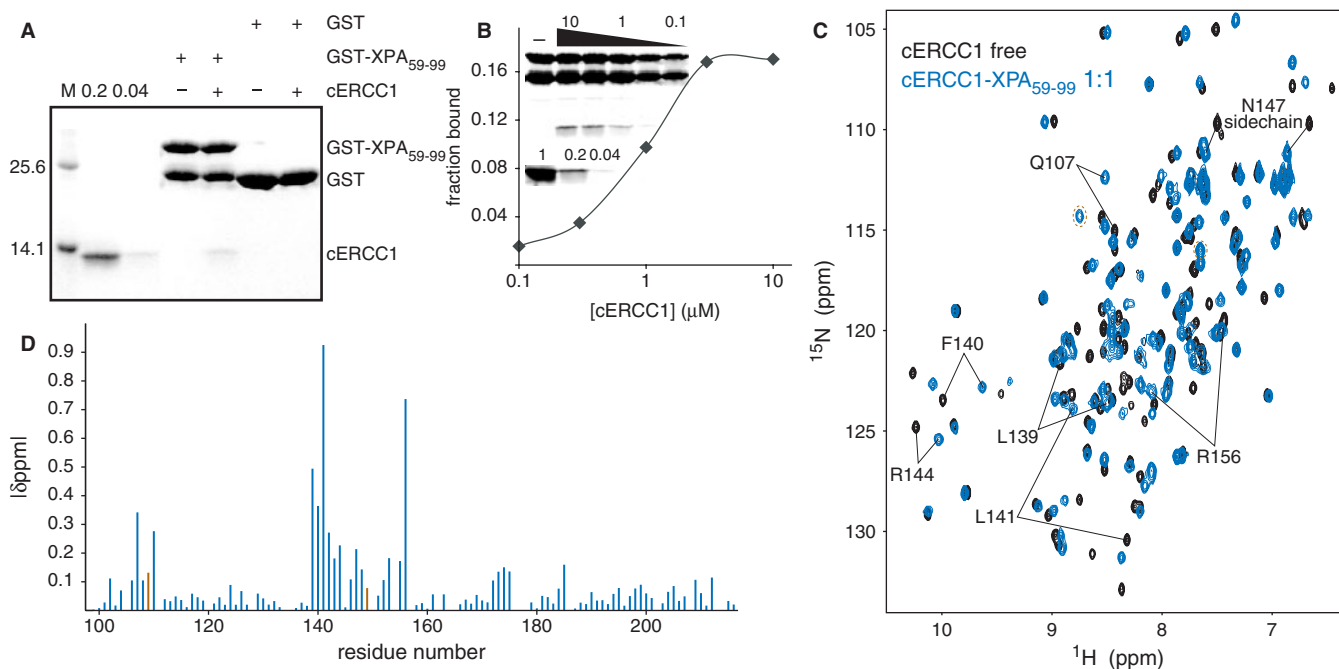


Figure 2. ERCC1–XPA interactions. (A) GST pull-down assay with 3 μg of GST or GST-XPA fusion proteins in the presence or absence of 2 μM cERCC1. Here, 0.2 and 0.04 refer to respectively 20 and 4% of the input cERCC1 protein present in the GST pull-down assay. (B) Semi quantitative GST pull-down assay showing the fraction of cERCC1 bound to 3 μg of GST-XPA at the indicated [cERCC1] (μM). The upper part of inset shows a representative GST pull-down assay, where the various cERCC1 concentrations used are depicted above (10, 3, 1, 0.3 and 0.1 μM). The lower panel shows respectively 100, 20 and 4% of cERCC1 added to the assay. (C) Chemical shift perturbation of the cERCC1 ¹H-¹⁵N HSQC upon complex formation with XPA. Free cERCC1 spectrum is in black and XPA-bound spectrum in blue, while brown circles in the bound spectrum indicate G109 and H149 resonances. (D) Normalized chemical shift changes between free and XPA-bound forms versus the cERCC1 sequence. The p.p.m. difference for G109 and H149 was calculated by their resonances in the free cERCC1 spectrum at pH 5.5, and are colored brown.

changes in the cERCC1 ¹H-¹⁵N HSQC NMR spectra. The titrations show slow exchange regime with respect to the NMR chemical shift timescale, in agreement with the determined affinity (Figure 2C). We have been able to assign all the amide resonances of cERCC1 in the bound form (1:1 complex stoichiometry) by using 3D NOESY-(¹⁵N, ¹H)-HSQC spectra. The pattern of the backbone amide NOEs does not change substantially compared to the free cERCC1 protein. This indicates that the overall structure is maintained, and led us to conclude that the observed chemical shift perturbations arise from specific contacts with the XPA peptide.

The most significant perturbations map to the surface of the V-shaped groove of the cERCC1 structure and involve many residues that compose the corresponding DNA cleavage site of the archaeal XPF endonucleases (Figures 2D and 3). Interestingly, among the largest chemical shifts of cERCC1 upon XPA binding are those of L139, F140 and L141. These ERCC1 residues correspond to the characteristic strongly conserved ERK catalytic triad of the XPF nuclease domain (Figure 1). The rest of the significantly perturbed residues are polar or positively charged (Q107, N110, S142, R144, N147 and R156), are competent for hydrogen bonding with the XPA peptide and are more conserved than the surrounding regions of the cERCC1 V-shaped groove (Figure 3). Additionally, of special interest are the amide signals of G109 and H149, which are not observed in the free protein spectrum due to fast exchange with the solvent at pH 8

(these amide resonances are observable at pH 5.5). However, both give sharp signals at the end of the titration, an indication that the solvent exchange in the free protein is quenched in the complex. These residues reside at the rim of the V-shaped binding site and may be involved in hydrogen bonding with the XPA peptide (Figure 3).

Although ERCC1 residues forming the XPA-binding site are different from the acidic and basic residues which form the active site in XPF (5), there are no significant changes in the backbone fold and the spatial side-chain orientations in the structure remain unaffected. However, their different chemical properties result in a distinct charge and hydrophobicity distribution on the structure surface (Figure 3). While XPF displays negative charge at the DNA cleavage site (important for divalent cation coordination), the equivalent XPA-binding site in ERCC1 is neutral or slightly positively charged. Therefore, the ERCC1–XPA interaction is a combination of hydrophobic and electrostatic interactions possibly involving the glutamic acid stretch of XPA (19) and two conserved positively charged residues (R106 and R156) of cERCC1.

DNA binding by the ERCC1 central domain

Given the proposed common origin of ERCC1 and XPF (33), it can be expected that the degenerated catalytic site of ERCC1 could still bind DNA in a similar fashion as the archaeal XPF active site (12). Because our ERCC1–XPA

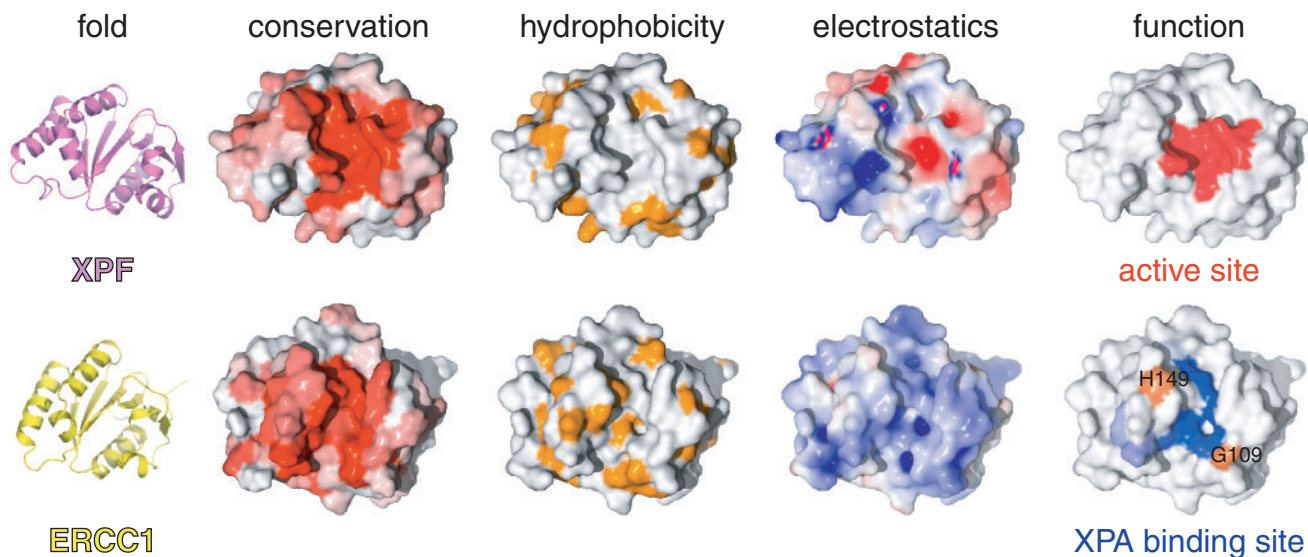


Figure 3. Common fold, different properties and distinct functions for XPF and ERCC1. Sequence conservation either for XPF or ERCC1 proteins is colored from white (non-conserved) to red (highly conserved). The opposite face has no significant conservation for either protein. Hydrophobic side chains are colored orange. Electrostatic surface potentials were calculated using APBS (39) and colored blue for positive or red for negative charge potential. Active site residues of XPF were chosen based on previous mutational studies (5) and the XPA-binding site of ERCC1 was identified by our NMR titrations.

interaction data render this unlikely, we explored whether cERCC1 is able to bind DNA as has been suggested previously (12).

We performed an electrophoretic mobility shift assay (EMSA) where increasing amounts of cERCC1-HIS protein were added to ssDNA. While we saw a loss of free probe, we failed to see any complex formation under these conditions. If the same protein–DNA complex was loaded on a 3.5% agarose gel, we detected a weak smeary complex at the highest protein concentration (Figure 4A). However, by using a HIS-GST-cERCC1 protein, we observed formation of the complex with ssDNA (Figure 4B). This suggests that the C-terminal his-tag interferes with DNA binding. Therefore, we used the HIS-GST-cERCC1 protein, which we cleaved with thrombin to obtain the untagged protein. DNA binding with the untagged cERCC1 demonstrated a faster migrating complex with various ssDNA substrates (data not shown). HIS-GST-cERCC1/DNA complex formation is prevented by incubation of the reaction mixture with MagneHIS magnetic beads or glutathione agarose beads, which depletes HIS-GST-cERCC1 from the reaction mixture either prior to or after the addition of DNA. If beads were added after complex formation, a small but significant decrease in the amount of free probe was observed. Depletion of the HIS-GST tagged cERCC1 protein from the binding reaction confirms that the protein–DNA complex is indeed formed by the HIS-GST-cERCC1 protein (Figure 4C).

In agreement with earlier studies, under these conditions no binding was observed for dsDNA (12), while fairly comparable binding affinities were found for both ssDNA and bubble substrates with 10 or 20 unpaired bases. We calculated an apparent K_d of $2.5 \pm 0.7 \mu\text{M}$ for the HIS-GST-cERCC1 protein bound to a bubble10 (b10)

substrate in the presence of 100 mM NaCl (Figure 4B). Corroborating our results, an equilibrium binding titration experiment using fluorescence anisotropy has demonstrated ssDNA binding for the same domain with an apparent K_d of 10 μM (12). The relatively small difference in affinity can be well explained by different salt conditions or substrate used.

For the NMR titrations we performed a thrombin cleavage on HIS-GST-cERCC1, to remove the his-tag. The untagged cERCC1 ^1H - ^{15}N HSQC spectrum is identical to that of cERCC1-HIS, except for the two amides preceding the artificial his-tag (Supplementary Figure 1). Addition of the b10 substrate to this cERCC1 protein caused a limited number of specific perturbations that were unambiguously assigned. Again, the perturbed resonances exhibit slow exchange behavior in the NMR titrations and indicate contacts with the DNA. The affected resonances include the backbone amides of N99, I102, L132, K213, A214 and the side-chain amide of Q134 (Figure 4D and E). Compared to the free-protein spectrum we only miss T211, which cannot be identified. The perturbation induced by the b10 DNA was complete at equimolar concentration with cERCC1. Chemical shift mapping reveals that all perturbations are in the vicinity of the C-terminus of the cERCC1 construct, consistent with the presence of a flexible tag interfering with DNA binding. The DNA-binding region resides at the surface of the structure where the N- and C-termini meet (Figure 4F). The perturbed residues belong to well-defined structure elements. The only positively charged residue we identified (K213) is fully conserved in the ERCC1 proteins and may explain the dependence of DNA binding on salt concentration. All the other affected residues are less conserved than the residues involved in XPA binding. The composition of the affected residues together with the

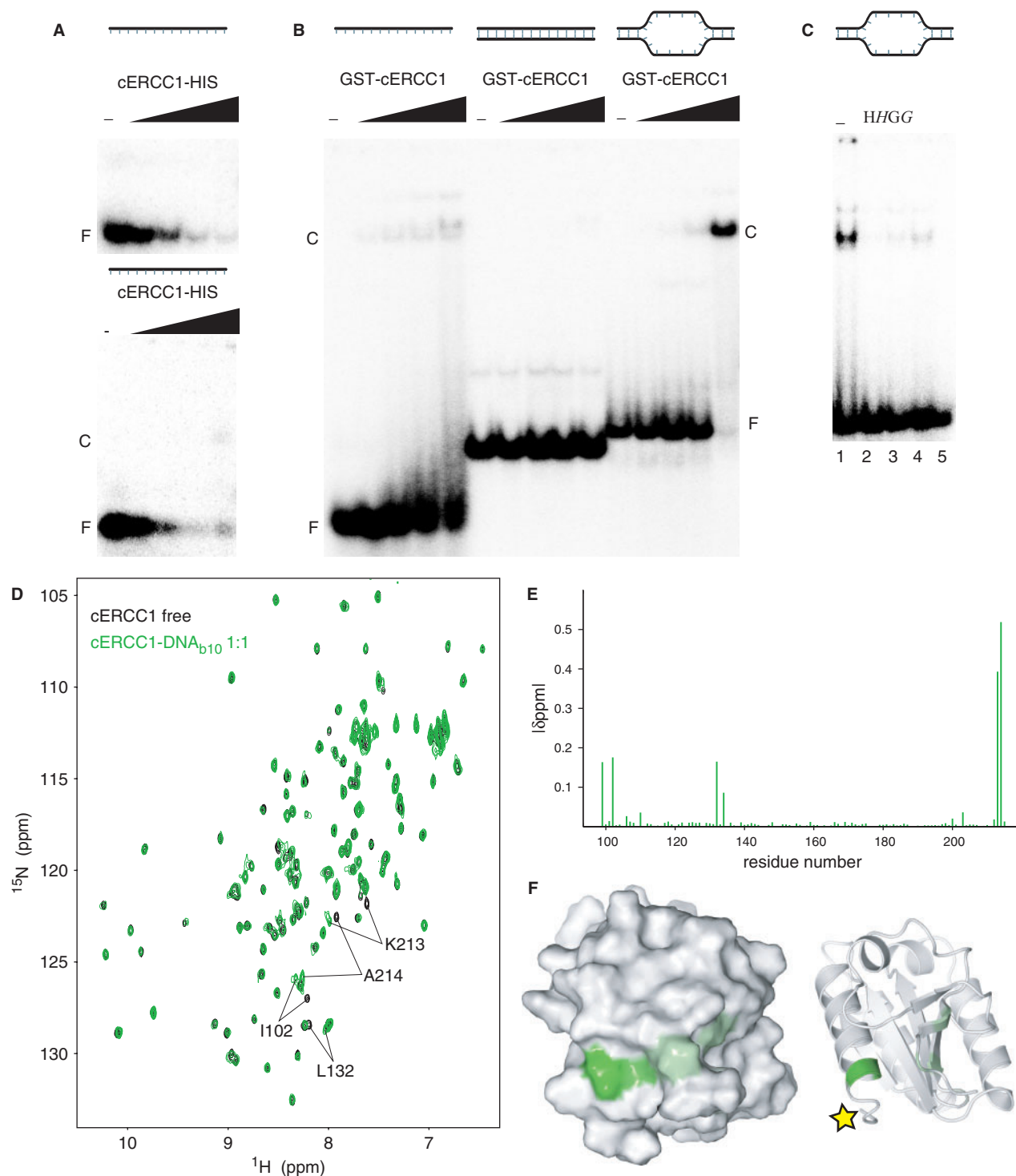


Figure 4. DNA binding by cERCC1. (A) EMSA with a 20-mer ssDNA substrate using increasing concentrations of cERCC1-HIS protein (0, 0.04, 0.2, 1, and 5 μ M), loaded on a 7.5% acrylamide gel (upper panel) or a 3.5% agarose gel (lower panel). (B) EMSA using ssDNA (20-mer), dsDNA (30bp) and bubble10, a dsDNA with 10 unpaired bases as substrate, in the presence of 0, 0.125, 0.25, 0.5 and 1 μ M HIS-GST-cERCC1. (C) Inhibition of cERCC1-b10 DNA complex formation ([cERCC1] = 0.5 μ M) by depletion of HIS-GST-cERCC1 from the EMSA reaction by the addition of MagneHis beads (H) or GST agarose beads (G) prior to or after addition of DNA (italic). F: free DNA, C: cERCC1-DNA complex. (D) Chemical shift perturbation after addition of b10 DNA. The spectrum of the free protein is shown in black and the spectrum after addition of equimolar amount of b10 in green. (E) Normalized chemical shift changes between free and b10-bound forms versus the cERCC1 sequence. (F) Surface and cartoon representations of cERCC1 colored according to normalized chemical shifts. Yellow star indicates the position of the C-terminal his-tag that was interfering with DNA binding.

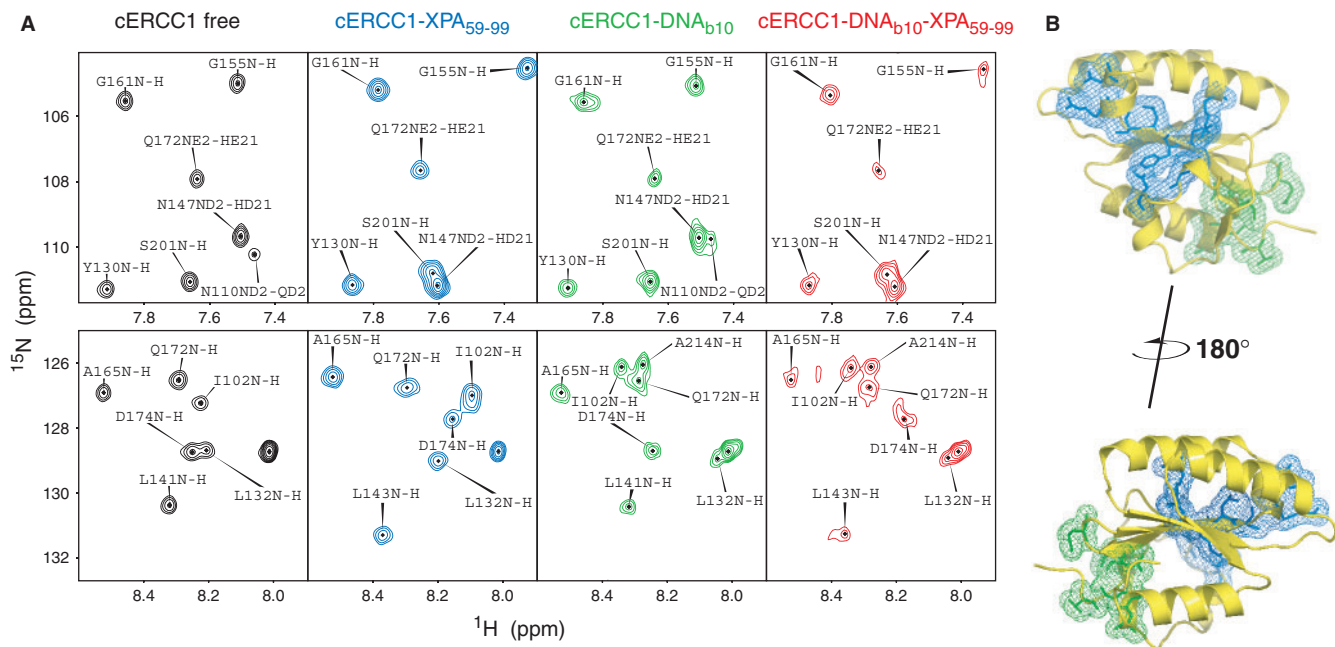


Figure 5. Dual function of cERCC1. (A) Indicative portions of the ^1H - ^{15}N HSQC spectrum for the free cERCC1, bound to either XPA (1:1) or DNA (1:1) separately, and the ternary complex (1:1:1). (B) Views of the two binding sites on cERCC1 structure.

slow exchange behavior suggests a large contribution to the binding by hydrophobic interactions with the DNA bases.

Summarizing, both the biochemical and the NMR experiments show that the ERCC1 central domain binds to DNA. The DNA-binding site we identified on cERCC1 by the NMR titrations suggests that cERCC1 contacts only a small part of the ssDNA, probably three to four bases. The situation is different in the full-length heterodimer, where other DNA-binding surfaces, such as previously established for the C-terminal ERCC1/XPF domains (12,14), assist in DNA recognition.

cERCC1 can bind simultaneously to XPA and ssDNA

Chemical shift perturbation experiments revealed distinct binding surfaces for ssDNA and XPA, and therefore, it is probable that both can bind simultaneously to cERCC1. We performed EMSA experiments in the presence of an equimolar or 5-fold excess of the HIS-XPA peptide. Addition of XPA did not influence binding of cERCC1 to ssDNA, while the binding affinity of XPA for the cERCC1 domain under these conditions would suggest that the vast majority of cERCC1 is in complex with XPA. We did not observe formation of a slower migrating HIS-GST-cERCC1/ssDNA/XPA super complex, possibly because the increase in mass by the addition of the HIS-XPA peptide was too small to discern (data not shown).

To independently confirm the formation of a ternary cERCC1/ssDNA/XPA complex, we added the HIS-XPA peptide to the cERCC1-bubble10 protein-DNA complex and followed the chemical shift perturbations in the ^1H - ^{15}N HSQC spectra. As shown in Figure 5, addition of an equimolar amount of the XPA peptide causes the same

amide displacements as in the titration with the free protein. On the contrary, most amides influenced by the DNA binding remain unaffected by the presence of XPA (Figure 5A). I102 and N110 were perturbed by both XPA and b10 DNA when the titrations were performed independently. In the ternary complex, however, I102 remains close to the position as in the DNA bound form. Conversely, the side-chain amide of N110, although affected by the DNA, adopts the distinct XPA bound position upon XPA addition. These data strongly indicate the formation of the ternary ERCC1/DNA/XPA complex. DNA and XPA-binding sites are distinct, and the two interactions can happen simultaneously (Figure 5B).

Functional role of the ERCC1 central domain in the heterodimer

Since the catalytic activity has been preserved in the human XPF protein through strong conservation of the nuclease signature (5), the human nuclease fold should be identical to that of the archaeal counterparts. In that sense, human XPF nuclease and human ERCC1 central domains are expected to exhibit the same fold. Moreover, the ERCC1 and XPF HhH₂ domains feature a common architecture and come into tight association exactly in the same way that the HhH₂ domains of both archaeal species do (10,11,14). Therefore, in structural terms the human ERCC1 and XPF proteins share the essential architectural subunits observed in the short XPF homodimer of *A. pernix*. The structural similarities within the XPF family provide additional evidence for the proposed common origin of the human ERCC1 and XPF proteins (33). Because ERCC1 is absent in archaea, this gene is thought to have been acquired from an ancient XPF gene

duplication in the eukaryal lineage. The ERCC1 and XPF genes have subsequently evolved by the process called subfunctionalization (36,37). This model suggests that after gene duplication, both copies may be reciprocally preserved through the fixation of complementary loss-of-subfunction mutations, which results in a partitioning of the tasks of the ancestral gene. From the functions present in the ancestral XPF protein (archaea), human ERCC1 retained the canonical HhH₂ domain that acts as a DNA-binding domain (14), while XPF retained the catalytic activity (5). Once the separation of the ancestral subfunctions occurred in ERCC1 and XPF genes, only the heterodimeric protein complex could restore the original function. Additionally, due to genetic alterations, the second helix-hairpin-helix motif of human XPF degenerated, yet the fold remained crucial for stabilizing the corresponding intact domain of human ERCC1 (14). Similarly, the central domain of ERCC1 lost the catalytic activity by sequence drift, but despite adoption of a novel dual function (XPA interaction and DNA binding), the 3D fold was preserved.

We suggest a mechanistic model for the heterodimeric function based on the model for the homodimeric archaeal homolog (Figure 6) (10,38). Human ERCC1 corresponds to the archaeal protomer that binds with the HhH₂ domain to the upstream duplex. Human XPF corresponds to the archaeal protomer that recognizes the downstream duplex. Since it cannot make contacts with the minor groove (14), the specificity of the human heterodimer has shifted to splayed-arms substrates consisting of only one duplex. This does not exclude contribution of the XPF HhH domain to ssDNA interactions, as reported before (12). We show here that the central domain of ERCC1 is also involved in ssDNA binding. Additional DNA interactions are likely for the nuclease of XPF, analogous to the archaeal case (38). Furthermore, ssDNA binding by cERCC1 will be stimulated by specific interactions with XPA, already present in the NER pre-incision complex and possibly bound to the DNA via its own DNA-binding site. Most importantly, through these coordinated interactions of the ERCC1 domains the XPF protein will be positioned to cleave the 3' protruding strand (limon), thereby retaining the polarity present in the archaeal homodimer.

Our model underlines the significant role of ERCC1 in the context of the full-length heterodimer. XPF is the catalytic module but the ERCC1 domains guarantee that the enzymatic activity is targeted properly. The presence of multiple distinct DNA-binding surfaces within the ERCC1/XPF/XPA/RPA repair protein intermediate coordinates cleavage to occur only when the DNA damage is recognized correctly by the NER machinery. The duplication of the ancestral XPF gene within the eukaryal kingdom resulted in an obligate heterodimer through loss of function (with an altered HhH₂ domain of XPF and a degenerated catalytic domain of ERCC1) and adoption of novel functions (ssDNA and XPA binding of cERCC1). This permitted additional quality control mechanisms through a more complicated molecular interaction network, mediated by the novel functional

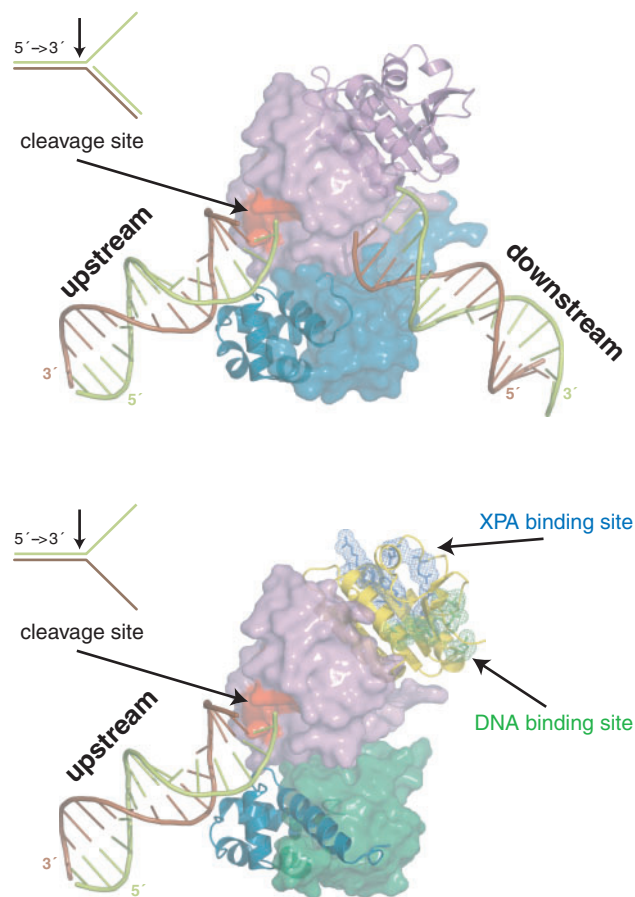


Figure 6. Models for the function of the archaeal homodimeric XPF (top) and human ERCC1/XPF heterodimer (bottom) constructed from the structure of the XPF homodimer bound to dsDNA (2bgw), the free structure of ERCC1/XPF C-terminal interacting domains (1z00) and the current structure of the ERCC1 central domain (2jpd). For the homodimer, one protomer is shown in a cartoon and the other in a surface representation. Accordingly, for the heterodimer ERCC1 is in a cartoon and XPF in a surface representation. The protein domains in both cases are colored as in Figure 1A.

domains, and thereby improving the fidelity of DNA damage repair.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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