



The periplasmic transaminase PtaA of *Pseudomonas fluorescens* converts the glutamic acid residue at the pyoverdine fluorophore to α -ketoglutaric acid

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The periplasmic conversion of ferribactin to pyoverdine is essential for siderophore biogenesis in fluorescent pseudomonads, such as pathogenic *Pseudomonas aeruginosa* or plant growth-promoting *Pseudomonas fluorescens*. The non-ribosomal peptide ferribactin undergoes cyclizations and oxidations that result in the fluorophore, and a strictly conserved fluorophore-bound glutamic acid residue is converted to a range of variants, including succinamide, succinic acid, and α -ketoglutaric acid residues. We recently discovered that the pyridoxal phosphate-containing enzyme PvdN is responsible for the generation of the succinamide, which can be hydrolyzed to succinic acid. Based on this, a distinct unknown enzyme was postulated to be responsible for the conversion of the glutamic acid to α -ketoglutaric acid. Here we report the identification and characterization of this enzyme in *P. fluorescens* strain A506. *In silico* analyses indicated a periplasmic transaminase in fluorescent pseudomonads and other proteobacteria that we termed PtaA for “periplasmic transaminase A.” An in-frame-deleted *ptaA* mutant selectively lacked the α -ketoglutaric acid form of pyoverdine, and recombinant PtaA complemented this phenotype. The *ptaA/pvdN* double mutant produced exclusively the glutamic acid form of pyoverdine. PtaA is homodimeric and contains a pyridoxal phosphate cofactor. Mutation of the active-site lysine abolished PtaA activity and affected folding as well as Tat-dependent transport of the enzyme. In pseudomonads, the occurrence of *ptaA* correlates with the occurrence of α -ketoglutaric acid forms of pyoverdines. As this enzyme is not restricted to pyoverdine-producing bacteria, its catalysis of periplasmic transaminations is most likely a general tool for specific biosynthetic pathways.

Iron ions play important roles in all living organisms, but in many habitats their availability is limited due to low solubility of Fe(III) oxide hydrates (1). Pyoverdines are siderophores that permit growth of many pseudomonads under iron limitation. Examples include pathogenic *Pseudomonas aeruginosa* as well

as non-pathogenic or even plant-growth promoting *Pseudomonas fluorescens* strains (2). All pyoverdines originate from non-ribosomally synthesized peptides that are translocated into the periplasm where a quinoline fluorophore is formed (3, 4). The many known “fluorescent pseudomonads” produce a multitude of distinct pyoverdine variants that differ generally in the sequence of the peptide chain and the nature of a fluorophore-attached residue that is synthesized as a glutamic acid and usually converted to succinamide, succinic acid, or α -ketoglutaric acid and in rare cases to malamide and malic acid (4). While the formation of the fluorophore is known to depend on the activity of the periplasmic enzyme PvdP (5), the chemical modification of the glutamic acid that is amide-bonded to the fluorophore via its γ -carboxylic acid has long been puzzling.

We recently revealed that the periplasmic enzyme PvdN is responsible for the direct conversion of the glutamic acid to succinamide by a novel PLP²-catalyzed reaction (6). From these findings, it was concluded that there had to exist a yet unknown transaminase in the periplasm that should, in competition with the PvdN-catalyzed succinamide formation, convert the glutamic acid residue to α -ketoglutaric acid (6). Here we demonstrate that this enzyme indeed exists. Initially, we identified *in silico* a unique candidate that was predicted to possess all required characteristics. The cofactor content, transport, and quaternary structure of this enzyme were experimentally confirmed. We inactivated the corresponding gene by scarless in-frame deletion and showed that the α -ketoglutaric acid variant was indeed absent in the mutant strain, a phenotype that was fully complemented in *trans*. Further analyses indicate that in pseudomonads the occurrence of this transaminase correlates with α -ketoglutaric acid-containing pyoverdines. However, there is strong evidence for additional functions of this transaminase in the periplasm, which is why we term this enzyme “periplasmic transaminase A” (PtaA).

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This article contains supplemental Figs. S1 and S2.

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² The abbreviations used are: PLP, pyridoxal phosphate; CAS, chrome azurol S; EDDHA, ethylenediamine di(*o*-hydroxy)phenylacetic acid; IEF, isoelectric focusing; ONC, overnight culture; Tat, twin-arginine translocation; UPLC, ultraperformance liquid chromatography; PVD, pyoverdine; DSC, differential scanning calorimetry; SEC, size-exclusion chromatography; MALS-RI, multiangle light scattering/refractive index; CAA, casamino acid; a.m.u., atomic mass units.

Table 1**Potential Tat substrates in *P. fluorescens* A506**

Potential Tat substrates were identified with TATFIND v.1.4 (7) and classified using the InterPro web service (8, 9). ABC, ATP-binding cassette.

Locus tag	Classification
PflA506_0002 ^a	DNA polymerase III, β -chain
PflA506_0050	Gluconate 2-dehydrogenase subunit 3
PflA506_0159	Pectin lyase/hemagglutinin
PflA506_0418	Flavin-dependent halogenase
PflA506_0574	Leu/Ile/Val-binding protein/urea ABC transporter, substrate-binding protein UrtA-like
PflA506_0673	Acid phosphatase, AcpA
PflA506_0796	Alkaline phosphatase D-related/metallo-dependent phosphatase-like
PflA506_0832	Rieske iron--sulphur protein
PflA506_0959	Mannose-1-phosphate guanylyltransferase/mannose-6-phosphate isomerase
PflA506_1047	Multicopper oxidase (CumA)
PflA506_1182	Glucan biosynthesis, MdoD
PflA506_1897	Alkaline phosphatase D-related/metallo-dependent phosphatase-like
PflA506_2032 ^a	Hypothetical protein/no classification possible
PflA506_2213	Oxidoreductase molybdopterin-binding subunit, LorB-related
PflA506_2582	Deferochelatase/peroxidase EfeB
PflA506_2836	Periplasmic binding protein-like II/Aliphatic sulfonate-binding protein-related
PflA506_2902	Dienelactone hydrolase
PflA506_3071	Copper chaperone SCO1/SenC
PflA506_3083	PvdP (uncharacterized domain, dicopper center, tyrosinase family protein)
PflA506_3085	PvdN (aminotransferase class V domain)
PflA506_3359	Leucine-binding protein domain
PflA506_3490	Aldehyde oxidase/xanthine dehydrogenase, molybdopterin-binding
PflA506_3831	Copper-resistance protein CopA
PflA506_4010 ^a	3-Oxoacyl-[acyl-carrier-protein] synthase 2
PflA506_4424	Aminotransferase, class I/class II
PflA506_4509	Oxidoreductase, molybdopterin-binding domain (YedY)
PflA506_4922	Six-bladed β -propeller, TolB-like/protein of unknown function, DUF839
PflA506_5086	Hypothetical protein/no classification possible
PflA506_5390	N-Acetylmuramoyl-L-alanine amidase

^a This hit might represent a false positive in the opinion of the authors.

Results***In silico* prediction of a pyoverdine-modifying periplasmic transaminase in *P. fluorescens* A506**

Recent genetic and biochemical analyses of our model strain *P. fluorescens* A506 indicated the existence of a yet unknown periplasmic enzyme that converts the fluorophore-attached glutamic acid residue of its pyoverdine (PVD_{A506}) into α -keto-glutaric acid (6). We approached the identification of this enzyme initially by combining *in silico* tools, anticipating that the conversion should be carried out by a PLP-containing transaminase, and like the PLP-containing PvdN (6), this enzyme should be Tat-dependently transported into the periplasm.

A TATFIND v.1.4 analysis (7) predicted the presence of 29 putative Tat substrates in the proteome of *P. fluorescens* A506 of which three are most likely false positives. The functional annotation based on the InterPro web service (8, 9) indicated that two of these predicted Tat substrates were sequence-related to transaminases, namely PvdN and PflA506_4424 (Table 1). PvdN has been structurally and functionally characterized (6, 10). It contains a PLP cofactor and catalyzes the unusual amine-retaining oxidative decarboxylation of the fluorophore-attached glutamic acid residue of pyoverdine to succinamide. PvdN thus has already been demonstrated not to be involved in the searched-for transamination reaction. Therefore, the only remaining candidate was PflA506_4424, which we will hereafter term PtaA to avoid confusion and to facilitate reading.

To gain a better understanding of PtaA's properties, we searched all genomes contained in the *Pseudomonas* Genome Database (11) for homologs of PtaA. To reduce the bias by over-represented sequences within the identified homologs, we performed a redundancy reduction with an identity threshold of

0.95 utilizing the CD-HIT web service (12). Thereafter, the sequences were aligned using the T-Coffee web service (13), and the multiple sequence alignment was visualized as sequence logo with WebLogo 3 (14). From the results of these analyses, it can be inferred that Tat signal peptides are common in PtaA homologs (Fig. 1A). Additionally, the initial classification of PtaA by the InterPro web service (8, 9) allowed us to identify the active-site lysine residue Lys-224 in a highly conserved sequence pattern (TFSK(I/L)YG(M/L)AGAR), which is highlighted in the sequence logo (Fig. 1B). Furthermore, the InterPro predictions suggested that PtaA likely forms a homodimer.

PtaA is a periplasmic homodimeric enzyme that requires cytoplasmic cofactor assembly for folding and transport

To identify the predicted PLP cofactor, PtaA was overproduced in its mature form in *Escherichia coli* and purified as described under "Experimental procedures." By means of a PLP binding assay, we removed the PLP cofactor from PtaA and detected the PLP oxime via its fluorescence (Fig. 2). The predicted periplasmic localization was addressed in complemented *P. fluorescens* mutant strains that were constructed for subsequent functional analyses (described below). We carried out an in-frame deletion of the gene encoding PtaA in the wild-type *P. fluorescens* A506 strain as well as in a Δ pvdN deletion strain that was generated previously (6). In *P. fluorescens* A506, the *ptaA* gene is flanked by genes encoding a predicted DNA-binding protein upstream in the opposite direction and a putative autoinducer-binding transcriptional regulator downstream in the same direction. The in-frame deletion removed the sequence from codons 5 to 357 of the coding region to avoid any potential polar effects.

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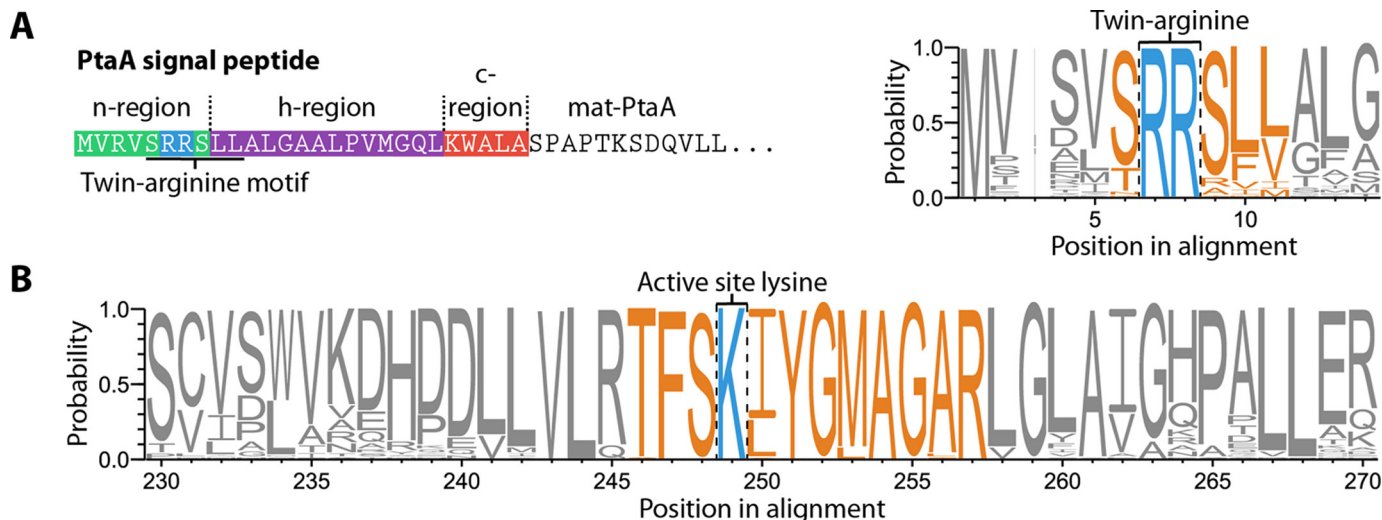


Figure 1. Analysis of the N-terminal Tat signal peptide of PtaA and its conserved PLP-binding site, the lysine residue Lys-224. *A*, depiction of the N-terminal Tat signal peptide of PtaA containing the eponymous twin-arginine motif (ZRRXΦΦ; Z denotes a polar amino acid, R is an L-arginine residue, X is any amino acid, and Φ is a hydrophobic amino acid; Ref. 30). Furthermore, a subsection of the sequence logo of the signal peptide region demonstrates the conserved twin-arginine motif. *B*, sequence logo of the active-site lysine, Lys-224, and the sequence in its vicinity. The process to generate the sequence logo is detailed under “Experimental procedures.”

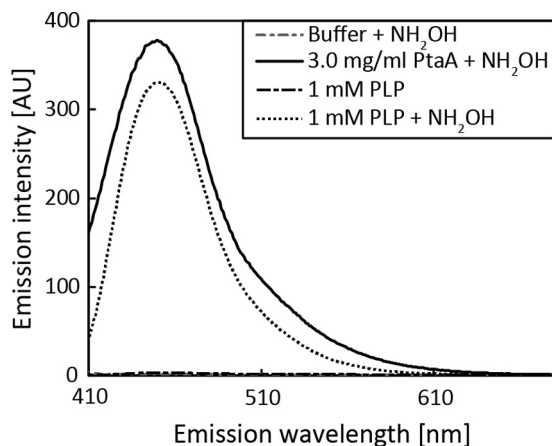


Figure 2. PLP binding assay. Shown are the fluorescence spectrum of the PLP oxime as generated by transamination with hydroxylamine with mature PtaA purified from *E. coli* (continuous line), the reference spectrum of the PLP oxime (dotted line), and buffer as well as PLP negative control spectra (dashed lines), indicating that PtaA binds PLP. Due to different sensitivity modes, spectra are to be qualitatively compared. AU, arbitrary units.

With the $\Delta ptaA$, $\Delta pvdN$, and $\Delta ptaA/\Delta pvdN$ strains in our hands, we constructed complementation vectors for recombinant production of functional PtaA or a PtaA(K224A) variant that is expected to abolish the PLP-dependent activity. The constructs were C-terminally tagged with a Strep-tag II to facilitate detection and purification.

Western blot detection in subcellular fractions proved that PtaA is indeed localized in the periplasmic fraction. Interestingly, the PtaA(K224A) variant accumulated in the membrane fraction, suggesting folding defects that are known to influence the efficiency and accuracy of Tat-dependent translocation (Ref. 15 and Fig. 3A). The precursor form still carrying the N-terminal signal peptide could be distinguished from the mature form in which the signal peptide was cleaved off. These observations were made with both the single- and the double-mutant strains. To examine whether folding defects were caused by the K224A exchange that could influence transport,

the protein was analyzed by differential scanning calorimetry (DSC) (Fig. 3C). The proteins used for this study were purified via affinity chromatography and subsequent size-exclusion chromatography (SEC). A Coomassie-stained SDS-PAGE gel of the purified proteins is depicted in Fig. 3B. The DSC data indicated that covalent PLP binding stabilized the overall protein structure to some extent since the unfolding transition temperature of the PtaA(K224A) variant was lowered from 70.8 to 69.0 °C in comparison with the wild-type PtaA. More importantly, a folded subdomain that gives rise to a minor peak at approximately 60 °C was almost absent in the PtaA(K224A) variant, which can be explained by a stabilization of the binding pocket in wild-type PtaA by the covalently attached PLP cofactor. To assess whether PtaA forms a homodimer, we analyzed the molecular size via SEC coupled with static multiangle light scattering/refractive index (MALS-RI) detectors as detailed under “Experimental procedures.” The experiment allowed us to calculate the approximate molecular mass from the data obtained (Fig. 3D). Since the size of mature monomeric PtaA and PtaA(K224A) (with the Strep-tag II) amounts to 37.3 and 37.2 kDa, respectively, the measured molecular masses of 73.2 ± 1.0 kDa for PtaA and 71.4 ± 1.1 kDa for PtaA(K224A) correlate well with the calculated molecular masses of the dimers when the influence of protein shape on MALS-derived values is considered. From these data, it can be concluded that both the wild-type PtaA and the PtaA(K224A) variant form homodimeric structures.

PtaA is not essential for the overall pyoverdine production and physiological function

The influence of the $\Delta ptaA$ deletion on formation/export of PVD_{A506} was assessed by means of a semiquantitative pyoverdine production assay (Fig. 4A). Furthermore, in the same assay, we included the analysis of $\Delta pvdN$ single- and $\Delta ptaA/\Delta pvdN$ double-deletion strains and the above mentioned in *trans* complementation systems in the analyses. The results obtained (Fig. 4A) demonstrate that the deletions do not affect

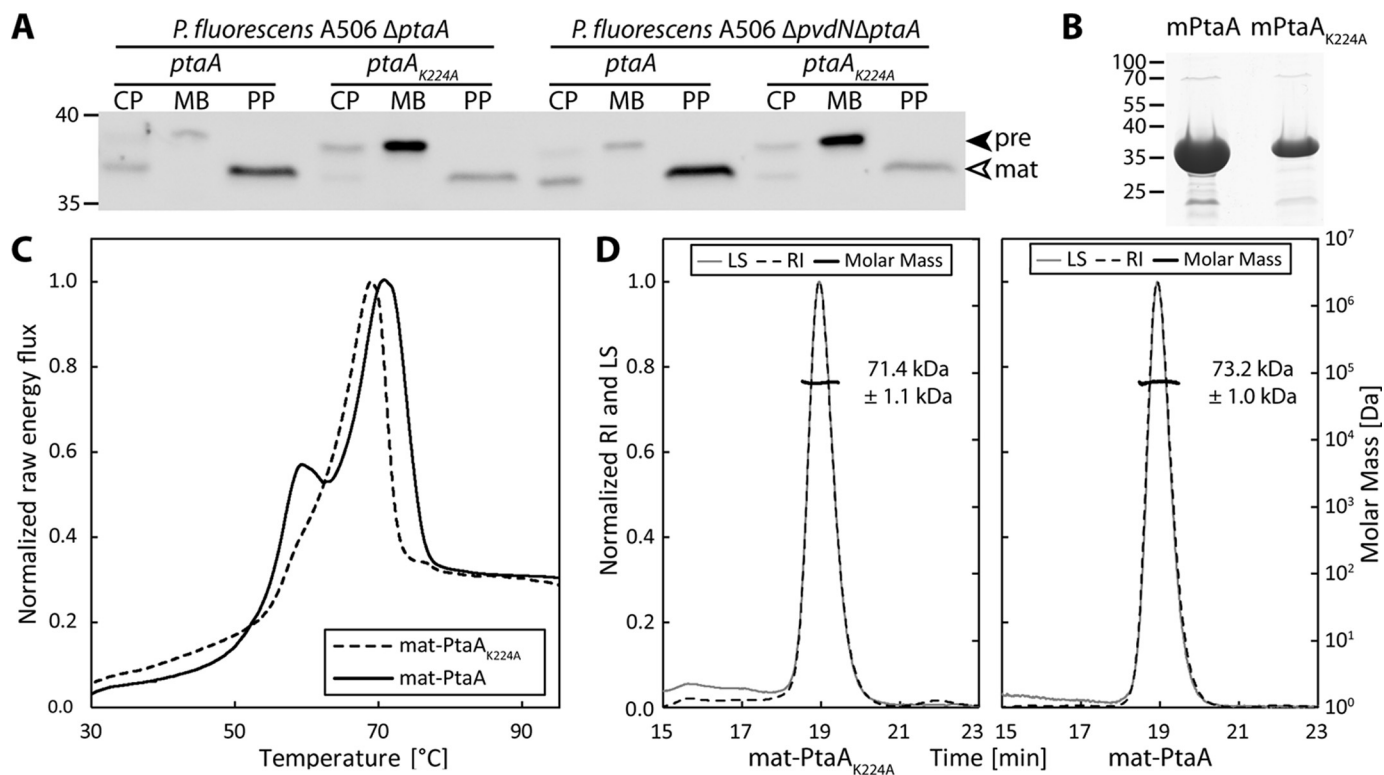


Figure 3. PtaA is periplasmically localized and forms a homodimer that is stabilized by covalent PLP binding. A, SDS-PAGE/Western blot analysis of subcellular fractions of *P. fluorescens* A506 deletion strains carrying plasmids for the production of the indicated C-terminally Strep-tagged PtaA variants. PtaA was detected using tag-specific antibodies. CP, cytoplasm; MB, membrane; PP, periplasm; pre, PtaA precursor; mat, mature PtaA. B, Coomassie stained SDS-PAGE gel of the purified mature PtaA (mPtaA) variants used for the DSC experiment depicted in C. C, DSC analysis of PtaA and PtaA(K224A). D, SEC-MALS-RI molecular mass determination of the indicated PtaA variants, indicating the dimeric structure of the enzyme. LS, light scattering; RI, refractive index. Data in C and D are normalized to the global maximum of the respective data set. In A and B, masses (in kDa) of standard proteins are indicated on the left.

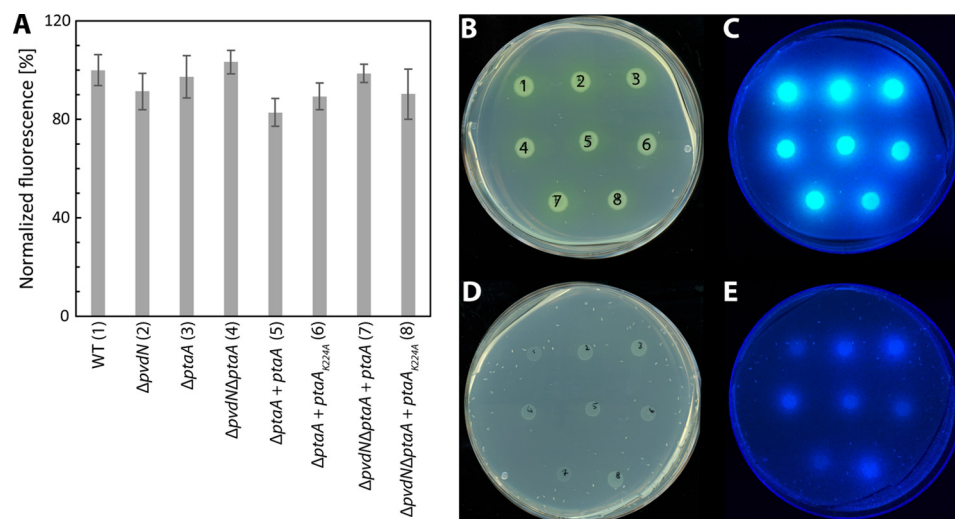


Figure 4. Semiquantitative pyoverdine production and iron binding assay for all *P. fluorescens* A506 strains used in this study. A, relative quantification of pyoverdine production in liquid culture, normalized to *P. fluorescens* A506 (WT) levels. The error bars indicate the standard deviation calculated from triplicates. B–E, droplet assays of all *P. fluorescens* A506 strains used in this study as indicated by numeric notation (1–8) in A. The incubation was performed either in the absence (B and C) or presence of EDDHA (D and E). The same plates were scanned (B and D), or fluorescence was detected on a UV table (C and E).

the overall pyoverdine production. Certainly, this type of assay did not differentiate between predicted differences in the side chain at the chromophore. To exclude that potential side-chain modifications impair iron affinity or pyoverdine import, iron binding and import were examined qualitatively by growth experiments (Fig. 4, B–E). In agreement with the

formation of functional siderophores, the mutant strains were able to grow similarly well on pyoverdine-inducing casamino acid (CAA) medium (Fig. 4, B and C), and even in the presence of 0.5 g/liter iron-depleting chelator ethylenediamine di(o-hydroxy)phenylacetic acid (EDDHA) (Fig. 4, D and E), growth was normal, indicating physiological sidero-

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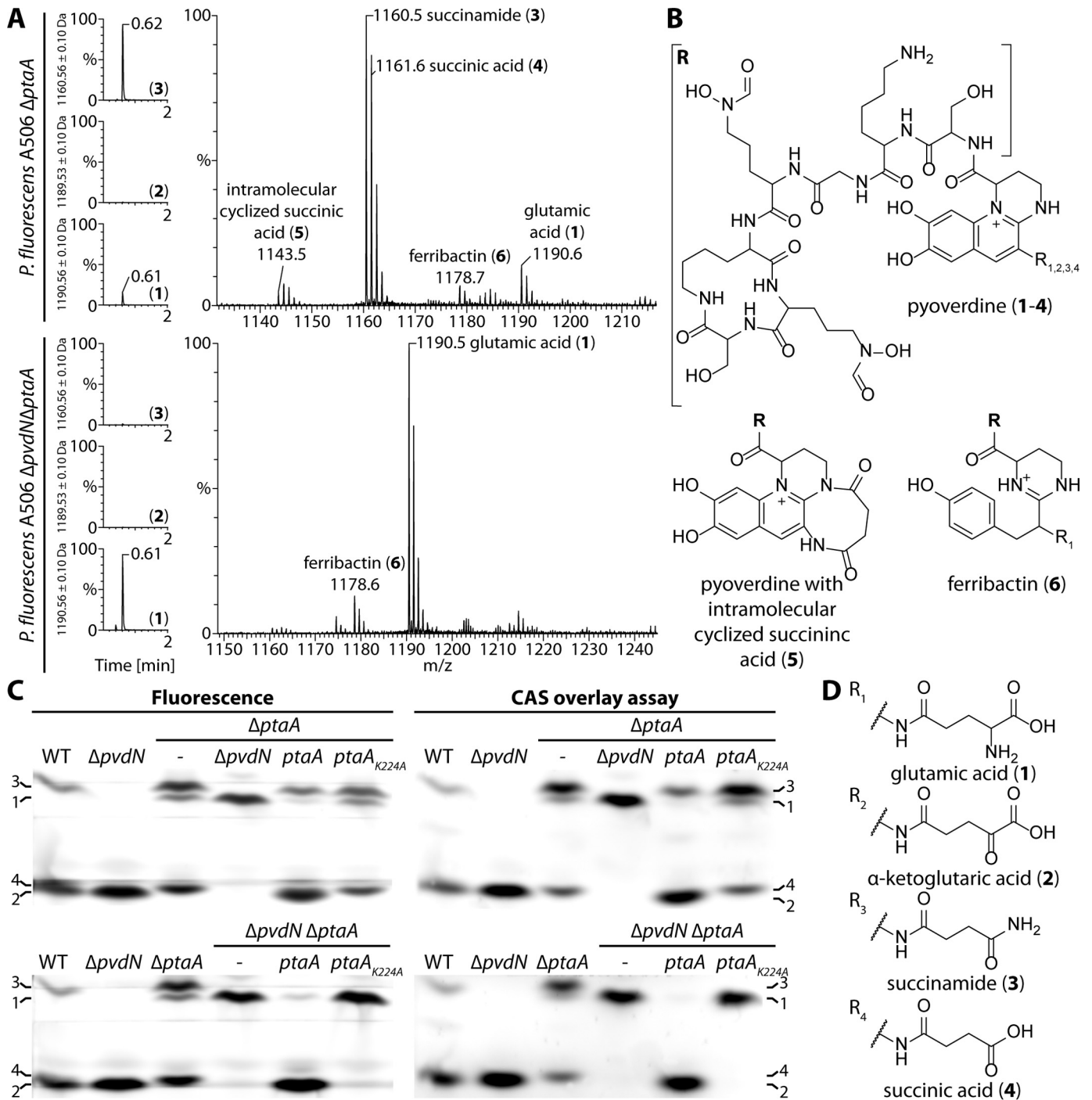


Figure 5. The enzyme PtaA is the periplasmic transaminase that converts the glutamic acid side chain of pyoverdine into the α -ketoglutaric acid variant. A, mass spectrometry results showing the UPLC chromatograms filtered for the masses of the succinamide, α -ketoglutaric acid, and glutamic acid variants on the left. On the right, the corresponding mass spectra of the peak at 0.6 min are depicted. The mass peaks were assigned to the structures shown in B and D. B, structure of the pyoverdine backbone, the intramolecular cyclized succinic acid variant, and the non-cyclized pyoverdine precursor ferribactin. The residues R_{1-4} correspond to the substructures shown in D. C, IEF gels visualized by pyoverdine fluorescence (left) and a CAS overlay assay (right). Numbers denote the identified pyoverdine variants, whose structures are represented in B in conjunction with D.

phore functionality of the pyoverdines produced in the mutant strains.

PtaA is the searched-for transaminase

To examine whether PtaA is responsible for the predicted side-chain modification, we isolated the pyoverdines from all strains and analyzed the molecular composition via ultraperfor-

mance liquid chromatography (UPLC)-coupled high-resolution mass spectrometry (HR-MS) and isoelectric focusing (IEF) gels in combination with the chrome azurol S (CAS) overlay assay. The results verified our hypothesis about the nature of the modification (Fig. 5, A–D). For convenience, the structures of all pyoverdine variants identified in this study are denoted in Fig. 5, B and D, and the assigned numbers 1–6 will be used

unambiguously attribute the compounds discussed. The masses of compounds **1–6** were determined by HR-MS and compared with the calculated molecular masses (denoted in parentheses) of the respective molecules: **1**, 1190.544 atomic mass units (a.m.u.) (1190.544 a.m.u.); **2**, 1189.510 a.m.u. (1189.513 a.m.u.); **3**, 1160.534 a.m.u. (1160.534 a.m.u.); **4**, 1161.520 a.m.u. (1161.518 a.m.u.); **5**, 1143.507 a.m.u. (1143.507 a.m.u.); **6**, 1178.581 a.m.u. (1178.581 a.m.u.).

To analyze whether the identified structures were truly distinct molecules and not artifacts of MS (e.g. dehydrations), we used a shallower gradient for elution and could thereby prove that all compounds have a distinct elution profile (supplemental Fig. S1). From the MS results, it was obvious that, in the absence of PtaA, no α -ketoglutaric acid variant (**2**) of PVD_{A506} was produced (Fig. 5A). The equilibrium between the side-chain modifications was clearly shifted in favor of the succinamide variant (**3**) of PVD_{A506}. Additionally, the unmodified glutamic acid precursor (**1**) of PVD_{A506} was detectable, indicating an incomplete turnover by PvdN. It should be noted that further pyoverdine species could be detected by MS, including the hydrolysis product of the succinamide variant, namely the succinic acid variant (**4**); traces of the intramolecular cyclization product thereof (**5**); and the non-cyclized pyoverdine precursor ferribactin (**6**). As expected from our previously published analysis of PvdN (6), the Δ *ptaA*/ Δ *pvdN* double-deletion strain produced neither the succinamide variant (**3**) or further derivatives thereof nor the α -ketoglutaric acid variant (**2**) of PVD_{A506}. Consequently, the only detectable variants were the fluorophore-containing precursor with the original glutamic acid residue (**1**) and its non-cyclized precursor ferribactin (**6**). To examine the validity of our MS data interpretation by an independent method, we performed IEF experiments in combination with the CAS overlay assay (Fig. 5C). The assignments of the bands are denoted on the sides of the images using the definitions given in Fig. 5D. The results obtained by IEF correlate well with the data obtained by MS. Noticeably, the wild-type strain produced an excess of the α -ketoglutaric acid variant (**2**), whereas the succinamide variant (**3**) was a minor side product. The Δ *pvdN* deletion strain did not produce the succinamide variant (**3**) anymore, and all pyoverdine was converted into the α -ketoglutaric acid variant (**2**) by PtaA. The Δ *ptaA* deletion strain in turn produced at least three variants of pyoverdine, namely the unmodified glutamic acid precursor (**1**), the succinamide variant (**3**), and the hydrolysis product thereof, the succinic acid variant (**4**). It should be noted that the lower band in the Δ *ptaA* strain could easily be mistaken for the α -ketoglutaric acid variant (**2**) due to its nearly identical pI, but the succinic acid variant (**4**) migrated slightly higher. Furthermore, in the Δ *ptaA*/ Δ *pvdN* double-deletion strain, no succinic acid variant (**4**) was present due to the lack of its precursor, the succinamide variant (**3**). This lane unambiguously clarifies the assignment of the closely migrating α -ketoglutaric acid (**2**) and succinic acid variants (**4**) of PVD_{A506}.

The Δ *ptaA* phenotype was entirely complemented in *trans* by the *ptaA* complementation strain, whereas the PtaA(K224A) variant did not complement the phenotype and was therefore inactive. This conclusion can be deduced from the comparison of the IEF pyoverdine band patterns of these strains with the

patterns obtained for the wild-type and the Δ *ptaA* deletion strains. The same holds true for the complementation of the Δ *ptaA*/ Δ *pvdN* double-deletion strains as the in *trans* complementation with *ptaA* resulted in the quantitative turnover of the glutamic acid precursor (**1**) to the α -ketoglutaric acid variant (**2**), whereas the strain with PtaA(K224A) produced only the unmodified glutamic acid precursor (**1**). Moreover, the CAS overlay assay indicated that the side-chain modifications did not alter the iron affinity of the pyoverdine variants as can be inferred from the identical band patterns in comparison with fluorescence imaging of the IEF gels (Fig. 4C).

Discussion

More than 30 years ago, an α -ketoglutaric acid was identified for the first time as an amide-bonded residue at the fluorophore of a pyoverdine in strains of *P. aeruginosa* and *P. fluorescens* (16, 17). Also the at that time already known succinamide (18) and succinic acid residues were detected at this position in those studies. Interestingly, pyoverdines of *P. aeruginosa* strains greatly differ in their peptide moiety but generally have the three above mentioned residues in common that are amidically attached to their fluorophore (19). Also a glutamic acid residue bound to the fluorophore via its γ -carboxylic acid was discovered at this position (20) as well as malamide (21), malic acid (22), and an intramolecular cyclized succinate (23). These variations of the pyoverdine are made in a species- and strain-specific manner (24). Based on the large variability of amino acid sequences and fluorophore-attached side chains, IEF-based siderotyping methods have been developed that use the strain-specific characteristics for systematic purposes (25).

The glutamic acid residue is always the first residue in ferribactins, initially discovered by Hohlneicher *et al.* (26). This glutamic acid residue is myristoylated for transport into the periplasm where it is deacylated by PvdQ (27–29). After formation of the fluorophore, it thus must be the glutamic acid residue that is initially present at this position, and the found succinamide, succinic acid, α -ketoglutaric acid, malamide, and malic acid residues must result from glutamic acid modifications.

Biochemistry of PtaA

It was clear from our recent study on PvdN (6) that an unknown enzyme had to be responsible for the formation of the α -ketoglutaric acid. Using *P. fluorescens* A506 as model organism, we have now identified PtaA as the periplasmic transaminase responsible for the formation of the α -ketoglutaric acid residue in PVD_{A506} (**2**). PtaA is a PLP-containing enzyme (Fig. 2) that is transported together with its covalently bound cofactor via the Tat pathway in a folded conformation (Fig. 3). The folded state is influenced by cofactor binding to position Lys-224 (Fig. 3C), which is most likely the reason for a partial mislocalization of the mutated protein in the membrane (Fig. 3A). A similar effect had been already observed with PvdN, which also requires PLP binding for Tat-dependent transport (6). In the case of PvdN, the exchange of the active-site Lys resulted even in a complete mislocalization of the enzyme in the mem-

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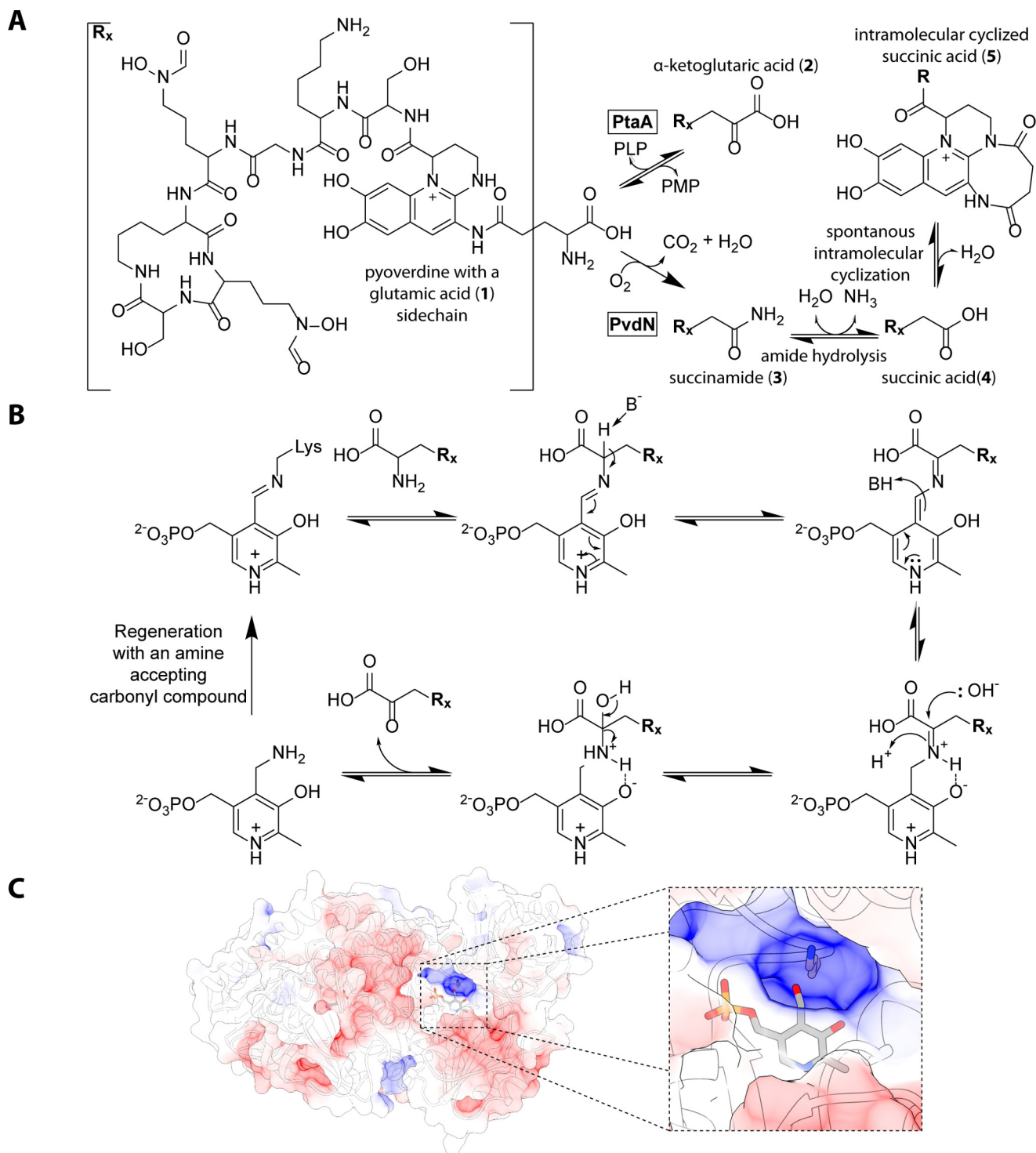


Figure 6. Role and mechanism of PtaA in periplasmic pyoverdine tailoring. A, beginning from the glutamic acid variant (1) of PVD_{A506r}, two competing tailoring pathways are present. Transamination by PtaA results in the α -ketoglutaric acid variant (2) PVD_{A506r}, whereas the PvdN modification results in the succinamide variant (3), which can be partially hydrolyzed to the succinic acid (4) and then intramolecularly cyclized (5). B, proposed mechanism of transamination as catalyzed by PtaA in the periplasm, postulating a carbonyl compound for regeneration of PLP. C, structure of PtaA as calculated by homology modeling, highlighting the Lys-224 residue and the PLP cofactor. The model was visualized with UCSF Chimera.

brane fraction (6). PvdN and PtaA are both homodimeric enzymes (Ref. 6 and Fig. 3D) with their PLP most likely bound close to the dimer interface (Ref. 6 and Fig. 6C). This class of enzymes includes therefore good examples for typical Tat substrates that need to be folded prior to translocation (30). They possess twin-arginine signal peptides that can be used for iden-

tification purposes in addition to transaminase sequence specifications (Fig. 1).

Fig. 6A summarizes the now established modifications of the fluorophore-attached glutamic acid residue of pyoverdines. The transamination probably proceeds via a standard transaminase mechanism, which is interesting to exist in the periplasm

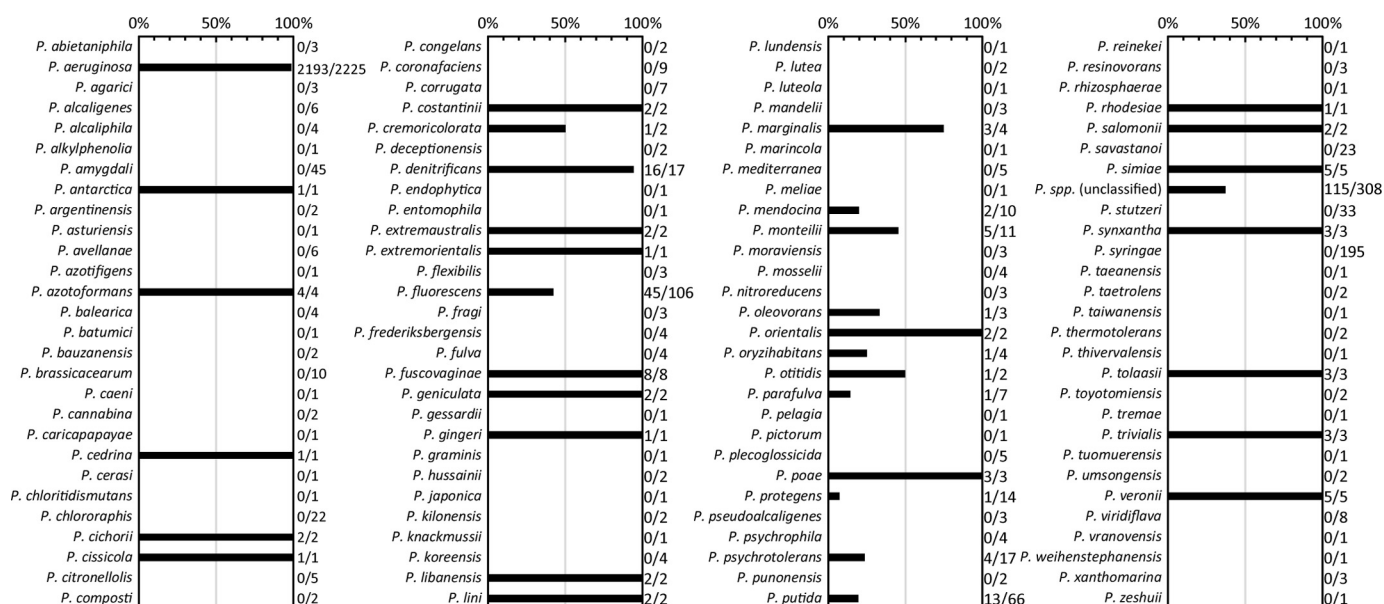


Figure 7. Distribution of PtaA homologs within the genus *Pseudomonas*. Shown is an evaluation of the distribution of PtaA within the genus *Pseudomonas*. Numbers to the right of the bar graph indicate the ratio between genomes positive for the presence of PtaA and the total number of analyzed genomes of the respective species.

as it requires a yet unknown periplasmic carbonyl compound that we postulate to exist (Fig. 6B). As the Tat transport and its PLP requirement indicate that PtaA assembles PLP inside the cytoplasm to fold to an active enzyme, PLP must be regenerated from PtaA-bound pyridoxamine inside the periplasm. The enzymes PvdN and PtaA are responsible for the initial conversions of the glutamic acid pyoverdine variant. The existence of these competing enzyme reactions explains the reported regulatory differences for the formation of succinamide and α -ketoglutaric acid forms of pyoverdine (31). As a side aspect, the glutamic acid variant of pyoverdine that is produced by the $\Delta ptaA/\Delta pvdN$ double mutant can be very helpful for drug delivery by “Trojan horse” antibiotics (32) because bulky side chains may be attached to this residue without affecting uptake (33). Malamide is likely formed by a hydroxylating enzyme acting on succinamide. The acids succinic acid and malic acid are hydrolysis products of their respective amides. A non-enzymatic hydrolysis has been observed and postulated to be responsible for the formation of these acids in cultures (34). It certainly might be that amidases contribute to the rates of these reactions.

PtaA, a transaminase for pyoverdines and other periplasmic compounds

In *P. aeruginosa* PAO1, the homolog of PtaA (53% sequence identity) is encoded by the locus PA2531, which is PvdS-dependently up-regulated in response to iron (35); hence this gene belongs to the regulon of PvdS, the σ factor that mediates pyoverdine production (36). An “iron starvation box” for PvdS binding (37) has been identified in the promoter region of PA2531 (35). We found the consensus box as described for *P. aeruginosa* PAO1 by Ochsner *et al.* (35) also in the *ptaA* promoter region of *P. fluorescens* A506, namely “TAAATN₁₆CGT.” However, the genetic context of *ptaA* is not conserved between *P. aeruginosa* PAO1 and *P. fluorescens*

A506. While the iron-responsive regulation of *ptaA* correlates with its function in pyoverdine modification, its usually *pvd*-unrelated genomic environment is suggestive for additional functions. This interpretation is further strengthened by a previous study on chorismate mutases that included the detection of transaminase activity in the periplasm of *P. aeruginosa* PAO1 (38). In that study, the periplasmic transaminase activity could be assigned to PA2531, the above mentioned PtaA homolog. PA2531 is dimeric just like PtaA from *P. fluorescens* in our study (Fig. 3D). At that time, the authors could not know about its role in pyoverdine modification, which is now clarified by our study. They used α -ketoglutaric acid and phenylalanine as substrates for the transaminase assay. Therefore, PtaA homologs cannot be highly specific for pyoverdine substrates and are likely to be involved in other conversions as well. In support of this conclusion, *ptaA* homologs are also present in some acidobacteria and α -proteobacteria, such as *Zymomonas mobilis* (supplemental Fig. S2), that do not produce any pyoverdines. Therefore, this periplasmic transaminase must be able to exert its function for distinct physiological purposes, possibly in conjunction with the generation of distinct secondary metabolites.

Distribution of PtaA among pseudomonads

As summarized in Fig. 7, we searched for *ptaA* homologs in all *Pseudomonas* genomes available in the *Pseudomonas* Genome Database (11) (see “Experimental procedures” for details). In agreement with the known occurrence of α -ketoglutaric acid forms of pyoverdines, we found this gene in almost all sequenced *P. aeruginosa* strains (2193 out of 2225 genomes; ~99%). The absence of hits in the ~1% of strains without this gene might be due to the inclusion of draft genomes in the analysis.

In the more diverse groups of *P. fluorescens* and *Pseudomonas putida* strains, only 45 out of 106 genomes and 13 out of 66 genomes, respectively, encode PtaA, suggesting that the presence of the α -ketoglutaric acid variant of pyoverdine is not

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Table 2

Primers used in this study

oePCR, overlap extension PCR.

Name	Sequence	Restriction site	Purpose
PfA506-4424-F1-MR PfA506-4424-R1-MR	ATAGCCGGATCC TAGACAGGTAGCGCCAAATCAGC AGGAGTAGTCACCATGGTGCCTGTACCGCAGGTGGTCTGA TCAGGCGGCGCTATAGCTG	BamHI	Forward primer for <i>ptaA</i> left flanking region Reverse primer for <i>ptaA</i> left flanking region
PfA506-4424-F2-MR PfA506-4424-R2-MR	GACACGCACCATGGTACTACTCCTTTG GGCCCGCAATTC TTTGTACAGATTCCTGATTTCCATCTTG	EcoRI	Forward primer for <i>ptaA</i> right flanking region Reverse primer for <i>ptaA</i> right flanking region
PfA506-4424-DF-MR PfA506-4424-DR-MR PfA506-4424-F-MR	TTTGAGTGTGCGCCTATTG CGCGCAATGCTGGTGAAGATATAG TCATCGCATATGGTGCCTGTAGTCGTCGATCC	NdeI	<i>ptaA</i> genomic deletion control primer <i>ptaA</i> genomic deletion control primer Forward primer for cloning PtaA-coding region into pEXH5
PfA506-4424-strep-R-MR	GGCCGCAAGCTTTTACTTTTCGAACTGCGGGTGGCTCCAG ACCACCTGCGTCGCAAAGGCCCTCGC	HindIII	Reverse primer for cloning PtaA-coding region into pEXH5
PfA506-4424_K224A-F-MR PfA506-4424_K224A-R-MR PfA506-mat4424-F-MR	CTGGTGCTGCGCACCTTCTCCGCCATCTAC CCGGCCATGCCGTAGATGGCCGAGAAAGGTG GATATACATATGAGCCAGCGCCGACAAAATCTGACC	NdeI	Primer for K224A exchange by oePCR Primer for K224A exchange by oePCR Forward primer for cloning mature PtaA-coding region into pEXH5
pEXH5-RBS-F-MR	GGCGCGGGATCCGTTTAACTTTAAGAAGGAGATATAC	BamHI	Forward primer for subcloning from pEXH5 into pME6010
pEXH5-strep-term-HindIII-R-MR	CCCCCTTAAGCTTAAAAAAAACCCCGCCCTGTACAGGGGCGG GGTTTTTTTTTTTACTTTTCGAACTGCGGGTGGCTCC	HindIII	Reverse primer for subcloning from pEXH5 into pME6010

strictly conserved in these species. In agreement with this prediction, *P. fluorescens* Pf0-1 has no PtaA and, as expected, has been shown not to contain the α -ketoglutaric acid (24). On the contrary, there are strains that form α -ketoglutaric acid pyoverdine variants but not any succinamide/succinate variants (24), such as the genome-sequenced *P. putida* strain H8234. We found that this strain has a *pvdMO-ptaA* operon instead of the commonly found *pvdMNO* operon. Therefore, in the rarely found strains that do not possess the succinamide/succinic acid modification pathway, *ptaA* can be associated with the *pvd* gene cluster that encodes the periplasmic pyoverdine maturation enzymes. In conclusion, some *Pseudomonas* species with at least the exception of *P. aeruginosa* show the interesting characteristic of containing either only the PtaA pathway or only the PvdN pathway (39). Strains with both pathways acting in parallel also exist. We did not find a strain that produces pyoverdine without modifying its glutamic acid. The reason for the apparent requirement for such modifications is unknown and must be unrelated to iron affinity and uptake (Ref. 40 and Fig. 4). Future studies will have to clarify the exact functions of these modifications as well as the additional roles that PtaA can have in the periplasm of proteobacteria.

Experimental procedures

Strains and growth conditions

For physiological studies, *P. fluorescens* A506 was used. For cloning *E. coli* DH5 α λ *pir*⁺ and for expression *E. coli* Rosetta 2 (DE3) pLysSRARE2 were utilized. *P. fluorescens* A506 was cultivated at 30 °C, whereas *E. coli* strains were cultivated at 37 °C unless noted otherwise. The standard cultivation medium was LB (1% (w/v) tryptone, 1% (w/v) NaCl, 0.5% (w/v) yeast extract). If necessary, the appropriate antibiotics were added to the cultivation media at the following final concentrations: 100 μ g/ml ampicillin, 25 μ g/ml chloramphenicol, 50 μ g/ml kanamycin, and 20 μ g/ml tetracycline.

The production of pyoverdine and the pyoverdine plate assay, pertaining to *P. fluorescens* A506, were carried out as described previously (6). For relative pyoverdine quantification, 5-ml LB overnight cultures (ONCs) of *P. fluorescens* A506 were inoculated from cryocultures and incubated at 30 °C at 180 rpm

overnight. Thereafter, 50 of ml CAA medium (41) (5 g/liter casamino acids, 5 mM K₂HPO₄, 1 mM MgSO₄) precultures in 100-ml Erlenmeyer flasks with one baffle were inoculated with 50 μ l of the respective ONC and incubated for ~16 h at 30 °C and 180 rpm. 2 ml of the respective preculture were sedimented at 16,000 \times g for 2 min at room temperature and subsequently washed twice with 1.5 ml of CAA medium. Afterward, the OD₆₀₀ of the cell suspensions was adjusted to 1.0 with CAA medium, and for each sample three 4.5-ml CAA cultures (without antibiotics) were inoculated with 0.5 ml of the respective adjusted cell suspension. The cultures were then incubated overnight at 30 °C and 180 rpm. These cultures were sedimented by centrifugation at 3,260 \times g and 4 °C for 10 min in 15-ml screw-top plastic tubes. 2.7 ml of the supernatant were mixed with 300 μ l of a 1 M HEPES buffer, pH 8.0, and transferred into 1-cm acrylic cuvettes. The samples were measured with a Jasco FP-6500 spectrofluorometer using the following settings for acquisition: excitation at 405 nm, emission at 460 nm, bandwidth of 3 nm for both excitation and emission, and 0.5 s response time in “low-sensitivity” mode.

Genetic methods and plasmids

The construction of all scarless and markerless deletions in *P. fluorescens* A506 were performed as described previously (6). Furthermore, complementation and production vectors as well as constructs with single-point mutations (pME6010 and pEXH5 derivatives) were generated according to our previously published procedure (6). All primers used for cloning are listed in Table 2. Constructs were verified by restriction analysis and sequencing. *E. coli* DH5 α λ *pir*⁺ cells were rendered competent and transformed as described previously (42).

Biochemical methods

For analysis of subcellular fractionations and protein overexpression by SDS-PAGE, successive Western blotting, or in-gel colloidal Coomassie staining, standard protocols were used (43–46). Western blots were developed according to the manufacturer’s instructions using StrepMAB-Classic (IBA, Göttingen, Germany) as primary antibody and anti-mouse-HRP con-

jugate (Carl Roth, Karlsruhe, Germany) as secondary antibody. Images were acquired with the MF-ChemiBIS 4.2 imaging system (DNR Bio-Imaging Systems, Jerusalem, Israel).

The overproduction of mature PtaA and mature PtaA(K224A) was performed using *E. coli* Rosetta 2 (DE3) pLysSRARE2 as described previously (47) with minor modifications. Briefly, the respective construct was transformed into *E. coli* Rosetta 2 (DE3) pLysSRARE2 by the transformation and storage solution method (48). After recovery, the cells were spread on MDAG-11 plates (47) and incubated at 37 °C overnight. A 5-ml MDAG-135 (47) ONC was then inoculated with a single colony and incubated at 180 rpm and 37 °C. For overproduction, two 0.5-liter ZYM-5052 preheated (37 °C) autoinducer medium cultures (47) in 3-liter Erlenmeyer flasks with four baffles were each inoculated with 0.5 ml of the ONC. The cultures were incubated for 3 h at 200 rpm and 37 °C. Subsequently, the temperature was decreased to 30 °C, and the cultures were incubated for approximately 19 h at 200 rpm. The cells were harvested, and the respective protein was purified by means of four 1.5-ml gravity-flow Strep-Tactin®-Sepharose columns (IBA) as described previously (6). The purified proteins were concentrated using Vivaspin® 6 concentrators with a cutoff of 10 kDa (Sartorius, Göttingen, Germany). For molecular weight determination, a Pharmacia FPLC (LKB Pump P-500, V-7 valve, 50- μ l sample loop) was connected to a size-exclusion chromatography column (Superdex 200 Increase 10/300 GL, GE Healthcare) coupled to MALS (miniDAWN TREOS, Wyatt Technology Europe GmbH, Dernbach, Germany) and refractive index detectors (Shodex RI-101, Showa Denko Europe GmbH, Munich, Germany). The flow rate of the mobile phase (PBS; 10 mM phosphate buffer, pH 7.4, 140 mM NaCl) was set to 0.75 ml/min. The chromatograms were recorded and analyzed with ASTRA 6.1 software (Wyatt Technology). For DSC, a NANO DSC in conjunction with a degassing station from TA Instruments (Lindon, UT) was used. For preparation of DSC samples, the protein elution peak (19-min retention time) from the SEC-MALS-RI system was collected by hand and subsequently concentrated with Vivaspin 6 concentrators (cutoff, 10 kDa). The protein concentration was determined using Roti®-Nanoquant according to the instruction manual (Carl Roth) using a SpectraMax M3 spectrophotometer (Molecular Devices, Biberach an der Riss, Germany). The protein of interest was diluted with degassed PBS to a concentration of ~1.5 mg/ml prior to analysis. The DSC reference cell was filled with degassed PBS. Analyses covered the temperature range from 20 to 110 °C at a heating rate of 1 °C/min with a 900-s pre-equilibration time at a constant pressure of 3 atm.

The subcellular fractionation of *P. fluorescens* A506 cultures was performed as described previously (49) with slight modifications. 50-ml cultures were grown to an OD₆₀₀ of 1 and fractionated into 1-ml fractions that were further analyzed without precipitation.

For PLP cofactor detection, a method modified from that of Ojha *et al.* (50) was used. The purified mature PtaA enzyme in 100 mM potassium phosphate buffer, pH 7.2, was treated with 5 mM hydroxylamine and incubated for ~72 h at 4 °C. The sample was loaded onto a Vivaspin 6 concentrator with 5-kDa cutoff, and the PLP oxime in the flow-through was detected at

446-nm emission (scan from 400 to 700 nm) with an excitation of 353 nm, 3-nm bandwidth for both excitation and emission, 0.2-s response time, and three accumulations in “high-sensitivity” mode using a Jasco FP-6500 spectrofluorometer. Reference solutions with the buffer only or with 1 mM PLP in the same buffer were treated identically. PLP- or PLP oxime-containing reference spectra were recorded with the same settings but in “medium-sensitivity” mode.

The extraction of pyoverdine, IEF analysis in conjunction with the CAS overlay assay, and UPLC-MS analysis were performed as described previously (6). To prove that the identified compounds are not artifacts of the MS, an adjusted linear gradient profile was utilized as follows: solvent A (double-distilled water with 0.1% (v/v) formic acid) and solvent B (acetonitrile with 0.1% (v/v) formic acid) at a flow rate of 0.6 ml/min; 2% B (0.0 min), 10% B (10.0 min), 90% B (10.5 min), 90% B (13.0 min), 2% B (13.5 min), and 2% B (20.0 min). The electrospray ionization voltage was set to 100 V, and the injection volume was 5 μ l. All other instrument parameters and all instrument hardware were identical to the method described previously (6) except that an ACQUITY UPLC Column Manager (Waters) was additionally installed on the instrument.

Bioinformatics methods

To perform an initial distribution analysis of PtaA within the genus *Pseudomonas*, we downloaded all amino acid sequences of all complete and draft genomic sequences from the *Pseudomonas* Genome Database (11) and used TATFIND v.1.4 (7) to search for potential Tat substrates. Subsequently, phmmer v3.1b (51) was used to search for PtaA homologs (cutoff, 1e–45) within all potential Tat substrates. Then a taxonomy table reduced to species level was constructed by means of in-house developed software. Thereafter, the ratio between the total number of genomes of one species and the number of genomes of the same species carrying a potential PtaA homolog was calculated.

To construct the sequence logos, the sequence redundancy of all PtaA homologs was reduced by means of the CD-HIT algorithm contained in the CD-HIT Suite (12) using a similarity threshold of 0.95. Then the resulting sequences were aligned with the T-Coffee web server (13). The sequence alignment was subsequently visualized with WebLogo 3 (14) by plotting the probability of each amino acid at each alignment position. For further investigations into the distribution of PtaA homologs across other bacterial species, a profile hidden Markov model (pHMM) was generated from the multiple sequence alignment with hmmbuild v.3.1b (51). Then all available reference sequences were searched with the generated model (E-value cutoff, 1e–100) using the HMMER web server (52, 53). The resulting sequences were then analyzed regarding their phylogenetic relationship using the Phylogeny.fr³ web service (54) in conjunction with the ETE toolkit (55) for visualization.

To initially predict the enzyme class and active-site lysine residue, Lys-224, the InterPro web service (8, 9) was used. The SWISS-MODEL web service (56) was used to model the struc-

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ture of PtaA (template, Protein Data Bank code 3LY1), and UCSF Chimera (57) in conjunction with APBS (58) was used to visualize the structure of PtaA.

Author contributions—M. T. R. performed the experiments, prepared the figures, and analyzed the data together with T. B. G. D. performed the MS analyses. T. B. conceived and coordinated the study. T. B. and M. T. R. wrote the paper. All authors reviewed the results and approved the final version of the manuscript.

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