



Rare Earth Element (REE)-Dependent Growth of *Pseudomonas putida* KT2440 Relies on the ABC-Transporter PedA1A2BC and Is Influenced by Iron Availability

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In the soil-dwelling organism *Pseudomonas putida* KT2440, the rare earth element (REE)-utilizing, and pyrroloquinoline quinone (PQQ)-dependent ethanol dehydrogenase PedH is part of a periplasmic oxidation system that is vital for growth on various alcoholic volatiles. Production of PedH and its Ca²⁺-dependent counterpart PedE is inversely regulated in response to lanthanide (Ln³⁺) bioavailability, a mechanism termed the REE-switch. In the present study, we demonstrate that copper, zinc, and in particular, iron availability influences this regulation in a pyoverdine-independent manner by increasing the minimal Ln³⁺ concentration required for the REE-switch to occur by several orders of magnitude. A combined genetic and physiological approach reveals that an ABC-type transporter system encoded by the gene cluster *pedA1A2BC* is essential for efficient growth on 2-phenylethanol with low (nanomolar) Ln³⁺ concentrations. In the absence of *pedA1A2BC*, a ~100-fold higher La³⁺-concentration is needed for PedH-dependent growth but not for the ability to repress growth based on PedE activity. From these results, we conclude that cytoplasmic uptake of lanthanides through PedA1A2BC is essential to facilitate REE-dependent growth on 2-phenylethanol under environmental conditions with poor REE bioavailability. Our data further suggest that the La³⁺/Fe²⁺/Fe³⁺ ratio impacts the REE-switch through the mismetallation of putative La³⁺-binding proteins, such as the sensor histidine kinase PedS2, in the presence of high iron concentrations. As such, this study provides an example for the complexity of bacteria-metal interactions and highlights the importance of medium compositions when studying physiological traits *in vitro* in particular in regard to REE-dependent phenomena.

Keywords: lanthanides, rare earth elements, ABC-transporter, *Pseudomonas putida*, pyrroloquinoline quinone, PedH, mismetallation, metal homeostasis

INTRODUCTION

Metal ions are essential for all living organisms as they play important roles in stabilizing macromolecular cellular structures, by catalyzing biochemical reactions or acting as cofactors for enzymes (Gray, 2003; Merchant and Helmann, 2012). They can, however, also be toxic to cells at elevated levels through the generation of reactive oxygen species or by aspecific interactions such as mismetallation (Cornelis et al., 2011; Dixon and Stockwell, 2014; Foster et al., 2014). Bacteria have hence developed a sophisticated toolset to maintain cellular metal homeostasis (Andrews et al., 2003; Schalk and Cunrath, 2016; Chandrangsu et al., 2017; Semrau et al., 2018). Common mechanisms include release of metal-specific scavenger molecules, the activation of high-affinity transport systems, the production of metal storage proteins, and the expression of specific efflux pumps.

As it is the case for all strictly aerobic bacteria, the soil-dwelling organism *Pseudomonas putida* KT2440 has a high demand for iron. However, the bioavailability of this metal is very poor under most oxic environmental conditions due to the fast oxidation of Fe^{2+} - and the low solubility of Fe^{3+} -species (Andrews et al., 2003). One strategy of many bacteria to overcome this challenge is to excrete self-made peptide-based siderophores (such as pyoverdines) into the environment that bind Fe^{3+} with high affinity, and thereby increase its bioavailability (Salah El Din et al., 1997; Cornelis and Andrews, 2010; Baune et al., 2017). A second adaptation of *P. putida* cells to iron-limitation is a change in the proteomic inventory to limit the use of Fe-containing enzymes, exemplified by the switch from the Fe-dependent superoxide dismutase (SOD) to a Mn-dependent isoenzyme or by re-routing of entire metabolic pathways (Kim et al., 1999; Sasnow et al., 2016). In contrast, when Fe bioavailability is high, the production of the bacterioferritins Bfr α and Bfr β is increased to enable intracellular storage and thereby improve cellular fitness under potential future conditions of iron starvation (Chen et al., 2010). The regulatory mechanisms for metal homeostasis of *P. putida* cells in response to other essential metal ions such as Co, Cu, Mg, Mo, Ni, and Zn are less well explored. Genes encoding for transport systems associated with the uptake and efflux of these metals can, however, be found in its genome (Nelson et al., 2002; Belda et al., 2016), and some of these have been studied in more detail (Miller et al., 2009; Ray et al., 2013).

We have recently reported that *P. putida* KT2440 is capable of using rare earth elements (REEs) of the lanthanide series (Ln^{3+}) when growing on several alcoholic substrates (Wehrmann et al., 2017, 2019). Under these conditions, the cells use the pyrroloquinoline quinone (PQQ)-dependent ethanol dehydrogenase (EDH) PedH, to catalyze their initial oxidation within the periplasm. Like many other organism, *P. putida* harbors an additional, Ln^{3+} -independent functional homolog of PedH termed PedE that depends on a Ca^{2+} ion as metal cofactor (Takeda et al., 2013; Wehrmann et al., 2017). Depending on the availability of REEs in the environment, *P. putida* tightly regulates PedE and PedH production (Wehrmann et al., 2017, 2018). In the absence of Ln^{3+} , growth is solely dependent on PedE whereas PedH transcription is repressed. The situation immediately changes in the presence of small amounts of Ln^{3+} (low nM

range) leading to a strong induction of the Ln^{3+} -dependent enzyme PedH and repression of its Ca^{2+} -dependent counterpart PedE. For *P. putida* KT2440 the PedS2/PedR2 two component system (TCS) is a central component of this inverse regulation (Wehrmann et al., 2018). Notably, the REE-switch in *P. putida* was also found to be influenced by environmental conditions, as the critical La^{3+} concentrations required to support PedH-dependent growth differ dramatically depending on the medium used, ranging from 5 nM up to 10 μM (Wehrmann et al., 2017).

Lanthanides are only poorly available in natural environments (often picomolar concentrations) due to the formation of low soluble hydroxide and/or phosphate complexes (Meloche and Vrátný, 1959; Firsching and Brune, 1991). The presence of active uptake systems to facilitate REE-dependent growth in bacteria has thus been favored by many researchers (Markert, 1987; Tyler, 2004; Aide and Aide, 2012; Gu et al., 2016; Gu and Semrau, 2017; Cotruvo et al., 2018; Picone and Op den Camp, 2019). A transcriptomic study of *M. trichosporium* OB3b cells observed that multiple genes encoding for different active transport systems were among the most regulated in the presence of cerium (Gu and Semrau, 2017). In addition, it has been found that a specific TonB-dependent receptor protein as well as a TonB-like transporter protein are highly conserved in bacteria that carry genes encoding for Ln^{3+} -dependent MDHs (Keltjens et al., 2014; Wu et al., 2015). Only very recently, different studies indeed identified both an ABC-transporter and TonB-dependent receptor proteins that are needed for REE-dependent growth of methano- and methylotrophs on methanol, strongly suggesting the existence of an uptake system that specifically transports a Ln^{3+} -chelator complex in these organisms (Groom et al., 2019; Ochsner et al., 2019; Roszczenko-Jasińska et al., 2019).

With the present study, we show that a homologous ABC-transporter system, encoded by the gene cluster *pedA1A2BC*, is essential for lanthanide-dependent growth on 2-phenylethanol in the non-methylotrophic organism *P. putida* KT2440 under low (nanomolar) concentrations of REEs. Notably, no homolog of the TonB-dependent receptor proteins found in methanotrophic or methylotrophic strains could be identified within the genome of *P. putida* KT2440 indicating either a lack or substantial differences in the chemical nature of such a Ln^{3+} -specific chelator system. Finally, we show that the siderophore pyoverdine plays no essential role for growth on 2-phenylethanol under low REE concentrations but provide compelling evidence that in addition to Cu^{2+} and Zn^{2+} the $\text{Fe}^{2+/3+}$ to Ln^{3+} ratio significantly alters the REE-switch most likely through mismetallation.

MATERIALS AND METHODS

Bacterial Strains, Plasmids, and Culture Conditions

The *Escherichia coli* and *Pseudomonas putida* KT2440 strains and the plasmids used in this study are described in **Table 1**. Maintenance of strains was routinely performed on solidified (1.5% agar) LB medium (Maniatis et al., 1982). If not stated otherwise, strains were grown in liquid LB medium (Maniatis et al., 1982) or a modified M9 salt

TABLE 1 | Strains and plasmids used in the study.

Strains	Relevant features	Source or References
KT2440*	KT2440 with a markerless deletion of <i>upp</i> . Parent strain for deletion mutants	Graf and Altenbuchner, 2011
$\Delta pedE$	KT2440* with a markerless deletion of <i>pedE</i>	Mückschel et al., 2012
$\Delta pedA1A2BC$	KT2440* with a markerless deletion of <i>pedA1A2BC</i> (PP_5538, PP_2669, PP_2668, PP_2667)	This study
$\Delta pedE \Delta pedA1A2BC$	$\Delta pedE$ with markerless deletion of gene cluster <i>pedA1A2BC</i>	This study
$\Delta pedE \Delta pvdD$	$\Delta pedE$ with a markerless deletion of <i>pvdD</i> (PP_4219)	This study
$\Delta pedH$	KT2440* with a markerless deletion of <i>pedH</i>	Mückschel et al., 2012
$\Delta pedH \Delta pedA1A2BC$	$\Delta pedH$ with a markerless deletion of gene cluster <i>pedA1A2BC</i>	This study
$\Delta pedE \Delta pedH$	KT2440* with a markerless deletion of <i>pedE</i> and <i>pedH</i>	Mückschel et al., 2012
$\Delta pedE \Delta tatC1$	$\Delta pedE$ with a markerless deletion of <i>tatC1</i> (PP_1039)	This study
$\Delta pedE \Delta pedH \Delta pedA1A2BC$	$\Delta pedE \Delta pedH$ with a markerless deletion of gene cluster <i>pedA1A2BC</i>	This study
$\Delta pedE \Delta tatC2$	$\Delta pedE$ with a markerless deletion of <i>tatC2</i> (PP_5018)	This study
<i>E. coli</i> BL21 (DE3)	$F^- ompT gal dcm lon hsdS_B(r_B^- m_B^-) \lambda[DE3 (lacI lacUV5-T7 gene 1 ind1 sam7 nin5)]$	Studier and Moffatt, 1986
<i>E. coli</i> TOP10	$F^- mcrA \Delta(mrr-hsdRMS-mcrBC) \phi 80lacZ \Delta M15 \Delta lacX74 nupG recA1 araD139 \Delta(ara-leu)7697 galE15 galK16 rpsL(Str^R) endA1 \lambda^-$	Invitrogen
<i>E. coli</i> HB101	$F^- mcrB mrr hsdS20(r_B^- m_B^-) recA13 leuB6 ara-14 proA2 lacY1 galK2 xyl-5 mtl-1 rpsL20(Sm^R) gln V44 \lambda^-$	Boyer and Roulland-Dussoix, 1969
<i>E. coli</i> PIR2	$F^- \Delta lac169 rpoS(Am) robA1 creC510 hsdR514 endA reacA1 uidA(\Delta MluI);pir$	Invitrogen
<i>E. coli</i> CC118 λ pir	$\Delta(ara-leu) araD \Delta lacX74 galE galK phoA20 thi-1 rpsE rpoB argE(Am) recA1 \lambda_{pir}$ phage lysogen	Herrero et al., 1990
KT2440*:Tn7M-pedH-lux	KT2440* with insertion of Tn7-M-pedH-lux	This study
$\Delta pedA1A2BC$:Tn7M-pedH-lux	$\Delta pedA1A2BC$ with insertion of Tn7-M-pedH-lux	This study
KT2440*:Tn7M-pedE-lux	KT2440* with insertion of Tn7-M-pedE-lux	This study
Plasmids		
pJOE6261.2	Suicide vector for gene deletions	Graf and Altenbuchner, 2011
pMW10	pJeM1 based vector for rhamnose inducible expression of <i>pedH</i> with C-terminal 6x His-tag	Wehrmann et al., 2017
pMW50	pJOE6261.2 based deletion vector for gene <i>pvdD</i> (PP_4219)	This study
pMW57	pJOE6261.2 based deletion vector for gene cluster <i>pedA1A2BC</i>	This study
pTn7-M	Km ^R Gm ^R , <i>ori R6K</i> , <i>Tn7L</i> and <i>Tn7R</i> extremities, standard multiple cloning site, <i>oriT</i> RP4	Zobel et al., 2015
pRK600	Cm ^R , <i>ori ColE1</i> , Tra ⁺ Mob ⁺ of RK2	Keen et al., 1988
pTNS1	Ap ^R , <i>ori R6K</i> , <i>TnSABC</i> + <i>D</i> operon	Choi et al., 2005
pSEVA226	Km ^R , <i>ori RK2</i> , reporter vector harboring the <i>luxCDABE</i> operon	Silva-Rocha et al., 2013
pSEVA226-pedH	pSEVA226 with a <i>pedH-luxCDABE</i> fusion	This study
pSEVA226-pedE	pSEVA226 with a <i>pedE-luxCDABE</i> fusion	This study
pTn7-M-pedH-lux	pTn7-M with a <i>pedH-luxCDABE</i> fusion	This study
pTn7-M-pedE-lux	pTn7-M with a <i>pedE-luxCDABE</i> fusion	This study

* is part of the strain name KT2440* which is strain KT2440 with a *upp* deletion, representing the parental strain for all mutant strains.

medium (Wehrmann et al., 2017) supplemented with 5 mM 2-phenylethanol, 5 mM 2-phenylacetaldehyde, 5 mM phenylacetic acid, or 25 mM succinate (pre-cultures) as carbon and energy source at 28°C to 30°C and shaking. 40 $\mu\text{g mL}^{-1}$ kanamycin or 15 $\mu\text{g mL}^{-1}$ gentamycin for *E. coli* and 40 $\mu\text{g mL}^{-1}$ kanamycin, 20 $\mu\text{g mL}^{-1}$ 5-fluorouracil, or 30 $\mu\text{g mL}^{-1}$ gentamycin for *P. putida* strains was added to the medium for maintenance and selection, if indicated.

Liquid Medium Growth Experiments

Liquid growth experiments were performed in biological triplicates by monitoring the optical density at 600 nm (OD₆₀₀) during growth in modified M9 medium supplemented with the corresponding carbon and energy sources (see above). For all experiments, washed cells from overnight cultures grown with succinate at 30°C and 180 rpm shaking were utilized to inoculate fresh medium with an OD₆₀₀ of 0.01 to 0.05. Depending on the

culture vessel, the incubation was carried out in 1 ml medium per well for 96-well 2 ml deep-well plates (Carl Roth) at 350 rpm shaking and 30°C or 200 μL medium per well for 96-well microtiter plates (Sarstedt) at 180 rpm shaking and 28°C. If needed, different concentrations of LaCl₃ (Sigma-Aldrich) were added to the medium.

Construction of Plasmids

The 600 bp regions upstream and downstream of gene *pvdD*, gene cluster *pedA1A2BC* and genes *tatC1* and *tatC2* were amplified from genomic DNA of *P. putida* KT2440 using primer pairs MWH56/MWH57 and MWH58/MWH59, MWH94/MWH95 and MWH96/MWH97, PBtatC1.1/PBtatC1.2 and PBtatC1.3/PBtatC1.4 and PBtatC2.1/PBtatC2.2 and PBtatC2.3/PBtatC2.4 to construct the deletion plasmids pMW50, pMW57, pJOE-tatC1 and pJOE-tatC2 (Table 2). The *Bam*HI digested pJOE6261.2 as well as the two up- and

TABLE 2 | Primers used in the study.

Name	Sequence 5' → 3'	Annealing temperature (°C)
MWH56	GCCGCTTTGGTCCCGGCCACCGGCGAGTTGCA	60
MWH57	CCCGAAAGCTTGAACATCTCCTACCAGGGC	60
MWH58	ATGTTCAAGCTTTGCGGGGCCG	60
MWH59	GCAGGTGCGACTCTAGAGCTTACAGATGCTGCTGCAGTGC	60
MWH94	GCCGCTTTGGTCCCGCAACAACGCCAGGCCAC	60
MWH95	GCCAGGTTTAAACACTCCACGGCAGATGG	60
MWH96	AGTGTGTTAAACCTGGCGTGAACCCG	60
MWH97	GCAGGTGCGACTCTAGAGCCAGGGAGGTTGCTATGC	60
PBtatC1.1	CGATGGCCGCTTTGGTCCCGCCCATCCGTGCATGCCTC	66
PBtatC1.2	CGATGGCCGCTTTGGTCCCGCCCATCCGTGCATGCCTC	66
PBtatC1.3	AAAATGCTTCGGCCCTTTCGCGGGCGGTG	72
PBtatC1.4	CCTGCAGGTGCGACTCTAGAGGCCATGCCAGTTCGCG	72
PBtatC2.1	CGATGGCCGCTTTGGTCCCGGGAGTACGAAATGGGTATCTTTGACTGGAAACAC	72
PBtatC2.2	AGCAACAGGTGGGGCTCGCGCGGTTGA	72
PBtatC2.3	CGCGAGCCCCACCTGTTGCTTCTGAAGAGG	62
PBtatC2.4	CCTGCAGGTGCGACTCTAGAGATCACCCAGCTGTACCGG	62

downstream fragments were therefore joined together using one-step isothermal assembly (Gibson, 2011) and subsequently transformed into *E. coli* BL21(DE3) or TOP10 cells. Sanger sequencing confirmed the correctness of the plasmids.

For measuring promoter activity of *pedE* and *pedH* *in vivo*, plasmids pTn7-M-*pedH-lux* and pTn7-M-*pedE-lux* were constructed. The DNA regions encompassing the promoters from *pedE* and *pedH* genes were amplified by PCR using the primer pairs p2674-FSac/p2674-RPst and p2679-FSac/p2679-RPst (Wehrmann et al., 2017). The PCR products were digested with *SacI* and *PstI* and inserted upstream the *luxCDABE* operon hosted by plasmid pSEVA226. The cargo module bearing the *pedE-lux* or *pedH-lux* fusion was then passed from the resulting pSEVA226-based constructs to pTn7-M as *PacI/SpeI* fragments.

Strain Constructions

For the deletion of chromosomal genes a previously described method for markerless gene deletions in *P. putida* KT2440 was used (Graf and Altenbuchner, 2011). In short, after transformation of the integration vectors carrying the up- and downstream region of the target gene, clones that were kanamycin (Kan) resistant and 5-fluorouracil (5-FU) sensitive were selected and one clone was incubated in liquid LB medium for 24 h at 30°C and 180 rpm shaking. Upon selection for 5-FU resistance and Kan sensitivity on minimal medium agar plates, clones that carried the desired gene deletion were identified by colony PCR.

Integration of the pTn7-M based *pedH-lux* and *pedE-lux* fusions into the chromosome of *P. putida* KT2440 was performed by tetraparental mating using PIR2/pTn7-M-*pedH-lux* or PIR2/pTn7-M-*pedE-lux* as the donor, *E. coli* CC118 λ pir/pTNS1 and *E. coli* HB101/pRK600 as helper strains and appropriate KT2440 strain as the recipient (Zobel et al., 2015). Briefly, cultures of the four strains grown under selective conditions were mixed, spotted on LB agar and incubated overnight at 28°C. Transconjugants were selected on

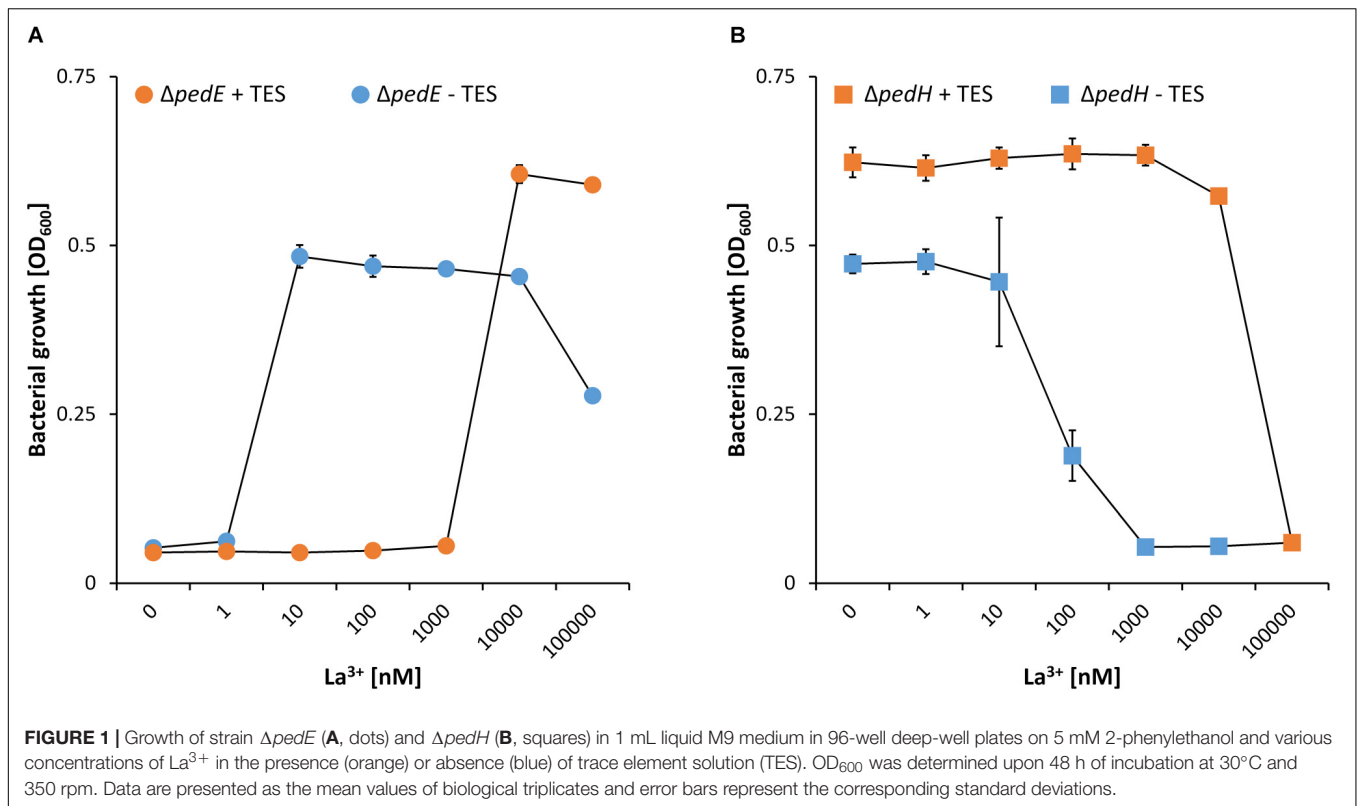
cetrimide agar (Sigma-Aldrich) containing gentamicin. Correct chromosomal integration of mini-Tn7 was checked by colony PCR using Pput-*glmSDN* and PTn7R primers as described elsewhere (Choi et al., 2005).

Promoter Activity Assays

Pseudomonas putida harboring a Tn7-based *pedH-lux* or *pedE-lux* fusion were grown overnight in M9 medium with 25 mM succinate, washed three times in M9 medium with no added carbon source and suspended to an OD₆₀₀ of 0.1 in the same medium with 1 mM 2-phenylethanol. For luminescence measurements, 198 μ l of cell suspension was added to 2 μ l of a 100-fold-concentrated metal salt solution in white 96-well plates with a clear bottom (μ Clear; Greiner Bio-One). Microtiter plates were incubated in a FLX-Xenius plate reader (SAFAS, Monaco) at 30°C with orbital shaking (600 rpm, amplitude 3 mm) and light emission and OD₆₀₀ were recorded after the indicated time periods. Promoter activity was expressed as relative light units (RLU) normalized to the corresponding OD₆₀₀. Experiments were performed in triplicates and data are presented as the mean value with error bars representing the standard deviation.

TABLE 3 | Trace element concentrations of M9 medium and MP medium.

	M9 medium	MP medium
Na ₃ -citrate	51 μ M	45.6 μ M
H ₃ BO ₃	5 μ M	–
CoCl ₂	–	2 μ M
CuSO ₄	4 μ M	1 μ M
FeSO ₄	36 μ M	18 μ M
MnCl ₂	5 μ M	1 μ M
Na ₂ MoO ₄ /(NH ₄) ₆ Mo ₇ O ₂₄	0.137 μ M	2 μ M
NiCl ₂	0.084 μ M	–
Na ₂ WO ₄	–	0.33 μ M
ZnSO ₄	7 μ M	1.2 μ M



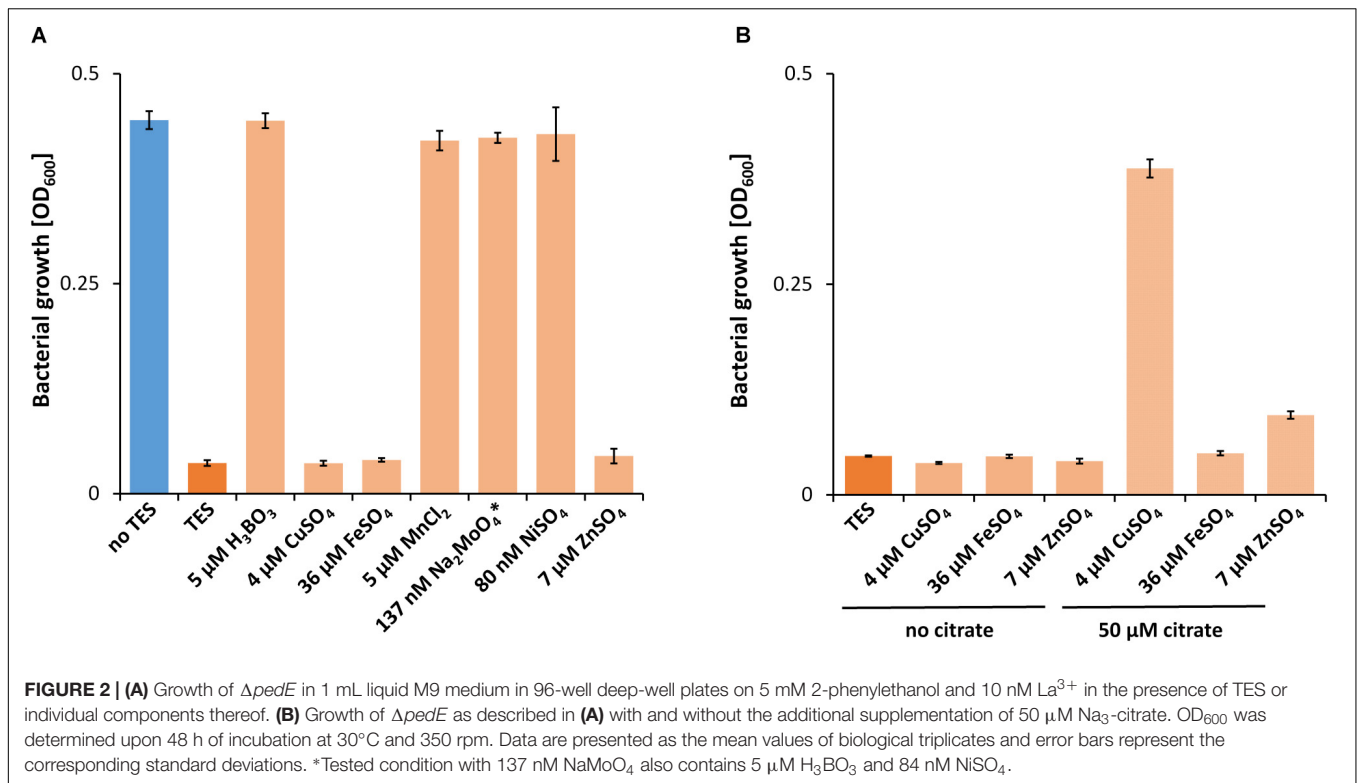
RESULTS

The REE-Switch in *P. putida* KT2440 Is Influenced by the Presence of Iron, Copper, and Zinc

Pseudomonas putida KT2440 makes use of a periplasmic oxidation system to grow on a variety of alcoholic substrates. Crucial to this system are two PQQ-dependent ethanol dehydrogenases (PQQ-EDHs), which share a similar substrate scope but differ in their metal cofactor dependency (Wehrmann et al., 2017). PedE makes use of a Ca^{2+} -ion whereas PedH relies on the bioavailability of different REE. During our studies, we found that the critical REE concentration that supports growth based on PedH activity differs dramatically depending on the minimal medium used. In a modified M9 medium, concentrations of about 10 μM of La^{3+} were necessary to observe PedH-dependent growth on 2-phenylethanol while only about 20–100 nM La^{3+} were required in MP medium (Wehrmann et al., 2017). One major difference between the two minimal media lies in their trace element composition and the respective metal ion concentrations (Table 3). The concentrations of copper, iron, manganese and zinc are between 2 \times and 7 \times higher in the modified M9 medium compared to MP medium, and other trace elements such as boron, cobalt, nickel or tungsten are only present in one out of the two media. To study the impact of the trace element solution (TES) on growth in the presence of La^{3+} , we used the $\Delta pedE$ strain growing

on 2-phenylethanol in M9 minimal medium in the presence and absence of TES.

While a critical La^{3+} concentration of 10 μM or higher was needed in the presence of TES to support PedH-dependent growth, this concentration dropped to as little as 10 nM La^{3+} in the absence of TES (Figure 1A). Similarly, inhibition of PedE-dependent growth by La^{3+} in strain $\Delta pedH$ differed dramatically depending on the presence of TES (Figure 1B). In the presence of TES, the addition of $\geq 100 \mu M$ of La^{3+} was required for growth inhibition in the $\Delta pedH$ strain within 48 h of incubation, whereas a minimum of only $\geq 1 \mu M$ La^{3+} was required in the absence of TES. From these experiments, we conclude that also a non-complemented minimal medium contains low, but sufficient, amounts of essential trace elements to allow growth even in the absence of TES. To find out whether the trace element mixture or a single trace element was causing the observed differences, we analyzed the growth of strain $\Delta pedE$ in more detail (Figure 2A). For concentrations of H_3BO_3 , Na_2MoO_4 , $NiSO_4$, and $MnCl_2$ similar to those found in complemented M9 medium, PedH-dependent growth on 2-phenylethanol in the presence of 10 nM La^{3+} was observed. In contrast, upon the individual supplementation with 4 μM $CuSO_4$, 36 μM $FeSO_4$ or 7 μM $ZnSO_4$ PedH-dependent growth could not be observed with 10 nM La^{3+} , as it was the case upon the supplementation with TES. Since citrate is used as a metal chelator in TES, we further tested the impact of citrate on growth inhibition of $CuSO_4$, $FeSO_4$ and $ZnSO_4$ when used as additional supplement (Figure 2B). The addition of 50 μM



of Na_3 -citrate restored growth of the $\Delta pedE$ strain in the presence of 4 μM $CuSO_4$ and 7 μM $ZnSO_4$, even though cell growth was still impaired for Zn-containing medium. However, in cultures containing 36 μM $FeSO_4$ the addition of citrate had no effect, strongly indicating that $FeSO_4$ is predominantly responsible for the inhibition of PedH-dependent growth of the $\Delta pedE$ strain under low La^{3+} concentrations in a TES complemented M9 medium.

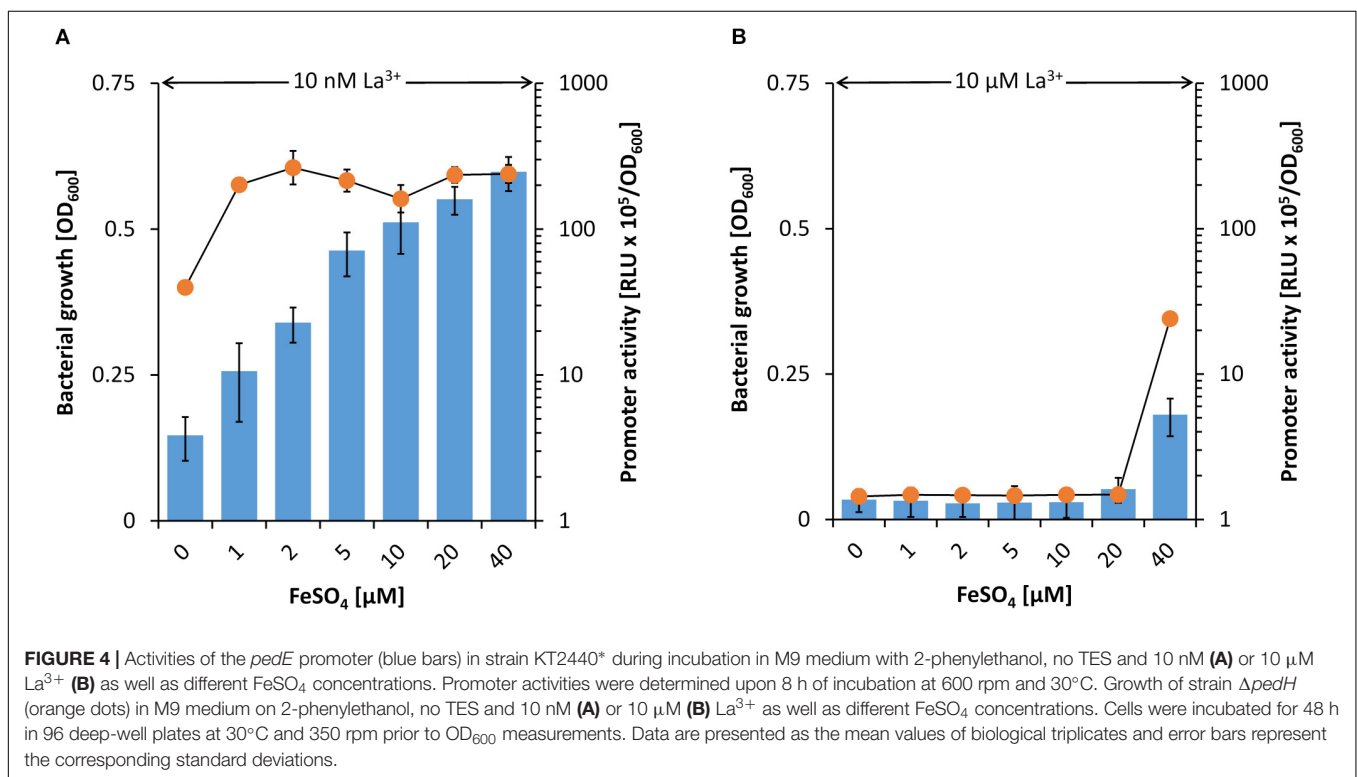
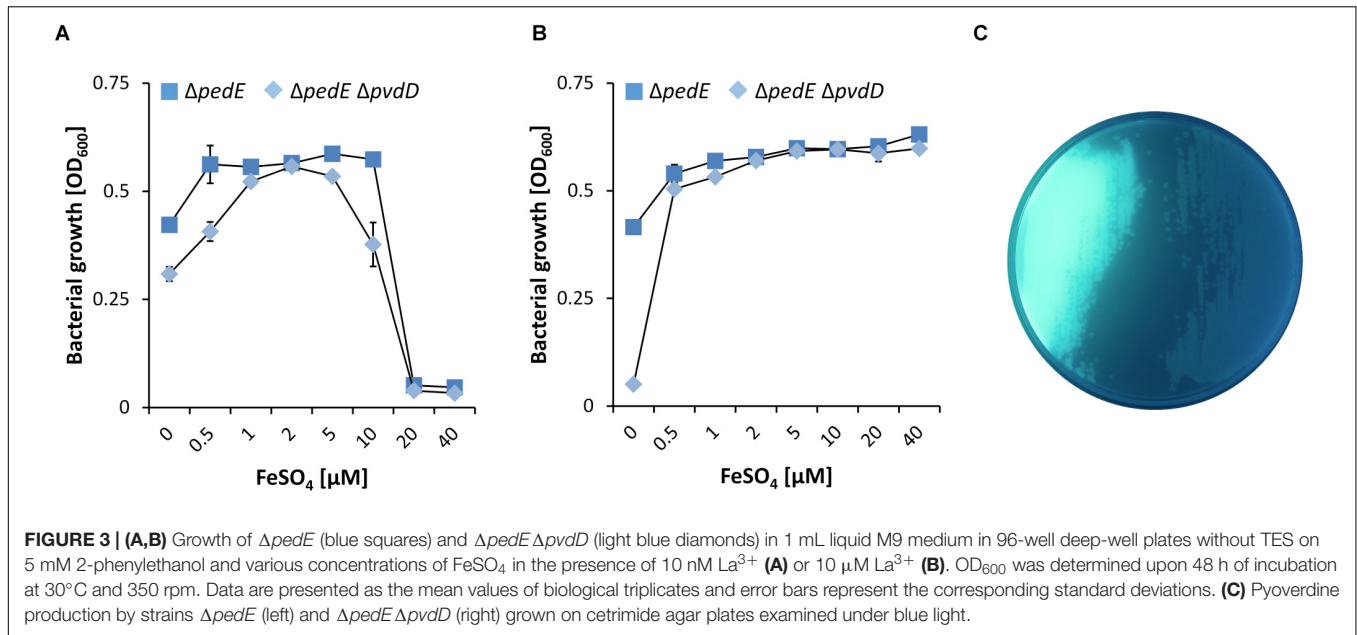
La^{3+} -Dependent Growth on 2-Phenylethanol Under Low La^{3+} Conditions Does Not Depend on Pyoverdine Production

To acquire iron under restricted conditions, *P. putida* KT2440 can excrete two variants of the siderophore pyoverdine (Salah El Din et al., 1997). Beside their great specificity toward Fe^{3+} , different pyoverdines can also chelate other ions including Al^{3+} , Cu^{2+} , Eu^{3+} or Tb^{3+} , although with lower affinity (Braud et al., 2009a,b). To test whether pyoverdine production in response to low iron conditions facilitates growth on 2-phenylethanol under low La^{3+} conditions, the mutant strain $\Delta pedE\Delta pvdD$ was constructed. This strain is no longer able to produce the two pyoverdines due to the loss of the non-ribosomal peptide synthetase *pvdD* (PP_4219; formerly known as *ppsD*) (Matilla et al., 2007), which was confirmed upon growth on agar plates (Figure 3C). In experiments with varying $FeSO_4$ supplementation, we found that PedH-dependent growth on 2-phenylethanol of strain $\Delta pedE$ was only observed for $FeSO_4$ concentrations $\leq 10 \mu M$

under low (10 nM) La^{3+} conditions (Figure 3A). With $\geq 20 \mu M$ $FeSO_4$ in the medium, no growth was observed. Strain $\Delta pedE\Delta pvdD$ exhibited the same $FeSO_4$ -dependent growth phenotype as the parental strain under low La^{3+} concentrations. Under high (10 μM) La^{3+} conditions, strain $\Delta pedE$ exhibited PedH-dependent growth under any $FeSO_4$ concentration tested (Figure 3B). Notably, strain $\Delta pedE\Delta pvdD$ showed nearly the same growth pattern as $\Delta pedE$ under high La^{3+} concentrations, with the exception of the condition where no $FeSO_4$ was added to the medium. Under this condition, no growth was observed.

The REE-Switch Is Influenced by the $Fe^{2+/3+}$ to La^{3+} Ratio

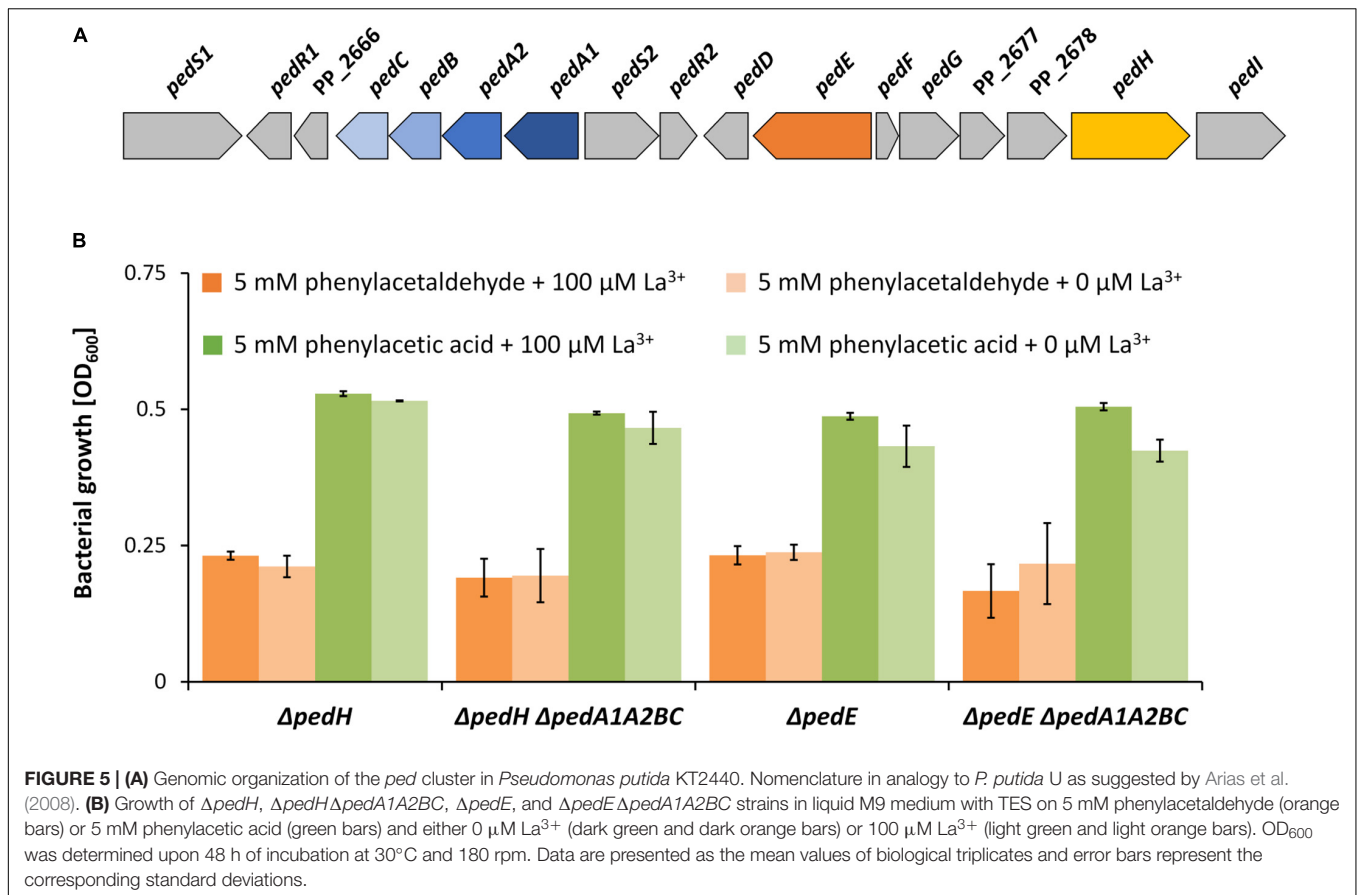
From the data described above, it can be speculated that beside the PedH-dependent growth also the inhibition of PedE-dependent growth on 2-phenylethanol is dependent on the $Fe^{2+/3+}$ to La^{3+} ratio. In the presence of 10 nM La^{3+} , *pedE* promoter activity was comparably high and increased with increasing $FeSO_4$ concentrations. In addition, strain $\Delta pedH$ grew readily on 2-phenylethanol under all these conditions even with no $FeSO_4$ supplementation (Figure 4A). When 10 μM La^{3+} was available, no growth of the $\Delta pedH$ mutant was observed in presence of $\leq 20 \mu M$ $FeSO_4$ and the *pedE* promoter activities were low (Figure 4B). However, when 40 μM $FeSO_4$ were present in the medium, representing a fourfold excess compared to La^{3+} , PedE-dependent growth and an increased *pedE* promoter activity was detected.



PedA1A2BC Is Required for La^{3+} -Dependent Growth on 2-Phenylethanol in Presence of Low La^{3+} Concentrations

Due to the very low concentrations of REEs (nM range) required for REE-dependent growth, it is commonly speculated that specific REE uptake systems must exist. From our previous

results, we can conclude that the produced pyoverdines do not contribute to such a system. A search of the genomic context of the *ped* gene cluster identified a putative ABC transporter system located nearby the two PQQ-EDHs encoding genes *pedE* and *pedH* (Figure 5A). The ABC-transporter is predicted to be encoded as a single transcript by the online tool “Operon-mapper” (Taboada et al., 2018). It consists of four genes encoding a putative permease [*pedC* (PP_2667)], an



ATP-binding protein [*pedB* (PP_2668)], a YVTN beta-propeller repeat protein of unknown function [*pedA2* (PP_2669)], and a periplasmic substrate-binding protein [*pedA1* (PP_5538)]. While efflux systems are usually composed of the transmembrane domains and nucleotide binding domains, ABC-dependent import system additionally require a substrate binding protein for functional transport (Biemans-Oldehinkel et al., 2006). As the gene *pedA1* is predicted to be such a substrate binding protein, it is very likely that this transporter represents an import system.

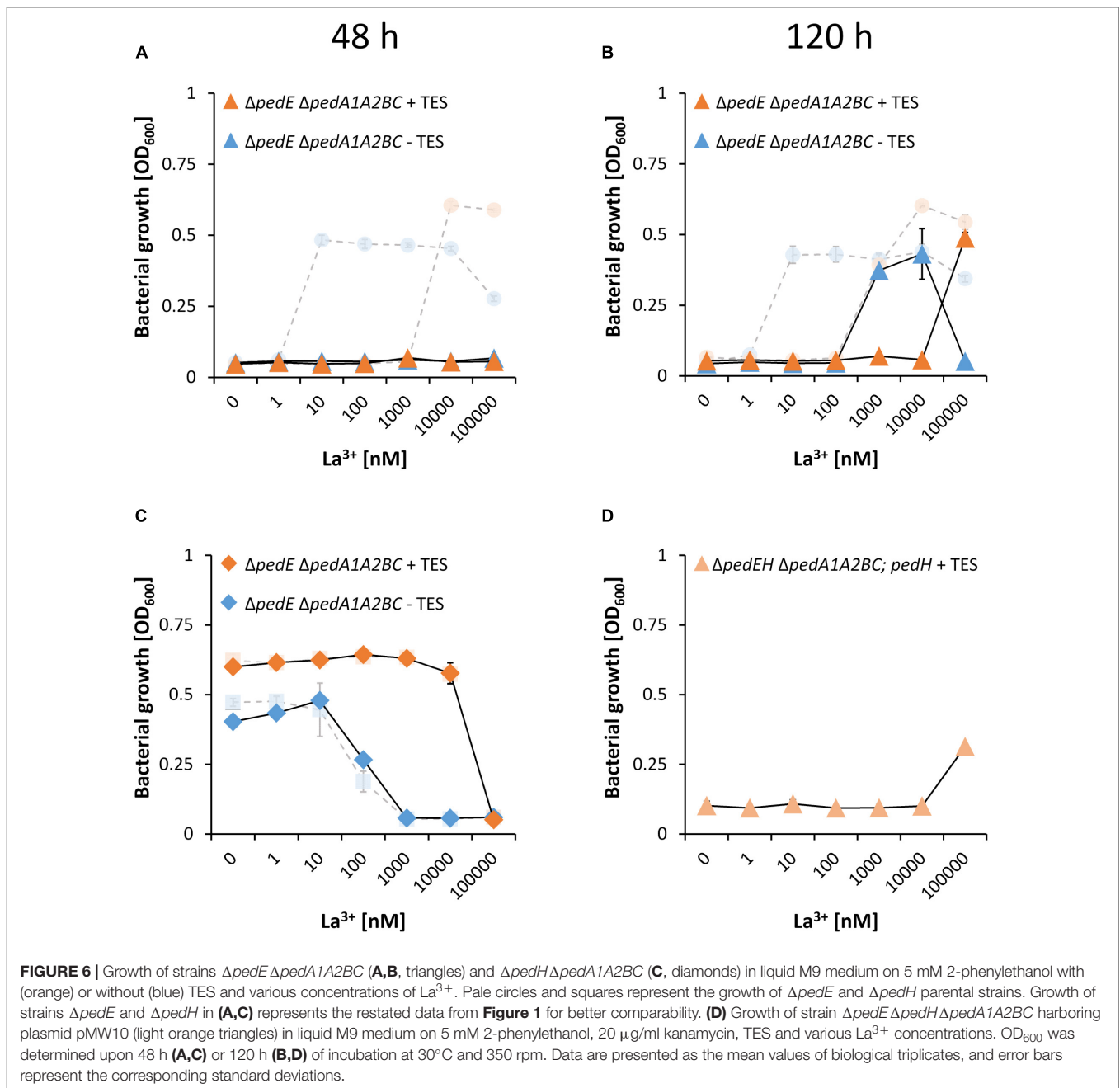
ABC-dependent importers can be specific for carbon substrates or metal ions. Growth experiments with $\Delta pedE$, $\Delta pedH$, $\Delta pedE \Delta pedA1A2BC$ and $\Delta pedH \Delta pedA1A2BC$ demonstrated that independent of La^{3+} (100 μM) availability, all strains were capable of growing on the oxidized degradation intermediates of 2-phenylethanol, namely 2-phenylacetaldehyde and phenylacetic acid, within 48 h of incubation (Figure 5B). This indicates, that the transport system is not involved in carbon substrate uptake.

When subsequently different La^{3+} concentrations were tested, we found that *PedH*-dependent growth on 2-phenylethanol of strain $\Delta pedE \Delta pedA1A2BC$ was inhibited for the first 48 h of incubation, irrespectively of the presence or absence of TES (Figure 6A). This was in contrast to the $\Delta pedE$ deletion strain, which grew in the presence of ≥ 10 nM La^{3+} or ≥ 10 μM La^{3+} depending on TES availability (Figures 1A, 6A pale symbols and lines). Upon an increased incubation time of 120 h, however,

strain $\Delta pedE \Delta pedA1A2BC$ eventually did grow with 1 and 10 μM La^{3+} or 100 μM La^{3+} depending on TES addition (Figure 6B). Notably, also the critical REE concentration for *PedH*-dependent growth of $\Delta pedE \Delta pedA1A2BC$ was increased by 100-fold compared to the $\Delta pedE$ strain under all conditions tested. Assuming that *PedA1A2BC* is specific for REE uptake, growth with the Ca^{2+} -dependent enzyme *PedE* should not be influenced by a loss of the transporter function. When we tested the $\Delta pedH$ and $\Delta pedH \Delta pedA1A2BC$ strain, we indeed could not find any difference in growth as both strains exhibited a similar inhibition pattern for concentrations ≥ 1 μM La^{3+} or ≥ 100 μM La^{3+} depending on the absence or presence of TES in the medium (Figure 6C).

PedA1A2BC Does Not Influence La^{3+} -Dependent Expression of *pedH*

ABC-transporter systems, or the transported compounds, can be involved in transcriptional regulation of specific target genes (Biemans-Oldehinkel et al., 2006). Thus, the impaired growth on 2-phenylethanol under low La^{3+} concentrations of the $\Delta pedE \Delta pedA1A2BC$ strain might be caused by the lack of transcriptional activation of the *pedH* gene. To test this hypothesis, strain $\Delta pedE \Delta pedH \Delta pedA1A2BC$ was complemented with a *pedH* gene independent of its natural promoter. Growth analysis of this strain on 2-phenylethanol

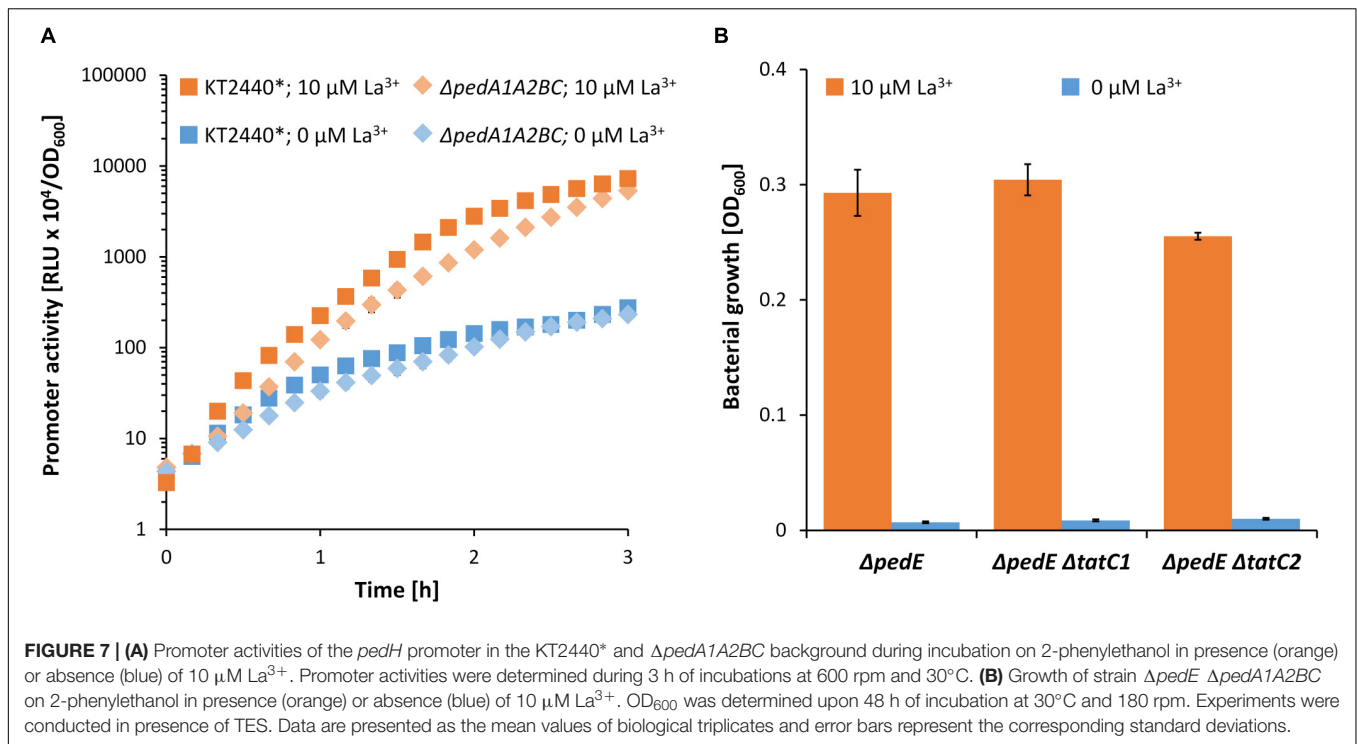


in the presence of TES and varying La³⁺ concentrations revealed no difference in the growth pattern when compared to strain $\Delta pedE \Delta pedA1A2BC$ (Figure 6D). This indicated that the impaired growth phenotype of the ABC-transporter mutant is not due to a lack of transcriptional activation of *pedH*. To further validate this conclusion, *pedH* promoter activities were measured during incubation with 2-phenylethanol in strain $\Delta pedA1A2BC$ and its parental strain in the absence and presence of 10 μ M La³⁺ (Figure 7A). Both strains showed a very similar induction of the *pedH* promoter in response to La³⁺ supplementation over time, which was about 20-fold higher at the end of the experiment (3 h) compared to the incubations in the absence

of La³⁺ (26-fold for KT2440*:Tn7-*pedH-lux* and 23-fold in $\Delta pedA1A2BC$:Tn7-*pedH-lux*).

La³⁺-Dependent Growth on 2-Phenylethanol of Strain $\Delta pedE$ Is Not Impaired Upon Interference With the Twin-Arginine Translocation (Tat) System

In contrast to PedE, the signal peptide of PedH contains two adjacent arginine residues, which is an indication that it might be transported to the periplasm in a folded state via the Tat (twin-arginine translocation) protein translocation system



(Berks, 2015). Therefore, one could argue that the transport of lanthanides into the cytoplasm might be beneficial as the incorporation into the active site of PedH could be more efficient during protein folding compared to the complementation of the apoenzyme in the periplasm. An initial analysis of the PedH signal peptide using different online software tools (TatP, PRED-TAT, SignalP 5.0, TatFind) could neither confirm nor refute this hypothesis (Rose et al., 2002; Bendtsen et al., 2005; Bagos et al., 2010; Almagro Armenteros et al., 2019). Therefore, we generated strains $\Delta pedE \Delta tatC1$ and $\Delta pedE \Delta tatC2$ in which the two individual TatC proteins [TatC1 (PP_1039) and TatC2 (PP_5018)] encoded in the genome of KT2440 are deleted. These strains should be restricted in the translocation of folded proteins into the periplasm, and if PedH would represent a Tat substrate, impaired growth on 2-phenylethanol in the presence of La³⁺ should be observable. However, neither a *tatC1* nor a *tatC2* mutation affected La³⁺-dependent growth on 2-phenylethanol (Figure 7B). Notably, various attempts to generate the double *tatC1/C2* mutant strain were unsuccessful.

DISCUSSION

In the present study, we reveal that iron availability severely affects the REE-switch in *P. putida* KT2440. This is evidenced by the reduction of the critical concentration of La³⁺ that is required both to promote PedH-dependent growth and for the repression of growth on 2-phenylethanol based on PedE activity. By using a $\Delta pvdD$ deletion strain, we demonstrate that the production of the iron chelating siderophore pyoverdine is not

required for PedH-dependent growth on 2-phenylethanol under low La³⁺ conditions. Our data suggest that the observed effects during high Fe^{2+/3+}/La³⁺ ratios are caused by mismetallation. In this scenario, the La³⁺-binding sites of proteins could be occupied by Fe^{2+/3+} ions that are in excess in the medium, and can also be present in the same 3+ oxidation state (Webb, 1970; Tripathi and Srivastava, 2006; Tottey et al., 2008; Foster et al., 2014). Transcriptional data show that *pedE* repression can be influenced by iron in a concentration dependent manner. Further, the impact of iron is not identical for PedE and PedH-dependent growth (100 fold vs. 1000 fold). Since PedE regulation is solely dependent on PedS2 (Wehrmann et al., 2017; 2018), these data are thus supportive of such a mismetallation hypothesis, assuming that the sensor histidine kinase PedS2 and PedH have different binding affinities to La³⁺ and/or Fe^{2+/3+}.

The same hypothesis might similarly explain why under high La³⁺ concentrations and in the absence of Fe^{2+/3+} supplementation, a pyoverdine-deficient strain is strongly impaired in growth. In this scenario the Fe^{2+/3+} binding sites of pyoverdine-independent Fe transporters, such as the ferrichrome, ferrioxamine and ferric citrate uptake systems, might be occupied by La³⁺ and prevent binding of Fe^{2+/3+} ions (Jurkevitch et al., 1992; Cornelis, 2010). Consequently, a pyoverdine deficient strain would be unable to take up enough of this essential element that is, most likely, present at trace levels in the medium even without additional supplementation.

It is further interesting to point out that also micromolar Cu²⁺ and Zn²⁺ inhibited growth on 2-phenylethanol in presence of La³⁺ in the nanomolar range, although these metals do not exist in the same 3+ oxidation state under natural conditions.

They are, however, the divalent transition metals that form the most stable complexes irrespective of the nature of the ligand, and as such also competitively bind non-cognate metal binding sites with high strength (Irving and Williams, 1953; Foster et al., 2014). Notably, Cu^{2+} has also been reported to interfere with REE-dependent regulation of PQQ-dependent methanol dehydrogenases in *M. trichosporium* OB3b (Gu et al., 2016; Gu and Semrau, 2017), and it is tempting to speculate that mismetallation might be involved in this process, too.

We provide compelling evidence that the predicted ABC-transporter PedA1A2BC is essential for PedH-dependent growth on 2-phenylethanol under low concentrations of La^{3+} . Based on the PedE-dependent growth phenotype, we can further show that PedA1A2BC is not involved in transcriptional repression of *pedE* under low La^{3+} conditions. The fact that a $\Delta pedE \Delta pedA1A2BC$ mutant strain can only grow with a 100-fold higher concentration of La^{3+} compared to the $\Delta pedE$ single mutant strongly indicates that PedA1A2BC functions as a La^{3+} -specific importer into the cytoplasm. Recently, it was demonstrated that in several *Methylobacterium extorquens* strains a similar ABC-transporter system is required for Ln^{3+} -dependent growth (Ochsner et al., 2019; Roszczenko-Jasińska et al., 2019). A BLAST analysis revealed that these ABC transporters show high similarities to all four genes of the *pedA1A2BC* operon (>43% sequence identity for *pedA2* and >50% for *pedA1*, *pedB* and *pedC*) and that all bacterial strains that have been reported to produce Ln^{3+} -dependent PQQ-ADHs thus far, carry homologs of this transporter system in their genome. Using a protein-based fluorescent sensor with picomolar affinity for REEs, Mattocks and colleagues were able to demonstrate that *M. extorquens* indeed selectively takes up light REEs into its cytoplasm (Mattocks et al., 2019) and it was later shown that cytoplasmic REE-uptake depends on the presence of the previously identified ABC-transporter system (Roszczenko-Jasińska et al., 2019).

Since the PedH enzyme, like all currently known Ln^{3+} -dependent enzymes, resides in the periplasm and since the purified apoenzyme of PedH can be converted into the catalytically active holoenzyme by Ln^{3+} supplementation *in vitro*, the question arises what the potential advantage of the postulated cytoplasmic Ln^{3+} uptake for *P. putida* would be. From our point of view, two different reasons can be imagined, namely that (i) the REE-dependent PedH protein is folded within the cytoplasm and the incorporation of the La^{3+} -cofactor is only possible or more efficient during the folding process; or (ii) La^{3+} binds to a cytoplasmic protein that either represents a so-far uncharacterized transcriptional regulator or another REE-dependent enzyme.

It has been demonstrated that the location of protein folding can regulate metal binding (Tottey et al., 2008). As such, Ln^{3+} insertion during folding in the cytoplasm, where metal concentrations are tightly regulated, could provide a means of preventing the Ln^{3+} binding site of PedH from mismetallation with potentially competitive binders such as Cu^{2+} , Zn^{2+} , or $\text{Fe}^{2+/3+}$ in the periplasm. However, we could not find evidence that PedH is a Tat substrate and consequently transported into the periplasm as a folded protein (Berks, 2015), as the individual *tatC1* or *tatC2* mutants both still showed PedH-dependent

growth. However, it cannot be excluded that the two Tat systems are functionally redundant since attempts to generate a *tatC1/C2* double mutant strain proved unsuccessful.

We can further conclude that the putative La^{3+} transport into the cytoplasm is not required to activate *pedH* transcription. It is, however, possible that additional genes/proteins required for PedH-dependent growth rely on, or are regulated by, the cytoplasmic presence of REEs. In this context, it is interesting to note that in a recent proteomic approach, we found that besides PedE and PedH, additional proteins of unknown function show differential abundance in response to La^{3+} availability (Wehrmann et al., 2019). It will hence be interesting to find out, whether any of these proteins is required for PedH function.

In *Methylobacterium extorquens* PA1 and *Methylobacterium extorquens* AM1, almost identical TonB-dependent receptor proteins (>99% sequence identity) were found to be crucial for REE-dependent growth suggesting a specific Ln^{3+} -binding chelator system in these organisms (Ochsner et al., 2019; Roszczenko-Jasińska et al., 2019). Interestingly, also in *Methylobacterium buriatense* 5GB1C a TonB-dependent receptor was identified that is crucial for the REE-switch to occur (Groom et al., 2019). The latter receptor only shows <20% sequence identity to those of the *M. extorquens* strains and thus could not have been identified by homology searches.

In *Pseudomonas putida* KT2440, no close homolog to any of the aforementioned TonB-dependent receptors can be identified (<30% sequence identity). *P. putida* preferentially resides in the rhizosphere whereas *M. buriatense* and *M. extorquens* PA1 were isolated from the sediment of a soda lake (pH 9 – 9.5) and the phyllosphere of *Arabidopsis thaliana*, respectively (Kaluzhnaya et al., 2001; Knief et al., 2010). One could speculate that a specific Ln^{3+} -chelator system is perhaps not relevant in the rhizosphere due to the large reservoir of REEs within the soil and the usually acidic environment near the plant roots caused by the secretion of organic acids (Raghothama and Karthikeyan, 2005; Ramos et al., 2016). The lack of a homologous TonB-dependent receptor could also be explained by structural differences in the REE-specific chelator system that might be employed by *P. putida* compared to that of the methylotrophic bacteria. Lastly, it is also possible that in *P. putida* the REE uptake across the outer membrane proceeds via the same chelator systems that are used for pyoverdine independent Fe-acquisition. This could further provide another explanation for the impact of the $\text{Fe}^{2+/3+}$ to La^{3+} ratio on the REE-switch.

Overall, the present study expands the crucial role of a conserved ABC-transporter system, which was very recently identified as Ln^{3+} -specific inner membrane transport system in methano- and methylotrophs, to non-methylotrophic organisms. It further provides new insight into the complexity of bacterial-metal interactions and demonstrates that Cu, Zn and in particular Fe ions can strongly interfere with the REE-switch in *P. putida*, most likely through mismetallation. The body of knowledge how REEs impact protein function, gene regulation and consequently physiology of different microorganisms is rapidly increasing.

As such, it will be very interesting to see when some of the most interesting questions, such as the cytoplasmic function of REEs or the nature and potential structural diversity of specific REE-chelator systems, will be resolved by future studies.

DATA AVAILABILITY STATEMENT

All datasets generated for this study are included in the article/supplementary material.

AUTHOR CONTRIBUTIONS

MW, PB, and JK designed the research. MW, CB, and PB performed the experiments. All authors analyzed

the data. MW and JK wrote the manuscript with contributions of PB.

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