The MobM relaxase domain of plasmid pMV158: thermal stability and activity upon Mn$^{2+}$ and specific DNA binding

Fabian Lorenzo-Diaz¹, Lubomir Dostal², Miquel Coli³, Joel F. Schildbach², Margarita Menendez⁵, and Manuel Espinosa¹,*

¹Centro de Investigaciones Biológicas, CSIC, Ramiro de Maeztu 9, 28040 Madrid, Spain, ²Department of Biology, Johns Hopkins University, 3400 North Charles Street, Baltimore, MD 21218, USA, ³Institut de Biologia Molecular de Barcelona, CSIC, ⁴Institute for Research in Biomedicine, Barcelona Science Park, Badiri Reixac 10, 08028 Barcelona and ⁵Instituto de Química Física Rocasolano, CSIC and CIBER of Respiratory Diseases (CIBERES), Serrano 119, 28006 Madrid, Spain

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ABSTRACT

Protein MobM, the relaxase involved in conjugative transfer of the streptococcal plasmid pMV158, is the prototype of the MOBV superfamily of relaxases. To characterize the DNA-binding and nicking domain of MobM, a truncated version of the protein (MobMN199) encompassing its N-terminal region was designed and the protein was purified. MobMN199 was monomeric in contrast to the dimeric form of the full-length protein, but it kept its nicking activity on pMV158 DNA. The optimal relaxase activity was dependent on Mn$^{2+}$ or Mg$^{2+}$ cations in a dosage-dependent manner. However, whereas Mn$^{2+}$ strongly stabilized MobMN199 against thermal denaturation, no protective effect was observed for Mg$^{2+}$. Furthermore, MobMN199 exhibited a high affinity binding for Mn$^{2+}$ but not for Mg$^{2+}$. We also examined the binding-specificity and affinity of MobMN199 for several substrates of single-stranded DNA encompassing the pMV158 origin of transfer (oriT). The minimal oriT was delimited to a stretch of 26 nt which included an inverted repeat located eight bases upstream of the nick site. The structure of MobMN199 was strongly stabilized by binding to the defined target DNA, indicating the formation of a tight protein–DNA complex. We demonstrate that the oriT recognition by MobMN199 was highly specific and suggest that this protein most probably employs Mn$^{2+}$ during pMV158 transfer.

INTRODUCTION

Bacterial plasmids are able to disseminate multiple resistance genes by conjugation. This horizontal gene transfer (HGT) is relevant because of the increase in infectious diseases provoked by dissemination of antibiotic-resistant bacterial pathogens (1,2). Among them, the Gram-positive (G+) bacterium Streptococcus pneumoniae (the pneumococcus) is responsible for nearly 2 000 000 human deaths annually, and this figure represents only 15–20% of the people infected (3). Thus, understanding HGT in pneumococcus and related bacteria may help in controlling the spread of medically important antibiotic resistance (4).

Conjugation between Gram-negative (G−) bacteria is a well-characterized HGT process (5). The mechanism involves the assembly of a stable multi-protein complex (the relaxosome) on a plasmid region termed origin of transfer (oriT), and the key player is a plasmid-encoded nicking-closing enzyme, the DNA relaxase (5). Initiation of transfer requires cleavage of the phosphodiester bond at a specific position within the oriT (the nick site) mediated by a Tyr residue of the relaxase, so that a covalent tyrosinyl-DNA adduct is formed (6,7). The cleaving reaction requires the target DNA to be in single-stranded configuration (ssDNA). Thus, like almost all cation-dependent nucleotidyl-transferase enzymes, relaxases leave a free 3′-OH end while remaining bound to the 5′-phosphate product. This complex is actively pumped to the recipient cell by a plasmid-encoded coupling protein and the transferosome, a type IV secretion system (8,9). Once in the recipient, the relaxase–ssDNA intermediate restores the original circular plasmid molecule after termination of transfer by means
of a reversion of the strand transfer reaction. Consequently, termination of DNA transfer resembles termination of rolling circle replication (RCR) (10). Finally, conversion of ssDNA molecules into double-stranded (ds) plasmids in the recipient cells is carried out by conjugative replication (11,12). Whereas self-transmissible plasmids encode the entire machinery for their transfer, many small plasmids only harbour an oriT and the relaxase gene (13,14). This second group, known as mobilizable plasmids, can be transferred when they co-reside in the cell with an auxiliary self-transmissible element that supply the coupling protein and the transferosome (14).

In spite of the knowledge accumulated during the past years on the structure and function of conjugative relaxases encoded by plasmids of G-bacteria (8), information on equivalent proteins encoded by plasmids from G+ bacteria is still scarce and limited to a small number of proteins, such as TraA_pIP501 (15,16), Mob proteins of pC221 (17,18) and MobM, encoded by the mobilizable plasmid pMV158 (19,20). In general, the C-terminal moiety of relaxases contains a DNA-helicase or primase domain and/or other functions related to membrane association and to protein–protein interactions, whereas the N-terminal moiety harbours the endonuclease (DNA nicking and closing) and DNA-binding domain. In many cases, the endonuclease domain exhibits, apart from the catalytic Tyr residue, a His-triad termed ‘3H motif’ (HxDxHxHxH, with ‘x’ being any residue and ‘u’ a hydrophobic residue) which is involved in the coordination of a single divalent cation required for DNA cleavage (21). In the architecture of the N-terminal nuclease domains of the four conjugative relaxases whose structures have been solved, this cation was shown to be Mg$^{2+}$ or Mn$^{2+}$ in TraI_F (6), Ni$^{2+}$, Cu$^{2+}$, or Zn$^{2+}$ in TrwC_R388 (7), and Mn$^{2+}$ in MobA_R1162 (22) and TraI_pCU1 (23).

With respect to the DNA-binding activity, it has been demonstrated that the interactions between the relaxase and the oriT are sequence- and structure-specific. Plasmid oriTs are complex DNA regions that contain inverted repeats (IR) and A+T-rich tracts. In the plasmids R388 and F, both belonging to MOB_F family (24), an IR is located 8 and 9 bp upstream to the nick site, respectively. Binding analyses using different fluorescently-labelled DNA fragments suggested that the relaxase domain of TraI_F could bind to its oriT in two distinct manners with different sequence specificities (25). Nevertheless, in vitro TraI protein-binding affinities for several ssDNA mutants and in vivo transfer efficiencies of F plasmid derivatives did not always correlate (26). These latter results suggest that the essential function of some bases in the oriT may be to position the scissile phosphate bond for cleavage, rather than to directly contribute to binding affinity. In the TrwC relaxase-ssDNA structure, the IR forms a hairpin and several protein–DNA contacts are established (7). This conformation was reported to be important for the termination reaction in the recipient cell (27,28).

The MobM protein has been considered representative of a family of relaxases termed MOB_F, which encompasses nearly 100 members (24). However, little information on biochemical and structural parameters of MOB_F relaxases has been provided. For that reason, the objective of this work was to characterize the N-terminal relaxase domain of protein MobM. Full length MobM has 494 residues. We have cloned a region of the mobM gene encoding the first 199 N-terminal residues (MobMN199). MobMN199 was purified as a native protein using a new protocol involving an affinity chromatography followed by a gel filtration step. In contrast to the dimeric form of the full length protein, the purified MobMN199 was a monomer in solution, but kept its nicking activity on supercoiled pMV158 DNA. Its secondary structure was determined and thermal denaturation studies showed that structural stability of MobMN199 strongly depended on Mn$^{2+}$ and on DNA binding. Although Mg$^{2+}$ ions could replace Mn$^{2+}$ for DNA cleavage, the former did not stabilize the MobMN199 structure, at least in the absence of its DNA substrate. MobMN199 binding affinities for Mn$^{2+}$ and for several oligonucleotides containing the oriT sequence were also measured. The ssDNA-binding experiments allowed us to define a minimal oriT sequence that contains an IR upstream the nick site. The results presented here demonstrate that the oriT recognition by the MobM relaxase is highly specific and suggest that this protein most probably employs Mn$^{2+}$ during pMV158 plasmid transfer.

MATERIALS AND METHODS

**Bacterial strains, plasmids and oligonucleotides**

*Escherichia coli* strains were grown on TY (Pronadisa, Spain) medium supplemented with 30 μg/ml kanamycin or 100 μg/ml ampicillin. *E. coli* BL21(DE3) was used for expression of MobM and MobMN199. The plasmid used for controlled MobM expression was pLGM2 (based on pET5 vector) (20). Plasmid used for cloning the *mobM* variant *mobMN199* was pET24(b) (Novagen). Purified pMV158 DNA was prepared from the *S. pneumoniae* 708 strain (29) by two consecutive CsCl gradients as described (30).

Oligonucleotides used for binding studies were synthesized and HPLC-purified by the Integrated DNA Technologies (Coralville, IA, USA); ORIT 42-mer GCA CACACTTTATGAATATAAAGTATGTG/TTAT ACTTTA; IR1 18-mer ACTTTATGAATATAAAGT; IR2 24-mer TAAAGTAGGT/TTATATCTTTA; IR3 32-mer GCACACACTTTATGAAATATAAAGT AGTGTG/; IR1+8 26-mer ACTTTATGAATATAAAG TATAGTG; IR1-R 11-mer GAATATAAAGT; IR3-L 15-mer GCACACACATTTATGAATATAAAGT AGTGTG/; 10-NIC 10-mer GTATAGTG/; where ‘?’ denotes the oriT nick site. The IR3, IR2 and ORIT oligonucleotides were also purchased with the Cy5 fluorophore label.

**DNA amplification and cloning**

To obtain plasmid pMobMN199, a 630-bp fragment that encodes the first 199 residues of MobM protein was amplified by PCR on pMV158 DNA (accession number X15669) with the following primers: NdeI-F (5’-AAGGA...
GGGAAACATATGAGTTACA-3′; coordinates 3718–3741) and XhoI-R/STOP (5′-GAAGTTCTCTCTCGAG TTACATATCGCCA-3′; coordinates 4347–4318). To facilitate cloning, forward and reverse primers included changes (underlined) to generate recognition sites for NdeI and XhoI restriction enzymes, respectively. Furthermore, a single change (boldface) was also introduced in the reverse primer to get an ochre stop codon instead of Glu200 (E200Stop). PCR amplifications were done in 50-μl reaction mixtures containing reaction buffer 1× [16 mM (NH₄)₂SO₄, 67 mM Tris–HCl (pH 8.8), 1.5 mM MgCl₂], 0.2 mM of each dNTP (Roche), 0.4 μM of each primer, 0.65 U of Phusion DNA polymerase (Finnzymes) and 1 ng of DNA template, under the following conditions: an initial denaturing step at 98°C (30 s); 30 cycles of denaturation at 98°C (10 s), annealing at 55°C (20 s) and extension at 72°C (30 s) followed by a final extension at 72°C (10 min). PCR products were purified with the QIAquick gel extraction kit (Qiagen) and after digestion with NdeI and XhoI enzymes, the fragments were ligated into pET24(b) previously digested with the same enzymes. The constructs were confirmed by nucleotide sequencing. Escherichia coli BL21(DE3) cells harbouring the desired plasmid were used for protein expression.

**Protein purification and N-terminal sequencing**

MobM and MobMN199 proteins were overproduced and purified by a newly developed protocol, with greater speed and higher yield than the one previously described for MobM (19,20). Briefly, cells were grown in 41 of TY-Km at 37°C to reach an OD₆₀₀ = 0.5. Expression of the plasmid-encoded genes was achieved by induction with 1 mM IPTG (30 min), followed by addition of rifampicin (200 μg/ml) and growth for an additional 90 min. Cells were sedimented by centrifugation (8000 rpm, 20 min, 4°C) and stored at −80°C. The cell pellet was thawed and resuspended (100× concentrated) in buffer A [20 mM Tris–HCl pH 7.6, 1 mM EDTA, 1 mM dithiothreitol, 5% (v/v) glycerol] plus 1 M NaCl and a tablet of protease inhibitor cocktail (Complete, Roche). Cells were then lysed by passage through a French pressure cell and the extract was centrifuged to remove cell debris. The clarified extract was treated with 0.2% (v/v) polyethyleneimine (Sigma) to precipitate nucleic acids, and proteins in the supernatant were precipitated at 70% (w/v) ammonium sulphate saturation. The proteins in the precipitate were collected by centrifugation and dissolved in buffer A with 300 mM NaCl. After equilibration in the same buffer by dialysis, the sample was loaded onto a 100 ml heparin–agarose (BioRad) column (flow rate of 50 ml/h). After washing with 5-column volumes of the same buffer, a 400 ml 0.3–0.8 M NaCl gradient was applied to elute the proteins retained. Fractions were analysed by 15% SDS–Tris–glycine polyacrylamide (PAAG) gel electrophoresis followed by staining with Bio-safe Coomassie (BioRad Laboratories). Fractions containing the peak of MobMN199 were pooled, dialysed against buffer A containing 500 mM NaCl, and concentrated by filtering through 3 kDa cut-off membranes (Pall) until the sample volume reached 1 ml. The protein sample was injected at 0.5 ml/min onto a HiLoad Superdex 200 gel-filtration column (Amersham) and subjected to fast-pressure liquid chromatography (FPLC; Biologic DuoFlow from BioRad). OD₃₀₀ values were monitored continuously and each recovered fraction (2 ml) was analysed as above. Fractions containing pure MobMN199 protein (>98%) were pooled and concentrated until the final concentration was 5 mg/ml protein and stored at −80°C. In these conditions, the protein retained full activity for at least 1 year. Edman’s sequential degradation was applied to analyse the N-terminal sequence of MobMN199 in a Procise 494 protein sequencer (Perkin Elmer) as reported (20). Concentration of MobMN199 protein was determined by spectrophotometric determinations and by determination of the amino acid composition of the protein (CIB-Protein Chemistry Service).

**Supercoiled DNA relaxation assays**

Nicking of supercoiled DNA by purified MobM or MobMN199 was performed essentially as reported (20). Standard reaction mixtures (20 μl) contained supercoiled pMV158 DNA (300 ng) in buffer N [20 mM Tris–HCl pH 7.6, 200 mM NaCl, 0.05 mM EDTA, 1% (w/v) glycerol, 1 mM dithiothreitol] to which different concentrations of divalent cations and purified protein were added. Samples were incubated at 30°C for 20 min, and reactions were stopped by addition of SDS (0.5%) and Proteinase K (100 μg/ml), followed by 30 min incubation. Generation of open circular forms (FII) by the nicking activity of MobMN199 was monitored by electrophoresis on 1.2% (w/v) agarose gels and quantified as described (20).

**Analytical ultracentrifugation**

Sedimentation equilibrium experiments were performed at 20°C in an Optima XL-A (Beckman-Coulter) analytical ultracentrifuge. A range of MobMN199 concentrations, from 2 to 20 μM protein, were examined in buffer UA [20 mM Tris–HCl pH 7.6, 500 mM NaCl, 1 mM EDTA, 1% (v/v) glycerol]. Samples (100 μl) were centrifuged at two successive velocities (20,000 and 35,000 rpm) and absorbance was measured at 3 h intervals to determine that samples had reached equilibrium. The equilibrium scans were monitored at the most appropriate wavelength (230, 280 or 290 nm), depending upon the MobMN199 concentration employed. Baseline signals were measured after high-speed centriufugation. Apparent average molecular masses (Mₓ,app) of MobMN199 were obtained using the program Heteroanalysis (www.biotech.uconn.edu). The partial specific volume of MobMN199, 0.723 ml/g, was calculated from its amino acid composition with the programme SEDNTERP (31). Sedimentation velocity experiments were performed at 48,000 rpm and 20°C with 400 μl samples of MobMN199 (2–20 μM, in buffer UA) that were loaded into double-sector cells. The sedimentation coefficient distribution for MobMN199 was calculated by least-squares boundary modelling of sedimentation velocity data using the sedimentation coefficient distribution method, as implemented in the SEDFIT program (32). The coefficient was corrected to standard conditions to get the corresponding s₂₀,w value using the SEDNTERP
program. The translational frictional coefficient of MobMN199 ($f$) was determined from the molecular mass and sedimentation coefficient of the protein (33), whereas the frictional coefficient of the equivalent hydrated sphere ($f_0$) was estimated using a hydration of 0.4506 g H$_2$O per g protein (34).

**Sequence-based secondary structure prediction**

Secondary-structure predictions of MobMN199 were carried out with programs PSIPRED (35), Jpred (36), SABLE (37), NPS@ (38) and PredictProtein (39) (http://www.expasy.ch/). The average number of residues involved in alpha helix, extended strand or random coil was calculated, and a secondary structure model of MobMN199 was constructed.

**Circular dichroism**

CD spectra were acquired from 185 to 260 nm in a J-710 spectropolarimeter (Jasco Corp.) fitted with a Peltier temperature control accessory using 0.02–0.1 cm optical path length quartz cells. Each spectrum was the average of four cumulative measurements, performed automatically, that were recorded at a scanning speed of 20 nm/min and 1 nm spectral bandwidth. Measurements were performed at 15 μM MobMN199 and the spectra were corrected by subtracting the buffer contribution. To obtain structural information, spectra were acquired at 10°C in buffer CD1 (20 mM sodium phosphate, pH 7.6, 50 mM ammonium sulphate), converted to mean residue ellipticity ($\Theta_{MRW}$) or to differences in molar absorbance ($\Delta\varepsilon$) and then deconvolved using CONTINLL (40) and SELCON3 (41) algorithms, with the reference data set #6, available at DichroWeb site (42). To analyse the temperature-associated changes in secondary structure with different Mn$^{2+}$ or Mg$^{2+}$ concentrations (‘Results’ section), MobMN199 was dialysed against buffer CD2 (20 mM HEPES pH 7.6, 50 mM ammonium sulphate), unless otherwise stated, and changes in ellipticity at 218 nm were recorded in 0.1 cm optical path-length quartz cells while increasing the temperature from 10 to 90°C at two different rates (20 and 50°C/h). In addition, CD spectra obtained from 205 to 260 nm were recorded between 10 and 90°C with temperature increments of 5°C and equilibration times of 1 min before acquiring each spectrum. Finally, the sample was cooled again to the initial temperature (10°C) and the protein spectrum was recorded under renaturing conditions. Data acquisition and processing were carried out using the Jasco Spectra-Manager software, and thermal-denaturation profiles were analysed in terms of two-state transitions [Equation (1)] using the Origin v6.0 software (Microcal Inc.).

$$\Delta\Theta(T)/\Delta\Theta_{\text{max}} = \frac{1}{\sum_{i=1}^{2} f_i \left[ \exp\left[-\Delta H_i(T_{mi} - T)/R \times T_{mi} \times T \right]/\left[1 + \exp\left[-\Delta H_i(T_{mi} - T)/R \times T_{mi} \times T \right]\right] \right]}$$

where $\Delta\Theta(T)$ is the ellipticity change at 218 nm at temperature $T$, $\Delta\Theta_{\text{max}}$ the maximum variation in the ellipticity at this wavelength, $R$ the gas constant, $f_i$ the relative contribution of transition $i$ to $\Delta\Theta_{\text{max}}$, and $T_{mi}$ and $\Delta H_i$ are the half-transition temperature and the enthalpy change of this transition, respectively.

**Calorimetric studies**

 Isothermal titration calorimetry (ITC) was performed using a MicroCal MCS-ITC calorimeter (Microcal Inc.). Before measurements, MobMN199 samples were dialysed against buffer ITC (20 mM HEPES, 400 mM NaCl, pH 7.6), which was also used to prepare the cation solutions. The protein was loaded into the calorimetric cell at a concentration of 75 μM and titrated by stepwise injection of 1.5 mM MnCl$_2$ or 20 mM MgCl$_2$ solutions. Typically, twenty 3–8 μl injections were performed while stirring at 300 rpm, and the heat of the dilution of the ligand was determined in separate runs. The experiments were carried out at 17°C and the binding isotherms were analysed using the Microcal Origin ITC software package. Differential scanning calorimetry (DSC) measurements were performed at a heating rate of 50°C/h in a VP-DSC microcalorimeter (Microcal Inc.), at a constant pressure of 2 atm. MobMN199 was equilibrated in ITC buffer supplemented with the required Mn$^{2+}$ concentration. Experiments performed in the presence of DNA (oligonucleotide IR1-8) were carried out at a MobMN199/DNA ratio of 2.0 using protein concentrations of 40 and 20 μM. Microcal Origin DSC software was used for data acquisition and analysis. Excess heat capacity functions were obtained after subtraction of the buffer–buffer base line and transformed into molar heat capacities dividing by the number of moles of MobMN199 in the DSC cell.

**DNA-binding affinity measurements**

 Titration of MobMN199 with 5'- or 3' labelled (Cy5) oligonucleotides was performed by electrophoretic mobility shift assays (EMSA), as reported (43). Purified MobMN199 was mixed, at different concentrations, with Cy5-labelled DNA (2 μM) in 10 μl of buffer A with 300 mM NaCl. Reaction mixtures were incubated at 25°C for 30 min. Free and bound DNAs were separated by electrophoresis on native 10% PAA gels. Fluorescent DNA bands were detected with a Typhoon scanner system (Molecular Dynamics) and quantified with QuantityOne software (Bio-Rad). Competitive EMSA were used to determine the relative affinities of MobMN199 for different DNA sequences. Mixtures of Cy5'-IR3 oligonucleotide (2 nM) and DNA fragments at differing concentrations were incubated simultaneously with 80 nM MobMN199 protein. The samples were incubated at 25°C for 30 min, loaded on native 10% PAA gels and the DNA species were quantified as above.

**RESULTS**

**Characterization of the MobM relaxase domain**

The full-length MobM protein has a predicted size of 57 874 Da (494 residues) and exhibits three conserved
motifs located in the N-terminal moiety (Figure 1a): (i) motif I (HxxR), of unknown function; (ii) motif II (NYD/EL), which contains the putative catalytic tyrosine; and (iii) motif III (HxDExxPHuH), also known as the 3H motif, probably involved in coordination of a divalent metal (13). To uncouple the N- and C-terminal domains of MobM, a strategy to obtain a protein containing only the first 199 N-terminal residues (MobMN199) was designed. The strategy was based on alignment of the relaxases belonging to the MOBV family (24), of which pMV158 is the prototype, revealed that roughly the first 200 amino acids were highly conserved among all the relaxases. Inspection of the DNA sequence around this region indicated that generation of a stop codon after the first 199 residues was relatively simple. Thus, the 5'-region of the pMV158 mobM gene (encoding the first 199 residues followed by a stop codon) was cloned into a suitable expression vector. To purify MobMN199, the protocol employed to obtain the full-length MobM protein (19) was improved by introducing one step of mechanical shearing (covered by lane 7). Fractions containing the peak of MobMN199 were pooled, dialysed against buffer A and concentrated. M indicates the molecular weight standards (in kDa). (c) Purified MobMN199 was injected onto a gel filtration column, and its elution profile was recorded; the inset shows the SDS–PAGE gel with the purified protein and the molecular size markers. (d) Relaxation assays with wild-type MobM and the MobMN199 protein. Supercoiled pMV158 DNA samples (300 ng; 8 nM) were incubated with and without (−) full-length MobM (left) or with the short MobMN199 fragment (right) in the presence of 15 mM MnCl2 at 30°C, 20 min. Protein concentrations used were 120, 240 and 480 nM. Generation of relaxed forms (FII) from supercoiled DNA forms (FI) was tested at several protein concentrations (from 120 to 480 nM) and reactions were performed in the presence of 15 mM MnCl2.

The ability of MobMN199 to convert supercoiled DNA (form FI) into open circular species (form FII) was tested at several protein concentrations (from 120 to 480 nM) and reactions were performed in the presence of 15 mM MnCl2. The samples were incubated at 30°C for 20 min, as reported for the full-length protein (19,20). The results showed that MobMN199 was able to relax pMV158 DNA with the same efficiency as the entire MobM,
converting up to 60% of supercoiled plasmid into form FII (Figure 1d). The failure to reach 100% cleavage could be explained as the result of the equilibrium between the nicking and closing activities of MobMN199; a behaviour observed for several relaxases (44–46). Selective precipitation by KCl and SDS of FII forms cleaved by a relaxase is the procedure used to detect relaxosome formation (46). We have previously shown that the full-length MobM protein was able to form stable complexes with pMV158 DNA, most likely through a covalent linkage (20). Similar experiments performed with MobMN199 showed that the FII forms generated by the truncated protein were also precipitated by KCl and SDS (not shown). This result indicates that the N-terminal domain of MobM contains all the information needed to generate a relaxosome with its target DNA.

Analytical ultracentrifugation analyses showed that the entire MobM protein behaved as an elongated dimer in solution (19). Since the shorter version MobMN199 lacks the last 295 C-terminal amino acids, we wanted to determine its oligomeric state and hydrodynamic properties. The elution volume of the truncated protein in the size-exclusion chromatography used for purification (Figure 1c) suggested that MobMN199 was a monomer. To corroborate this, analytical ultracentrifugation assays were performed at three protein concentrations (2, 8 and 20 µM). The results of sedimentation equilibrium and velocity assays are shown in Figure 2. Sedimentation velocity profiles fit well to a model of single sedimenting species (98.1% of total concentration loaded), with an average molecular mass determined by MALDI-TOF. No improvement in the best-fitting parameters was obtained considering more sedimenting species, an indication of sample homogeneity. At 8 µM, the experimental sedimentation equilibrium data fitted best to a single species (98.1% of total concentration loaded), with an average molecular mass (MW,app) of 22,944 ± 2,820 Da, which agrees with the monomer mass determined by MALDI-TOF. No improvement in the best-fitting parameters was obtained considering more sedimenting species, an indication of homogeneity. At 8 µM, the experimental sedimentation equilibrium data fitted best to a single species (98.1% of total concentration loaded), with an average molecular mass (MW,app) of 23,000 ± 700 Da, which agrees with the monomer mass determined by MALDI-TOF. No improvement in the best-fitting parameters was obtained considering more sedimenting species, an indication of sample homogeneity. At 8 µM, the experimental sedimentation equilibrium data fitted best to a single species (98.1% of total concentration loaded), with an average molecular mass (MW,app) of 23,000 ± 700 Da, which agrees with the monomer mass determined by MALDI-TOF. No improvement in the best-fitting parameters was obtained considering more sedimenting species, an indication of sample homogeneity.
at the higher Mn$^{2+}$ concentrations tested. A very different picture was observed when the cation used was Mg$^{2+}$: even at 15 mM, the metal concentration typically used in DNA-nicking assays (19), Mg$^{2+}$ addition only induced a small thermal up-shift in the final part of the denaturation profile (Figure 4a; orange triangles).

The reversibility of MobMN199 denaturation was examined by recording the protein spectra after cooling the heated samples to the initial temperature of the experiments. Under renaturing conditions, the spectrum of the native protein was substantially recovered (Figure 4b), indicating that MobMN199 was able to refold, at least to a large extent (~60–70%), both in the absence and in the presence of added cations. In addition, the functionality of refolded MobMN199 samples was addressed by performing a cleavage assay on supercoiled pMV158 DNA (FI forms) at 30°C in the presence or absence of the two tested divalent cations. Activity assays showed that MobMN199 was active after the slow heating and cooling steps (Supplementary Figure S1). The specific appearance of open circle forms (FII) as well as relaxed covalently closed forms (FI') demonstrated the nicking‐closing activity of the refolded protein. Although the closing activity of MobMN199 samples renatured in the absence of cation was significantly lower (~16% FI' forms) than the one shown by the native protein (~50% FI' forms), it was almost completely restored when the protein refolded in buffers containing Mg$^{2+}$ (33% FI' forms) or Mn$^{2+}$ (52% FI' forms).

The analysis of the CD denaturation profiles showed that they could be described as the superimposition of two apparently independent processes whose thermodynamic parameters are summarized in Table 2. The values of their relative contributions to the total ellipticity change at 218 nm with and without Mn$^{2+}$ could indicate that the first transition observed in the absence of metal cations ($F_1$~0.36) would correspond to the highest temperature in Mn$^{2+}$-bound samples. The assumed model was further supported by the CD denaturation experiments carried out in ITC buffer (see below).

The influence of Mn$^{2+}$ (and of DNA binding; see below) in MobMN199 structural stability was also examined by DSC. At 124 μM Mn$^{2+}$ (Mn$^{2+}$:MobMN199 ratio 1.5), denaturation apparently takes place in a single step with an enthalpy change, $\Delta H_D$, of 41 kcal/mol and a $T_m$ of 45°C (Figure 5a), and can be described in terms of a two-state transition (red dashed line). However, the heat capacity profile was significantly affected by the cation concentration and, at 8 mM Mn$^{2+}$, MobMN199 denaturation proceeded with a substantial increase of the total enthalpy change ($\Delta H_D = 93$ kcal/mol). In addition, deconvolution of the endotherm indicated that, at this cation concentration, MobMN199 unfolding might take place in three steps (Figure 5a). The two first conform to the two-state transition model ($T_{mB1} = 43.1°C$, $\Delta H_{B1} = 34$ kcal/mol, $T_{mB2} = 50.2°C$, $\Delta H_{B2} = 55$ kcal/mol) and would take place, approximately, in the same temperature interval as the first step observed in the CD profile recorded at this cation concentration. In addition, the shoulder appearing at high temperature ($T_{mA} \approx 63°C$) might

Figure 3. Secondary-structure analysis of MobMN199. (a) Computation analyses. The amino acid sequence of MobMN199 is depicted as well as a summary of the most reliable predictions of secondary structure (using PSIPred, Jpred, NPS@, SABLE and PredictProtein programs). Boxes and arrows below the amino acid sequence of MobMN199 correspond to helices and β-strands, respectively. Boldface letters indicate the conserved residues in the MobM family (Figure 1a). (b) CD spectra of MobMN199 (15μM) protein in buffer CD1, at 10°C. The solid line represents the fit of the experimental curve by the CONTIN method. Experimental data (diamonds) were acquired using 0.02-cm optical path-length quartz cells.

Table 1. MobMN199 secondary-structure content

| Method used   | α-Helix | β-Strand | Turns | Unordered |
|---------------|---------|----------|-------|-----------|
| Deconvolution | CONTIN  | 41.1     | 14.8  | 20.2      | 24        |
| Selcon3       | 47.4    | 7.9      | 22    | 24.1      |
| Average       | 44 ± 4  | 11 ± 5   | 21 ± 1| 24.0 ± 0.1|
| Prediction    | PSIPred | 41.9     | 19.2  | ND        | 38.9      |
| Jpred         | 38.4    | 16.2     | ND    | 45.5      |
| NPS@          | 47.5    | 10.1     | ND    | 42.4      |
| SABLE         | 37.9    | 18.2     | ND    | 43.9      |
| PredictProtein| 36.9    | 15.7     | ND    | 47.5      |
| Average       | 40 ± 4  | 16 ± 4   | ND    | 44 ± 3    |

Data and standard deviation are given in percentages. ND, not determined.
correspond to the last step of CD curves (Figure 4a). As DSC studies were performed in the ITC buffer that contained NaCl instead of (NH₄)₂SO₄, the loss of secondary structure was also monitored in the former buffer following the ellipticity change at 218 nm. Substitution of ammonium sulphate by sodium chloride modified the thermal denaturation profile that became clearly biphasic, even in the absence of Mn²⁺ (Figure 4a, inset). Its deconvolution in terms of two two-state transitions showed an increase of about 5°C in the Tₘ value of the second transition (Table 2 and inset in Figure 4) whereas the first one remained unchanged. These observations support the analyses of the CD profiles monitored in the absence of Mn²⁺ and at low cation concentration in buffer CD2 in terms of two transitions. Moreover, the shift of both transitions to higher temperatures upon addition of 8 mM MnCl₂ was comparable in both buffers (Table 2; inset Figure 4a). However, comparison of DSC and CD results suggested that the first process observed in CD denaturation profiles might actually comprise two transitions at high Mn²⁺ concentrations. Indeed, a better fit of the CD curve recorded at 8 mM MnCl₂ in ITC buffer was obtained using the DSC-derived thermodynamic parameters (Table 2).

Figure 4. Temperature-associated changes in the secondary-structure of MobMN199. (a) Thermal denaturation profiles of the protein (15 μM) measured at 218 nm in presence or absence of divalent metals (free protein, black; 15 mM MgCl₂, orange; 0.015 mM MnCl₂, dark blue; 0.2 mM MnCl₂, red; 1.5 mM MnCl₂, green; 8 mM MnCl₂, purple; and 15 mM MnCl₂, blue). Data fit (continuous lines) assumed the superimposition of two apparently independent transitions whose thermodynamic parameters are shown in Table 2. The inset compares the thermal denaturation profiles of MobMN199 in CD2 (solid symbols) and ITC (open symbols) buffers in the absence (black) and in the presence (purple) of 8 mM MnCl₂. (b) Far-UV CD spectra registered before denaturation (10°C, solid lines) or under renaturing conditions (heated samples cooled to 10°C, dashed lines) without metal (black), and in the presence of 15 mM Mg²⁺ (orange) or 15 mM Mn²⁺ (blue). A spectrum of MobMN199 at 85°C (solid grey line) is also depicted.
The affinity of MobMN199 for Mn$^{2+}$ and Mg$^{2+}$ was examined by ITC. Titrations were performed at 17°C to preserve the native state of MobMN199 during the experiment. The analysis of the binding isotherms (Figure 6) showed that MobMN199 binds Mn$^{2+}$ with high affinity and one-to-one stoichiometry. The equilibrium constant ($K_b$) for the complex formation was $2.3 \pm 0.4 \times 10^6$ M$^{-1}$ and the enthalpy change ($\Delta H_b$) was $27.3 \pm 0.3$ kcal mol$^{-1}$ (average of three experiments). These results evidenced that binding of Mn$^{2+}$ to MobMN199 was enthalpically driven since the entropy change ($\Delta S_b = 65.2 \pm 0.7$ cal mol$^{-1}$ K$^{-1}$) was largely unfavourable. The high affinity of Mn$^{2+}$ for MobMN199 is consistent with the strong stabilization observed in the CD profiles at high cation concentrations (Figure 4). Further, the high values calculated for the enthalpic and entropic contributions indicate the existence of significant conformational rearrangements in the structure of MobMN199 upon Mn$^{2+}$ binding. In contrast, no evidence of Mg$^{2+}$ binding to MobMN199 was found neither from direct binding assays nor from the protein titration with Mn$^{2+}$ in the presence of 10 mM Mg$^{2+}$ (metal competition for the same site should have decreased the apparent affinity of MobMN199 for Mn$^{2+}$).

The effect of metal concentration on MobMN199 nicking-activity

MobM relaxation activity, measured by conversion of FI to FII forms, was previously tested in presence of different metal ions, being Mn$^{2+}$ and Mg$^{2+}$ the more efficient (19). Here we analysed the effect of Mn$^{2+}$ and Mg$^{2+}$ dosage on the activity of MobMN199, while keeping protein–DNA ratio (250 and 8 nM, respectively) and reaction conditions fixed. As shown in Figure 7, the nicking activity of MobMN199 strongly depended on the divalent cation concentration. Relaxed plasmid forms were not observed in the presence of EDTA (10 mM) or in the cation-free control. Maximum activity was found above 8 mM of divalent cation, with conversion levels to FII forms of 46 or 62% in the presence of saturating concentrations of MgCl$_2$ or MnCl$_2$, respectively.

Table 2. Apparent thermodynamic parameters for MobMN199 CD thermal transitions

| Buffer | [MnCl$_2$]$^a$ | $T_m1$ | $\Delta H_{1,app}$ | $F_1$ | $T_m2$ | $\Delta H_{2,app}$ | $F_2$ |
|--------|----------------|--------|-------------------|-------|--------|-------------------|-------|
| CD2    | NA             | 21 ± 2 | 44 ± 8            | 0.33 ± 0.07 | 37.8 ± 0.8 | 48 ± 4 | 0.67 ± 0.07 |
| 0.2    | 37 ± 2         | 34 ± 5 | 0.6 ± 0.1         | 51.0 ± 0.6 | 110 ± 40 | 0.4 ± 0.1 |
| 1.5    | 41 ± 3         | 30 ± 4 | 0.6 ± 0.1         | 55.8 ± 0.6 | 100 ± 40 | 0.4 ± 0.1 |
| 8      | 42 ± 1         | 29 ± 2 | 0.60 ± 0.04       | 64.0 ± 0.4 | 87 ± 10 | 0.41 ± 0.04 |
| 15     | 42 ± 1         | 52 ± 10| 0.48 ± 0.06       | 63.5 ± 0.9 | 68 ± 15 | 0.52 ± 0.06 |
| 15 (Mg$^{2+}$) | 22 ± 2 | 50 ± 20| 0.3 ± 0.1         | 40 ± 2 | 37 ± 7 | 0.7 ± 0.1 |
| ITC    | NA             | 22 ± 1 | 44 ± 7            | 0.32 ± 0.04 | 42.8 ± 0.6 | 41 ± 3 | 0.68 ± 0.04 |
| 8      | 46.5 ± 0.4     | 28 ± 2 | 0.6 ± 0.1         | 64.1 ± 0.6 | 48 ± 5 | 0.4 ± 0.1 |
|        | (43.1 ± 0.9)   | (34 ± 4) | (0.39 ± 0.02) | (65.1 ± 0.7) | (56 ± 7) | (0.43 ± 0.03) |
|        | (50.2 ± 0.1)   | (55 ± 1) | (0.18 ± 0.05)     |      |        |            |

$^a$Concentrations are given in millimolar units. Addition of Mg$^{2+}$ instead Mn$^{2+}$ is indicated.

$^b$Data in parenthesis correspond to the fit based on DSC parameters (see Figure 5 and text for details).

NA, not added.

Figure 5. Thermal denaturation of MobMN199 relaxase measured by DSC. (a) Influence of Mn$^{2+}$ binding. Continuous grey and black lines are the thermograms of DNA-free protein monitored at 124 μM and 8 mM Mn$^{2+}$ (83 μM MobMN199), respectively. Dashed red line is the fit of MobMN199 thermogram at low Mn$^{2+}$ concentration to the two-state model. Dashed–dotted black lines are the elementary transitions under the thermogram at 8 mM MnCl$_2$ assuming that MobMN199 denaturation takes place in three steps and the dashed–dotted green line is the theoretical envelope ('Results' section). (b) Influence of DNA binding. The presence of oligonucleotide IR1+8 shifted the unfolding endotherms of MobMN199 toward higher temperatures. Curves monitored at fixed protein:DNA ratio of 2:1 are in dark yellow (13.3 μM:6.5 μM) and blue (40 μM:20 μM). Mn$^{2+}$ concentrations were fixed to have a free metal concentration of 82 μM at half denaturation.

MobMN199 affinity for cations

The affinity of MobMN199 for Mn$^{2+}$ and Mg$^{2+}$ was examined by ITC. Titrations were performed at 17°C to preserve the native state of MobMN199 during the experiment. The analysis of the binding isotherms (Figure 6) showed that MobMN199 binds Mn$^{2+}$ with high affinity and one-to-one stoichiometry. The equilibrium constant ($K_b$) for the complex formation was $2.3 \pm 0.4 \times 10^6$ M$^{-1}$ and the enthalpy change ($\Delta H_b$) was $27.3 \pm 0.3$ kcal mol$^{-1}$ (average of three experiments). These results evidenced that binding of Mn$^{2+}$ to MobMN199 was enthalpically driven since the entropy change ($\Delta S_b = -65.2 \pm 0.7$ cal mol$^{-1}$ K$^{-1}$) was largely unfavourable. The high affinity of Mn$^{2+}$ for MobMN199 is consistent with the strong stabilization observed in the CD profiles at high cation concentrations (Figure 4). Further, the high values calculated for the enthalpic and entropic contributions indicate the existence of significant conformational rearrangements in the structure of MobMN199 upon Mn$^{2+}$ binding. In contrast, no evidence of Mg$^{2+}$ binding to MobMN199 was found neither from direct binding assays nor from the protein titration with Mn$^{2+}$ in the presence of 10 mM Mg$^{2+}$ (metal competition for the same site should have decreased the apparent affinity of MobMN199 for Mn$^{2+}$).

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forms, followed by a further increase of ~48% at higher cation concentrations, was required to fit well the experimental data (Figure 7c). These results indicate that MobMN199 may contain two or more different classes of Mn$^{2+}$ binding sites, at least in the presence of DNA, and that full nicking activity would be observed when the high and low affinity sites become saturated. Finally, based on this and on previously reported results (19), we can establish that the hierarchy of preference for cation usage by MobM is: Mn$^{2+}$ > Mg$^{2+}$ > Ca$^{2+}$ > Zn$^{2+}$ > Ba$^{2+}$.

**MobMN199 interactions with ssDNA**

The oriT of pMV158 was proposed to be included within a DNA region that contains IR1 and IR2 which encompassed the MobM-mediated nick site (located between coordinates 3591 and 3592) (20). A closer inspection of the sequence allowed us to find a third 31-bp IR (IR3) that includes IR1 plus 5 and 8 bases up- and downstream, respectively (Figure 8a). To reveal the requirements of the DNA target for recognition by MobMN199, we designed a set of Cy5-labelled oligonucleotides that contained part or all of the oriT sequence (Figure 8a). EMSA analysis using 2 nM of DNA showed that MobMN199 recognized specifically the oriT sequence, since it generated single complexes with the IR3-Cy5 and ORIT-Cy5 oligonucleotides (Figure 8b and c, respectively). However, MobMN199 did not bind to IR2-Cy5,
even at high protein concentration (Figure 8d). After quantification of the bound DNA fraction, it was possible to estimate the dissociation constants ($K_d$) of MobMN199 as 60±7 and 58±6 nM for IR3 and ORIT oligonucleotides, respectively. These results indicate that the region encompassing IR1/3 was able to bind MobMN199 with high affinity. However, loss of the 5'-region of the oriT sequence (represented by oligonucleotide IR2) resulted in, at least, a 5-fold decrease in binding affinity ($K_d > 320$ nM). We also analysed the stoichiometry of MobMN199 binding to IR3 oligonucleotide. Graphic representation of the percentage of bound DNA showed that saturation was reached at an approximately 2:1 molar ratio of MobMN199:DNA (Supplementary Figure S2), which suggested the presence of two monomers of MobMN199 per DNA molecule in the complex. The same result was obtained with the ORIT oligonucleotide (data not shown).

The minimal oriT sequence includes IR1 hairpin

Competition binding assays between IR3-Cy5 (2 nM) and different amounts of the same unlabelled DNA (Figure 9a) or the ORIT oligonucleotide (Figure 9b) were performed. The results obtained at a fixed MobMN199 concentration (80 nM) showed a comparable competition ability of both oligonucleotides. This indicated that the recognition sites of MobMN199 for oligonucleotides IR3 and ORIT were equivalent. Thus, the minimal oriT region able to bind MobMN199 with high affinity could be located within the IR3 sequence. To test this hypothesis, new competition experiments were done using DNA fragments shorter than IR3. The results showed that a DNA including the IR1 hairpin sequence plus 8 nt downstream (IR1 + 8), just up to the nick site, was able to compete for binding of MobMN199 to IR3 with, at least, the same efficiencies than IR3 or ORIT (Figure 9c). However, other oligonucleotides harbouring only the IR1 sequence (IR1), the right-arm of IR3 (IR3-R), or 10 nt upstream the nick site (10-NIC), did not compete effectively with IR3-Cy5 for binding to MobMN199 at the concentrations employed (Figure 9d). The same behaviour was observed with oligonucleotides containing the left-arm of IR3 (IR3-L), right-arm of IR1 (IR1-R), or IR2 sequences (not shown).

To further examine the interactions between MobMN199 and IR1 + 8, the influence of DNA binding on the protein thermal stability was evaluated using DSC (Figure 5b) at low Mn2+:MobMN199 ratio (ITC buffer). Addition of IR1+8 oligonucleotide at a protein:DNA molar ratio of 2:1 increased the $T_m$ of the peak observed under similar conditions without DNA (Figure 5a) by >10°C (57.2 and 58.4°C at 13.3 and 40 μM MobMN199, respectively), and also induced a substantial increase in the enthalpy change ($\Delta H_D = 100$ kcal/mol). A small shoulder was also visible at lower temperatures that might be due to DNA-free protein since no transitions were observed in the runs performed with IR1 + 8 in the absence of MobMN199 (not shown). The strong stabilization of the protein structure derived from the DNA–MobMN199 interaction was indicative of a tight complex formation. The dissociation rate constants of IR3 and IR1+8 oligonucleotides were also analysed. These measurements were
done by combining MobMN199 and labelled IR3-Cy5 oligonucleotide and then adding a 100-fold excess of IR3 or IR1+8 unlabelled oligonucleotides to prevent the re-association of labelled oligonucleotide after dissociation from MobMN199. The same behaviour was observed in both the cases, and the fit of the dissociation curves required two exponentials whose average rate constants were 0.030 min⁻¹ and 0.001 min⁻¹ (Supplementary Figure S3). All these results indicated that the minimal oriT target for MobMN199 binding on ssDNA is represented by IR1+8.

DISCUSSION

Here we present the first report on the location, at the N-terminal moiety, of the DNA-binding and DNA-nicking domain of MobM, the representative relaxase of the MOBV family (24). We have separated the relaxase domain from the C-terminal moiety of the protein which, in turn, would harbour the regions involved in protein–protein interactions. Such interactions may include those with the coupling protein provided by either the host chromosome or the auxiliary plasmid participating in the transfer. Further, since MobMN199 was shown to be a monomer, rather than the dimer formed by the full-length protein (19), we propose that the C-terminal moiety would be also involved in MobM dimerization, which would be probably mediated by a putative leucine-zipper motif located between residues 317 and 338 (LENHKSLEAKICLDESNQLQ). The truncated MobMN199 was unable to promote conjugal transfer of pMV158, as well as a MobM-derivative lacking the last 18 residues (results not shown), indicating the relevance of the C-terminal moiety for the in vivo function of MobM in pMV158 transfer.

Activity and stability of protein MobMN199

Spectroscopic and functional results evidenced the strong stabilization of MobMN199 structure provided by Mn²⁺ binding, which correlates with the high affinity of the cation for MobMN199 measured by ITC. The equilibrium constant (2.3 ± 0.3 × 10⁶ M⁻¹) was comparable to Mn²⁺ affinities for the minMobA (the relaxase domain of MobA; 186 residues) and TraI_F relaxases (Kᵇ of 5 × 10⁶ and 2 × 10⁶ M⁻¹, respectively) (48,49), and demonstrated that MobM exhibits high affinity binding for Mn²⁺ ions. Based on previous results obtained with the protein TraI_F (50), this site most likely corresponds to the 3H-motif of MobM (Figure 1a). Interestingly, despite its high binding affinity for Mn²⁺, the metal ion concentration required for MobM optimal activity was >8 mM. This observation indicated that MobM protein may require the uptake of additional cations for an efficient DNA nicking activity than just the one bound to the active centre. In the case of the relaxase domain of minMobA, crystallographic and biophysical studies revealed that, apart from the active centre, there were two more Mn²⁺ binding sites of lower affinities (Kᵇ = 2.5 × 10⁴ and 2.2 × 10⁴ M⁻¹) (22,49). Besides, variation of minMobA activity with cation concentration also supports the notion that saturation of metal ion low-affinity sites was required to reach the full activity (49). These observations indicate that this feature might be shared by other relaxases. Thus, cation binding might
Interactions of MobMN199 with DNA

Employment of fluorophore-labelled oligonucleotides (43) allowed us to study the interactions between MobMN199 and the oriT sequence in single-strand conformation. The oriT structure of pMV158 is different from the other well-known transfer systems because it exhibits three IRs instead of one. Recognition of oriT by MobMN199 was specific and the minimal oriT region defined here was constituted by the IR + 8 oligonucleotide (Figure 8). Similarly to other transfer systems, ssDNA substrates lacking the IR1 left arm of pMV158-oriT (represented by IR2, IR1-R, IR3-R and 10-NIC oligonucleotides; Figure 8a) showed very low affinities for MobMN199. In the case of TraI_F and TrwC, in vitro binding assays indicated that these proteins recognized their cognate oriT-containing oligonucleotides with higher affinities when they included a 5'-segment corresponding to an IR (7, 25). The crystal structure of the TrwC relaxase domain bound to its cognate DNA showed that the IR generates a hairpin, which is involved in the strand-transfer reaction during the termination of conjugation (7, 54). A recent report has corroborated this finding, showing that there are two distinguishable recognition-binding and cleavage-sites for TrwC, both required for efficient conjugation of R388 (28). Furthermore, the relaxase TraI_F could bind with high affinity to the same or similar DNA sequences in two different conformations (25). The two dissociation rate-constants observed in the kinetic analysis performed here suggest that there might be two different modes for the binding of oriT-containing oligonucleotides (IR3) to the MobMN199 protein. Furthermore, our results demonstrated that the average protein:IR3 stoichiometry was 2:1 as experimentally determined using EMSA. We propose that this may correlate with the dimeric nature of the native MobM protein (19), where each monomer could recognize two sites in the oriT during the plasmid transfer. Although the role of IR2 is still unknown, conservation of the oriT sequence among the MOBV plasmid family (not shown) suggests that any one of the three IRs could be involved in the Mob-recognition of the oriT at the initiation of relaxosome formation in the donor cell and/or termination reaction to close the T-strand, in the recipient cell.

Relevance of Mn²⁺ in pMV158 and related plasmids

The only available information on metal ion recognition in MOBV family is represented by Mob_pBBR1 from Bordetella bronchiseptica (55) and BmpH_Tn5520 from Bacteroides fragilis (56) systems; both are active with Mg²⁺. It would be interesting to know whether the selection of the cation reflects a preference for proteins of the MOBV family, or a preference in Mn²⁺ utilization by S. pneumoniae. In this sense, it is worth pointing out that the second nucleotidyl-transferase encoded by pMV158, the RepB initiator of RCR, also required Mn²⁺ for optimal activity (57). Furthermore, this cation was present in the active pocket of the protein as observed in the three-dimensional (3D) structure of RepB (47). The discovery of the pneumococcal mntE Mn²⁺ efflux system...
as involved in pathogenesis and invasive response of this bacterium (58), supports the relevance of this cation in vivo and, perhaps, in transactions involved in horizontal transfer of genetic material (conjugation and transformation being the two most relevant processes in S. pneumoniae). Nevertheless, further experiments, including the solution of the structure of the MobM protein bound to its DNA target, will allow us a deeper biochemical and structural characterization of the MOBv family of relaxases, as well as a better understanding of the mechanisms involved in transfer of the pMV158 promiscuous replicon.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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