Role of phenazine-enzyme physiology for current generation in a bioelectrochemical system

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Introduction
Energy is the key factor, which drives metabolism in all organisms (Alexandre et al., 2004). Microorganisms produce energy in the form of ATP mainly by the systematic transfer of electrons through their electron transport chains to a terminal electron acceptor such as oxygen, nitrate or sulphate (Kracke et al., 2015; Kuyper et al., 2018). Electroactive microorganisms are in addition able to transport electrons through their cellular membrane to the extracellular environment to reduce solid-state electron acceptors such as Fe(III) and Mn(IV) (Kato, 2015; Shi et al., 2016). This discharge of electrons can be harnessed in a bioelectrochemical system (BES), where electrons are channelled to an electrode (anode) as an alternative electron acceptor (Pant et al., 2012; Logan et al., 2019). BES technology currently holds potentials in various fields, the most renowned being the microbial fuel cell (MFC) used for wastewater treatment (Pandey et al., 2016). Additionally, a fairly new field applies BES in biotechnology for new oxygen-limited bioprocesses, as they can provide an oxygen-independent alternative for discharging excess electrons and thereby maintaining the metabolic flux of the organism (Schmitz et al., 2015; Askitosari, 2019).

Electrons are transferred to the electrodes of a BES either by direct contact of the microorganisms or by using soluble redox mediators (Shi et al., 2016; Beblawy et al., 2018; Lovley and Walker, 2019; Saunders et al., 2020). For biotechnological applications, the use of the latter has several advantages. Redox mediators are in principle non-exhaustible (Brutinel and Gralnick, 2012) and based on the three-dimensional culture broth, there is no limit to the extent mediators can travel. This is in contrast to direct electron transfer, which is limited to the 2D surface area of the electrode. The natural mediators produced by Pseudomonas aeruginosa are termed phenazines, with phenazine-1-carboxylic acid (PCA) and pyocyanin (PYO) being the ones dominantly produced (Mavrodi et al., 2001). Although these colourful secondary metabolites are mostly known for the different roles they play as a pathogenicity factor in P. aeruginosa infections (Hall et al., 2016; Yang et al., 2016), they also have been shown to enhance the anaerobic survival of the organism by shuttling electrons to distant oxygen (Wang et al., 2010). Phenazines may receive electrons from NAD(P)H, as shown in vitro, and from some

Summary
Pseudomonas aeruginosa produces phenazine-1-carboxylic acid (PCA) and pyocyanin (PYO), which aid its anaerobic survival by mediating electron transfer to distant oxygen. These natural secondary metabolites are being explored in biotechnology to mediate electron transfer to the anode of bioelectrochemical systems. A major challenge is that only a small fraction of electrons from microbial substrate conversion is recovered. It remained unclear whether phenazines can re-enter the cell and thus, if the electrons accessed by the phenazines arise mainly from cytoplasmic or periplasmic pathways. Here, we prove that the periplasmic glucose dehydrogenase (Gcd) of P. aeruginosa and P. putida is involved in the reduction of natural phenazines. PYO displayed a 60-fold faster enzymatic reduction than PCA; PCA was, however, more stable for long-term electron shuttling to the anode. Evaluation of a Gcd knockout and overexpression strain showed that up to 9% of the anodic current can be designated to this enzymatic reaction. We further assessed phenazine uptake with the aid of two molecular biosensors, which experimentally confirm the phenazines’ ability to re-enter the cytoplasm. These findings significantly advance the understanding of the (electro) physiology of phenazines for future tailoring of phenazine electron discharge in biotechnological applications.
cytoplasmic enzymatic conversions of central metabolic pathways (Price-Whelan et al., 2007; Glasser et al., 2017; Jo et al., 2020). Reduced phenazines mediate these electrons to distant oxygen or other feasible electron acceptors such as the anode of a BES and with this, enable the survival of *P. aeruginosa* in an oxygen-limited environment (Pham et al., 2008; Glasser et al., 2014). Recently, biotechnologically important strains such as *P. putida* KT2440 and *E. coli* have been engineered to produce these phenazines, thereby boosting the prospects of using these redox-active compounds for electron transfer in electrochemical bioprocesses (Schmitz et al., 2015; Feng et al., 2018; da Silva et al., 2021).

However, the energetic use of phenazines for electron discharge, both in native and biotechnologically relevant *Pseudomonas* species, is limited, and only a small fraction of electrons provided by the substrate, for example glucose, is discharged to the extracellular electron acceptor (Bosire et al., 2016). Additionally, this electron discharge currently does not seem to directly yield ATP for the cell (Glasser et al., 2014). Since phenazine biosynthesis and all identified phenazine reduction pathways occur in the cytoplasm (Blankenfeldt and Parsons, 2014; Glasser et al., 2017; Jo et al., 2020), a central question is if phenazines can in fact re-enter the cell once they have been pumped outside. This has not been experimentally verified yet, but is a requirement to enable a potentially broader enzymatic reduction. We here hypothesize that a significant part of the phenazine reduction takes place already in the periplasm (Fig. 1) and tested this hypothesis with two approaches: (i) specific phenazine reduction by periplasmic reductases and (ii) the development of bioassays to verify phenazine entry into the cytosol.

*Pseudomonas* species commonly utilize glucose via a direct glucose oxidation pathway, which occurs in the periplasm (Daddaoua et al., 2010; Kohlstedt and Wittmann, 2019). This pathway involves the oxidation of glucose to 2-ketogluconic acid via gluconate, and both sugar acids are commonly found as accumulated intermediates in *Pseudomonas* BES cultivations (Schmitz et al., 2015; Bosire et al., 2016). The first step of this reaction, the oxidation of glucose to gluconate, is catalysed by a membrane-bound pyrroloquinoline quinone (PQQ)-dependent glucose dehydrogenase (Gcd), which transfers two electrons via PQQ to ubiquinone, localized in the membrane and eventually to the electron transport chain (Van Schie et al., 1985; Kobayashi et al., 2005). Here, we evaluate this enzyme for its capacity of phenazines reduction.

To assay the uptake of phenazines into the cytosol, we developed two molecular biosensors. One of them utilized the fact that PYO is synthesized intracellularly from PCA as a precursor (Mentel et al., 2009). We provided external PCA to a strain exclusively harbouring the genes to turn PCA into PYO, and measured the cytoplasmic transformation into PYO. The second biosensor tracked PYO access to the cell by harnessing the ability of endogenous phenazines to oxidize the (2Fe-2S) transcription factor SoxR, which in turn activates the promoter of the MexGHI-OpmD efflux pump operon involved in phenazine export (Palma et al., 2005; Dietrich et al., 2006; Sakhtah et al., 2016).

**Results**

To test our first hypothesis that phenazine reduction can take place in the periplasm, we evaluated the interaction of the phenazines with the first enzyme of the periplasmic glucose oxidation pathway *in vitro* and *in vivo*.

**Membrane-bound glucose dehydrogenase (Gcd) of Pseudomonas reduces phenazines in vitro**

We first tested, if phenazines could interact with the membrane-bound glucose dehydrogenase (Gcd) of *Pseudomonas* and as such mediate the electrons generated through the oxidation of glucose in the periplasmic space to extracellular terminal electron acceptors. The *gcd* genes of *P. aeruginosa* PA14 (PA14_34970) and *P. putida* KT2440 (PP_1444) were cloned and expressed in *E. coli*. Both proteins, each made up of 803 amino acid residues, are PQQ-dependent and require divalent ions such as calcium ions as an enhancing co-factor. The initial 138 to 140 amino acids at the N-terminus form a membrane anchor with five transmembrane helices, while the remaining part of the protein, including the catalytic site, flanks the periplasmic space. The final construct of the Gcds used for protein expression contained C- and N-terminal His-tag sequences and excluded the first 8 native amino acids, since the construct with the full native sequence showed retarded growth and a low protein expression. The membrane proteins were purified for the enzyme assay using affinity and size permeation chromatography (Fig. S1).

If the Gcds can interact with phenazines, they should be able to reduce these mediators by transferring electrons from the glucose oxidation through the POQ co-factor to the phenazines (Fig. 1). An enzyme assay was performed with the purified proteins using PCA and PYO as prospective electron acceptors. The standard electron acceptors of an established dehydrogenase enzyme assay (Jahn et al., 2020), phenazine methanosulfate (PMS) and dichlorophenolindophenol (DCPIP), were used as control reaction to verify activity of the enzyme. When either PCA or PYO was used as electron acceptor in this reaction, the Gcd of *P. aeruginosa* PA14 was able...
to catalyse the transfer of electrons from glucose to these phenazines, which resulted in their measurable reduction (Fig. 2A and B). A control in which water was used to replace either the substrate or the enzyme did not show any phenazine reduction. We also performed the same assay with Gcd obtained from *P. putida* KT2440, a prospective organism for limited-oxygen BES processes (Askitosari *et al.*, 2019). Comparison of the Gcds of *P. aeruginosa* PA14 and *P. putida* KT2440 showed 81.64% and 78.93% sequence identities in their nucleotide and amino acid sequences respectively. The enzyme assay with the Gcd of *P. putida* KT2440 equally showed reduction of PCA and PYO utilizing electrons from glucose oxidation (Fig. 2C and D). No reduction of the phenazines was observed for both enzymes when PQQ or calcium ions were excluded from the reaction (results not shown). The reduction rate of PYO was about 60 times faster than that of PCA for both Gcds (Fig. 2). The experiments clearly confirmed the membrane-bound glucose dehydrogenases of *P. aeruginosa* and *P. putida* as potential phenazine reducing enzymes.

**Confirming Gcd-phenazine reduction in vivo using bioelectrochemical system (BES)**

Having established that a periplasmic enzyme can interact with phenazines, we tested if this is a major pathway for phenazine reduction in vivo. To ascertain the proportion of electrons generated from the periplasmic glucose oxidation pathway versus downstream cytoplasmic pathways, a BES was set up using three *Pseudomonas putida* KT2440 strains: a wild-type strain (KT2440 wt), a glucose dehydrogenase deletion mutant (KT2440 Δgcd), in which the native Gcd was removed, and an overexpression strain, which was supplied with a plasmid for expressing the glucose dehydrogenase of *P. aeruginosa* PA14 (KT2440 Δgcd–PA14gcd). The Gcd of PA14 was used for the overexpression because PA14 is a natural producer of phenazines and had an overall better performance in the enzyme assay (Fig. 2). The wild-type and overexpression strains are able to metabolize glucose using the periplasm as well as the cytoplasmic pathways, while the Gcd mutant strain can only use the cytoplasmic pathway. PCA (80 mg l⁻¹) or PYO (18 mg l⁻¹)
were added to the system 23 h after inoculation (time point ‘a’ in Fig. 3A and B). A higher concentration of PCA was used because it showed a lower Gcd reduction rate (Fig. 2) and to be in consistency with the amounts typically produced by *Pseudomonas* in a BES (Askitosari et al., 2019). As expected, no electrons were discharged to the anode during the active aeration growth period (first 31 h) as electrons from glucose oxidation were preferably moved to oxygen (Fig. 3A and B). The current, however, increased for the three strains once the supply of oxygen was turned off (passive aeration, time point ‘b’ in Fig. 3A and B), indicating that phenazines mediated electrons to the anode (Fig. 3A and B). The KT2440 Δ*gcd*–*PA14* gcd strain attained the highest maximum current density (25.9 µA cm⁻²) in the PCA-BES, which was a 55% and 63% increase compared to the KT2440 wt and the KT2440 Δ*gcd* respectively (Fig. 3A). Similarly, in the PYO-BES, KT2440 Δ*gcd*–*PA14* gcd had a peak current density of 12.2 µA cm⁻², while the KT2440 wt and KT2440 Δ*gcd* showed a maximum of 8.5 µA cm⁻² (~ 70% of former) and 4.9 µA cm⁻² (~ 40% of former) respectively (Fig. 3B). Thus, the current density increase for the overexpression strain was somewhat lower in the PCA-BES, which supports our earlier finding that the PYO enzymatic reduction was faster and more efficient than that with PCA (Fig. 2). The decrease in current density observed in all the BES after the peak current was reached, was primarily due to a slower substrate conversion rate (Table 1) under sustained passive aeration, but also for PYO because of degradation of this phenazine over time. PCA with a half-life of more than 10 days appeared more stable than PYO, which half-life was about one day (Fig. 4).

More detailed analyses of glucose metabolism and the products of the direct glucose oxidation showed a correlation with the peak currents. The specific glucose consumption was highest for the first two days of active aeration, with rates being the lowest for the Gcd mutant and highest for the overexpression strain (Table 1). The initial higher glucose conversion rate especially for the overexpression but also the wild-type strains in both PCA- and PYO-BES could be linked to the utilization of the periplasmic pathway for glucose oxidation. This was

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Fig. 2. Phenazine reduction by periplasmic membrane-bound PQQ-dependent glucose dehydrogenase (Gcd) of *P. aeruginosa* PA14 (PA14 Gcd) and *P. putida* KT2440 (KT2440 Gcd). (A) increase in concentration of reduced phenazine-1-carboxylic acid (PCA) and (B) decrease in concentration of oxidized pyocyanin (PYO) by the interaction with PA14 Gcd; (C) increase in concentration of reduced PCA and (D) decrease in concentration of oxidized PYO by the interaction with KT2440 Gcd. To measure the phenazines, decrease in the absorbance of oxidized PYO was followed at 690 nm and the concentration (mM) was calculated using an extinction coefficient of 4.5 mM⁻¹cm⁻¹, while the formation of the reduced PCA was followed at 440 nm and the concentration (mM) was calculated using an extinction coefficient of 2.05 mM⁻¹cm⁻¹. The reaction mixture contained in 200 µl: 100 mM Tris-HCl pH 7.0, 1 mM CaCl₂, 0.45 mM of the phenazines, 65 µg ml⁻¹ enzyme or water, 5 µM PQQ and 20 mM glucose or water was used to complete the reaction. Results show the average of three replicates; error bars indicate standard deviation (SD).
conferred by the presence of gluconate and 2-ketogluconic acids in the supernatant (Fig 4). The amount of gluconate and 2-ketogluconic acids were expectedly low (about 0.2 g l\(^{-1}\)) in the wild type (Fig. 4A and B), since the rate of gluconate uptake in \(P.\ putida\) \(KT2440\) is well matched to glucose oxidation with a negligible amount going all the way to 2-ketogluconate (Nikel et al., 2015). The initial difference in the current density (after turning down the aeration) for both phenazines between the wild type and the mutant confirmed a contribution of the periplasmic pathway (Fig. 4C and D). It seems like phenazines can only access electrons and enable current production with a delay in the absence of Gcd. Meanwhile, the products of the periplasmic pathway were found accumulated in a high amount in the overexpression strain (Fig. 4E and F). In the PCA-BES, almost 95% of the glucose converted by the overexpression strain was found as gluconate and 2-ketogluconic acid after 2 days, showing that the sugar oxidation pathway was greatly accelerated by the overexpression; however, sugar acid uptake into the cytosol could not match their production. The effect was similar for this strain in the PYO-BES, although only 42% of the glucose was found as intermediate products of direct glucose oxidation. The correlation between the metabolic flux increase of the overexpression strain in the glucose oxidation pathway and the increased electron discharge with this strain to the BES anode strongly suggests that the direct glucose oxidation pathway indeed is an important interaction point for phenazines to mediate electrons to the anode. However, if electrons are discharged at this point, no energetic gain for the cell is possible, since the electrons will be diverted away from the electron transport chain, which drives ATP-production. To quantitatively compare the efficiency of the electron discharge via phenazines, the coulombic efficiency was calculated for the three strains. It gives the fraction of the electrons from glucose oxidation, which was mediated to the anode for current generation via the phenazines. For the PCA-BES, this value is 4.1%, 5.0% and 8.8% for the wild-type; Gcd mutant, and the overexpression strain, respectively, while it was expectedly lower in PYO-BES.

Table 1. Specific glucose consumption and maximum biomass yield of \(P.\ putida\) \(KT2440\) strains in BES using PCA or PYO as electron mediators.

| Strain          | PCA- BES                      | PYO- BES                      |
|-----------------|-------------------------------|-------------------------------|
|                 | Coulombic efficiency (%)      | Specific glucose consumption (g l\(^{-1}\) h\(^{-1}\)\(^{a}\)) | Maximum biomass yield (gDCW gGlc\(^{-1}\)) | Coulombic efficiency (%) | Specific glucose consumption (g l\(^{-1}\) h\(^{-1}\)\(^{a}\)) | Maximum biomass yield (gDCW gGlc\(^{-1}\)) |
| KT2440 wt       | 4.06                          | 0.09 ± 0.01                   | 0.17 ± 0.01                           | 0.76                      | 0.14 ± 0.03                     | 0.16 ± 0.04                           |
| KT2440 Δgcd     | 5.02                          | 0.05 ± 0.001                  | 0.16 ± 0.01                           | 0.81                      | 0.1 ± 0.02                      | 0.18 ± 0.01                           |
| KT2440 Δgcd-PA14 gcd | 8.84                      | 0.139 ± 0.02                  | 0.16 ± 0.02                           | 1.27                      | 0.17 ± 0.06                     | 0.18 ± 0.02                           |

\(\text{a}\). Specific glucose consumption was calculated for the first 2 days of the BES experiment.
with 0.8%, 0.8% and 1.3%, because the current was not sustained due to the instability of PYO. A higher coulombic efficiency of the overexpression strain is in agreement to the high current density obtained for this strain in comparison with the other strains and to the less complete oxidation of the glucose with accumulated intermediates (Fig. 4E and F). It should be highlighted that a coulombic efficiency of almost 9% is the highest ever achieved with P. putida.
observed for phenazine-based electron discharge with *Pseudomonas* (Askitosari et al., 2019).

Assessing the transfer capacity of phenazines across the cell membrane into the cytosol

The finding that the Gcd mutant could still generate current using the phenazines, even though it is not able to utilize the periplasmic direct glucose oxidation pathway, made it likely that phenazines also have access to electrons released in cytoplasmic metabolic pathways. Therefore, we developed two molecular biosensors to ascertain if phenazines, after initial release when synthesized or when externally added to the cell, could (re-)enter the cytoplasm (Fig. 5). The first biosensor was made to track PCA and was based on the two reaction steps that modify PCA to PYO, catalysed by the two enzymes PhzM, a methyltransferase, and PhzS, a monoxygenase (Mavrodi et al., 2001). The genes encoding PhzM and PhzS were expressed via a plasmid in KT2440 and a PA14 mutant strain, lacking the PCA biosynthetic genes (*P. aeruginosa* PA14 *Δphz1,2*) (Schmitz and Rosenbaum, 2020). Since neither organism can produce phenazines, but now house the cytosolic enzymes PhzM and PhzS, it is expected that externally added PCA would be transformed to PYO, only if PCA has access to the cytoplasm (Fig. 5A). Both biosensors (PA14 *Δphz1,2-Ms* and KT2440-Ms) converted externally added PCA (40 mg l⁻¹) to about 8 mg l⁻¹ PYO after 3 h (Fig. 6A). Less than 2 mg l⁻¹ PYO was detected in an in vitro PCA reduction assay using just the supernatant of the biosensor culture, which had been induced but was never supplied with PCA. This confirmed that majority of PCA conversion occurred in the cytoplasm and was not a result of secreted enzymes or lysed cells.

Next, we designed a biosensor, which could enable us to evaluate whether PYO has access to the cytoplasm (Fig. 5B). PYO has been shown to activate the constitutively expressed transcription factor SoxR by a reversible one-electron oxidation of its [2Fe-2S] cluster (Dietrich et al., 2006; Sheplock et al., 2013). Since neither organism can produce phenazines, but now house the cytosolic enzymes PhzM and PhzS, it is expected that externally added PCA would be transformed to PYO, only if PCA has access to the cytoplasm (Fig. 5A). Both biosensors (PA14 *Δphz1,2-Ms* and KT2440-Ms) converted externally added PCA (40 mg l⁻¹) to about 8 mg l⁻¹ PYO after 3 h (Fig. 6A). Less than 2 mg l⁻¹ PYO was detected in an in vitro PCA reduction assay using just the supernatant of the biosensor culture, which had been induced but was never supplied with PCA. This confirmed that majority of PCA conversion occurred in the cytoplasm and was not a result of secreted enzymes or lysed cells.

Biosensors induced with PYO showed more Gfp fluorescence compared to the un-induced or those induced with PCA (Fig. 6B). Un-induced biosensors, however, showed some fluorescence (Figs 6B and 7), which might be a result of superoxide ions generated during aerobic growth, which are able to activate SoxR although such oxidation is less effective in *P. aeruginosa* (Fujikawa et al., 2012). Inability of PCA to yield significant fluorescence compared to the un-induced biosensor could be a result of its low redox potential, since SoxR is only activated by phenazines of high redox potential (Sheplock et al., 2013; Sakhtah et al., 2016). Although SoxR in *P. aeruginosa* and *P. putida* have a similar DNA binding domain (Park et al., 2006; Sheplock et al., 2013), the PYO-induced KT2440 biosensor shows a comparatively lower fluorescence in respect to its PA14 counterpart. This may be due to the fact that KT2440 SoxR was unable to completely activate the pMexG promoter, since the respective operon is not part of its native target regulon (Park et al., 2006).

Overall, both biosensors independently show that PCA and PYO, once released from the cell after synthesis, are able to re-enter the cytoplasm and therefore are able to access the diverse enzymatic reactions, for which an interaction with phenazines has already been described in vitro (Glasser et al., 2017; Jo et al., 2020).

Discussion

The low electron discharge efficiency of phenazines to the anode of a BES and the poor understanding of the physiology of phenazine electron transport presents a challenge for further developing an oxygen-limited bio-process that would be supported by phenazine electron mediation. We anticipated that if periplasmic redox enzymes interact with phenazines, improving such interaction could increase the electron flux to the anode, as it has been observed for the synthetic redox mediator ferri-cyanide (Yu et al., 2018) with which highly efficient glucose oxidation to 2-ketogluconate was coupled to anodic electron discharge. Here, we have confirmed that the periplasmic PQQ-dependent glucose dehydrogenase (Gcd) is able to reduce PCA and PYO. PCA, the precursor of PYO, had an about 60-fold lower enzymatic reduction rate than PYO (Fig. 2). The reason might be their structural difference or their different redox properties (E°[PCA]=−0.24 V; E°[PYO]=−0.18 V vs. Ag/AgCl sat.) (Bosire et al., 2016). The different functional groups surrounding the core phenazine structure have been shown to affect both, the chemical properties such as partitioning coefficients and the redox potentials of the individual derivatives (Bellin et al., 2014). Nevertheless, the longer half-life exhibited by PCA in BES operations resulted in a better performance over time compared to PYO (Fig. 3). Hence, PCA is better suited if long-term phenazine electron mediation is desired. Since phenazine electron discharge in *Pseudomonas* is not

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limited to glucose as a substrate, we also investigated other PQQ-dependent periplasmic enzymes. The alcohol substrates 2,3-butanediol and ethanol also show anodic current production (Bosire et al., 2016) and can be metabolized by the periplasmic PQQ-dependent ethanol dehydrogenase (Edh). However, the in vitro assay of this

Fig. 5. Schematic illustration of the biosensor for evaluating phenazine ability to re-enter the cell once they are released. A. detects PCA ability to enter the cytoplasm through the conversion of externally added PCA to PYO by the gene products of phzM and phzS. B. detects phenazine (especially PYO) ability to enter the cytoplasm via the oxidation of SoxR by the phenazines, which subsequently activates the promoter of the MexGHI efflux pump (pMexG) inducing gfp expression.

Fig. 6. Detection of phenazine ability to enter the cytoplasm of P. aeruginosa PA14 and P. putida KT2440 using cellular biosensors. A. Biosensors which convert PCA to PYO if the former enters the cytoplasm. (a) P. aeruginosa PA14 Δphz1,2 (phenazine knockout, control strain) and PA14 Δphz1,2-MS (biosensor); (b) P. putida KT2440 (wild type, control strain) and KT2440-MS (biosensor) were tested. Cells were grown to OD 1 before addition of PCA (40 mg l⁻¹) and induced with 0.1 mM sodium salicylate and were further cultivated for 3 h. (+) added PCA, (*) induction by sodium salicylate, (+*) added PCA and induction by sodium salicylate, and (+**) PCA was added to the supernatant of induced biosensors strain and incubated for 1 h at 37°C. (B) Biosensors for the detection of intracellular PYO – based on a PYO-regulated promoter controlling Gfp fluorescence. The gfp gene was cloned behind the promoter of the P. aeruginosa PA14 efflux protein MexGHI-OpmD (pMexG), which is activated when phenazines oxidize the redox regulator SoxR. P. aeruginosa PA14 Δphz1,2-pMexG-gfp (biosensor) and P. putida KT2440-pMexG-gfp (biosensor) and their respective controls were induced after 3 h for the expression of Gfp with 4 mg l⁻¹ (20 µM) PYO or 9 mg l⁻¹ (40 µM) PCA and cultivated for 16 h. The fluorescence of the control strains shows the auto-fluorescence of the organism. Fluorescence of the Gfp construct without phenazine addition shows untargeted induction of the pMexG promoter, for example by other redox-active compounds. Unpaired t-test was used to compare the fluorescence of Gfp and is represented as (**P < 0.01; ***P < 0.001 and n.s.-no statistical significance-P > 0.05). Results are average of three replicates, and error bars represent the SD.
enzyme showed no reduction of PCA or PYO with either of the alcohols (data not shown). In contrast, the artificial electron acceptor DCPIP was reduced in this assay, showing functionality of the Edh enzyme. The PQQ-dependent enzyme reaction is believed to use the same (hydrid) transfer mechanism for both alcohol (Edh) and aldehyde (Gcd) dehydrogenases (Anthony, 1996; Oubrie et al., 1999). Nonetheless, factors such as differences in the charged amino acid residues around the substrate entrance pocket might still influence electron acceptor preference (Hiraka et al., 2020). The mode of extracellular electron transfer achieved in vivo using phenazines or other redox mediators, however, does not only depend on the enzymes, which directly catalyse the reduction of these mediators. Other respiratory components of the cell, such as menaquinone, cytochrome reductase and terminal oxidases, were recently shown to play a critical role in the reduction of these mediators (Feng et al., 2020; Jo et al., 2020; Lai et al., 2020). However, the delay experienced in attaining peak currents with the Gcd mutant strain in addition to the improved current from the overexpression strain point to a central role of periplasmic Gcd for phenazine-based electron discharge. Further research is now required to elucidate a possible connection of the Gcd-phenazine interaction to the reported components of the electron transport chain.

While our results from the periplasmic enzyme investigation in the BES showed a strong involvement of the glucose oxidation pathway in phenazine charging, they also showed that even when this pathway is abolished in the Gcd mutant strain, electrons are transferred to phenazines. If glucose cannot be oxidized in the periplasm, it is directly taken up into the cytoplasm and phosphorylated by a glucokinase (Nikel et al., 2015). Although several cytoplasmic enzymes have been shown to be able to reduce phenazines (Glasser et al., 2020;...

Fig. 7. Microscopy images of the parent and biosensor strains for evaluating phenazine cytoplasmic access. Bright field and the corresponding fluorescent microscopy of the strains used to track phenazine access to the cytoplasm through the expression of Gfp. Scale bar: 10 µm.

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2017; Jo et al., 2020), it was not clear if phenazines could re-enter the cytoplasm once they were released after their synthesis. Since phenazines themselves are also toxic, _Pseudomonas aeruginosa_ uses different mechanisms such as a MexGHI-OpmD efflux pump and PumA, a monoxygenase, for self-resistance (Sakhtah et al., 2016; Sporer et al., 2018). While the efflux pump is speculated to be involved in phenazine expulsion, no putative entry pathway has been proposed yet. A direct observation of the phenazine transfer across the cell membrane is difficult due to the lack of labelling strategies for phenazines. Instead, with the aid of molecular biosensors, we here show that both phenazines can have access to the cytoplasm once released and consequently access the electron pool in this part of the cell.

However, taking all presented experiments together; the sustainability of extracellular current production in the _gcd_ deletion mutant, the established uptake of PCA and the probable uptake of PYO into the cytoplasm; it is not clear yet, why this access to metabolic electrons is so limited, that is does not translate to higher coulombic efficiencies. Potentially, several central enzymes of the glucose catabolism should be able to reduce phenazines, diverting electrons away from the native electron transport chain for oxygen reduction. Moreover, the fact that phenazine reduction is influenced by components of the electron transport chain (Jo et al., 2020), gives further indication that there must be mechanism to prevent cytosolic enzymes from reducing phenazines. This makes sense from an energetic point of view, because phenazine reduction by catabolic enzymes would result in the loss of reducing equivalents without energy conservation. Therefore, a better understanding of the control or synergy between catabolic enzyme catalysed phenazine reduction and reduction of phenazine by components of the respiratory chain would provide a wider view to understand and improve phenazine extracellular electron transfer. With this next level of insight, finally an answer could be found to the question if and how phenazine-based electron discharge can or could be coupled to cellular ATP generation.

**Experimental procedures**

*Microorganisms, plasmids and culture conditions*

Details on all strains and plasmids used for this study are listed in supporting Table S1.

All _E. coli_ strains were grown in lysogeny broth (LB) containing 50 \( \mu \)g ml\(^{-1} \) kanamycin (km) at 37°C. For enzyme purification, _E. coli_ expressing the gene for _gcd_ were grown in a 4-L batch using a Biostat® B-DCU bioreactor (B. Braun Biotech International GmbH, Melsungen Germany) as detailed in the Data S1. _P. aeruginosa_ PA14 and _P. putida_ KT2440 were initially cultivated in LB at 37°C and 30°C respectively. For all experiment with _P. aeruginosa_ PA14, MOPS medium was prepared according to (Neidhardt et al., 1974) with 3.5 g l\(^{-1} \) glucose as carbon source. 50 \( \mu \)g ml\(^{-1} \) km was used as antibiotic and 0.1 mM sodium salicylate was used to induce the plasmids when required. For the biosensor experiment, a 10 ml culture was cultivated until an optical density of 1, after which PCA was added and cells were induced when necessary. MOPS medium was also used for the _P. putida_ KT2440 biosensor experiment converting PCA to PYO because it was easier to keep the pH stable. All other experiments with _P. putida_ KT2440 were performed in a Delft mineral salt medium, which contained per litre: 10 g glucose, 3.88 g K_2HPO_4, 1.63 g NaH_2PO_4, 2 g (NH_4)_2SO_4, 0.1 g MgCl_2 \_6 H_2O, 10 mg EDTA, 2 mg ZnSO_4 \_7H_2O, 1 mg CaCl_2 \_2H_2O, 5 mg FeSO_4 \_7H_2O, 0.2 mg Na_2MoO_4 \_2 H_2O, 0.2 mg CuSO_4 \_5 H_2O, 0.4 mg CoCl_2 \_6 H_2O and 1 mg MnCl_2 \_2 H_2O. The pH of the BES bioreactors was maintained above 6.0 using 2 N NaOH.

**Heterologous enzyme expression and purification**

The molecular constructs for the overexpression of the _gcd_ gene of _P. aeruginosa_ PA14 and _P. putida_ KT2440 in _E. coli_ BL21 (DE3), the _gcd_ gene of _P. aeruginosa_ PA14 in _P. putida_ KT2440 \( \Delta gcd \), the enzyme production and purification protocols as well as the molecular construction of the biosensors are explained in detail in the Data S1.

**Enzyme assay**

Activity of the _gcd_ enzymes for phenazine reduction was assessed with a modified PQQ-dependent reductase assay as described in Mennenga et al. (2009). In the original assay, the electrons released during glucose oxidation are transferred from the PQQ co-factor to the synthetic redox mediator PMS and finally to the non-reversible chromophore electron acceptor DCPIP. In our modified version, the synthetic electron acceptor system (PMS + DCPIP) was replaced by either PCA or PYO and their reduction was followed spectrometrically. The concentration of the protein used for activity measurement was determined using the Bradford method (Bradford, 1976). Then, all components of the enzyme assay were first gassed with nitrogen for about 30 min to ensure anaerobic conditions, as oxygen easily re-oxidizes the phenazines and therefore hampers the measurement. The reaction mixture was prepared in an anaerobic chamber (Coy Lab products, Grass Lake, MI, USA) prior to measurement outside in a CLARIOstar plate reader (BMG Labtech, Orteberg, Germany), which was constantly flushed with nitrogen resulting in a
reduced oxygen environment of 1.2%. The assay mixture contained a final volume of 200 µl (100 mM Tris-HCl, pH 7.1; 1 mM CaCl2, 5 µM PQ, 0.45 mM phenazine (PYO or PCA), 65 µg purified enzyme and 20 mM glucose). All assay components, except glucose, were first mixed and incubated at 30°C for 30 min before glucose was added as a substrate or an equivalent volume of water as a control, to start the reaction. Remaining oxidized PYO was measured at 690 nm, while accumulated reduced PCA was measured at 440 nm. Since PCA is re-oxidized by oxygen very fast even though the enzymatic reduction is slow, the microtiter plate containing the assay mixture with PCA was kept in an anaerobic chamber and only brought out at intervals for plate reader measurements. Extinction coefficients used were ε440 = 2.206 mM⁻¹cm⁻¹ for reduced PCA (Glasser et al., 2017) and ε690 = 4.5 mM⁻¹cm⁻¹ for oxidized PYO (determined in this work, see Fig. S2).

Bioelectrochemical systems set-up and procedures

BES set-up (see Fig. S3) and operation were identical as reported previously (Askotsari et al., 2019). The BES reactor was made of a single-chamber 500-ml (working volume) glass reactor with a water jacket for temperature control (Laborglas Lammek, Moers, Germany). It contained a three-electrode set-up: a working electrode (anode) made of a carbon comb - 7.3 × 4.7 × 0.9 cm (geometrical surface area = 156.32 cm²) attached to a graphite rod, a counter electrode (cathode) - 7.3 × 2.2 × 0.9 cm carbon block (49.22 cm³) attached to a graphite rod and a reference electrode (RE) – Ag/AgCl, saturated with KCl (192 mV vs. SHE at 30°C). All carbon materials were of high-grade graphite (EDM-3, Novotec, Nettetal, Germany). Butyl rubber gaskets were used to seal all ports for the electrodes, gas-in/out and sampling.

The BES reactors were operated with a VMP-300 potentiosstat (BioLogic Science Instruments, Seyssinet-Pariset, France). Experiments were performed at 30°C and stirred at 200 rpm with a magnetic stirrer. Chronoamperometry measurement of the electric current was recorded at a set potential of 0.2 V against RE. Electric current of the blank medium was first recorded for 24 h before inoculation, and thereafter, the recording was interrupted every 22 h for a 2 h cyclic voltammetry measurement (data not shown). Sampling for determination of the optical density at 600 nm, pH and metabolites production was performed at regular intervals. Experiments were performed under two oxygen-limited conditions: initially, active aeration (AA) of the medium with an oxygen flow rate of 30 ml min⁻¹ was supplied to the reactors for the first 31 hours through a sparger connected to a 0.2 µm sterile PTFE filter (DIA-Nielsen, Düren, Germany); thereafter, passive aeration (PA) was applied for the rest of the experiment that allowed air passively into the reactor headspace through a 0.2 µm sterile PTFE filter. Phenazines (80 mg l⁻¹ PCA or 18 mg l⁻¹ PYO) were added to the system 23 h after inoculation.

Analytical procedures

Analysis of central metabolites: Glucose was measured using a YSI 2900D Biochemistry Analyser (YSI Incorporated, Ohio, USA). Analysis of secreted metabolites was performed using a HPLC (Jasco Deutschland GmbH, Pfungstadt, Germany) with an Aminex HPX-87H 300 nm × 7.8 mm, 9 µm ion exclusion column (Bio-Rad, Hercules, CA, USA). Isocratic elution was achieved using 5 mM H₂SO₄ at a flow rate of 0.5 ml min⁻¹. Signals were detected with a UV detector (210 nm) and a refractive index (RI) detector.

Analysis of phenazines: A 1:2 dilution of the supernatant from the BES in acetonitrile was kept at 4°C overnight, to allow for protein precipitation. Thereafter, the samples were centrifuged, and the supernatant was used for measurement. Phenazines were analysed using a reverse-phase HPLC with a 250 × 4.0 mm, 5 µm column (Bischoff Chromatography, Leonberg, Germany). Elution was performed in a gradient of 0.1% trifluoroacetic acid (TFA) (v/v) in water plus 0.1% TFA (v/v) in acetonitrile with a flow rate of 1 ml min⁻¹. PCA and PYO were detected at 360 and 280 nm, respectively, with retention times of 18.1 min (PCA) and 9.0 min (PYO) and compared to standard solution of phenazines (PYO, Sigma Aldrich, Taufkirchen, Germany; PCA, Apollo Scientific, Cheshire, UK).

Calculating charge/energy balance

To determine the coulombic efficiency as the fraction of harvested electrons in relation to the charge (Q) contained in the substrate glucose in the form of reducing equivalents, the following equation was used to calculate the energy balance:

\[ \text{CE} \% = \frac{(Q_{\text{anode}})}{(Q_{\text{glucose utilized}}) - Q_{\text{sugar acid metabolites}})} \times 100. \]

With Q being the charge equivalent [C] accessible, which can be calculated using Faraday’s Law \( Q = n \times z \times F \), where: \( n \) = amount of consumed substrate [mol], \( z \) = number of transferable electrons per molecule of substrate (for glucose = 24), \( F \) = Faraday constant (96485.3 C mol⁻¹), or from the integrated anodic current \( Q_{\text{anode}} = \int I (t) \), where \( I \) = anodic current [A] and \( t \) = time [s]. Anodic charge was calculated using the EC-Lab software (Biologic Science instruments, Seyssinet-Pariset, France), while charge for other terms was determined from the obtained quantitative data.

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Conflict of interest

None declared.

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**Supporting information**

Additional supporting information may be found online in the Supporting Information section at the end of the article.

**Fig. S1.** SDS PAGE showing the purified membrane-bound glucose dehydrogenase (Gcd) of *P. putida* KT2440 and *P. aeruginosa* PA14.

**Fig. S2.** Determination of the extinction coefficient ($\varepsilon_{690}$) in mM$^{-1}$ cm$^{-1}$ of oxidized PYO.

**Fig. S3.** Photography of the BES setup.

**Table S1.** Strains and plasmids used for this study.

**Table S2.** Primers used in this study.

**Data S1.** Heterologous enzyme expression and purification.