Deep Trefoil Knot Implicated in RNA Binding Found in an Archaeabacterial Protein

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INTRODUCTION

(βα)₆ (TIM) barrel comprises one of the most abundant and versatile folds in nature. Although its structure is conserved, its primary sequence is highly divergent and gives rise to a plethora of distinct functions. Most structures deposited into the Protein Data Bank consist of the TIM barrel fold, and TIM barrel appears to be the most common fold in yeast. Nearly all TIM barrel proteins are enzymes, and so far, 15 distinct enzymatic functions have been assigned to TIM barrel containing proteins. The exception, narbonin, has no known function. Most commonly, their active sites are positioned within the β loops at the C-terminal end of the protein. TIM barrels are differentiated by auxiliary features such as location (and nature) of additional domains, identity of cofactor, number of (βα) units, location of the barrel major axis, and are based on strand and shear number. In general, in all TIM barrels and most other protein folds the main-chain does not cross over (to form a knot), although protein topologies involving formation of a knot have been reported in proteins.

The M. thermoautotrophicum MT1 gene is conserved in archaea, it lies in a ribosomal protein operon, and it codes for a 268 amino acid protein of unknown function. We report here the structure of MT1 that is novel from several standpoints: (i) the structure contains a novel topological unit—a deep C-terminal trefoil knot first observed in a TIM barrel-like fold, archaeabacterial proteins and rarely observed in other proteins; (ii) structurally, it contains only five (βα) units, and the arrangements of its hydrophobic and hydrophilic surfaces are opposite to that found in classical TIM barrel proteins; (iii) functionally, although it lacks typical features found in enzymes of the barrel family, it has strongly conserved residues clustered on the surface that form a potential catalytic site; (iv) the structure provides a first example of barrel-like fold linked to an RNA-binding domain, suggesting an extension of TIM barrel functionality to nucleic acid binding and/or catalysis.

MATERIALS AND METHODS

The MT1-coding region was cloned into pET-15b (Novagen) as a fusion with His₆ affinity tag and thrombin cleavage site. The protein was expressed in E. coli BL21 (DE3) containing plasmid that overexpresses rare E. coli tRNAs. Protein was purified by metal affinity chromatography on Ni-NTA superflow resin (Qiagen). The Se-Met derivative was produced as described previously.

Crystallographic statistics were performed by using the Screens I and II (Hampton Research). The best crystals were grown from a solution containing 5.7 mg/mL protein, 15% v/v Jeflamine M-600, 50 mM MES (pH 6.5), 25 mM CsCl in a drop equilibrated against 30% v/v Jeflamine M-600, 100 mM MES (pH 6.5), 50 mM CsCl at 23°C. The monoclinic C2 crystals (a = 101.307 Å, b = 51.353 Å, c = 109.25 Å, β = 94.69°) contain two molecules in the asymmetric unit related by a twofold noncrystallographic symmetry axis. A three wavelengths MAD data set was collected to 2.9 Å, and native data were collected to 2.3 Å at 100 K at SBC 19-ID beamline at the Advanced Photon Source of the Argonne National Laboratory by using a 3 × 3 mosaic CCD detector. All data were analyzed, indexed, and scaled by using HKL2000 (Table I).

The selenium substructure was solved by using the program SnB and refined with the program SHARP. Solvent flattening, histogram mapping, and twofold noncrystallographic symmetry averaging followed by electron density.

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density map calculation were performed by using the CCP4 suite\textsuperscript{15} (see references) and MAPMAN (Table II).\textsuperscript{16} The model was built manually by using O\textsuperscript{17} and refined against 2.3 Å native data using CNS.\textsuperscript{18} The final model has an R work\textsuperscript{22.1%} and R free\textsuperscript{27.7%}, and 120 water molecules. The Ramachandran plot calculated with the program PROCHECK (see CCP4 reference) shows all residues have favorable $\Phi$ and $\Psi$ angles (Table III). The electron density maps calculated from the final model refined at 2.3 Å were clear for all main-chain atoms except for the last six and the last four residues in the A and B monomers, respectively. The maps do not show density for residues N-terminal to the initiating methionine. The coordinates have been deposited in the Protein Data Bank with accession code 1K3R. Tertiary structure alignments were performed by using the DALI web server.\textsuperscript{19}

RESULTS AND DISCUSSION

The sequence homologues of MT1 are found only in archaea and eukaryotes (Fig. 1) and show no significant
sequence similarity with other proteins of known structure. The MT1 monomer consists of a large dimerization domain (MT1-DD) and a small β-barrel auxiliary domain (MT1-CSD). Two monomers dimerize via the helical face of their MT1-DDs, burying 3740 Å² (28%) of accessible surface area from each monomer (Fig. 2). The MT1-CSD is inserted in the second (αβ) loop region of the dimerization domain, and the two are connected by an α-helix and a 3_10 helix (Fig. 2). The β-barrel and a TIM barrel dimerization domains create a continuous 95 Å long positively charged surface for possible interaction with nucleic acid. Several strictly conserved residues are found on the dimerization as well as on MT1-DD/CSD interfaces.

The MT1-DD consists of 197 amino acids (residues 1–92 and 160–264) (Fig. 1). Analysis of structural homologs using the program DALI showed that the MT1-DD shares remarkable structural homology with TIM barrel enzymes, including *E. coli* methylenetetrahydrofolate re-
ductase (MTHFR) (PDB acc. no. 1B5T, Z score: 5.7, RMSD: 3.1 Å over 119 equivalenced residues), *M. kandleri* coenzyme F420-dependent tetrahydromethanopterin reductase (PDB acc. no. 1EZW, Z score: 5.4, RMSD: 3.8 Å over 124 equivalenced residues), and rabbit muscle pyruvate kinase (PDB acc. no. 1A49, Z score: 4.6, RMSD: 3.4 Å over 117 equivalenced residues). Despite missing half of the structural elements of a TIM barrel, the MT1-DD five strands and four helices superpose with \((\alpha/\beta)\) arrangement of the MTHFR TIM barrel with an RMSD of 3.1 Å (\(\beta\)-strands 1, 2, 3, 7, and 8 and \(\alpha\)-helices 1, 2, 7, and 8 are present in MT1-DD, using the numbering convention from MTHFR [Fig. 3]. Helix 3’s from MT1-DD and MTHFR do not overlap structurally because the MT1-DD \(\alpha\)-helix is dragged out of position to connect with the \(\beta\)-strand 7 (Fig. 4). Hence, the 4th, 5th, and 6th (\(\beta\alpha\)) units of MTHFR are missing in MT1-DD. Nevertheless, the MT1 dimer does not reconstitute complete TIM barrel structure. The existence of the MT1-DD stable partial TIM barrel strongly supports the notion that the ancestral TIM barrel was a half-barrel\(^{20}\) and suggests that the MT1-DD may be an early prototype of a TIM barrel. The \(\beta\) strand from the MT1 homolog from *S. pombe* is flanked by a glycine and proline with a spacing and internal \(\beta\)-strand sequence consistent with other TIM barrels\(^{21}\) (Fig. 1). The MT1 structure appears also like a classical nucleotide binding Rossmann fold.\(^{22}\) However, MT1-DD has no conserved nucleotide binding motifs.

At its C-terminus, the dimerization domain contains a knot. The 35 C-terminal residues are threaded through a loop on a surface that connects \(\alpha\)-helix 7 with \(\beta\)-strand 8 (Figs. 2, 3, and 5). The crossover involves residues Val233, Asn234, Ala193, and Ser194 (Figs. 1 and 5). The sequence that is threaded through the loop is not conserved with the exception of Asp230 and Pro237 on the C-side of the knot. There is also conserved proline residue within the loop (Pro195) that is located \(\sim 5\) Å from the Pro237. Several hydrophobic residues are
scattered throughout the sequence, and the length of the region varies among homologs (24–37 residues). The knot conformation is stabilized on N-site by Trp232, which appears to act as an anchor and on C-site by H-bond between Arg191 and Glu239. This region of MT1-DD is involved in dimer formation. Threading 35 residues through the loop region requires a major structural rearrangement (or cleavage and religation) of the protein main-chain.

Only a few other proteins have a knotted fold. 7–9 It is interesting that a very similar knot structure has been reported recently for the RrmA protein catalytic domain from Thermus thermophilus, which is predicted to be a 2'-O-ribose methyltransferase. 8 RrmA shares strong structural homology with MT1-DD including the knot region (PDB acc. no. 1IPA, Z score: 12.7, RMSD: 2.4 Å over 129 equivalenced residues). These proteins also show strikingly similar design. In MT1, a half-TIM barrel is fused with a putative cold-shock RNA-binding domain. In RrmA, the three-layer sandwich is fused with a eukaryotic ribosomal protein L30. 8 Despite the structural and perhaps functional similarities, MT1 and RrmA share virtually no sequence similarity, and the proposed catalytic residues in RrmA are not conserved in MT1 family. Present data suggest that the machinery responsible for creating the knot structure is present in bacteria, archaea, and eukaryotes.

The charge properties of the surface of the MT1-DD, with a polar interior and a hydrophobic exterior, are unlike those found in TIM barrels. Within the barrel, MT1-DD has an atypical abundance of charged and polar residues that point into the center of the barrel. The polar nature is further strengthened by the third α3 helix and the loop

![Fig. 5. The MT1 trefoil knot. Residues 1–190 and 199–229 are shown in solvent accessible surface representation (1.4 Å radius). The knot loop (residues 191–198) is in blue and the polypeptide chain threaded through the loop (residues 230–264) is labeled red. Crossover residues (Ser194 and Asn234) as well as Arg191 on the loop, Trp232 and Glu239 on threaded through chain are marked as a reference points. Carboxy terminus is labeled “C”. Figure was prepared with WebLabPro.](image)

### TABLE I. Data Collection Parameters

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<th>Wavelength (Å)</th>
<th>Resolution range (Å)</th>
<th>No. of reflections measured (unique)</th>
<th>% complete</th>
<th>Redundancy</th>
<th>R(merge) (%)</th>
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<td>78049 (11476)</td>
<td>92.4 (38.6)</td>
<td>6.80 (4.85)</td>
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<td>9.4 (53.6)</td>
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<td>26.77 (3.21)</td>
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*Rmerge = ∑|I − <I>|/|I|*, where I is the intensity of an individual measurement and <I> is the average intensity from multiple observations. Data in parentheses typically indicate values for last resolution shell.
TABLE II. Phasing Power Statistics

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<td>RMSD improper (°) (^d)</td>
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\(^a\) \(R = \sum_{hkl}||F_o|| - |F_c|/\sum_{hkl}|F_o|\), where \(F_o\) and \(F_c\) are the calculated and observed structure factor amplitudes for reflection hkl, respectively.

\(^b\) \(R_{free}\) is the same as \(R\) but calculated over a 10% fraction of the reflection data that were randomly selected and not included in the refinement.

\(^c\) Program PROCHECK.\(^c\)

\(^d\) Root-mean-square deviations from standard geometry values, as determined with the CNS\(^d\) program package.

linking strands \(\beta8\) and \(\beta9\), which shield several hydrophobic residues (data not shown). In contrast, its \(\alpha\)-helical face is unusually hydrophobic and drives dimerization.

MT1 has an auxiliary domain inserted into a loop of the TIM barrel [Figs. 1 and 4(b)]. The auxiliary domain is 67 amino acids long and contains residues 93–159 in the MT1 protein [Figs. 1 and 2]. The program DALI shows the this domain shares significant 3D structural similarity with TIM barrel proteins: *E. coli* major cold-shock protein CspA (PDB acc. no. 1MJC, Z score 5.6; RMSD: 2.5 Å over 58 equivalent residues), *E. coli* polyribonucleotide nucleotidyl transferase-S1 RNA-binding domain (PDB acc. no. 1SRO, Z score: 5.0; RMSD 2.3 Å over 52 equivalent residues), *Thermus thermophilus* S17 protein (PDB acc. no. 1FJF, Z score 5.1, RMSD: 2.6 Å over 54 equivalent residues). All four proteins are five-stranded antiparallel \(\beta\)-barrels that share the identical topology, known as the cold-shock domain\(^23\) [Fig. 4(b)]. This type of structure is also classified as an oligonucleotide-binding (OB) fold. Preliminary data suggest that MT1 is not a cold-shock protein (Giometti and Töllaksen, personal communication). The electrostatic potential surface map drawn by using GRASP\(^24\) also indicates a potential nucleic acid-binding role. MT1 has a distinct positively charged face (data not shown). Although the C-terminal \(\beta\)-sheet residues of the barrels contribute a small portion of the positive charge, most of the charge is located in the MT1-CSDs. This charge distribution is similar to other nucleic acid-binding proteins with this fold.

It is likely that the MT1 binds single- or double-stranded RNA for the following reasons. First, MT1-CSD has significant structural similarity with CspA, which is an RNA chaperone that binds RNA to prevent hairpin formation for transcription antitermination,\(^25,26\) and with the ribosomal protein S17, which binds double-stranded regions of the 16s rRNA. Second, the gene for MT1 is located within a cluster of genes related to ribosomal function. However, we cannot rule out the possibility that MT1 binds DNA because the OB fold is found in many ssDNA-binding proteins,\(^27\) including the product of BRCA2 oncogene that contains three such units and binds single-stranded DNA.\(^28\)

The MT1 structure, particularly the inserted domain, provides additional support to the observation that organisms accomplish complex tasks using modular protein design from a limited number of modules. There are at least two other examples of TIM barrel proteins that contain a similar insertion. Rabbit muscle pyruvate kinase contains a 9-stranded barrel inserted after the third \(\beta\alpha\) loop. In that protein, the \(\beta\)-barrel domain forms part of the active site with another structurally unrelated domain.\(^29\) The *Bacillus cereus* \(\beta\)-amylase TIM barrel contains a seven-stranded barrel, which forms the maltose-binding site and inserted at the C-terminus of the protein.\(^30\) In both instances, the inserted \(\beta\)-barrel domains are important for enzymatic catalysis.

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