Assembly dynamics and stability of the Pneumococcal epsilon zeta Antitoxin Toxin (PezAT) system from *Streptococcus pneumoniae*

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Running title: Dynamics of the PezAT system

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The Pneumococcal epsilon zeta Antitoxin Toxin (PezAT) system is a chromosomally encoded, class II toxin antitoxin system from the human pathogen *Streptococcus pneumoniae*. Neutralization of the bacteriotoxic protein PezT is carried out by complex formation with its cognate antitoxin PezA. Here we study the stability of the inhibitory complex in vivo and in vitro. We found that toxin release is impeded in *Escherichia coli* and *Bacillus subtilis* due to the proteolytic resistance of PezA once bound to PezT. These findings are supported by in vitro experiments demonstrating a strong thermodynamic stabilization of both proteins upon binding. A detailed kinetic analysis of PezAT assembly revealed that these particular features of PezAT are based on a strong, electrostatically guided binding mechanism leading to a stable toxin antitoxin complex with femtomolar affinity. Our data show that PezAT complex formation is distinct to all other conventional toxin antitoxin modules and a controlled mode of toxin release is required for activation.

Toxin-antitoxin (TA) systems were initially discovered on prokaryotic low-copy number plasmids, where they ensure stable inheritance in a growing population. By a postsegregational killing mechanism these systems trigger cell death of offspring that have lost the plasmid and were thus also designated as addiction or suicide modules. Most of those TA systems are transcribed from a single bicistronic operon, where the first gene product (antitoxin) inhibits the cytotoxic function upon binding to the product of the succeeding gene (toxin) (1). Addiction to the TA locus is achieved by differential proteolytic stabilities of the two proteins since antitoxic proteins are metabolically less stable and prone to faster degradation by intracellular proteases when compared with their cognate toxins. Hence, loss of the TA-loci results in a faster reduction of the pool of antitoxic proteins than that of toxins due to lack of de-novo synthesis, which eventually confronts the host with the freed and corrosive activity of the toxin.

In addition to plasmid encoded TA-systems, similar modules were also discovered to be encoded from chromosomes of free-living bacteria (2). However, the function of these chromosomally encoded TA-systems is currently under debate (3,4) and different functions such as growth control (5), stabilization of mobile genetic elements (6,7) or programmed cell death (8) have been suggested.

TA systems of the epsilon/zeta family are encoded on various plasmids or chromosomes of Gram-positive pathogens like *Streptococcus pyogenes* or *Streptococcus pneumoniae* as well as Gram-negative such as *Fusobacterium nucleatum*(9-12). The related zeta-toxins are ~30 kDa monomeric proteins owning a monophosphate kinase fold and are thought to phosphorylate a still enigmatic substrate (13). In addition to the toxin-inactivating function of the antitoxin epsilon, various homologues harbour a transcriptional...
repressor domain that regulates the operon (11,14).

The PezAT system (pneumococcal epsilon zeta) from <i>S. pneumoniae</i> encoded from the chromosomal Pneumococcal Pathogenicity Island 1 (PPI1) (15) is one of the only two epsilon/zeta systems characterized to date. <i>S. pneumoniae</i> strains containing PPI1 encode for various virulence factors that are highly beneficial for infection and these strains show increased pathogenicity. Strains lacking the toxin PezT gene displayed an attenuated infection progress and were out-competed by wildtype strains. This finding indicated an important role of PezT for pneumococcal pathogenicity as virulence factor (16).

Expression of PezT leads to growth arrest in <i>Escherichia coli</i> (11). The toxic activity of PezT is inhibited by complex formation with the dimeric three-helix bundle domain of PezA which is flexibly linked to the amino-terminal helix-turn-helix domain. The crystal structure of the inhibitory complex reveals a dumbbell-shaped heterotetrameric assembly, where the central dimeric three-helix bundle domain of the antitoxin binds two toxins and neutralizes them by blocking the conserved nucleotide binding site. Expression of PezAT is regulated by the transcriptional repressor domain of PezA, which binds to the σ70 like promotor of the bicistronic pezAT operon.

The formation of the inhibitory TA complex is the crucial step in toxin neutralization of bacterial TA systems. We investigated the proteolytic stability of the inhibitory PezAT complex under in vivo-like conditions in bacterial model systems. Furthermore we performed a detailed characterisation of the association / dissociation dynamics of the PezAT system and thereby can provide a general model for heterotetramer formation. Taken together, the results of this study indicate that PezT gets sequestered by PezA in a quasi-irreversible manner indicating that PezAT is a unique TA system.

**EXPERIMENTAL PROCEDURES**

Protein expression and purification – PezA, PezT and all of their variants were expressed from pET28b vectors using the <i>E. coli</i> BL21(DE3)/Codon Plus-RIL strain (Stratagene). Site directed mutagenesis was performed using the QuikChange protocol (Stratagene). For in vivo studies of protein stability the pezAT ORF was cloned into the pASK5+ vector (IBA) using Bsal (NEB). This vector results in an N-terminal Strep II tag fusion with the wildtype PezA. All constructs were verified by DNA sequencing. Overexpression of protein constructs from the pET system, as well as purification of PezT variants was carried out as described before (11). Note that all PezT constructs bore a non-cleavable C-terminal His<sub>6</sub>-tag. The initial step of PezA purification using Ni-affinity chromatography was the same as for PezT (11). Subsequently, PezA was dialyzed overnight against buffer A (all buffer compositions are listed in Table S1) in presence of thrombin to cleave the N-terminal His-tag, except for PezA preparations used in western blot analysis. The protein was then loaded onto a MonoQ column (GE Healthcare) equilibrated with buffer B and eluted in a gradient to 60 % buffer C. Fractions were analyzed by SDS-PAGE, pooled and further purified by using a Superose 6 (GE Healthcare) size exclusion column equilibrated with buffer D. All protein concentrations were determined by measuring the absorption at 280 nm. PezA, PezT and all of their variants were stored in buffer D.

Labelling of PezT(D66T) and PezA(M66C) with fluorescent dyes – PezT(D66T) was labelled at cysteine C209 with Alexa488-maleimide (Invitrogen) according to the dye manufacture’s protocol. Briefly, to a vial containing 1.5 mL of PezT(D66T) (3 mg/ml in buffer E), 140 µl of 10 mM Alexa488-maleimide was added dropwise under constant stirring and incubated overnight at 4°C. The reaction was quenched by adding 10 mM DTE. Precipitated protein was dissolved in buffer D containing 8 M urea and refolded by rapid dilution (1:15) in buffer D. Free fluorescent dye and residual urea were removed by buffer exchange in a 10 kDa cut-off spin column. The label efficiency was estimated spectrophotometrically to be ~100%. Proper refolding of the labelled PezT species was verified by circular dichroism and PezA binding experiments. In order to obtain the fluorescently labelled His-F-PezA variant, DTE free His<sub>6</sub>-PezA(M66C) was labelled with 1,5-IAEDANS...
(Invitrogen) by diluting 25 µM of His<sub>6</sub>-PezA(M66C) in buffer G. A 1,5-IAEDANS stock (20 mM in DMF) was added drop-wise under rigorous stirring to a final concentration of 380 µM. The mixture was further incubated for 1 hour at 4°C before the labelling reaction was quenched with 10 mM DTE. Free label was removed similar as described above. The label efficiency was estimated spectroscopically to be ~130% due to additional labelling of the second, poorly accessible cysteine at position 50.

**In vivo stability of PezAT** – To monitor degradation of PezA and PezT in vivo, *E. coli* DH5α cells bearing pASK5+(pezAT) or pASK5+(pezA) were grown in 200 mL LB at 37°C in presence of 100 µg/mL ampicillin. At an OD<sub>600</sub> of 0.5, protein expression was induced for one hour by addition of 0.2 µg/ml anhydrotetracycline. After that, chloramphenicol was added to a final concentration of 50 µg/mL to inhibit protein synthesis. Samples of 1 mL were taken in one hour intervals and cells were harvested at 16,000 x g for 1 min. Subsequently, the cells were resuspended in 100 µl buffer H and incubated at 95 °C for 5 minutes. Equivalents of 7.5 µl/OD<sub>600</sub> were analyzed by SDS-PAGE and subsequent Coomassie staining.

For the experiments probing PezT release, PezAT was expressed as described above. Both cultures expressing either PezA or PezAT were harvested at an OD<sub>600</sub> of 0.8 by centrifugation (5 min at 6,000 x g, 4°C). Cells were washed in LB-medium after centrifugation and finally resuspended in 100 mL LB, which was used to inoculate fresh LB medium supplemented with ampicillin 100 µg/mL to a final OD<sub>600</sub> of 0.05. The cell growth was monitored at 37°C by measuring the OD<sub>600</sub>.

**PezAT degradation in crude extract** – Cellular crude extract was obtained from *B. subtilis* YB886 overnight cultures grown in LB medium containing 5 µg/mL kanamycin. Cells were harvested by centrifugation (15 min at 6000 x g and 4°C) and 3 g of wet pellet was washed once in buffer I and finally resuspended in 30 mL of buffer I. Cell walls were opened by sonication for 20 minutes on ice. The resulting crude extract was supplemented with 10 mM MgCl<sub>2</sub>, 5 mM DTE and either 1 µM His-F-PezA, 1 µM PezT(D66T) or 1 µM His-F-PezA/PezT(D66T) complex. Individual samples were incubated at room temperature and aliquots were withdrawn at t = 0 h, 1 h, 2 h and 3 h. Degradation of His-F-PezA and PezT(D66T) was monitored by ECL western blot analysis using a Penta-His HRP Conjugate (Qiagen) following the supplier’s instructions. Additionally, degradation of His-F-PezA was monitored by visualizing IAEDANS fluorescence using an Imago compact imaging system (B&L systems).

**Urea denaturation of PezAT** – To monitor unfolding of PezA, 5 µM PezA or 5 µM PezA/PezT(D66T,W232Y) complex were incubated 15 h at 10°C and 5 h at 25°C in buffer D at different urea concentrations ranging from 0 M to 7.2 M. The fluorescence spectral centre of mass <λ> for each emission spectrum ranging from 335 nm to 400 nm was calculated by

\[
<\lambda> = \frac{\sum \lambda_i \cdot I_i}{\sum I_i}
\]

where \(I_i\) is the fluorescence intensity of the ith recorded point and \(\lambda_i\) the respective wavelength in nanometer. The mean change in residue ellipticity of the same samples was monitored in a J-810 CD Spectropolarimeter (Jasco) at 222 nm.

**Fluorescence titrations** – Toxin-antitoxin binding isotherms were obtained by titrating wildtype PezA or PezA(W111F) to a cuvette containing 3 µM PezT(D66T) in buffer D. The change in fluorescence intensity at 347 nm was recorded using a FluoroMax-3 Spectrofluorometer (HORIBA Scientific) with an excitation wavelength at 295 nm.

**Stopped-flow experiments** – PezAT association experiments were carried out in a SF61-DX2 stopped-flow instrument from TgK Scientific. The excitation wavelength was set to 295 nm, and the fluorescence was detected using a 320 nm low pass filter. Assembly of PezAT was monitored after rapid mixing of PezT(D66T) with PezA(W111F) in buffer D at 25 °C. The change in fluorescence intensity was measured at 0.25 µM, 0.5 µM, 0.75 µM, 1 µM, 2 µM, 3 µM and 5 µM toxin antitoxin concentrations for shots with 1:1 stoichiometry and 3 µM/6 µM, 6 µM/3 µM, 2 µM/1 µM, 1.5 µM/1 µM and 2 µM/1.5 µM antitoxin/toxin.
concentrations for asymmetric stoichiometries. 12-15 time traces of each measurement were averaged and analyzed using Dynafit 4.0 (Biokin) (17). In the final fit assuming a two-step model, the apparent rate constants were fitted globally, whereas the trace offsets and amplitudes were fitted locally.

Salt dependency of PezAT association was characterized by rapid mixing experiments of PezA and PezT (1 µM and 1:1 stoichiometry) at different NaCl concentrations and by fitting of the two-step model to individual time traces. Except for measurements at low ionic strength (below 100 mM NaCl), additional time traces for assembly experiments at 3 µM/1 µM antitoxin/toxin concentrations were included in the fit to support convergence into a well defined minimum. In order to account for any buffer contribution of 50 mM Hepes NaOH at pH 7.5, 23 mM were added to the overall ionic strength.

The dynamics of the PezA homodimer equilibrium were addressed by rapid dilution experiments where 5 µM PezA was mixed with buffer D to final ratios of 1:15, 1:12, 1:10, 1:8, 1:6, 1:5 and 1:4 in the absence and the presence of stoichiometric amounts of tryptophan free PezT(D66T,W232Y) using a SFM-400 stopped-flow device (BioLogic). Individual time traces from at least 20 shots were averaged per concentration to increase the signal to noise ratio of the intrinsically weak amplitude. These averaged time traces were fitted globally applying a mono-exponential assuming perturbation of a one-step dimerization equilibrium (18)

\[
F(t) = F_0 + \Delta F \cdot \text{Exp}\left(-\sqrt{8k_1[A]_0 + (k_{-1})^2} \cdot t\right)
\]

where \(F_0\) is the fluorescence at time zero, \(\Delta F\) the increase in fluorescence, \(k_1\) the association rate of dimer formation and \(k_{-1}\) the rate of dissociation.

Determination of PezAT dissociation rate – Displacement of PezT(D66T)Alexa488 from the PezAT complex with unlabelled PezT(D66T) was performed in buffer J at room temperature. 6 µM of unlabelled PezT(D66T) was incubated with 0.3 µM of a preformed PezA(W111F)/PezT(D66T)Alexa488 complex. The complex was separated from unbound PezT using a Superdex 200 (GE Healthcare) gel filtration column equilibrated in buffer J, connected to a Waters HPLC system with attached fluorescence detector and autosampler device. Elution volumes for either the complex or PezT alone were determined by calibration runs. 50 µL of sample were injected every 60 min onto the column and fluorescence was recorded at 525 nm with excitation set to 488 nm. Elution profiles of labelled PezAT and PezT were fit to a two-component skew-normal distribution function using CalcCenter 3.0 (Wolfram Research)

RESULTS

PezT remains inactive after inhibition of PezAT expression in E. coli - In genuine TA systems, toxin activation is caused by selective degradation of the antitoxin by AAA+ chaperone-proteases like Lon, ClpXP, ClpAP or ClpCP (5,19-22). To probe liberation of PezT by degradation of PezA in E. coli owning the entire K12 protease background, we expressed PezAT from a tetracycline-inducible promoter in E. coli DH5α. Cells harboring pASK5+(pezAT) were induced at an OD600 of 0.5. Subsequently, protein translation was inhibited by adding 50 µg/mL chloramphenicol after 1 h of expression. Inhibition of cell growth was verified by measuring the optical density in one hour intervals. In parallel, protein levels of PezA and PezT were monitored using Coomassie stained PAGE gels loaded with equivalent amounts of lysed cells (Fig. 1a). Intriguingly, we observed only a weak decrease of full-length PezA protein in presence of PezT during 4 hours of growth inhibition. Since PezA is co-expressed with PezT from a single bicistronic operon (11), intracellular PezA concentrations can be expected to be higher than those of PezT due to failure in translation initiation of the second open reading frame. Thus, the observed loss of PezA is probably caused by degradation of free PezA, which is in excess over PezT. This finding was somewhat unexpected, since Lon and ClpXP dependent degradation of the PezA homologue epsilon was reported to occur on a time-scale of minutes in the presence of zeta toxin when translation was inhibited in B. subtilis (23,24).

To probe if the observed initial loss of full-length PezA correlates with PezT release or
was simply due to terminal truncation and/or degradation of unbound excess PezA, respectively, we selectively switched off expression from the tet-promoter in order to provoke cell poisoning by PezT. Therefore, we induced PezAT expression from pASK5+ (pezAT) in DH5α. In contrast to the experiment described above, cells were harvested by centrifugation and washed extensively in order to remove any tetracycline after one hour of induction. The bacteria containing the expressed PezAT complex were then used to re-inoculate fresh LB medium and the cell growth was monitored (Fig. 1b). Similar to what was shown for the PezT homologue zeta (11,23,24) we expected a perceptible growth defect when expression from the tet promoter was impaired indicating a release of PezT upon PezA degradation. Intriguingly, we could not observe a difference between the cell cultures that had either expressed PezAT or, as control, PezA alone, suggesting that no significant release and accumulation of PezT occurred.

The PezAT core complex is protected from degradation in Bacillus subtilis crude extracts - Since we could not provoke PezT release in E. coli, we speculated about any host specific degradation and therefore performed a semi-in vivo approach in B. subtilis. In these experiments we could follow degradation of PezA and PezT over time using controlled and stoichiometric concentrations. As probe for the antitoxin, we used a PezA mutated variant His6-PezA(M66C) with a 1,5-IAEDANS fluorescence label attached to residue C66 (further simply referred to as His-F-PezA). Cysteine 66 is located in the peptide linker between the N-terminal helix-turn-helix motif and the C-terminal PezT binding domain; the three helix bundle. This modified variant enabled us to follow degradation of both domains independently. Whereas the western blot analysis using an Anti-His antibody indicated the presence of the N-terminus of the polypeptide chain, fluorescence imaging enabled detection of any proteolytic fragments that harbour the peptide linker. Degradation of PezT(D66T) was monitored by western blot analysis detecting its C-terminal His-tag. Both, His-F-PezA and PezT(D66T) were incubated separately and together at 1:1 molar ratio in highly concentrated, cytosolic extracts from freshly prepared B. subtilis.

We found that when His-F-PezA was incubated in B. subtilis cell extracts, the N-terminal His-tag became undetectable within the first hour of incubation (Fig. 1c). In parallel, we observed the occurrence of a faint band with higher electrophoretic mobility, indicating that to some extent N-terminal truncation to a smaller but stable domain occurred. On the other hand, PezT(D66T) turned out to be more stable than PezA and the C-terminal His-tag was still detectable after 1h of incubation (Fig. 1d).

However, when both proteins were incubated together, the His-tag of His-F-PezA was still detectable after 2h of incubation. Apparently, once bound to PezT, full-length His-F-PezA is protected to some extent. In contrast to the previous experiment, almost the entire His-F-PezA was subsequently truncated to the smaller stable domain containing the fluorophore, which was detected by the fluorescence imaging technique (Fig. 1e). Furthermore, PezT remained stable throughout the entire experiment suggesting that the smaller, proteolytically derived C-terminal fragment is sufficient to protect PezT from degradation. We can exclude that neither mutation nor the labelling of PezA have caused the observed stabilizing effect since similar results were obtained with unlabelled wild type PezA (data not shown).

Our experiments show that a PezAT core complex lacking the helix-turn-helix motif of PezA is extremely resistant towards proteolysis when compared to the individual proteins alone. Additional experiments using native gel electrophoresis monitoring IAEDANS fluorescence verified the identity and integrity of this PezAT core complex (Fig. S1). Thus, PezA undergoes a fast N-terminal truncation, leaving the remaining three-helix bundle associated to PezT. In the core complex the PezA three helix bundle and PezT are stably associated and both polypeptide chains are resistant towards any further proteolytic cleavage. This result is in good agreement with our stability studies in E. coli (see above) since we observed slight reduction of PezA levels in our PAGE analysis.
but the cells did not show impaired growth (Fig. 1a).

PezAT complex formation strongly increases the thermodynamic stability of the two proteins – Protein susceptibility to degradation depends on their global thermodynamic stability and, if they are substrates for ATP dependent proteases, on the accessibility and local stability of recognition motifs (25-27). In order to probe if the observed proteolytic stability of the PezAT complex is correlated with an increased thermodynamic stability, we performed unfolding studies of PezAT and both individual proteins using urea induced denaturation monitored by the red-shift in tryptophan fluorescence and circular dichroism. In these experiments a mutated tryptophan free PezT variant (PezT(D66T,W232Y)) was used, since both, wild type PezA and PezT contain a single tryptophan residue. Whereas W232 of PezT is solvent exposed, W111 of PezA is hidden in the hydrophobic core of the homodimer interface of the C-terminal three-helix bundle (Fig. 3b). Thus, a change in tryptophan fluorescence upon denaturation was expected to be more pronounced for PezA than for PezT and we could exclusively follow the equilibrium unfolding properties of this domain in absence and presence of PezT.

In absence of PezT(D66T,W232Y), the unfolding profile of the PezA three helix bundle shows a broad major transition, with a denaturation mid-point (D_{1/2}) at 4.1 M urea (Fig. 2a). Interestingly, the presence of PezT(D66T,W232Y) leads to a considerably steeper unfolding transition and the D_{1/2} increases to 4.9 M urea, indicating enhanced stability and a more pronounced cooperativity in unfolding of the PezA three-helix bundle (Fig. 2b). We note that we already observed a minor, protein concentration dependent red-shift of PezA tryptophan fluorescence at lower urea concentrations, indicating that the homodimer equilibrium of PezA is disturbed (Fig. S2).

When we monitored the global folding/unfolding equilibrium of PezA and PezAT by circular dichroism at 222 nm, we found a D_{1/2} at ~4.6 M urea for unbound PezA (Fig. 2b). Apparently, the fluorescence and CD response curves of PezA alone have different denaturation mid-points indicating the presence of a folding intermediate. Intriguingly, for the PezA/PezT(D66T,W232Y) complex we observe only a single but steep transition (D_{1/2} = 4.9 M) that overlays well with the major transition in tryptophan fluorescence of bound PezA as described above (Fig. 2c). This indicates that PezA shows different unfolding behaviour once bound to PezT and an unfolding intermediate occurs only for free PezA. However, both proteins unfold with high cooperativity once they are in complex. Additionally, free PezT(D66T,W232Y) already unfolds at much lower urea concentrations (D_{1/2} = 3.5 M) than the complex, indicating a reduced thermodynamic stability of the free toxin compared to the free antitoxin. This rather untypical finding for TA systems seem to be a distinct feature of the family of epsilon/zeta systems since it has also been reported for the zeta toxin from S. pyogenes (23). In summary, our results demonstrate that the proteolytic stability of the PezAT complex correlates with a strong increase in thermodynamic stability of both PezA and PezT once they are bound to each other. Consequently, the heterotetrameric assembly of the complex must provide both proteins with a strong increase in conformational rigidity.

Association of PezT and PezA leads to a conformational change of PezT - Our previous results prompted us to perform a detailed investigation of the dynamics of PezAT complex formation. Recent isothermal calorimetry experiments indicated a strong 1:1 interaction between PezA and PezT with an affinity in the sub-nanomolar range (11). In an initial tryptophan fluorescence titration experiment, we found that the tight binding event is accompanied by an overall increase in tryptophan fluorescence of the complex (Fig. 3a). We could confirm that the increased quantum-yield can be attributed to W232 of PezT, since titration experiments using the tryptophan free PezA(W111F) variant showed the same change as observed for wild type proteins. Additionally, our experiments could confirm the previously reported 1:1 stoichiometry, since the fluorescence only increased until equimolar concentrations were
reached (Fig. 3a). Thus, we can unambiguously assign the fluorescence increase to a conformational change around W232 of PezT. W232 is located in the loop between the two C-terminal helices of the toxin and, notably, not part of any protein contact interface. Therefore, the enhanced emission intensity observed can only be explained by a local conformational change in the environment of the fluorophore. It is worth mentioning that deletion of W111 in PezA decreased the total tryptophan fluorescence of the complex only marginally, indicating that this fluorophore is highly quenched.

PezT associates with PezA in a rapid two step mechanism - Since the absence of any curvature in the binding isotherm did not allow determination of a dissociation constant (Fig. 3a), we decided to measure binding kinetics using rapid mixing techniques. In these experiments, PezT(D66T) was mixed with PezA(W111F) at a set of different protein concentrations in the micromolar range using a stopped-flow device (Fig. 3c-e). This non-toxic PezT(D66T) variant was previously shown to have impaired toxic activity of PezT in vivo, however, the side chain of D66 is not involved in any PezA/PezT interaction (11).

Intriguingly, these experiments revealed an ultra-rapid formation of the complex, since the kinetic traces level-out within milliseconds at low micromolar concentrations. We collected a dataset of 13 time traces at stoichiometric and non-stoichiometric mixing-ratios and fitted them globally to different possible association pathways. Such global fitting procedures of different datasets have been shown to be adequate for elucidating sophisticated binding pathways (28,29). Surprisingly, the minimal association mechanism that describes the data sufficiently is a simplified, quasi-reversible two-step model (Scheme 1) that involves a conformational change of PezT (T) after binding to an empty PezA binding site (A_i)

\[ A_i + T \xrightarrow{\lambda_1} A_i \cdot T \xrightarrow{\lambda_2} A_i T^* \]

Scheme 1

where the apparent rate constant \( \lambda_1 \) describes formation of a transient complex \( A_i \cdot T \) and \( \lambda_2 \) describes a rapid conformational change of PezT upon formation of the stereospecific complex, leading to the observed increase in tryptophan fluorescence. The best fit values obtained for the experimental conditions are well defined (Fig. S3) and converge to 100 \( \mu M^{-1}s^{-1} \) for \( \lambda_1 \) and 410 \( s^{-1} \) for \( \lambda_2 \).

This association pathway is reminiscent to a generalized protein-protein association model for two binders, where formation of an initial transient complex, also often denoted as ‘encounter complex’, is followed by formation of the stereospecific complex (30,31). Notably, convergence of the fit to the data was already obtained using this simplified, quasi-reversible two-step model and including any additional putative rate constant accounting for the oligomeric state of PezA or a transiently occurring trimeric intermediate etc. was not required. This finding together with the fact that the initial binding rate \( \lambda_1 \) exceeds the expected value for diffusion controlled protein-protein association, suggested the presence of long-range electrostatic interactions between PezA and PezT.

The PezAT complex forms a stable heterotetramer in solution - Any possible disassembly factor which is capable of separating PezA from PezT would also need to interfere with the homodimer interface of PezA, since dimeric PezA is sandwiched between two PezT molecules in the heterotetrameric complex (see (11) and Fig. 3b). We therefore investigated the kinetics of PezA homodimerization and asked whether a PezAT heterotetramer or a PezAT heterodimer is the dominating oligomeric state in solution. In relaxation experiments we could monitor the change in tryptophan fluorescence of the highly quenched tryptophan W111 of PezA which is located in the homodimer interface. Briefly, either 5 \( \mu M \) of PezA or PezAT complex (but using the tryptophan free PezT variant) were rapidly diluted to various concentrations using a stopped-flow device (Fig. 4a, b). Global fit analysis of the resulting relaxation kinetics revealed that dimerization of PezA and PezAT heterodimers can be described by a simple one-step dimerization mechanism. We found that PezA alone forms homodimers with a \( K_d \) of 16 \( nM \) following moderate
association \( (k_{on} = 6.3 \, \mu M^{-1} s^{-1}) \) and dissociation \( (k_{off} = 0.1 \, s^{-1}) \) kinetics. In contrast, we observed a slightly decreased affinity of PezA homodimers once bound to PezT resulting in a \( K_d \) of 60 nM. Whereas, the dissociation rate constant increased to \( k_{off} = 0.3 \, s^{-1} \), the association rate constant was only reduced modestly to \( k_{on} = 5.3 \, \mu M^{-1} s^{-1} \). Apparently, the homodimer is slightly destabilized in the PezAT complex but assuming that PezA and PezT have similar cytosolic concentrations as reported for the related epsilon/zeta system (23), we speculate that a heterotetramer is the predominating oligomeric state \( \text{in vivo} \). Thus, any factor that can separate PezT from PezA needs to interfere with the heterotetrameric PezA/PezT arrangement.

PezAT association is electrostatically enhanced - Several protein-protein complexes have been described, where complementary electrostatic interactions strongly accelerate association kinetics (32-35). Since we observed association kinetics that are faster than those of a purely diffusion controlled process, we expected PezAT complex formation to be dependent on the ionic strength of the solute. Our assumption was further supported by calculations of electrostatic surface potentials of PezA and PezT using APBS (36-38), which showed that the PezT binding interface of PezA has an overall highly negative surface charge, while the cognate interface of PezT is complementarily positively charged. (Fig. 5a).

The existence of intermolecular electrostatic forces can be verified by measuring the rate dependencies of interactions on the ionic strength of the solute. To probe the PezAT toxin antitoxin association for such an electrostatic enhancement, we measured formation of 1 \( \mu M \) PezAT complex at increasing concentrations of NaCl. Plotting \( \lambda_1 \) versus the ionic strength of the solute verified the electrostatic interaction experimentally (Fig. 5b). Indeed, \( \lambda_1 \) is \( \sim 230 \, \mu M^{-1} s^{-1} \) at 0.05 M NaCl and decreases by more than an order of magnitude to less than 15 \( \mu M^{-1} s^{-1} \) at 1 M NaCl. On the other hand, the second observed rate constant \( \lambda_2 \) seems to be affected only mildly by varying the salt concentration. It retained close to a value of 400 \( s^{-1} \) at 0.1 to 0.4 M NaCl and remained fast but indeterminable at higher salt concentrations due to the increasing rate limitation by \( \lambda_1 \). This is reminiscent to the Colicin E9-Im9 complex, where the rate of the second association step is not grossly affected by the ionic strength of the solute indicating that \( \lambda_2 \) is mostly dependent on short-range interactions (35).

The salt dependency of the rate describing the formation of transient protein-protein complexes can be well explained on the basis of Debye’s treatment of the association rates of charged spheres (39-41)

\[
\ln(k(I)) = \ln(k(\infty)) - \frac{U}{kT} \left( \frac{e^{-\kappa(r-a)}}{1+\kappa a} \right)
\]

Equation 1

where \( k(\infty) \) is the rate at infinity ionic strength, \( U \) the interaction energy between the two proteins at zero ionic strength and \( \kappa \) the Debye-Hückel screening parameter directly dependent on the ionic strength of the solute. According to Gabdoulline and Wade, \( a \) represents the double distance between the protein charges relevant for association and the ion centers in the solvent (\( \sim 6 \) Å for NaCl) and \( r \) the relative separation of the protein charges in the transient complex (42). When fitting the salt dependency of \( \lambda_1 \) using Equation 1, we obtain a value of \( \sim 7 \) Å for \( r \) indicating that PezA and PezT are already in very close proximity prior to the second binding step. Thus, in the transient complex both proteins are most probably in close proximity and in a conformation which is similar to the final stereospecific complex. This is corroborated by the quasi-irreversibility of the first reaction step of the global fit.

The PezAT complex has femtomolar affinity at physiological ionic strength - Anticipated from the fast association kinetics, we were interested in the dissociation kinetics of the PezAT complex. Initial experiments hinted at a very slow toxin exchange rate from the complex. Since such kinetics are difficult to investigate by conventional stopped-flow measurements, we established a long-time displacement assay using Alexaass-labelled
Briefly, PezT(D66T) Alexa488 in complex with PezA(W111F) was incubated with a 20 fold excess of unlabelled PezT(D66T) over a time period of two days. Samples taken at individual time points were separated by size exclusion chromatography. The integrity of the complex was verified by SDS-PAGE analysis after long time storage in the buffer system used in the experiment (data not shown). Exchange kinetics were monitored by the observed decrease of the fluorescence signal of the PezAT complex due to displacement of labelled PezT(D66T) by unlabelled species and an increase of the fluorescence signal of labelled monomeric PezT (Fig. 6a). Both, occurrence of detectable free PezT(D66T) and disappearance of the complex could be fitted with a mono-exponential yielding an overall dissociation rate constant of PezT replaced from the PezAT heterotetramer with $\lambda_{\text{off}} = 6.5 \cdot 10^{-6}$ s$^{-1}$ resulting in an average half-life of $t_{1/2} \approx 30$ h for the PezAT interaction (Fig. 6b).

According to our findings, PezA forms homodimers with nanomolar affinity that are rapidly attracted to unbound PezT due to complementary electrostatic charges. After formation of an initial, transient complex, a fast rearrangement locks the PezAT assembly in a predominantly heterotetrameric state.

This outstanding femtomolar affinity of PezA and PezT is the strongest reported among all TA systems, which have been characterized so far. Intriguingly, other TA systems like PhD/Doc (44), RelB/RelE (45) and CcdB/CcdA (46) have been described to have affinities 3 to 8 orders of magnitudes lower than PezAT. Even for the PezAT homologue epsilon/zeta just a micromolar affinity has been reported (23).

In summary, PezAT is rather reminiscent to endotoxin-inhibitor complexes like barnase/bastar (34) or members of the colicin/immunoprotein family (33,35), which are not meant to dissociate in the host cell but kill susceptible target cells after secretion.

The difference in affinity within the family of epsilon/zeta TA systems is somewhat unexpected, because given the structural similarity of PezT and zeta (11,13), one can expect that both proteins have a similar toxic function. Furthermore, site directed mutagenesis studies have shown that conserved amino acids are involved in toxicity, although the target of PezT and zeta is still enigmatic. A probable explanation for these contradicting findings might be different cellular missions of PezAT and epsilon/zeta. Whereas disruption of the PezT open reading frame in S. pneumoniae resulted in impaired infectiousness of the pathogen, which implies that PezAT acts as a virulence factor (16), epsilon/zeta was reported to be a classical TA system ensuring stable inheritance of the plasmid pSM19035 (14). It seems plausible that toxin release rates have been fine tuned and adapted to perform different cellular tasks in their specific host.

DISCUSSION

In this study we investigated the overall stability of the chromosomally encoded PezAT TA-system from the human pathogen S. pneumoniae. Our data suggest an intriguing resistance of the inhibitory complex towards proteolysis in E. coli and B. subtilis preventing release of toxin under the conditions tested. Results from in vitro unfolding experiments further indicated that there is a tight interaction between toxin and antitoxin that has a great impact on the thermodynamic stability of both proteins. Figure 7 summarizes the detailed kinetic analysis of the PezAT interaction determined using rapid mixing methods and time-resolved size exclusion chromatography.
toxin release by simple dissociation. Additionally, it is unlikely that PezT release upon simple dissociation is caused by proteolytic removal of the transcriptional repressor domain of PezA, since the C-terminal domains binds to PezT with comparable affinity as full length protein (Fig. S4). Our results rather suggest that specific degradation of PezA from the complex either requires a host specific protease, which might be absent in the organisms used in this study, or a triggered activation of an adaptor molecule by an external queue required for proper recognition of PezA by a constitutive AAA+ protease.

It is tempting to speculate that the absence of “leakiness” in PezT inhibition renders the host-cell capable to actively control toxin release. This could be in contrast to other TA systems like epsilon/zeta, CcdA/CcdB, mazE/mazF, RelE/RelB or PemI/PemK, which have been shown to respond directly to loss of either the TA open reading frame or general stress conditions that impair protein translation (14,19,47-50). On the other hand, examples of TA systems have been reported to require bacterial signalling pathways in order to carry out sophisticated functions like programmed cell death upon fruit-body formation in Myxococcus xanthus or growth regulation by quorum sensing in E. coli (5,8). Moreover, TA systems are key players during persister cell formation and the development of bacterial biofilms (51-54). Given the observed lack of proteolytic degradation and the reported thermodynamic properties of the PezAT system, PezAT most likely is similar to the latter described type of TA systems.

In general, activation of any TA system will enable new therapeutic approaches in our antimicrobial defence. Thus, understanding of the assembly and disassembly pathways and kinetics is of crucial importance for such challenging goals. However, in case of PezAT activation of toxin by antitoxin-displacement by peptidic mimics as recently suggested for other TA systems (55,56) might not be feasible.
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**FOOTNOTES**

The abbreviations used are: TA, toxin-antitoxin; Pez, pneumococcal epsilon-zeta; DTE, dithioerythritol; CD, circular dichroism; ECL, enhanced chemiluminescence; HRP, horseradish peroxidase; 1,5-IAEDANS, 5-([idoacetyl]amino)ethyl]amino)naphthalene-1-sulfonic acid; AAA, ATPases associated with various cellular activities

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FIGURE LEGENDS

Figure 1. In vivo and semi-in vivo stability of PezAT (A) Degradation pattern of PezAT after PezAT expression and subsequent translation inhibition using chloramphenicol (Cm). E. coli DH5α cells bearing pASK5+(pezAT) (black) were induced with 0.2 µg/mL anhydrotetracycline at an OD$_{600}$ of ~ 0.5 (blue arrow). After one hour Cm was added leading to inhibition of protein expression (red arrow). The growth profiles of a control culture bearing pASK5+(pezA) demonstrated that growth inhibition was caused by Cm only are shown in light grey. Samples with equivalent amounts of cells were analyzed using SDS-gel electrophoresis to monitor PezA and PezT degradation (lower panel). (B) Growth profiles subsequent to removal of anhydrotetracycline after 1 hour of expression of PezAT. E. coli DH5α cells bearing either pASK5+(pezAT) (black) pASK5+(pezA) (gray) were induced with 0.2 µg/mL at an OD$_{600}$ of ~ 0.5. Fresh LB-medium was inoculated to a final OD$_{600}$ of ~ 0.1 and cell growth was monitored. Cell cultures in which PezAT expression was abolished after removal of inducing agent show similar growth profiles the control. (C) Degradation of 1 µM of the antitoxin His-F-PezA in B. subtilis raw extract. His-F-PezA was either monitored by ECL western blot analysis using Anti-His$_5$ (left) or imaging of IAEDANS fluorescence (right). (D) Degradation of 1 µM of the toxin PezT(D66T) in B. subtilis raw extract monitored by western blot analysis as in (C). (D) Degradation of 1 µM His-F-PezA/PezT(D66T) complex in B. subtilis raw extract similar as in (C).

Figure 2. Urea induced denaturation experiments (A) Unfolding of 5 µM PezA in absence (white circles) and presence (grey circles) of 5 µM PezT(D66T,W232Y) monitored by the red-shift of the spectral centre of mass of tryptophan fluorescence. (B) PezA denaturation (5 µM) monitored either by tryptophan fluorescence (white circles) or by the mean residue ellipticity at 222 nm (white triangles). (C) Denaturation of PezA (5 µM) in presence of equimolar concentrations of PezT(D66T,W232Y) monitored by tryptophan fluorescence (grey circles) and denaturation of the complex by mean residue ellipticity at 222 nm (grey triangles). (D) Denaturation of 5 µM PezT(D66T,W232Y) alone (white diamonds) and in presence of 5 µM PezA (gray diamonds) monitored by mean residue ellipticity at 222 nm. Note that individual values have been normalized in panel D.

Figure 3. Tryptophan fluorescence experiments of PezA and PezT upon complex formation. (A) Fluorescence titrations of 3 µM PezT(D66T) with wildtype PezA (dark grey circles) or tryptophan free PezA(W111F) (light grey circles) or buffer (white circles). Excitation was set to 295 nm and emission recorded at 347 nm. (B) Ribbon representation of the PezAT heterotetramer of PezA (red) and PezT (blue). Wild type tryptophan residues W232 of PezT (orange) and W111(green) are highlighted as stick models. Whereas W232 enables monitoring binding of PezA, W111 of PezA was shown to be sensitive to changes in the homodimer interface of the antitoxin. (C) Exemplary real time trace of binding of 1 µM PezT(D66T) to 1µM PezA(W111F) (grey) measured in a stopped-flow device. Tryptophan fluorescence of PezT(D66T) was excited at 296 nm and emission recorded with a 320 nm long-pass filter. The corresponding fit according to the two-step model is shown as black line. (D) Similar as (C) but 5 µM PezT(D66T) and 5 µM PezA(W111F) (E) Similar as (C) but 3 µM PezT(D66T) / 1.5 µM PezA(W111F). Note the difference in scaling of the time-axis.

Figure 4. Relaxation kinetics of the PezA homo-dimerization measured by tryptophan fluorescence. (A) Exemplary time trace of a rapid 1:15 buffer dilution at 10 µM PezA (grey) and the monoexponential fit describing a one step dimerization mechanism. (B) Exemplary relaxation time traces of the PezAT heterodimer / heterotetramer equilibrium measured by rapid 1:15 buffer dilution at 10 µM PezA / PezT(D66T,W232Y) complex (grey) and the corresponding fit (black). Protein fluorescence was excited at 280 nm and emission recorded with a 320 nm long-pass filter.

Figure 5. PezA and PezT binding is accelerated by electrostatic interactions. (A) Open-book representation of the PezA and PezT binding interface showing complementary charged surface potentials.
at moderate ionic strength. Charge distributions have been calculated using the Pymol APBPS/PDB2PQR at 0.2 M NaCl. 

(B) Dependency of the apparent rate constants $\lambda_1$ (dark grey) and $\lambda_2$ (light grey) on the ionic strength. The black line corresponds to the fit of $\lambda_1$ using Equation 1 with a fixed value of 6 Å for $a$. Best fit values were -9.5 for $k/\kT$ and $1 \cdot 10^6$ M$^{-1}$ s$^{-1}$ for $k(\infty)$.

**Figure 6.** Dissociation kinetics of the PezAT complex measured by exchange kinetics. 0.3 µM of preassembled PezA(W111F)/PezT(D66T)-Alexa$_{488}$ was incubated with 3 µM unlabelled PezT(D66T). 

(A) The decrease of fluorescent PezAT species and increase of free, labelled PezT was monitored by analytical gel filtration. (B) Data after peak integration and normalization were fit in parallel with a mono-exponential, yielding an apparent dissociation constant of $6.5 \cdot 10^{-6}$ s$^{-1}$.

**Figure 7.** Model of the PezAT assembly dynamics. The onset of PezAT heterotetramer formation is driven by rapid association of PezT with either PezA monomers or dimers. This initial association step is based on long-range electrostatic interactions resulting in a transient PezAT complex in which both proteins are still separated by ~5 Å. A subsequent rapid rearrangement eventually leads to the final stereospecific inhibitory PezAT complex, which is predominantly a heterotetramer due to the nanomolar affinity of the central PezA homodimer interface.
Figure 1

A

B

C

D

E
Figure 2
Figure 3

A) Graph showing relative fluorescence change against molar ratio.

B) Molecular structure with labeling of W232 and W111.

C) Graphs showing fluorescence signal over time.

D) Graph showing time course of fluorescence.

E) Graph showing time course of another signal.
Figure 4
Figure 5

A

-3 kT/e to +3 kT/e

B

 Ionic strength (M)

$\lambda_1$ (μM$^{-1}$ s$^{-1}$) and $\lambda_2$ (s$^{-1}$)
Figure 6
Assembly dynamics and stability of the Pneumococcal epsilon zeta Antitoxin Toxin (PezAT) system from Streptococcus pneumoniae.
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