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2376



Conjugative transfer through type IV secretion multiprotein complexes is the most important means of spreading antimicrobial resistance. Plasmid pIP501, frequently found in clinical Enterococcus faecalis and Enterococcus faecium isolates, is the first Gram-positive (G+) conjugative plasmid for which self-transfer to Gram-negative (G-) bacteria has been demonstrated. The pIP501-encoded type IV secretion system (T4SS) protein TraN localizes to the cytoplasm and shows specific DNA binding. The specific DNA-binding site upstream of the pIP501 origin of transfer (oriT) was identified by a novel footprinting technique based on exonuclease digestion and sequencing, suggesting TraN to be an accessory protein of the pIP501 relaxase TraA. The structure of TraN was determined to 1.35 Å resolution. It revealed an internal dimer fold with antiparallel β -sheets in the centre and a helixturn-helix (HTH) motif at both ends. Surprisingly, structurally related proteins (excisionases from T4SSs of G+ conjugative transposons and transcriptional regulators of the MerR family) resembling only one half of TraN were found. Thus, TraN may be involved in the early steps of pIP501 transfer, possibly triggering pIP501 TraA relaxase activity by recruiting the relaxosome to the assembled mating pore.

1. Introduction

As the major contributor to horizontal gene transfer, conjugative spread of plasmid-encoded antibiotic-resistance and pathogenicity genes represents a serious threat to hospitalized and immunosuppressed patients (Williams & Hergenrother, 2008). In the process of conjugation, DNA is transported from a donor to a recipient cell, requiring direct contact between the cells (Cascales & Christie, 2003; Alvarez-Martinez & Christie, 2009). The transfer is mediated by a plasmidencoded, multi-protein complex that is large enough to span the bacterial cell wall: the type IV secretion system (T4SS; Llosa et al., 2002). T4SSs have been studied in detail in bacteria of Gram-negative (G-) origin, such as Escherichia coli and Agrobacterium tumefaciens (Llosa et al., 2009; de la Cruz et al., 2010; Hayes et al., 2010; Rêgo et al., 2010; Smillie et al., 2010; Wallden et al., 2010; Thanassi et al., 2012; Zechner et al., 2012; Christie et al., 2014). For systems of Gram-positive (G+) origin, most knowledge of the T4SSs was originally based on their similarity to their G- counterparts (Grohmann et al., 2003; Abajy et al., 2007), but recently significant progress has been made towards their functional and structural characterization. The findings of Chen et al. (2008) support a model in which the putative coupling protein PcfC of the Enterococcus faecalis sex pheromone plasmid pCF10 initiates the conjugative transfer of the substrate through a pCF10encoded T4S channel. As with G- T4SSs, this mechanism is NTP-dependent and requires the presence of the peptidoReceived 12 March 2014 Accepted 17 June 2014

PDB references: TraN, 4p0z; 4p0y; 4pm3



glycan hydrolase PrgK (Laverde Gomez *et al.*, 2014). Only recently has structural information on several G+ T4SS proteins become available: a proposed scaffolding factor TcpC from plasmid pCW3 (Porter *et al.*, 2012), a VirD4-like ATPase domain from *Thermoanaerobacter pseudethanolicus* (Walldén *et al.*, 2012), the C-terminal VirB8-like domain of the pIP501 T4S protein TraM (Goessweiner-Mohr, Grumet *et al.*, 2013) and the pIP501 T4S protein TraK, which exhibits a novel fold (Goessweiner-Mohr *et al.*, 2014). A broad overview of the current G+ T4SS knowledge base is provided in a recent review (Goessweiner-Mohr, Arends *et al.*, 2013).

pIP501, a multiple antibiotic-resistance plasmid frequently encountered in clinical E. faecalis and E. faecium isolates (Rosvoll et al., 2010), is the conjugative plasmid with the broadest host range in G+ bacteria (Kurenbach et al., 2003). 15 putative T4SS genes are organized in a single operon. Significant sequence similarity to the T4SS from A. tumefaciens has only been detected for three of these proteins. TraE, an ATPase related to VirB4, interacts with itself as well as with several other potential pIP501 T4SS proteins (Abajy et al., 2007). This motor ATPase is most likely to drive the conjugation process. The hexameric coupling protein TraJ (E.-K. Celik, W. Keller & E. Grohmann, unpublished data), related to VirD4, lacks the transmembrane region shown for other coupling proteins (Gomis-Rüth et al., 2001; Atmakuri et al., 2004). Coupling proteins connect the macromolecular complex of single-stranded plasmid DNA and relaxosomeassociated proteins, which are transported to the recipient cell with the T4SS secretory conduit (Gomis-Rüth et al., 2004). Another pIP501 T4SS protein, TraI, might be responsible for recruitment of the coupling protein TraJ to the cell membrane (Alvarez-Martinez & Christie, 2009). To assemble the conjugative core complex in G+ bacteria, the thick peptidoglycan layer needs to be opened locally. In the case of the pIP501 T4SS this role is performed by the muramidase TraG (Arends et al., 2013), a protein related to VirB1. Recently, we published the 2.5 Å resolution structure of the C-terminal domain of the pIP501 T4SS protein TraM, with similarity to VirB8-like proteins (Goessweiner-Mohr, Grumet et al., 2013), as well as the 3.0 Å resolution structure of TraK_{pIP501} (Goessweiner-Mohr et al., 2014). Despite these emerging structural data and the above-described insight into some of the 15 potential T4SS proteins, we still lack information on most of the individual components.

Here, we present the biophysical and structural characterization of the T4SS protein TraN, a 17.6 kDa protein (formerly ORF14; GenBank CAD44394.1) encoded by the conjugative plasmid pIP501. The cytoplasmic protein binds to doublestranded (ds) DNA. Using various $oriT_{pIP501}$ -flanking DNA constructs, we were able to determine the preferred binding site as a 34 bp sequence positioned upstream of $oriT_{pIP501}$, which suggests TraN to be an accessory or regulatory factor within the pIP501 transfer system. We report the 1.35 Å resolution TraN structure, which has been solved by selenomethionine single-wavelength anomalous dispersion (SAD) methods. TraN comprises an HTH fold with an internal dimer composition. The TraN domains resemble the fold of excisionases (Xis) from transposons of G+ origin and bacteriophages as well as transcriptional regulators of the MerR family. A search for structurally related proteins in putative T4SSs from conjugative plasmids, transposons, ICEs and genomic islands (GI) revealed only a few mobile genetic elements which possess a TraN-like protein.

2. Materials and methods

Detailed information on the expression, purification and crystallization of native TraN has been published in Goess-weiner-Mohr *et al.* (2012).

The final coordinates and structure-factor amplitudes of the TraN crystal structures have been deposited in the PDB as entries 4p0z (crystal form I), 4p0y (crystal form II) and 4pm3 (crystal form III).

2.1. Expression of selenomethionine-labelled TraN

For the expression of selenomethionine-labelled TraN, pQTEV-*traN* plasmid DNA was transformed into the methionine-deficient *E. coli* strain B834 (DE3) (Novagen, Merck, Darmstadt, Germany) using standard protocols. The cells were resuspended in M9 minimal medium at an OD₆₀₀ of ~0.6, grown for 1 h at 37°C and induced with 1 mM IPTG. 25 mg selenomethionine was added and overexpression was continued for 3 h. The cells were harvested and frozen at -20° C. Selenomethionine-labelled TraN expression levels were monitored by SDS–PAGE. Protein extraction was performed as described in Goessweiner-Mohr *et al.* (2012).

2.2. Biophysical characterization of TraN

TraN was extracted and His-affinity purified in 50 m*M* Tris, 100 m*M* ammonium sulfate pH 7.45. TraN-containing Hisaffinity fractions were pooled and concentrated *via* centrifugation in Amicon tubes (Millipore Amicon, 3000 MWCO). TraN was further purified by size-exclusion chromatography with a Superdex 200 HR 10/30 column (GE Healthcare, Chalfont St Giles, England). A gel-filtration standard (670, 158, 44, 17 and 1.35 kDa; Bio-Rad, Hercules, California, USA) was used to calculate the molecular weight of TraN. TraNcontaining size-exclusion fractions with a maximal concentration of 1.58 mg ml⁻¹ were stored at -80 °C for subsequent experiments.

Circular-dichroism (CD) measurements were performed on a Jasco J715 spectropolarimeter (Jasco, Gross-Umstadt, Germany) equipped with an external thermostat. Spectra were measured from 260 to 190 nm in a 0.01 cm cuvette with a protein concentration of 0.89 mg ml⁻¹. Ten individual spectra were accumulated and the standard deviation was calculated from the repeated measurements. Temperature scans were performed in a 0.02 cm temperature-controlled cuvette from 25 to 95°C using a step–scan procedure with a constant wavelength of 208 nm. Spectra were recorded in triplicate from 260 to 190 nm every 5°C. The temperature slope was set to 1°C min⁻¹. TraN was applied at a concentration of 0.42 mg ml⁻¹. CD data were evaluated using the online service *DichroWeb* (Whitmore & Wallace, 2008) and reference data set No. 4. For the DLS measurements, a size-exclusion fraction containing 0.9 mg ml⁻¹ TraN was measured in a 45 µl cuvette. Ten measurements with constant baseline were merged and the monodispersity was assessed.

For the SAXS measurements on the X33 beamline at DESY, Hamburg, Germany, TraN was suspended in 100 mM ammonium sulfate, 100 mM NaCl, 50 mM HEPES pH 7.0. Size-exclusion purified protein was concentrated and TraN was measured at three different concentrations: 3.18, 1.57 and 0.96 mg ml⁻¹. *PRIMUS* (Konarev *et al.*, 2003) was applied to subtract the buffer from the protein data. The maximum intensity (I_0) and radius of gyration (R_g) were extracted from the Guinier plot generated from the data at 1.57 mg ml⁻¹. I_0 was used to calculate the molecular weight of TraN in solution, GNOME (Svergun, 1992) was used to generate the output file for subsequent ab initio modelling with GASBOR (Svergun et al., 2001), and CRYSOL (Svergun et al., 1995) was used to generate a scattering curve from the X-ray-derived model. The BUNCH software (Petoukhov & Svergun, 2005) was used to fit the X-ray data to the SAXS scattering curve of TraN. This was carried out by ab initio and rigid-body modelling of the N-terminal His tag of the construct, which was present in solution but was lost or not visible in the X-ray model.

2.3. Subcellular fractionation of *E. faecalis* JH2-2 (pIP501) and immunolocalization of TraN

Subcellular fractionation of E. faecalis JH2-2 (pIP501) was performed according to Buttaro et al. (2000) with modifications. An exponentially growing culture ($OD_{600} = 0.5$) of E. faecalis JH2-2 (pIP501) was chilled on ice, washed twice in an equal volume of 50 mM potassium phosphate buffer pH 7.0 and resuspended [1:50(v:v)] in lysis buffer $(50 \text{ m}M \text{ KH}_2\text{PO}_4/\text{ m})$ K_2 HPO₄ pH 7.0, 1 mM EDTA, 1 mM MgCl₂, 100 µg ml⁻¹ DNase, $100 \ \mu g \ ml^{-1}$ RNase). The cells were broken by FastPrep-24 (MP Biomedicals, Illkirch, France) using lysing matrix E (1.4 mm ceramic spheres, 0.1 mm silica spheres, 4 mm glass beads; MP Biomedicals, Illkirch, France). Unlysed cells were removed by low-speed centrifugation. The cell-wall fraction was harvested by high-speed centrifugation at 17 000g for 20 min at 4°C and the membrane fraction was obtained by ultracentrifugation of the supernatant at 45 000 rev min⁻¹ for 2 h at 4°C (OTD Combi ultracentrifuge, Thermo Fisher Scientific GmbH, Dreieich, Germany). The remaining supernatant contained the soluble proteins.

TraN was localized in the fractions (cell wall, membrane, cytoplasm) by immunostaining with primary polyclonal anti-TraN antibodies (BioGenes, Berlin, Germany) and a secondary horseradish-peroxidase conjugated anti-rabbit IgG antibody (Promega GmbH, Mannheim, Germany).

2.4. EMSA with unlabelled DNA fragments

Eight unlabelled DNA fragments of variable length were tested (fragments A–H; Supplementary Fig. $S3^{1}$). Except for fragments D and G, all fragments contained the pIP501 *oriT*

sequence and a sequence upstream and downstream of the *oriT*. Fragments A–E mostly comprised an upstream *oriT* sequence. 3 μ l of the respective fragments, corresponding to 1–2 pmol dsDNA, were mixed with a 0/1/2.5/5/10/25/50/100-fold molar excess of TraN protein, 3 μ l glycerol, 1 m*M* EDTA, 15 m*M* NaCl, 20 m*M* Tris–HCl pH 7.8, 20 μ g BSA and H₂O to a final volume of 20 μ l. The samples were incubated at room temperature (RT) for 30 min. 1 μ l DNA loading buffer (R0611; ThermoFisher Scientific, Waltham, Massachusetts, USA) was added and the samples were run on 1% agarose gels at a constant voltage of 80 V for 35 min.

2.5. Exonuclease digestion and sequencing-based footprinting assay

10 µl of DNA fragments C and L (see Supplementary Fig S3; up to 100 ng ml^{-1}) were incubated with a tenfold molar excess of TraN protein and without protein (negative control), respectively. The 20 µl incubation mixture consisted of 15 mM NaCl, 20 mM Tris-HCl pH 7.8, 20 µg BSA. The samples were incubated at RT for 30 min. 1.1 µl H₂O, 0.5 µl T7 exonuclease (New England Biolabs, Frankfurt, Germany) and 2.4 μ l 10 \times NEB4 buffer were added and the samples were mixed and incubated at RT for 10, 15 or 20 min. 5 μ l 20 mg ml⁻¹ proteinase K (New England Biolabs, Frankfurt, Germany) was added to stop the exonuclease reaction. The samples were incubated at RT for 30 min, followed by purification of the digested DNA fragments using the Wizard SV Gel and PCR Clean-Up System (Promega GmbH, Mannheim, Germany). The DNA was dissolved in distilled H₂O and the respective forward and reverse primers were added to allow Sanger sequencing of both DNA strands. Samples were sent to either the LGC Genomics GmbH (Berlin, Germany) or the Microsynth AG (Wolfurt-Bahnhof, Austria) sequencing services. The respective chromatograms were analyzed for the presence of a significant deterioration in the base-calling signal and the corresponding position in the DNA sequence was identified for all forward and reverse sequences. The sequences were aligned and the matching (protected) area was identified as the specific binding site of TraN.

2.6. Thermofluor assay

10 μ l of TraN (0.85 mg ml⁻¹) was mixed with random or specific 34-mer dsDNA oligonucleotides in a 1:1 ratio, distilled water was added to 20 μ l, 5 μ l of 50× SYPRO Orange (Sigma-Aldrich, St Louis, Missouri, USA) was added and the solution was incubated at RT for 10 min. In samples without oligonucleotides, the DNA was substituted by an equal volume of H₂O. The melting curve was measured from 25 to 95°C in increments of 0.1°C and with a 6 s incubation time at every step. The mean and the standard deviation of the resulting melting temperatures were individually calculated for all three samples (TraN alone, TraN plus random 34-mer and TraN plus 34 bp binding site).

¹ Supporting information has been deposited in the IUCr electronic archive (Reference: CB5056).

2.7. ITC experiments

ITC was performed on a MicroCal VP ITC System (GE Healthcare, Uppsala, Sweden). The calorimeter was calibrated according to the manufacturer's instructions. DNA oligonucleotides containing either the 34 bp binding site (CGGAAATGTCAGGTTAAACATATTTACTTTTATA) or the $oriT_{pIP501}$ (nonspecific DNA; AGGGCGCACTTATAC-GCAGTAACTTCGTTACTTC) sequence, as well as their complementary strands, were ordered from Life Technologies. The DNA was dissolved in TraN purification buffer to minimize artifacts owing to minor differences in buffer composition. Equal amounts of the complementary ssDNA oligonucleotides were mixed and heated to 95°C to anneal the two strands. The concentration was determined prior to the measurements. The sample cell (1.421 ml) was loaded with 19- $26 \,\mu M$ protein. A typical titration experiment consisted of 25 injections of DNA solution (205-221 µM), 12 µl volume and 4 s duration with a 5 min interval between additions. The titration was performed at constant temperature of 25°C. The stirring rate was set to 270 rev min^{-1} . For the competitive titration, the protein was saturated with nonspecific DNA (molar ratio of 1:1.2) and injected with specific DNA accordingly. For the reverse titration, the sample cell was filled with $12.9 \,\mu M$ specific binding-site DNA and the titration consisted of 25 injections of protein with a concentration of 259 µM. TraN protein and DNA ligands were titrated against buffer to measure nonspecific heat effects.

The raw data were integrated, corrected for nonspecific heats, normalized for concentration and analyzed according to a single-binding-site interaction model utilizing the *Origin* v.7.0 scientific plotting software (supplied with the ITC instrument). For the reverse titration with specific binding-site DNA, a two-binding-site interaction model was used to fit the data utilizing the results of the single titrations as starting values for the fitting.

2.8. Mass spectrometry of TraN crystals

Several crystals of TraN were dissolved in $10 \,\mu l$ distilled water and investigated by MALDI-TOF spectrometry (ultra-fleXtreme, Bruker, Vienna, Austria).

2.9. Crystallization of selenomethionine-derivatized TraN

TraN crystallization and optimization have been described in Goessweiner-Mohr *et al.* (2012). Since no significant sequence similarity with available structures was found, we generated a selenomethionine derivative of TraN. Optimized conditions were used for the following setups with selenomethionine-containing TraN. A seeding stock was prepared from TraN crystals and seeding-optimization plates were produced with the microbatch method and Index condition 42 only. For some setups TEV-cleaved TraN was used. Plates had a drop volume of 1 µl with a drop ratio of 50%(v/v) in the nonseeding conditions and a drop ratio of 40%(v/v) protein, 40%(v/v) buffer and 20%(v/v) seeding stock in the seeding conditions. Promising crystals were harvested from two conditions: (i) 0.1 *M* bis-tris pH 5.5, 25%(w/v) PEG 3350, no seeding, TEV-cleaved protein, stock concentration 0.54 mg ml⁻¹ and (ii) 0.1 *M* bis-tris pH 5.5, 25%(w/v) PEG 3350, nondiluted seeding stock, protein stock concentration 1.52 mg ml⁻¹.

2.10. Data collection, processing and refinement

Data collection was performed at 100 K on the synchrotron beamline X06DA at SLS, Villigen, Switzerland. No cryoprotectant was required. The selenomethionine-containing crystals diffracted to a resolution of about 1.8 Å on our home source and to 1.35–1.5 Å resolution at SLS. Several crystals showed isotropic diffraction with clear spots and nontwinned patterns. A fluorescence scan was performed to validate the presence of selenomethionine within the crystals and to define the optimal setup for anomalous data collection at the selenium f'' inflection, peak and high-end remote wavelengths (0.9794, 0.9796 and 0.9714 Å, respectively). A crystal-todetector distance of 120 mm, an oscillation range of 1.0° and an exposure time of 1 s per image were chosen. Multiwavelength anomalous dispersion (MAD) data sets at the mentioned wavelengths were collected for several crystals.

The crystals that were used for data processing and refinement all belonged to space group $P2_1$, with unit-cell parameters a = 33.15, b = 55.09, c = 35.52 Å, $\beta = 113.05^{\circ}$ for crystal form I, a = 33.10, b = 62.82, c = 55.87 Å, $\beta = 90.03^{\circ}$ for crystal form II and a = 32.88, b = 54.94, c = 57.71 Å, $\beta = 91.89^{\circ}$ for crystal form III, with one (form I) or two (forms II and III) molecules per asymmetric unit. The Matthews coefficient (Kantardjieff & Rupp, 2003) was calculated to be 2.04, 1.99 and 1.83 Å³ Da⁻¹ for crystal forms I, II and III, respectively, and the solvent content was 39.9, 38.2 and 32.9%, respectively. Details are presented in Table 2.

The first data set from crystal form I was used to solve the preliminary structure of TraN at the synchrotron. The resulting model was utilized in the data processing of further data sets (data 1 from crystal form I, data 2 from crystal form II and data 3 from crystal form III) using molecular replacement. Data 3 represents the previously published data set for the TraN protein (Goessweiner-Mohr et al., 2012). Although not stated in Goessweiner-Mohr et al. (2012), the crystal was soaked in K₂PtBr₄ solution for 1.5 h and the data set was obtained at 1.0615 Å, corresponding to the high-end remote wavelength of the Pt L_{III} absorption edge. As the anomalous signal was weak and the heavy-atom position could not be unambiguously determined, at the time the data were treated as a native data set. In the evaluation of the data the Pt and Br atom positions of the heavy-atom compound were placed in the residual electron density and were refined with a partial occupancy of 0.5. The resulting structures were refined to final $R/R_{\rm free}$ values of 0.1685/0.2022 (data 1), 0.1833/0.2233 (data 2) and 0.1693/0.2134 (data 3). In the models from data 1 and 2, 100% of the residues were found in the favoured Ramachandran areas. The final MolProbity scores were 1.30 (93rd percentile), 1.65 (69th percentile) and 1.13 (99th percentile) for data 1, 2 and 3, respectively.

The data sets were processed and scaled together using XDS and XSCALE (Kabsch, 2010), *iMosflm* (Battye et al.,

2011) and SCALA (Evans, 2006). The programs SHELXC/D/ E (Sheldrick, 2010) were used to define the selenium heavyatom sites and to build an initial model (SHELXE chaintracing function). The resulting model was completed manually in Coot (Emsley et al., 2010) and refined with REFMAC5 (Murshudov et al., 2011). For further data sets, the refined TraN SAD structure was the basis for molecular replacement with MOLREP (Vagin & Teplyakov, 2010) to generate the initial models. Data set 3 was also refined with phenix.refine (Afonine et al., 2012) from the PHENIX software suite (Adams et al., 2010). The refined X-ray models were validated by MolProbity (Chen et al., 2010). Three-dimensional alignments of the TraN structure with structural homologues were performed with DALI (Holm & Rosenström, 2010) and MATRAS (Kawabata, 2003). Structural alignment of TraN internal monomers was conducted with MASS (Dror et al., 2003). PyMOL (Schrödinger) was used to prepare structural representations and to calculate the r.m.s.d.s of TraN monomer alignments. Structural and sequential similarities between the two halves of TraN and structurally related proteins were examined using the pairwise structural alignment feature of MATRAS and the SSM algorithm (Krissinel & Henrick, 2004) included in Coot.

2.11. Sequencing-based comparison and characterization

HMMTOP (Tusnády & Simon, 2001), *MemsatSVM* (Nugent & Jones, 2009) and *Memsat3* (Jones *et al.*, 1994) were used to search for transmembrane motifs in the TraN sequence and potential homologous proteins.

PSIpred (Jones, 1999) was used to predict the secondarystructure content of TraN and of structurally related proteins, but where known, the secondary structure was derived from the crystal structure. The general features of the His-tagged TraN construct were assessed with *ProtParam* (Gasteiger *et al.*, 2003).

3. Results

3.1. TraN localizes to the cytoplasm

To localize the TraN protein *in vivo*, an exponentially growing culture of *E. faecalis* JH2-2 (pIP501) was fractionated into cell-wall, membrane and cytoplasmic fractions as described by Buttaro *et al.* (2000). TraN was exclusively found in the cytoplasmic fraction (Fig. 1*a*). We exclude contamination of the cytoplasmic fraction with cell-envelope proteins and *vice versa* as TraM_{pIP501}, which is predicted by *PSORTb* to localize to the cell envelope, was exclusively found in the cytoplasm under the same conditions (Goessweiner-Mohr, Grumet *et al.*, 2013).

3.2. Biophysical characterization

The expression and purification of a His-tagged TraN construct (17.6 kDa) have recently been reported (Goess-weiner-Mohr *et al.*, 2012). The protein eluted from the gel-filtration column as a single peak (Supplementary Fig. S1*a*) with an apparent molecular weight of 15.4 kDa, indicative of

a homogenous monomeric protein. Dynamic light-scattering (DLS) measurements yielded a single peak with a calculated polydispersity of 26.3% and a hydrodynamic radius (R_h) of 2.7 nm (Supplementary Fig. S1*b*).

SAXS measurements were performed to obtain further information about the oligomeric state and shape of TraN in solution. Data collection was performed on the X33 beamline (DESY, Hamburg, Germany) for three different protein concentrations. An I_0 of 13.7 was calculated for the data at medium concentration (1.57 mg ml⁻¹), resulting in a molecular weight of 14.1 kDa, which is in agreement with the theoretical molecular weight of TraN (17.6 kDa). A radius of gyration (R_{o}) of 2.28 nm was calculated from the Guinier plot and a D_{max} of 6 nm was determined from the p(r) function. The TraN monomers in the crystal showed a maximal diameter of only about 4.4 nm. An extended hydration shell may explain the significantly larger particle depicted in the SAXS measurements. On calculating ab initio models from the scattering function we observed an elongated particle, which may be owing to the flexible N-terminal end of the TraN construct containing the unstructured His tag.

TraN is folded in solution and possesses a mixed α - β composition as shown by CD spectroscopy (Supplementary Fig. S2*a*). The α -helical content exceeds that of β -sheets

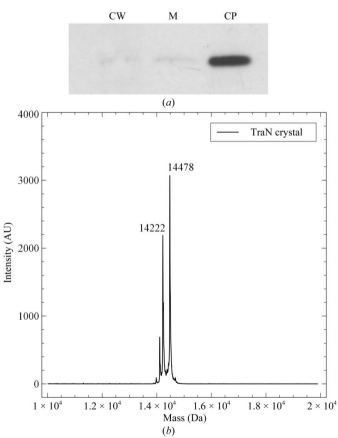


Figure 1

TraN localization and MS analysis of TraN crystals. (*a*) TraN localizes to the cytoplasm of pIP501-harbouring *E. faecalis* JH2-2 cells. TraN was localized in the cell fractions by Western blotting with anti-TraN antibodies. CW, cell wall; M, membrane; CP, cytoplasm. (*b*) MALDI-TOF analysis of TraN crystals.

(Supplementary Fig. S2*b*). The large proportion of unordered structure (27%) may result mainly from the N-terminal His tag. Temperature scans revealed that TraN is a stable protein with a melting temperature $T_{\rm m}$ of 55°C. Upon a thermal shift to 95°C the protein unfolded completely, but it returned to its original fold after cooling to RT (Supplementary Fig. S2*c*).

3.3. TraN binds to a sequence upstream of $oriT_{pIP501}$

As TraN co-purified with DNA, we tested its ability to interact with DNA molecules: the protein was incubated with radiolabelled ssDNA and dsDNA oligonucleotides, respectively, in electrophoretic mobility-shift assays (EMSA; data not shown). The oligonucleotides had a length of 42 bases and contained either the *oriT*_{pIP501} sequence (GenBank L39769.1, bp 1259–1296) or a random sequence with no sequence similarity to *oriT*_{pIP501}. TraN showed only a weak shift for the ssDNA oligonucleotides, whereas a significant shift was observed for the dsDNA fragments. There was no difference in the binding affinity between the random and the *oriT*_{pIP501}-containing oligonucleotide.

In order to check for a potential sequence-specific TraN binding site on the pIP501 plasmid, we performed gel-shift assays with dsDNA fragments containing the $oriT_{pIP501}$ sequence as well as sequences upstream and downstream of the $oriT_{pIP501}$ region (Supplementary Fig. S3). All tested DNA fragments were cooperatively shifted at higher TraN concen-

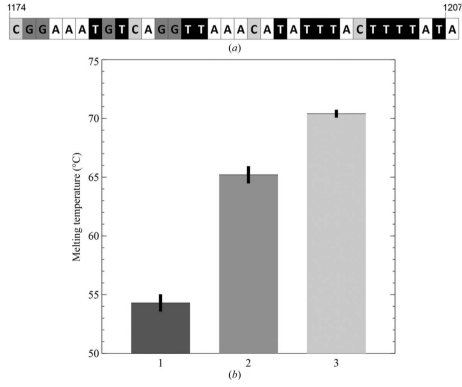


Figure 2

The TraN specific binding site and its stabilizing effect on the protein. (*a*) Sequence of the 34 bp fragment identified as the specific binding site of TraN upstream of $oriT_{pIP501}$. The position on the pIP501 plasmid sequence is indicated. (*b*) Thermofluor-based analysis of the TraN melting temperature without DNA (1), in the presence of a random 34-mer dsDNA oligonucleotide (2) and in the presence of the specific binding site (3). The standard deviation of each of the sets of 25 individual measurements is displayed.

trations, thus confirming the nonspecific binding of our initial EMSA experiments. A small but significant stepwise shift was visible for fragments A and D using an equimolar protein:DNA ratio. These two fragments both comprise a common 149 bp sequence upstream of the oriT sequence (Supplementary Fig. S3), which therefore has to contain a preferred binding site. Fragments which did not comprise this sequence (fragments B, G and H) did not exhibit a shift at equimolar protein and DNA concentrations. Upon close examination, fragments C, E and F also exhibited a defined, albeit small, shift at equimolar TraN and DNA concentrations. These three fragments contain the same 149 bp oriT upstream sequence as fragments A and D but are significantly larger. Thus, it appears that the smaller relative molecular mass change through TraN binding results in a less pronounced shift than for the smaller fragments A and D. From these experiments we conclude that the 149 bp sequence of fragment D contains a preferred high-affinity TraN binding site, as shown in Supplementary Fig. S3.

3.4. Identification of the specific TraN binding site

We developed a new footprinting protocol based on 5'-to-3' exonuclease digestion to identify the specific binding site of TraN within the previously identified pIP501 sequence. The new method was validated with three different DNA-binding proteins: the bZIP protein GCN4 (PDB entry 2dgc; Keller *et*

> *al.*, 1995), the ribbon-helix-helix protein ParD of the conjugative plasmid RP4 (PDB entry 2an7; Oberer *et al.*, 2007) and the helix-turn-helix protein ParR of the conjugative plasmid R1 (Breüner *et al.*, 1996; Møller-Jensen *et al.*, 2007). In all three cases, the published binding sites could be verified by this novel method (N. Goessweiner-Mohr & W. Keller, unpublished data available on request).

> In the case of TraN, the binding site was determined as a 34 bp sequence which is located 69 bp upstream of $oriT_{pIP501}$ (Fig. 2*a*; Supplementary Fig. S4 shows an example of the data obtained with the new method: positions 1174–1207 on plasmid pIP501; GenBank L39769.1). Notably, the TraN binding site contains no direct or inverted repeats but is heavily biased towards A/T bases.

3.5. TraN DNA-binding mode and characterization

We investigated the thermal stability of TraN in the presence and absence of DNA using a Thermofluor-based assay. As depicted in Fig. 2(b), the melting temperature (T_m) of TraN alone

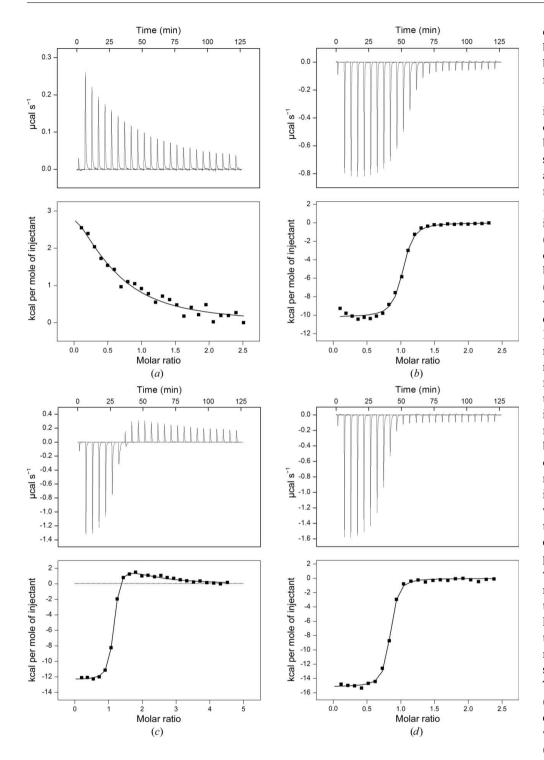


Figure 3

DNA-binding properties of TraN. Binding of TraN to a nonspecific 34-mer ds oligonucleotide (a) and the specific 34-mer ds binding site DNA with DNA as injectant (b) and with protein as injectant (c) were examined *via* ITC. The higher affinity of TraN for its specific binding site was tested in a competitive binding setup (d) in which protein was pre-incubated with nonspecific DNA and the specific DNA was used as the injectant.

amounted to 54.3 \pm 0.6°C; the binding of a nonspecific (random) 34-mer dsDNA oligonucleotide raised the $T_{\rm m}$ to 65.2 \pm 0.6°C, whereas addition of DNA containing the specific binding site increased the $T_{\rm m}$ to 70.4 \pm 0.2°C. The stabilizing

nonspecific DNA, respectively. We conclude that the second binding event of TraN to the specific DNA fragment is equivalent to the interaction of TraN with a random DNA sequence. To determine the difference in binding enthalpy

effect indicates an enhanced binding affinity for the specific binding site compared with the random DNA.

To determine whether there is a difference in the molar ratio of the TraN-DNA interaction between the random and the specific oligonucleotides, as well as to obtain information on the respective binding constants and ΔH values, we performed isothermal titration calorimetry (ITC) experiments with the oligonucleotide containing the binding site and the nonspecific (random) oligonucleotides which were used in the Thermofluor experiments (Fig. 3 and Table 1). In the titration setup with nonspecific DNA two TraN molecules bound to one dsDNA fragment (in a 2:1 ratio), whereas the stoichiometry for the specific interaction was equimolar (1:1 ratio) as expected for the specific binding site (Fig. 3b). We observed a mixed pattern of nonspecific and specific binding in the first steps of the titration with the fragment harbouring the binding site. As these effects could not be deconvoluted, we performed a reverse titration in which the DNA was in the measuring cell and the concentrated protein was added. The latter setup allowed a fit of a two-binding-site interaction model to the data. Owing to the significantly stronger binding of TraN to the specific binding site (exothermic reaction; binding constant of $10^7 M^{-1}$) compared with the nonspecific reaction (endothermic reaction; binding constant of $10^5 M^{-2}$ and a 2:1 binding ratio), it seems that the two sites are occupied consecutively. The resulting binding constants of the two calculated sites correspond well to the single titrations with specific and

Table 1
ITC-derived binding properties of TraN to nonspecific and specific 34-mer ds oligonucleotides.

			Reverse specific DNA		Competitive binding	
	Nonspecific DNA	Specific DNA	Specific binding	Nonspecific binding	Nonspecific versus specific DNA	
Molar ratio Binding constant† ΔH (cal mol ⁻¹)	$\begin{array}{c} 0.512 \pm 0.115 \\ 1.04 \times 10^5 \pm 3.02 \times 10^4 \\ 5380 \pm 1502 \end{array}$	$\begin{array}{c} 1.03 \pm 0.005 \\ 7.23 \times 10^6 \pm 7.72 \times 10^5 \\ -10500 \pm 108 \end{array}$	$\begin{array}{c} 1.08 \pm 0.004 \\ 3.38 \times 10^7 \pm 3.31 \times 10^6 \\ -12300 \pm 82 \end{array}$	$\begin{array}{c} 1.01 \pm 0.055 \\ 1.36 \times 10^5 \pm 2.46 \times 10^4 \\ 3811 \pm 262 \end{array}$	$\begin{array}{l} 0.797 \pm 0.004 \\ 1.45 \times 10^7 \pm 1.77 \times 10^6 \\ -15100 \pm 109 \end{array}$	

† The unit of the binding constant is M^{-1} for 1:1 stoichiometry and M^{-2} for 1:2 stoichiometry.

Table 2

Data-collection, phasing and refinement statistics.

Values in parentheses are for the highest resolution shell.

	Crystal form I	Crystal form II	Crystal form III
	Data 1	Data 2	Data 3
PDB code	4p0z	4p0y	4pm3
Data collection			
Beamline	X06DA (PXIII), SLS	X06DA (PXIII), SLS	X06DA (PXIII), SLS
Space group	$P2_1$	P21	P21
Detector	MAR CCD	MAR CCD	MAR CCD
Unit-cell parameters (Å, °)	a = 33.15, b = 55.09, $c = 35.52, \beta = 113.05$	a = 33.10, b = 62.82, $c = 55.87, \beta = 90.03$	a = 32.88, b = 54.94, $c = 57.71, \beta = 91.89$
Wavelength (Å)	0.9794	0.9794	1.0615
Resolution range (Å)	28.57-1.325 (1.40-1.33)	22.78-1.35 (1.48-1.40)	28.97-1.80 (1.90-1.80)
$R_{\text{meas}}(\%)$	9.9 (67.7)	9.1 (83.2)	14.1 (61.6)
$\langle I/\sigma(I) \rangle$	5.1 (1.1)	14.3 (2.7)	8.8 (3.7)
No. of molecules in asymmetric unit	1	2	2
Matthews coefficient ($Å^3 Da^{-1}$)	2.04	1.99	1.83
Solvent content (%)	39.9	38.2	32.89
Unique reflections	27168 (3914)	43358 (6179)	19191 (2804)
Measured reflections	199889 (27845)	333921 (47047)	69059 (10004)
Multiplicity	7.6 (7.1)	7.7 (7.6)	3.6 (3.6)
Completeness (%)	99.3 (98.3)	96.4 (94.2)	100.0 (100.0)
Refinement	99.5 (98.5)	90.4 (94.2)	100.0 (100.0)
Resolution range (Å)	32.68–1.35	21.35-1.40	28.97-1.80
	25731		
Unique reflections R	0.1685	43358 0.1833	19177 0.1693
R_{free}	0.2022	0.2233	0.2134
$R_{\rm free}$ test-set size (%)	5	5	5
No. of atoms	1011	10.10	1007
Protein	1014	1949	1886
Water	215	231	234
B factors $(Å^2)$			
Protein	14.96	14.73	15.15
Water	30.51	23.51	23.6
Pt			22.22 [2 atoms]
Br			43.72 [4 atoms]
Completeness (%)	99.4	96.1	99.95
Average B factor $(Å^2)$	17.72	15.70	16.14
R.m.s.d. from ideal values			
Bond lengths (Å)	0.025	0.024	0.007
Bond angles (°)	2.481	2.701	0.962
Ramachandran outliers	0	0	0
Ramachandran favoured (%)	98.33	99.12	99.10
Rotamer outliers	0/110	8/212	2/204
MolProbity score	0.96 [99rd percentile]	1.69 [65th percentile]	1.13 [99th percentile]

between the two modes of interaction, we performed a competitive binding experiment. Again, the values corresponded well to the results of the single titrations (Fig. 3d and Table 1).

3.6. TraN crystallization and data collection

Crystals of TraN were obtained as described in Goessweiner-Mohr *et al.* (2012), leading to two different crystal forms (using Index conditions 42 and 72). The integrity of TraN in the crystals was examined by matrix-assisted laser desorption/ionization time-of-flight (MALDI-TOF) mass spectrometry (MS), which showed that the protein in the crystals was significantly smaller than the original His-tagged construct (17.6 kDa). Two peaks of equal height with molecular masses of 14 222 and 14 478 Da, respectively (Fig. 1*b*) were visible, implying two cleavage sites with a distance of two amino acids. The His tag including the TEV cleavage site and linker amount to a molecular weight of

3213 Da (ExPASy; Gasteiger *et al.*, 2003), corresponding well to the difference between the full-length protein and the two species of TraN observed in the crystals. This indicates that the N-terminal tag of TraN is cleaved by *in situ* proteolysis during the crystallization process.

Details of the crystallization of selenomethionine-derivatized TraN as well as data collection, processing and refinement are described in §2.

3.7. The TraN crystal structure

Table 2 provides an overview of the data-collection, phasing and refinement statistics of the three data sets. Three different crystal forms with one (crystal form I) or two (crystal forms II and III) molecules per asymmetric unit were found. The N-terminal His tag and five residues at the C-terminal end of TraN were not observed in the electron-density map. This finding matches the loss of about 30 residues indicated by MS analysis of TraN crystals. Supplementary Fig. S5 provides an overview of the TraN construct as well as the predicted and actual secondary-structure content of the protein. The final coordinates and structure-factor amplitudes have been deposited in the PDB as entries 4p0z (crystal form I), 4p0y (crystal form II) and 4pm3 (crystal form III).

The crystal structure of TraN (amino acids 1–117) reveals an internal dimer fold (Fig. 4*a*) and shows an α – β composition with nine helices (h1–h9) and six β -strands (s1–s6). Structural

alignment of the two TraN halves yielded a nearly perfect superposition (Fig. 4b; Supplementary Table S1) except for the C-terminal helix h9, which is unique to the C-terminal half of TraN. The β -sheets of the halves (s1–3 and s4–6) form a β -barrel-like motif at the internal dimer axis. The nonpolar side chains of the following residues face towards the centre of this motif: Leu6 and Ile9 (h1), Ile46 (s2), Ile51 (s3), Leu63 (s4), Ile97 (s5) and Phe104 (s6). The side chains occupy the centre of the barrel-like structure, enabling a strong hydrophobic interaction that stabilizes the formation of the internal dimer (Fig. 4c).

Helices 3 and 7 are arranged in a parallel manner with a distance of approximately 33 Å (Fig. 5*a*). Furthermore, both helices are part of HTH motifs at the ends of TraN (h2 and h3, and h6 and h7, respectively) and each HTH motif together with the adjacent β -sheet resembles a winged-helix (WH) motif, which is a common variation of the HTH motif (Gajiwala & Burley, 2000). The distance and orientation of the HTH motifs is ideally suited for the recognition helices to enter two adjacent major grooves of a B-form dsDNA (Supplementary Fig. S6). In this scenario, the tip of the loops between β -strands 2 and 3 (residues Gly47 and Gly48) as well as strands 5 and 6 (residues Gly100 and Lys101) are positioned for a putative minor-groove interaction close to the internal twofold axis. Potential residues of the two HTH motifs that may be involved in the interaction with the major grooves of

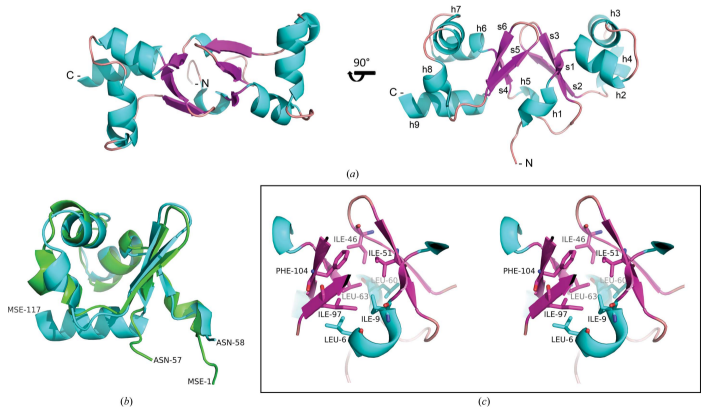


Figure 4

The structure of TraN_{1-117} . (a) Cartoon representation of TraN_{1-117} viewed from above and rotated by 90° about the threefold axis; secondary-structure elements are highlighted (helices in cyan, strands in purple) and labelled. (b) Structural alignment of the TraN internal halves (N-terminal half, green; C-terminal half, cyan); the respective start/end residues are labelled. (c) Wall-eyed three-dimensional representation of the hydrophobic residues facing towards the centre of the β -barrel-like motif in the centre of TraN.

DNA are Lys13, Asn22, Arg23, Asn24, Gln28, Arg31 and Asn32 of the N-terminal half and Lys67, Asn76, Asp77, Glu78, His82, Lys85 and Arg86 of the C-terminal half of TraN, respectively (Fig. 5b). We aligned the two halves of TraN, calculating the identity scores for the sequence and the secondary-structure composition, and compared the binding motifs. The sequence identity between the two halves is only about 30%, but the secondary-structure similarity is greater than 80% (Table S1). While the putative N-terminal binding motif (boxed amino acids in Fig. 5c) is mostly comprised of neutral or positively charged residues, the C-terminal motif contains two negatively charged residues, and five of the nine putative DNA-binding residues show non-conserved changes (Fig. 5c). This asymmetry in DNA-binding motifs suggests that different DNA half-sites should be recognized by the N- and C-terminal domains of TraN, which is in good agreement with the experimentally identified DNA-binding site of TraN, which actually does not contain inverted repeats.

3.8. TraN shows structural similarity to excisionases and the MerR family of transcriptional regulators

In searches for structurally similar proteins we only found hits that resemble one half of TraN. Consequently, we continued the search with only the larger C-terminal half of the protein. A closer examination of structural relatives (data not shown) revealed a relation to the structures of an excisionase from conjugative transposon Tn916 from E. faecalis (PDB entry 1y6u; Abbani et al., 2005) and an excisionase encoded by the Klebsiella pneumoniae genome (PDB entry 2kvv; Northeast Structural Genomics Consortium, unpublished work). Furthermore, excisionases from several bacteriophages such as phage λ (PDB entry 1rh6; Sam *et al.*, 2004) and phage HK022 (PDB entry 1pm6; Rogov et al., 2003) seem to share this fold. Excisionases (or Xis proteins) are required for the excision of bacteriophage DNA integrated in the genome of a host bacterium (Landy, 1989). Upon induction of a bacteriophage λ lysogen, the excision of the phage genome is performed via a site-specific recombination reaction (Sam et al., 2002). The Xis protein thus functions as a DNA architectural factor (Vargas & Landy, 1991; Abbani et al., 2007), which allows the integrase (Int) to switch to a second mode of action, namely the excision of the integrated DNA. Like bacteriophages, most of the time conjugative transposons are integrated into the bacterial host genome. To enable propagation, the integrated DNA is similarly excised from the host genome (Sam et al., 2002).

The TraN fold also resembles that of the N-terminal domain of transcriptional regulators (TR) of the MerR family, for

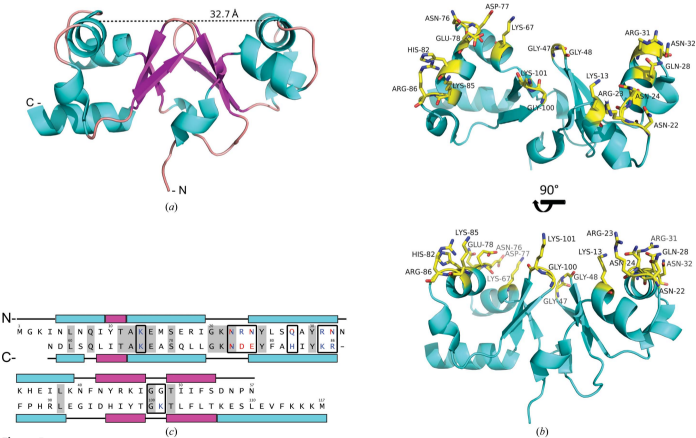


Figure 5

Structural basis for DNA binding of TraN. (a) Cartoon representation of TraN with the distance between the two parallel helices indicated; the length was measured in *Coot*. (b) Highlight of the TraN residues (yellow and labelled) putatively interacting with two succeeding major grooves of dsDNA viewed from above and rotated by 90° about the threefold axis. (c) Sequence and secondary-structure-based alignment of the two TraN internal halves [residues 1–57 (N-terminal half) and 58–117 (C-terminal half)]. Secondary-structure elements are highlighted (helices in cyan, β -strands in purple), identical amino acids are marked with a grey background and the putative DNA-binding residues are boxed and coloured according to their net charge.

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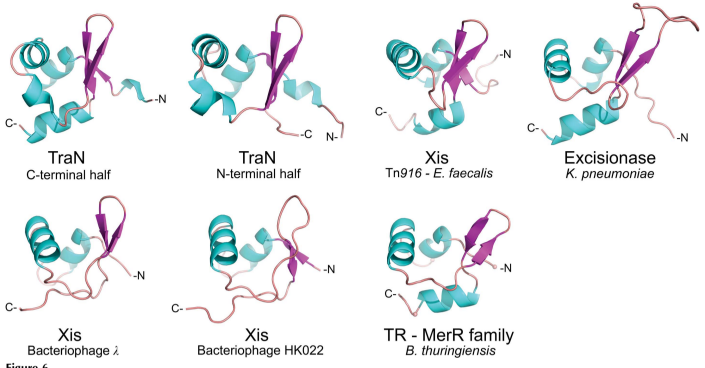


Figure 6

Structural comparison of TraN with itself and with related proteins. Cartoon representation of the TraN C-terminal and N-terminal halves, Xis from Tn916 from E. faecalis (PDB entry 1y6u), an excisionase from K. pneumoniae (PDB entry 2kvv), Xis from bacteriophage λ (1rh6), Xis from bacteriophage HK022 (PDB entry 1pm6) and a transcriptional regulator (TR) of the MerR family from B. thuringiensis (PDB entry 3gpv); secondarystructure elements are highlighted (helices in cyan, β -sheets in magenta).

example a transcriptional activator from Bacillus thuringiensis (PDB entry 3gpv; New York SGX Research Center for Structural Genomics, unpublished work). Transcriptional activators of the MerR family feature an N-terminal WH DNA-binding region and recognize the specific DNA site as a dimer where the recognition helices of the HTH motifs are inserted into two adjacent major grooves. The dimerization motif of MerR proteins is a coiled-coil arrangement between the C-terminal parts of two MerR proteins (Supplementary Fig. S6), which is completely different from the internal dimer configuration of TraN, which is based on hydrophobic interactions within a β -barrel-like motif in its centre. It is interesting to note that the three-stranded β -sheet of this motif corresponds to the 'wing' of the WH motif of MerR, which in the protein-DNA complexes of MerR family members is inserted into the minor grooves adjacent to the major groove-embedded recognition helices (Newberry & Brennan, 2004). In contrast to the MerR family proteins, which feature a C-terminal effector-binding region (Brown et al., 2003), neither TraN nor the TraN-like proteins of related T4SSs (see below) contain such a C-terminal extension.

Despite the low sequence identity of 8-13% between the TraN C-terminal domain and the respective domains of the compared proteins (Supplementary Table S1), the fold of all aligned proteins is highly conserved (Fig. 6). A distinct feature of the TraN fold is a short helix adjacent to the putative DNArecognition helix (h4 next to h3 in the N-terminal half and h8 next to h7 in the C-terminal half of TraN; Fig. 4a). This feature is not observed in the excisionase proteins or in the MerR family proteins (Fig. 6), suggesting that TraN-like proteins form a unique subgroup of the WH proteins.

We performed an extended search for TraN-like proteins (i.e. containing the internal dimer arrangement of the two HTH domains) in a broad spectrum of conjugative plasmids, transposons, ICEs and GIs from G- and G+ bacteria. All TraN-like proteins found are of enterococcal origin (Supplementary Fig. S7) and show a high sequence identity to TraN. All other proteins identified (excisionases and proteins belonging to the MerR family) contained only a single TraNlike domain.

4. Discussion

The research field of conjugative DNA transport has attracted increasing attention over the last decades. This is largely owing to its enormous importance in human healthcare: conjugative transfer greatly increases prokaryotic genome plasticity and has been identified as a major vehicle for the spread of antibiotic resistance among pathogens and commensal bacteria (Zechner et al., 2012).

The structural elucidation of the core complex of the pKM101 T4SS from E. coli by electron microscopy and X-ray crystallography was the most important recent contribution to the understanding of the assembly and core-complex architecture of the G- conjugation apparatus (Chandran et al., 2009; Fronzes et al., 2009; Rivera-Calzada et al., 2013). Unfortunately, less progress has been achieved for the corresponding systems of G+ origin, in spite of the occurrence of major multi-resistant human pathogens such as enterococci, streptococci or staphylococci in this group of bacteria (Burns, 2003). Only very recently has structural information become available for G+ T4SSs: the structure of the VirB8-like transfer protein TcpC from *C. perfringens* plasmid pCW3 (Porter *et al.*, 2012) was reported in early 2012, followed by the structure of VirB4 from *T. pseudethanolicus* (Walldén *et al.*, 2012), the structure of the VirB8-like protein TraM from *E. faecalis* conjugative plasmid pIP501 (Goessweiner-Mohr, Grumet *et al.*, 2013) and, most recently, the 3.0 Å resolution structure of the TraK_{pIP501} extracellular domain (Goessweiner-Mohr *et al.*, 2014).

In this study, structural and biophysical approaches were used to characterize TraN, a putative T4SS protein from the *E. faecalis* conjugative model plasmid pIP501. It is the third structure to be reported of a component of the pIP501 T4SS. TraN localizes to the cytoplasm and was found to bind preferentially to dsDNA, which points to a role in DNA processing prior to the actual transport of the single-stranded plasmid to the recipient cell. Based on the structural resemblance of TraN to the DNA-binding domains of prokaryotic transcription regulators, we predicted that TraN would be a specific DNA-binding protein and we were finally able to identify a 34 bp sequence upstream of $oriT_{pIP501}$ as the high-affinity binding site. These findings suggest several possible roles for TraN in the pIP501 transfer process.

Firstly, TraN may act as an accessory protein for the pIP501encoded relaxase TraA. The role of accessory proteins is well understood in G- T4S: detailed overviews of the conjugative DNA metabolism in G-bacteria have been published by de la Cruz et al. (2010) as well as by Zechner et al. (2012). In the case of IncF plasmids, TraY is responsible for the bending of DNA (Luo et al., 1994) and the subsequent relaxase binding and nicking in preparation for conjugative DNA transfer (Inamoto et al., 1994; Howard, 1995; Karl et al., 2001). A second protein, TraM, binds the DNA as a tetramer at preferred sites close to the oriT (Verdino et al., 1999; Wong et al., 2011). It was shown to promote relaxosome formation and to stimulate the niccleavage and oriT DNA-unwinding reactions of TraI (Ragonese et al., 2007; Sut et al., 2009). Moreover, it was found to promote interactions between the coupling protein TraD and the relaxosome (Beranek et al., 2004; Lu et al., 2008).

Our data suggest a functional role for $\text{TraN}_{\text{pIP501}}$ similar to that of the IncF auxiliary proteins TraM and TraY. This hypothesis is based on the preferred binding of TraN upstream of the pIP501 *oriT* DNA sequence, which implies an involvement in the activity of the relaxase TraA at the *nic* site. The two-domain HTH-like fold of the TraN crystal structure suggests a DNA-binding mode in which two adjacent major grooves of DNA are occupied, possibly including the bending or local unwinding of the plasmid DNA which are supposedly necessary for relaxase activity *in vivo* (Zechner *et al.*, 2012). Furthermore, TraN could be involved in connecting the relaxosome complex to proteins of the pIP501 secretion machinery. TraN has been reported to interact with the ATPase TraE, the muramidase TraG, the membrane-associated protein TraH and the polytopic protein TraL, as shown by yeast two-hybrid and pull-down assays (Abajy *et al.*, 2007) as well as with the coupling protein TraJ (Goessweiner-Mohr *et al.*, unpublished work). The latter resembles the interaction of the IncF auxiliary protein TraM with the respective coupling protein TraD (Beranek *et al.*, 2004; Lu *et al.*, 2008). However, we could not prove an interaction between TraN and the relaxase TraA using a yeast two-hybrid system (Abajy *et al.*, 2007).

Secondly, and surprisingly, TraN shows structural similarities to excisionases from bacteriophages and from the conjugative transposon Tn916. Via the bending of DNA and interaction with the integrase, the excisionase is responsible for a switch in the mode of operation of the integrase which leads to the excision of the mobile genetic element (Calì et al., 2004; Singh et al., 2014). A similar function for TraN is unlikely, since (i) pIP501 is not integrated in the chromosome and (ii) the excisionases only resemble the fold of one TraN domain. The proposed binding for a single TraN domain involves the interaction of the HTH motif with the major groove as well as of the loop between the two β -strands with the minor groove. Although this mode of binding is very similar to the observed mode of interaction of excisionases with their target sequence (Sam et al., 2002), the symmetric arrangement of the two HTH domains of TraN demands a distinctly different mode of DNA interaction.

Nevertheless, TraN could have an excisionase-like function by carrying out a role in altering the DNA architecture and stimulating relaxase activity. It could be proposed that TraN causes changes in the DNA structure at the oriT site on an already assembled TraA-DNA complex to stimulate cleavage of the nic site, similar to the mode of function of excisionases (Ramsay et al., 2006). This model is distinct from that for other relaxase accessory proteins described above, but is supported by our previous and current work and by the genetic organization of the traA-traO operon. The following model could be suggested. (i) Transcription of the tra operon initiates from the promoter upstream of traA and TraA immediately binds the oriT and represses further expression of the operon (Kurenbach et al., 2006), but is not able to immediately catalyze nicking of the oriT. (ii) Since TraN is encoded by the secondlast gene in the operon (orf14), nearly all putative mating pore proteins are translated before TraN. (iii) TraN is finally expressed, binds and bends the DNA upstream of the TraA-DNA complex and thereby stimulates nicking and subsequent transfer of the pIP501 plasmid.

A third possible function of TraN concerns its structural relation to transcriptional regulators of the MerR family. This class of transcriptional regulators have been reported to be involved in numerous metal-dependent stress-response mechanisms (Hobman, 2007; Kidd *et al.*, 2011; McEwan *et al.*, 2011; Liao & Sauer, 2012). Moreover, MerR-like regulators have also been reported to be involved in bacterial virulence (Stroeher *et al.*, 2007). Similar to the excisionases, MerR-like proteins only resemble the fold of a single TraN domain, but their mode of binding to DNA involves the formation of a homodimer (PDB entry 3gpv) which binds to two adjacent

major grooves of dsDNA, as proposed for TraN. Expression of the pIP501 transfer genes is already under auto-regulation by TraA relaxase (Kurenbach et al., 2006), thus TraN might fulfil a different role in the control of pIP501 plasmid transfer. We suggest that TraN may work as a transfer regulator by preventing the pIP501 plasmid DNA from being nicked by the relaxase without the establishment of cell-to-cell contact and formation of the conjugative core complex. Interestingly, TraN has been shown to interact with several key components of the pIP501 T4SS, despite being localized in the cytoplasm. Through these interactions, for example with the ATPase TraE, the muramidase TraG and the coupling protein TraJ, TraN could be prompted to release the plasmid DNA and would subsequently allow the relaxase to nick the plasmid DNA at the *oriT* site. The proposed mechanism would ensure that the relaxase-dependent processing of the conjugative plasmid is only started once the transfer apparatus has been assembled. A major question remains concerning the late expression of the TraN protein as the second-last gene in the polycistronic mRNA of the transfer region. This raises the question whether TraN is indeed capable of playing the proposed crucial regulatory role in the early steps of the transfer process.

TraN-like proteins are rarely found among T4SSs. In fact, only five other proteins with an equivalent internal dimer fold have been identified to date, all of which are of enterococcal origin (Supplementary Fig. S7). These findings imply that the corresponding conjugative systems might all have developed from an ancestral G+ T4SS and that the separation into distinct conjugative plasmids and ICEs has happened relatively recently. An explanation for the rare presence of TraNlike proteins in T4SSs could be the host range of the respective T4SSs. pIP501 has the broadest host range of known G+ T4SSs and has been even shown to stably replicate in bacteria of G- origin (Kurenbach et al., 2003). Consequently, TraNlike proteins might play an important role in the adaptation of their respective T4SSs to new hosts. This adaptation probably involves the regulation or the adjustment of the specific processing of the conjugative plasmid DNA.

The structure of TraN and the elucidation of its DNAbinding properties are milestones in the growing understanding of G+T4S. Nevertheless, further efforts are needed to decipher the specific function of TraN-like proteins.

5. Related literature

The following reference is cited in the Supporting Information for this article: van Dijk & Bonvin (2009).

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