

NAC domain, the one of *A. thaliana* ANAC019 (Ernst et al, 2004, EMBO Rep, 5:297-303), revealing a novel dimeric transcription factor fold. The structure is now the basis for mutagenesis studies to identify the DNA-binding mode (Olsen et al, 2005, Plant Science 169: 785-797) Structure determination of a new crystal form of this NAC domain shows that the dimer is likely to have some flexibility, but the solution structure as determined by small angle X-ray scattering is in good agreement with the original crystal structure. Based on ongoing crystallographic, small angle scattering and mutagenesis studies, our current model for NAC proteins binding to DNA will be presented.

Keywords: transcription factor, NAC protein, structure

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Structure of the topoisomerase IV from *S. pneumoniae* with a DNA target and quinolone drug

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Eighteen percent of all clinical bacterial infections are now treated with quinolone based antibiotics [1], which target the decatenating enzyme topoisomerase IV (a Class II topo) in gram-positive bacteria such as *S. pneumoniae*. Topoisomerase IV consists of both ParE and ParC domains. We have crystallised the complex of ParC55 (55 kDa) and ParE (30kDa) with a 32 base-pair DNA target and quinolone drug. Crystals were grown in space group P3₂ with cell dimensions $a=b=118.30$ Å, $c=177.90$ Å $\alpha=90^\circ, \beta=90^\circ, \gamma=120^\circ$ both by conventional hanging drop vapour diffusion in 24-well limbro plates and by sitting drop in 96 well MRC crystallisation plates. The structure has been solved by molecular replacement (CNS) using as search models our ParC55 structure [2] and a ParE domain homology modelled on the basis of the structure of the TOPRIM domain of the yeast type IIa [3]. The DNA has been positioned from difference Fourier maps following refinement using CNS. The veracity of the model was confirmed by the ability of this phase set to determine the correct Pt sites for a K₂PtCl₄ heavy atom derivative. Subsequently multiwavelength anomalous diffraction data were collected at the SOLEIL synchrotron about the Pt absorption edge in order to calculate a MAD map. We should like to thank the beamline personnel headed by K. McAuley at the DIAMOND synchrotron, Chilton, Oxford for their help in collecting native and fixed Pt edge synchrotron data on station IO3.

References:

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RNA splicing related proteins; Crystal structure of RNA 3'-terminal phosphate cyclase

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RNA 3'-terminal phosphate cyclase (Rtc) is an enzyme related to RNA splicing to convert the 3'-terminal hydroxyl group of truncated RNA to 2',3'-cyclic phosphate which is required just before its ligation. This reaction may occur in the following two steps: (i) $Rtc + ATP \rightarrow Rtc-AMP + PP_i$ and (ii) $RNA-N3'p + Rtc-AMP \rightarrow RNA-N>p + Rtc + AMP$. To reveal the reaction mechanism, Rtc overexpressed in *E.coli* was crystallized in the following states, Rtc, Rtc-AMP, Rtc+Mn and Rtc+ATP, and their structures have been determined at 2.25, 2.25, 3.2 and 2.4 Å resolutions, respectively. Rtc is a single protein folded into two domains, the large domain being composed of three $\beta\alpha\beta\alpha\beta\beta$ motifs arranged by pseudo three-fold symmetry, and the small domain being formed with a $\beta\alpha\beta\beta\alpha$ motif inserted into the third motif of the large domain. The overall structures of other derivatives are almost the same as that of Rtc. At the catalytic site of Rtc-AMP, the α -phosphate group of AMP is covalently bound to the N ϵ atom of His307, and the adenine moiety of AMP is stacked between the side chains of Pro126 and His283. The two hydroxyl groups at 2' and 3' positions of the ribose are bound to the side chain of Asp286 through hydrogen bonds. These structure features suggest the following reaction scheme. A Mg²⁺ ion bound to Glu10 induces conformational changes of the α and β phosphate groups of ATP which is trapped by Arg17 and Arg39, so that the N ϵ atom of His307 easily attacks to the α -phosphate group to form a P-N bond in the first reaction. In the second reaction, when a truncated RNA is bound, its 3'-phosphate group might be forced to react with the phosphate group of AMP, and the activated 3'-phosphate group is attacked by the 2'-hydroxyl group to generate the 2',3'-cyclic phosphodiester.

Keywords: RNA 3'-terminal phosphate cyclase, *Sulfolobus tokodaii*, X-ray crystal structure analysis

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Crystal structure of human DGCR8 core

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A complex of Drosha with DGCR8 (or its homolog Pasha) cleaves primary microRNA (pri-miRNA) substrates into precursor miRNA and initiates the microRNA maturation process. Drosha provides the catalytic site for this cleavage, whereas DGCR8 or Pasha provides a frame for anchoring substrate pri-miRNAs. To clarify the molecular basis underlying recognition of pri-miRNA by DGCR8 and Pasha, we determined the crystal structure of the human DGCR8 core (DGCR8S, residues 493 – 720). In the structure, the two double-stranded RNA – binding domains (dsRBDs) are arranged with pseudo two-fold symmetry and are tightly packed against the