Biochemical characterization of three putative ATPases from a new type IV secretion system of Aeromonas veronii plasmid pAC3249A

Ashraf Y Rangrez1,2, Mohammad Y Abajy3, Walter Keller3, Yogesh Shouche1*, Elisabeth Grohmann2*

Abstract

Background: Type four secretion systems (TFSS) are bacterial macromolecular transport systems responsible for transfer of various substrates such as proteins, DNA or protein-DNA complexes. TFSSs encode two or three ATPases generating energy for the secretion process. These enzymes exhibit highest sequence conservation among type four secretion components.

Results: Here, we report the biochemical characterization of three ATPases namely TraE, TraJ and TraK (VirB4, VirB11 and VirD4 homologs of the Agrobacterium tumefaciens transfer system, respectively) from the transfer system of Aeromonas veronii plasmid pAC3249A. ATPases were expressed as His-tag fusion proteins in E. coli and purified by affinity chromatography. ATP binding and ATP hydrolysis experiments were performed with the purified ATPases. TraE and TraK showed strong binding to TNP-ATP and TNP-CTP (fluorescent analogs of ATP and CTP respectively) whereas TraJ showed weak binding. The optimum temperature range for the three ATPases was between 42°C and 50°C. Highest ATP hydrolysis activity for all the ATPases was observed in the presence of Mg2+ and Mn2+. However, TraJ and TraK also showed activity in the presence of Co2+. TraJ exhibited the highest specific activity of all the three ATPases with \( v_{\text{max}} \) 118 ± 5.68 nmol/min/mg protein and \( K_M \) 0.58 ± 0.10 mM.

Conclusions: This is the first biochemical characterization of conjugative transport ATPases encoded by a conjugative plasmid from Aeromonas. Our study demonstrated that the three ATPases of a newly reported TFSS of A. veronii plasmid pAC3249A are functional in both ATP hydrolysis and ATP binding.

Background

TFSS are promiscuous macromolecular transporters of Gram-negative and Gram-positive bacteria that mediate intercellular transfer of various substrates, e.g. proteins, DNA or protein-DNA complexes between bacteria or between bacteria and eukaryotic cells [1-5]. The bacterial conjugation machines form a subgroup of TFSS. They mediate the spread of antibiotic resistance genes and virulence traits among bacterial populations. Agrobacterium tumefaciens uses an archetypal TFSS composed of VirD4, also called coupling protein and VirB1-VirB11 mating pair formation proteins [6]. TFSS encode two or three ATPases belonging to the P-loop NTPase family [1,7]. These ATPases exhibit the highest sequence conservation among TFSS components [1]. They are also characterized by highly conserved nucleotide binding Walker A (GxxGxGKT/S) and Walker B (hhhhDE) motifs [8].

By transfer DNA immunoprecipitation assay, Atmakuri et al. demonstrated that the three energetic components, VirD4, VirB11 and VirB4, mediate successive early steps of the postulated T-DNA translocation pathway [6]. They also suggested that VirD4, VirB11 and VirB4 interact with each other, and that they promote substrate transfer by both ATP-independent and -dependent mechanisms. R388 TrwK (VirB4 homolog), A. tumefaciens VirB11 and Brucella suis VirB11 homolog BsB11 self-assemble into hexamers and show ATPase activity [9-11]. VirB4 has a role in substrate export, whereas for VirB11 a possible chaperone/morphogenetic function was postulated [12]. Evidence for VirB4 self-association [13] and structural contribution...
to channel formation independent of ATPase activity has been shown [10,14,15]. ATPases are also considered as potential drug targets to prevent the spread of disease. Hilleringmann et al. (2006) showed that inhibitors of *Helicobacter pylori* ATPase CagAlfa (VirB11 homolog) block CagA transport and cag virulence [16]. VirD4-like proteins, so called coupling proteins (CP) are considered to link the DNA transfer intermediate to, and perhaps lead it through the mating channel [17]. Coll and coworkers proposed an elegant model based upon the crystal structure of TrwB, the CP of plasmid R388 [17–19]. The strong structural resemblance of TrwB with ring helicases suggests that the transferred ssDNA might pass through the central channel of the TrwB hexamer, thereby entering the translocation apparatus. ATP hydrolysis would provide the energy to pump the ssDNA through the TrwB channel [18].

Recently, we reported a TFSS encoded by plasmid pAc2349A in *A. veronii* consisting of twelve ORFs including three ATPases [20]. The genetic organization of the *A. veronii* TFSS is represented in Figure 1. The *Aeromonas* species used in this study was initially proposed as a novel bacterium, *A. culicicola* [21], which was later defined as a strain belonging to the species *A. veronii* [22]. *A. veronii* is a Gram-negative, rod-shaped bacterium. In humans *A. veronii* can cause diseases ranging from wound infections and diarrhoea to septicaemia in immune compromised patients [23-25]. In this study, we have biochemically characterized the three ATPases TraE, TraJ and TraK (VirB4, VirB11 and VirD4 homologs of *A. tumefaciens* respectively) by ATP binding and ATP hydrolysis experiments. We defined the optimum pH and temperature range of ATP hydrolysis and the requirement of divalent cations.

**Figure 1** Organization of the pAc2349A tra region. Different fillings of the ORFs are indicative of the putative role of the deduced protein. traB is the gene coding for a putative lytic transglycosylase (homologous to VirB1 of the Agrobacterium T-DNA transfer system). Dotted segments represent putative components of pilus assembly whereas thin lined segments represent proteins forming the core complex. Thick lined segments code for three putative ATPases; TraE and TraJ show the conserved nucleotide binding site motif A (Walker A box) and a motif B (Walker B box) of proteins belonging to the VirB4 and VirB11 family of nucleoside triphosphate binding proteins respectively. TraK is a member of the pfam02534 family of TraG/TrwB/VirD/VirD4 coupling proteins. It shows the P-loop motif (Walker A box) and a Walker B motif for nucleotide binding.

**Results**

**Overexpression and purification of TFSS ATPases**

TraE, TraJ and TraK were overproduced in *E. coli* XL10 as His-tagged recombinant proteins and purified by affinity chromatography. Purity of the proteins was checked by SDS-PAGE (Figure 2). TraE and TraK migrate at 90 kDa and 70 kDa in the denaturing PAGE according to their expected molecular mass of 89.97 kDa and 71.21 kDa, respectively. However, the TraJ band migrates at approximately 35 kDa, significantly faster than expected for its calculated molecular mass of 40.63 kDa (Figure 2). Expression of TraE, TraJ and TraK resulted in 2.3 g, 2.1 g and 1.8 g of cells per liter culture. The yield of purified TraE, TraJ and TraK was approximately 2.8 mg, 2.5 mg and 2 mg per liter culture respectively. Highly pure eluted fractions of TraE, TraJ and TraK from Ni-NTA affinity column were concentrated to 0.5-1 mg/mL for enzymatic characterization.

**TraE, TraJ and TraK show ATP binding activity in vitro**

Purified TraE, TraJ and TraK were used for nucleotide binding assays using TNP-ATP and TNP-CTP. The study revealed that all three ATPases show in vitro ATP binding. The ATP binding activity of TraE was higher than that of TraJ and TraK (Figures 3, 4 and 5). In case of TraE and TraK, there was a shift in the absorption maximum after binding to TNP-ATP (Figures 3 and 5). Binding studies with TNP-CTP gave similar results. TNP-CTP binding affinity of TraE was also higher than that of TraJ and TraK. However, the binding affinity for all the ATPases was higher for ATP than for CTP (Figures 3 and 5).

**TraE, TraJ and TraK exhibit ATP hydrolysis activity**

Further characterization revealed that TraE, TraJ and TraK exhibit in vitro ATP hydrolysis activity. The ATP hydrolysis activity was highest for TraJ (118.0 ± 5.68...
nmol/min/mg protein), lower for TraK (82.76 ± 6.82 nmol/min/mg protein) and lowest for TraE (53.35 ± 3.64 nmol/min/mg protein) (Figure 6). As derived from a Michaelis-Menten plot, the $K_M$ of TraE, TraJ and TraK was 0.55 mM, 0.58 mM and 0.92 mM, respectively. The estimated $K_M$ value for TraK is significantly higher than that observed for TraE and TraJ showing its lower affinity for ATP. Utilization of other nucleotides such as CTP, GTP, TTP and ADP as substrate in the hydrolysis assay was also studied. Maximum hydrolyzing activity for TraE was found to be associated with both ATP and TTP. TraJ hydrolysis activity was two times higher with CTP as compared to ATP whereas TraK hydrolyzed TTP more efficiently than other nucleotides (Figure 7).

To eliminate the possibility of the presence of any other contaminant ATPase in the purified TraE/TraJ/ TraK fractions, the purified relaxase (TraL) (which was overproduced and purified following the procedure applied for TraE, TraJ and TraK) was used as a control in ATP binding and hydrolysis experiments (AYR and WK, unpublished data). Neither ATP binding nor ATP hydrolysis activity was observed for the relaxase, ruling out the possibility of any ATPase contamination.

**Mg$^{2+}$ is the best cofactor for TraE, TraJ and TraK ATPase activity**

We analyzed seven different divalent metal ions (Mg$^{2+}$, Cu$^{2+}$, Co$^{2+}$, Zn$^{2+}$, Ca$^{2+}$, Mn$^{2+}$ and Ni$^{2+}$ 10 mM each) in an ATPase assay to define the cofactor requirement of TraE, TraJ and TraK. Mg$^{2+}$ was observed to be the best cofactor followed by Mn$^{2+}$ for ATPase activity of all the three ATPases (Figure 8). Surprisingly, ATPase activity of TraK in presence of Mg$^{2+}$ and Co$^{2+}$ was nearly the
same (Figure 8). However, Cu$^{2+}$, Zn$^{2+}$, Ca$^{2+}$ and Ni$^{2+}$ were less effective in activating the ATP hydrolysis activity of the ATPases.

**TraE, TraJ and TraK are active between pH 6.5 - 8.0**

The effect of different buffers on ATPase activity was studied to determine the optimum pH for the ATP hydrolysis activity of TraE, TraJ and TraK. A set of buffers ranging from pH 5.0 to 9.5 (sodium citrate (pH 5.0 - 5.5), sodium phosphate (6.0 - 6.5), HEPES-NaOH (pH 7.0), Tris-HCl (pH 7.5 - 8.5), and sodium borate-NaOH (pH 9.0 - 9.5)) were used to determine the pH optimum. The optimal ATPase activity of TraE, TraJ and TraK was observed within a pH range of 6.5 to 8.0, with highest activities for TraE and TraK at pH 7.0 and for TraJ at pH 7.5 (Figure 9). Notably, ATP hydrolysis activity for TraE declined gradually beyond the optimum pH range. However, ATP hydrolysis by TraJ and TraK dropped sharply outside the optimum pH range (Figure 9).

**Optimum temperature range for TraE, TraJ, and TraK**

ATPase activity of TraE, TraJ and TraK was measured over the temperature range of 10 - 80°C (Figure 10). Approximately 60-80% of the total ATPase activity for all the ATPases was observed within a temperature range of 36 - 50°C. Maximum activity for TraJ and TraK occurred at 42°C and for TraE the optimum temperature observed was 45°C. At lower temperatures (below 36°C), only 10 - 15% activity was measured. However, at temperatures between 60°C - 70°C, 15-20% of the total activity was retained. It sharply dropped to less than 5% at 80°C (Figure 10).

**Discussion**

TraE, TraJ and TraK (VirB4, VirB11 and VirD4 homologs of the *A. tumefaciens* TFSS respectively) are components of the first putative conjugative TFSS of *Aeromonas*. Here we present a functional characterization of these ATPases of *A. veronii* plasmid pAc3249A. Like other transport ATPases, TraE, TraJ and TraK
and TraK displayed Mg2+ dependent ATPase activity towards Tides effectively acting as an ATPase. Purified TraE, TraJ and TraK bind and hydrolyze nucleotides with v_max of 53.35, 118.0 and 82.76 nmol/min/mg protein respectively, ATP is the likely substrate utilized intracellularly due to the relatively low abundance of other nucleotides in vivo [29].

As expected, Mg2+ was the preferred divalent cation for the ATPase activity of TraE, TraJ and TraK [29]. However, TraJ retained approximately 35% and TraK 95% activity in the presence of Co2+ when compared to Mg2+. This observation was surprising considering the low concentration of Co2+ and its toxic effect on bacterial cells [30,31]. Ca2+ was less effective whereas Cu2+, Zn2+ and Ni2+ were not effective in activating ATP hydrolysis as was the case with the A. hydrophila ATPase [29]. The optimum pH for TraE and TraJ ATPase activity is around 7.0. This is comparable with the pH optimum of their respective homologs R388 TrwK (pH 6.5) and R64 PilQ (pH 6.5) [11,32]. ExeA hydrolyzed GTP and CTP more efficiently than ATP. Although highest hydrolytic rates of TraJ and TraK were obtained with CTP and TTP, respectively, ATP is the likely substrate utilized intracellularly due to the relatively low abundance of other nucleotides in vivo [29].

The ATPase activity identified for TraE (53.35 nmol/min/mg protein) is comparable with the recently reported activity for its homolog R388 TrwK (48.4 nmol/min/mg protein) [11]. The ATPase activities observed for TraJ (VirB11 homolog) and TraK (VirD4 homolog) are approximately 10 fold higher than the values reported for their respective homologs: plasmid pTiC58 VirB11 and plasmid R388 TrwD presented weak ATPase activity of 1 - 15 nmol/min/mg protein [27,28]. The weak activity of these proteins may be attributed to denaturing-renaturing conditions used during protein purification [27,28]. In this study, we maintained native conditions for the proteins throughout the extraction and purification steps which likely explain the higher ATPase activities for TraE, TraJ and TraK.

The ATPase activity measured for TraJ is approximately two fold higher than that for TraE and TraK. Though TraE, TraJ and TraK were able to utilize both purine and pyrimidine nucleotides as substrate, we observed major differences in hydrolysis activities with respect to the different nucleotide substrates. TraE exhibited approximately the same hydrolysis activity in the presence of ATP and TTP but only 50 - 60% activity (compared to ATP) in the presence of ADP, CTP and GTP. Hydrolysis activity of TraJ was almost twofold higher for CTP than for ATP. However, TraJ could not hydrolyze GTP. TraK utilized TTP most efficiently, followed by ATP and ADP. Interestingly, similar results on substrate selectivity were reported for ExeA, an ATPase involved in the type II secretion pathway of Aeromonas hydrophila [29]. ExeA hydrolyzed GTP and CTP more efficiently than ATP. Although highest hydrolytic rates of TraJ and TraK were obtained with CTP and TTP, respectively, ATP is the likely substrate utilized intracellularly due to the relatively low abundance of other nucleotides in vivo [29].

As expected, Mg2+ was the preferred divalent cation for the ATPase activity of TraE, TraJ and TraK [29]. However, TraJ retained approximately 35% and TraK 95% activity in the presence of Co2+ when compared to Mg2+. This observation was surprising considering the low concentration of Co2+ and its toxic effect on bacterial cells [30,31]. Ca2+ was less effective whereas Cu2+, Zn2+ and Ni2+ were not effective in activating ATP hydrolysis as was the case with the A. hydrophila ATPase [29]. The optimum pH for TraE and TraJ ATPase activity is around 7.0. This is comparable with the pH optimum of their respective homologs R388 TrwK (pH 6.5) and R64 PilQ (pH 6.5) [11,32]. We noticed moderate ATPase activity for TraE, TraJ and TraK also above 50°C. The optimal temperature range for all the ATPases was between 36°C - 50°C. Interestingly, for all of them, the ATP hydrolysis activity observed at 30°C was lower than the activity at 50°C.

The involvement of the pAc3249A TFSS in conjugative plasmid transfer was confirmed by mating experiments (AYR and EG, unpublished data). In triparental mating experiments with A. veronii as a donor, we could show mobilization of plasmid pDL277-
oriT\textsubscript{pAc3249A} (a derivative of plasmid pDL277 containing the oriT region of plasmid pAc3249A) to \textit{E. coli} XL1 blue cells. Further studies on putative key factors of the \textit{Aeromonas} TFSS are in progress. We are currently characterizing the putative relaxase encoded by the pAc3249A transfer region. In conjunction, these data will elucidate the molecular mechanism of the first TFSS encoded by a plasmid from pathogenic \textit{Aeromonas}.

Conclusions
This is the first biochemical characterization of conjuga-
tive transport ATPases encoded by a conjugal plas-
mid from \textit{Aeromonas}. This study illustrated that the
three ATPases, TraE, TraJ, and TraK of the newly
reported TFSS of the \textit{A. veronii} plasmid pAc3249A bind and hydrolyze ATP. Ongoing efforts to solve the three
dimensional structure of these ATPases will help explain
the substrate preferences and nucleotide binding activ-
ities of the respective proteins.

Methods
Cloning of \textit{traE}, \textit{traJ} and \textit{traK}
The \textit{traE}, \textit{traJ} and \textit{traK} genes of plasmid pAc3249A (GenBank\textsuperscript{*} accession number DQ890522) were ampli-
cified by PCR using Phusion DNA polymerase (New
England Biolabs, Frankfurt, Germany) and the specific
primers given in Table 1 (Eurofins MWG Operon,
Ebersberg, Germany). The PCR products were cut with
restriction enzymes \textit{Bam}HI and \textit{Hind}III and purified by PCR cleanup kit (Qiagen, Hilden, Germany).
Expression vector pQTEV (Qiagen, Hilden, Germany)
was also cut with \textit{Bam}HI and \textit{Hind}III. The ligation
mixture was incubated at 16°C overnight with T4
DNA ligase (Roche Diagnostics, Mannheim, Germany)
and used for transformation of \textit{Escherichia coli} XL1
Gold cells (Stratagene, Amsterdam, The Netherlands).
Plasmid DNA of selected clones of pQTEV-\textit{traE},
pQTEV-\textit{traJ}, and pQTEV-\textit{traK} was sequenced by SMB
(Rüdersdorf, Germany) to confirm the open reading
frames.

| Primer | Sequence (5’-3’) | Target gene | Nucleotide position (in pAc3249A) |
|--------|-----------------|-------------|----------------------------------|
| \textit{traE\textsubscript{f}} | GCCGGATTCATGAAAGCAGATAAAAGCA | \textit{traE} | 1250 - 1266 |
| \textit{traE\textsubscript{r}} | GCCAAGCTTAACTCCTCTCAGTTTG | | 3605 - 3623 |
| \textit{traJ\textsubscript{f}} | GCCGGATTCATGTCGCCGAAATATT | \textit{traJ} | 8007 - 8023 |
| \textit{traJ\textsubscript{r}} | GCCGAAGCTTAAATTGCATCTCGAGC | | 9066 - 9085 |
| \textit{traK\textsubscript{f}} | GCCGGATTCATGAAAAATAAACGGG | \textit{traK} | 9099 - 9115 |
| \textit{traK\textsubscript{r}} | GCCGAAGCTTATGATATGTTATAGGC | | 10965 - 10983 |

Nucleotides shown in bold indicate the restriction sites, GGATCC - \textit{Bam}HI and AAGCTT - \textit{Hind}III.

Expression of \textit{TraE}, \textit{TraJ} and \textit{TraK}
5 mL overnight cultures of the corresponding expression
strains (\textit{E. coli} XL1 Gold cells harbouring pQTEV-\textit{traE},
pQTEV-\textit{traJ}, and pQTEV-\textit{traK} respectively) were har-
vested by centrifugation, 6000 g at 4°C for 15 min and
resuspended in 100 mM Tris-HCl (pH 7.6), 100 mM
NaCl and 10% glycerol (buffer A). Cell lysis was carried
out by incubating the suspension with 1 mg/mL lyso-
zyme at room temperature for 30 min followed by five
cycles of ultrasonication, 1 min each. Cell debris was
removed by centrifugation at 25000 g at 4°C for 30 min.
The supernatant was loaded onto a Ni\textsuperscript{2+} charged Ni-
NTA (Qiagen, Hilden, Germany) affinity column equili-
brated with buffer A. The column was first washed with
buffer A followed by two washes with buffer A contain-
ing 30 mM and 40 mM imidazole respectively. The
respective protein (TraE/TraJ/TraK) was eluted in buffer
A with 250 mM imidazole. The purification process was
monitored by 12% SDS-PAGE. Eluted fractions of high
purity were pooled and concentrated by centrifugation
with the Amicon CentriPrep system (molecular mass
cut-off of 10 kDa) (Millipore, Vienna, Austria). The con-
centrated proteins were used for nucleotide binding and
hydrolysis studies.

NTP hydrolysis assay
The NTP hydrolysis assay was performed in 50 \textmu l reac-
tions as described by Lanzetta \textit{et al}. (1979) with minor
modifications [33]. The standard reaction mixture con-
taining 2.5 \textmu g of \textit{TraE}, \textit{TraJ} or \textit{TraK} protein, 2.5 mM
ATP and 10 mM MgCl\textsubscript{2} in buffer A (100 mM Tris-HCl

Table 1 Primers used for the amplification of \textit{traE}, \textit{traJ} and \textit{traK} genes of plasmid pAc3249A (GenBank accession number DQ890522).

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(pH 7.6), 100 mM NaCl and 10% glycerol) was incubated at 37°C for 30 min. Furthermore, the effect of different ATP concentrations (0.025 mM - 3.5 mM), different nucleotides (2.5 mM each), different divalent cations (10 mM each), different pH (pH 5 - pH 9.5) and different temperatures (10°C - 80°C) on hydrolysis activity was analyzed. The reactions were stopped by the addition of 800 μL of a colour reagent (3:1 ratio of 0.045% malachite green and 4.2% ammonium molybdate in 4 N HCl) and 100 μL of 34% citric acid solution. After incubation at room temperature for 30 min, the absorbance was measured at 660 nm. One unit of ATPase activity was defined as the hydrolysis of 1 nmol of ATP/min/mg of the respective protein. Three independent experiments were performed for each parameter and graphs were plotted using mean values and standard deviations wherever appropriate.

Nucleotide binding assay

The nucleotide binding assay was performed following the method described in Schröder and Lanka [26]. The fluorescent nucleotide analogues TNP-ATP (Invitrogen Biosciences, Jena, Germany) were used as substrates. Briefly, 0.5-1 mg/mL protein (TraE, TraJ or TraK) in buffer A (100 mM Tris-HCl (pH 7.6), 100 mM NaCl and 10% glycerol) was incubated for 20 s with TNP-ATP (15 μg) or TNP-CTP (15 μg) at room temperature. Fluorescence spectra were taken at room temperature by using a Varian Cary Eclipse spectrophluorometer (Varian Inc., Palo Alto, USA) with excitation at 410 nm and emission scanning in the range of 470 to 630 nm. The fluorescence maxima were determined graphically.

Abbreviations Used

TFSS: type IV secretion system; CP: coupling protein; ORF: open reading frame; TNP-ATP: 2',3'-O-(2,4,6-trinitrophenyl) adenosine 5'-triphosphate; TNP-CTP: 2',3'-O-(2,4,6-trinitrophenyl) cytidine 5'-triphosphate; ss: single stranded; NTP: nucleotide tri-phosphate.

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Author details

1Molecular Biology Unit, National Centre for Cell Science, Pune 411007, India.
2Department of Environmental Microbiology, Technical University of Berlin, Franklinstr. 29, FRI-2, D-10587 Berlin, Germany.
3Institute for Molecular Biosciences, Karl-Franzens-University Graz, Humboldtstrasse 50/3, A-8010 Graz, Austria.

Authors’ contributions

AYR, YS, WK and EG designed the experimental strategy for this study. AYR and MYA were involved in standardization of the experimental conditions.

AYR was involved in acquisition of the data. AYR, WK and EG analyzed and interpreted the data. AYR drafted the manuscript and other authors made corrections to the manuscript. All authors read and approved the final manuscript.

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