The Metabolic Redox Regime of Pseudomonas putida Tunes Its Evolvability toward Novel Xenobiotic Substrates

Akkaya, Özlem; Pérez-Pantoja, Danilo R.; Calles, Belén; Nikel, Pablo I.; de Lorenzo, Victor

Published in:
mBio (Online)

Link to article, DOI:
10.1128/mBio.01512-18

Publication date:
2018

Document Version
Publisher's PDF, also known as Version of record

Link back to DTU Orbit

Citation (APA):
Akkaya, Ö., Pérez-Pantoja, D. R., Calles, B., Nikel, P. I., & de Lorenzo, V. (2018). The Metabolic Redox Regime of Pseudomonas putida Tunes Its Evolvability toward Novel Xenobiotic Substrates. mBio (Online), 9(4), [e01512-18]. https://doi.org/10.1128/mBio.01512-18
The Metabolic Redox Regime of *Pseudomonas putida* Tunes Its Evolvability toward Novel Xenobiotic Substrates

Özlem Akkaya,¹ Danilo R. Pérez-Pantoja,¹ Belén Calles,² Pablo I. Nikel,³ Victor de Lorenzo³

¹Department of Molecular Biology and Genetics, Faculty of Sciences, Gebze Technical University, Kocaeli, Turkey
²Programa Institucional de Fomento a la Investigación, Desarrollo e Innovación, Universidad Tecnológica Metropolitana, Santiago de Chile, Chile
³Systems and Synthetic Biology Program, Centro Nacional de Biotecnología, Madrid, Spain

**ABSTRACT** During evolution of biodegradation pathways for xenobiotic compounds involving Rieske nonheme iron oxygenases, the transition toward novel substrates is frequently associated with faulty reactions. Such events release reactive oxygen species (ROS), which are endowed with high mutagenic potential. In this study, we evaluated how the operation of the background metabolic network by an environmental bacterium may either foster or curtail the still-evolving pathway for 2,4-dinitrotoluene (2,4-DNT) catabolism. To this end, the genetically tractable strain *Pseudomonas putida* EM173 was implanted with the whole genetic complement necessary for the complete biodegradation of 2,4-DNT (recruited from the environmental isolate *Burkholderia* sp. R34). By using reporter technology and direct measurements of ROS formation, we observed that the engineered *P. putida* strain experienced oxidative stress when catabolizing the nitroaromatic substrate. However, the formation of ROS was neither translated into significant activation of the SOS response to DNA damage nor did it result in a mutagenic regime (unlike what has been observed in *Burkholderia* sp. R34, the original host of the pathway). To inspect whether the tolerance of *P. putida* to oxidative challenges could be traced to its characteristic reductive redox regime, we artificially altered the NAD(P)H pool by means of a water-forming, NADH-specific oxidase. Under the resulting low-NAD(P)H status, catabolism of 2,4-DNT triggered a conspicuous mutagenic and genomic diversification scenario. These results indicate that the background biochemical network of environmental bacteria ultimately determines the evolvability of metabolic pathways. Moreover, the data explain the efficacy of some bacteria (e.g., pseudomonads) to host and evolve with new catabolic routes.

**IMPORTANCE** Some environmental bacteria evolve with new capacities for the aero-
bic biodegradation of chemical pollutants by adapting preexisting redox reactions to novel compounds. The process typically starts by cooption of enzymes from an available route to act on the chemical structure of the substrate-to-be. The critical bottleneck is generally the first biochemical step, and most of the selective pressure operates on reshaping the initial reaction. The interim uncoupling of the novel sub-
strate to preexisting Rieske nonheme iron oxygenases usually results in formation of highly mutagenic ROS. In this work, we demonstrate that the background metabolic regime of the bacterium that hosts an evolving catabolic pathway (e.g., biodegrada-
tion of the xenobiotic 2,4-DNT) determines whether the cells either adopt a genetic diversification regime or a robust ROS-tolerant status. Furthermore, our results offer new perspectives to the rational design of efficient whole-cell biocatalysts, which are pursued in contemporary metabolic engineering.

Received 18 July 2018  Accepted 23 July 2018  Published 28 August 2018

Citation Akkaya O, Pérez-Pantoja DR, Calles B, Nikel PI, de Lorenzo V. 2018. The metabolic redox regime of *Pseudomonas putida* tunes its evolvability toward novel xenobiotic substrates. mBio 9:e01512-18. https://doi.org/10.1128/mBio.01512-18

Editor Caroline S. Hanwood, University of Washington

Copyright © 2018 Akkaya et al. This is an open-access article distributed under the terms of the Creative Commons Attribution 4.0 International license.

Address correspondence to Pablo I. Nikel, pabnik@biosustain.dtu.dk, or Víctor de Lorenzo, vdlorenzo@cnb.csic.es.

This article is a direct contribution from a Fellow of the American Academy of Microbiology. Solicited external reviewers: Jim Spain, University of West Florida; Maia Kivisaar, University of Tartu, Estonia.
Environmental bacteria that catabolize xenobiotic pollutants (existing only since the onset of synthetic chemistry) offer a unique opportunity to inspect the rules that govern the evolution of metabolic networks (1, 2). Unlike resistance to antibiotics, which can be caused by mutations modifying the target or by evolving just one protein (3–6), new catabolic phenotypes require multiple changes in the protein complement of the pathway along with other functions in the host that tune the activity of the novel route according to the background biochemical network (7–10). Among the microorganisms known to host aerobic routes for catabolism of typical industrial pollutants, e.g., chloroaromatic (11, 12) and nitroaromatic chemicals (13, 14), *Pseudomonas* species stand out as recurrent hosts of catabolic routes that enable growth on such unusual chemicals (15–20). This state of affairs raises two related questions. The first question is why new pathways evolve preferentially in this bacterial domain and not so much in other species. The second question is how an evolutionary solution to the novel metabolic challenge remains in the same bacterial domain rather than propagation into other prokaryotic hosts (15). The biochemical reactions at stake often involve redox modifications on the substrate, such as mono- or dioxygenations executed by Rieske nonheme iron oxygenases (21–23). Owing to their mechanism of action, when such enzymes act on substrates-to-be that do not fit well in the active enzyme center, reactive oxygen species (ROS) are released (24–27). This phenomenon is due to a suboptimal kinetic scenario in which the substrate leaves the active pocket unscathed before the Fe-activated oxygen molecule can attack the aromatic structure (28, 29). Uncoupled reactions of this sort result in considerable redox stress in the host (30) and—at least in some cases—DNA damage and acquisition of a DNA mutagenic regime (31, 32). The interplay between faulty redox reactions, ROS formation, and DNA damage has been previously characterized in strain *Burkholderia* sp. R34. This environmental bacterium degrades (if with difficulties) 2,4-dinitrotoluene (2,4-DNT), an archetypal xenobiotic compound (33). The first enzyme of the *dnt* pathway is a Rieske nonheme iron oxygenase that evolved from a precursor protein that acts on naphthalene (34). The extant catabolic pathway is still evolving, as its substrate profile and regulation have features of the antecedent route. When *Burkholderia* sp. R34 is exposed to 2,4-DNT, the substrate is indeed degraded, but cells undergo a massive intracellular production of ROS (35) stemming from the first reaction (i.e., dioxygenation of the nitroaromatic compound to yield 4-methyl-5-nitrocatechol). While ROS formation results in killing most of the microbial population, this DNA-damaging and protein-perturbing agent causes the surviving cells to diversify genetically (35). At least part of the ROS-triggered DNA mutagenesis can be traced to misincorporation of 8-hydroxy-2′-deoxyguanosine (8-oxoG) to DNA, although other mechanisms of ROS-mediated inhibition of the DNA mismatch repair system could be at play (36–38). The example of 2,4-DNT degradation illustrates how stress arising from an abortive metabolic reaction can paradoxically promote evolution of novel routes, as genetic diversification fosters exploration of the solution space by the whole bacterial population, plausibly leading to an optimized catabolic outcome that ultimately becomes fixed in the genome. One consequence is that the evolution of aerobic degradation pathways for xenoaromatic compounds can occur only in bacterial hosts able to cope with intracellular ROS generation to a level that allows genetic diversification without surpassing a deadly threshold. The most common mechanism to counter oxidative stress involves the action of detoxifying enzymes (e.g., catalases, peroxidases, and hydroperoxide reductases) that inactivate ROS (39). The corresponding reactions are ultimately fed by metabolic NADPH (40, 41), which provides the reductive currency to counteract the noxious effects of ROS, e.g., via reduced glutathione (42, 43). In this work, we addressed the effect of the background redox metabolism on the evolvability of environmental bacteria hosting a new biodegradation/biotransforma-
tion pathway. We implanted the dnt route for 2,4-DNT catabolism in the genome of the model soil bacterium *Pseudomonas putida* and inspected the effects of metabolizing this compound on intracellular ROS production, redox stress, and genetic variability resulting from DNA mutagenesis. *P. putida* is a frequent host of pathways for aerobic degradation of aromatics, and it is a habitual carrier of both evolving routes and naturally optimized pathways (44–46). These qualities are generally attributed to the distinct core metabolic network of this bacterium, geared to maintain high NADPH levels (47, 48), which is further reinforced through the action of stress-induced pyridine nucleotide transhydrogenases (49). The results below indicate that ROS, resulting from faulty reactions of the dnt pathway on 2,4-DNT, are translated into genetic diversification of the host in a fashion that depends on its redox status—and therefore that evolvability of new traits is ultimately tuned by the background metabolism of the bacterial host. Consequently, some bacteria seem to be more suitable in hosting evolution of new pathways and delivering their activities in sustained form, compared to others in which the background metabolic network cannot cope with the massive formation of ROS.

**RESULTS**

**Construction of a stable 2,4-DNT–degrading *P. putida* strain.** The 2,4-DNT degradation pathway of *Burkholderia* sp. R34 converts the aromatic substrate into propionyl-coenzyme A and pyruvate through the sequential action of six enzymes (DntA to DntE) (Fig. 1a). The genes encoding the entire pathway (i.e., *dntAaAbAcAd dntB dntD dntE dntG*), along with an open reading frame (ORF) encoding a native regulatory protein (i.e., DntR) (50), were cloned by PCR from *Burkholderia* sp. R34, assembled in a synthetic Tn7-based vector (Fig. 1b), and delivered into the chromosome of *P. putida* EM173. This strain is a derivative of wild-type KT2440 devoid of four prophages and the endogenous Tn7 transposase (51) and displays enhanced genetic stability, a trait exploited when manipulating enzymes involved in harsh biochemical reactions (52). After checking the proper insertion of the genes into the target chromosome by PCR, the capability of the resulting strain (termed *P. putida* EM-DNT) to degrade 2,4-DNT was tested. For this, cells were grown overnight at 30°C and then diluted in fresh M9 minimal medium with 0.4% (wt/vol) succinate. When cultures reached an optical density at 600 nm (OD600) of 0.5, 2,4-DNT was added at 0.5 mM. After 3 h of incubation—and similarly to the original 2,4-DNT–degrading *Burkholderia* strain (35)—*P. putida* EM-DNT secreted distinctively colored metabolites (Fig. 1c), which indicated the activity of the dnt route in the surrogate *P. putida* host.

As a quantitative measure of the activities implanted in *P. putida*, both the consumption of 2,4-DNT and the appearance of key metabolic intermediates in the route were assessed by gas chromatography coupled to mass spectrometry (Fig. 1d). After 24 h of incubation, *P. putida* EM-DNT processed 64% of the aromatic substrate, with a specific consumption rate of 0.78 ± 0.04 μmol 2,4-DNT/g cell dry weight (gCDW)/h. The observed rate of 2,4-DNT consumption was comparable to that previously shown in cultures of *Burkholderia* sp. R34, the original host of the pathway, in which 70% of the substrate was consumed after 24 h of incubation (8, 35). As expected, no substrate consumption was detected in cultures of the parental strain EM173. Both 4-methyl-5-nitrocatechol and 2-hydroxy-5-methylquinone, the metabolic products of DntA and DntB, respectively (53, 54), were detected in supernatants of *P. putida* EM-DNT cultures with added 2,4-DNT but not in control experiments. The specific formation rates of 4-methyl-5-nitrocatechol and 2-hydroxy-5-methylquinone in these cultures were 0.24 ± 0.03 and 0.19 ± 0.05 μmol/gCDW/h, respectively. The appearance of the colored species in culture supernatants indicated that at least the first two steps of the biotransformation of 2,4-DNT into the corresponding metabolic intermediates are catalytically active in the surrogate host of the degradation pathway. Taken together, these results show that the 2,4-DNT degradation route grafted into *P. putida* (and, in particular, the ROS-generating DntA Rieske nonheme iron oxygenase that processes 2,4-DNT) was active under the culture conditions tested.
2,4-DNT degradation by *P. putida* EM-DNT results in ROS generation and activation of the cellular response to oxidative stress. Despite having the necessary genetic and biochemical complement, *Burkholderia* sp. R34 grows poorly on 2,4-DNT as the sole carbon source. This difficulty that can be traced to the formation of ROS upon exposure of the cells to the substrate of the biodegradation pathway (35). Against this background, we evaluated the generation of ROS in *P. putida* EM-DNT when exposed to 2,4-DNT at either 0.25 or 0.5 mM by assessing the fluorescence brought about by the ROS-sensitive dye 2',7'-dichlorodihydrofluorescein diacetate (H2DCF-DA) in single cells by using flow cytometry (Fig. 2a). The amount of ROS proportionally increased with respect to the concentration of 2,4-DNT, indicating that substrate consumption generates oxidative stress. The proportionality between the activity of the degradation pathway and ROS generation was qualitatively assessed in these experiments, since the...
amount of 2-hydroxy-5-methylquinone produced by the cells increased with the concentration of 2,4-DNT in the same fashion as ROS formation did. Quantification of ROS indicated that the mere exposure of the cells to 2,4-DNT is not the main cause of ROS generation; instead, the biochemical transformation of the substrate leads to endogenous oxidative stress. Indeed, *P. putida* EM·DNThad a 10-fold increase in ROS formation upon exposure to 2,4-DNT (Fig. 2b). In contrast, strain EM173 had a lower accumulation of ROS (less than 4-fold increase) than strain EM·DNT when challenged with 2,4-DNT, even at the highest substrate concentration tested. While biotransformation intermediates 4-methyl-5-nitrocatechol and 2-hydroxy-5-methylquinone (55) could be mutagenic, we previously demonstrated that increasing turnover of metabolites downstream of 4-methyl-5-nitrocatechol by controlled overexpression of *dntB* in *Burkholderia* sp. R34 does not result in a decrease of ROS formation (35). This indicates that the activity of the first enzyme of the pathway (and not the metabolic intermediates therein) is responsible for the bulk of the oxidative stress observed upon addition of 2,4-DNT.

Yet, is the observed ROS formation connected to the activation of stress responses in the engineered *P. putida* strain? To answer this question, the transcriptional activities of genes involved in the oxidative stress response (i.e., *ahpC* [PP_2439, alkyl hydroperoxide reductase] and *katA* [PP_0481, a catalase]) were studied by fusing the corresponding promoter region of these two genes to the reporter monomeric superfolder green fluorescent protein (msf·GFP) (Fig. 2c). The ROS reporter plasmids or pSEVA237M, the promoterless, msf·GFP-containing vector (56), were individually transformed into *P. putida* EM·DNT, and the msf·GFP signal was evaluated in cultures exposed to 2,4-DNT by using flow cytometry. The output signal qualitatively followed the same trend as observed for ROS accumulation (Fig. 2b): the induction of the two oxidative stress-
responsive promoters significantly increased in the presence of the aromatic compound (i.e., 10- and 42-fold increase in the transcriptional activities of \( P_{ahpC} \) and \( P_{katA} \), respectively, when cells were exposed to 2,4-DNT at 0.25 mM). In accordance with previous observations in \( Burkholderia \) sp. R34 (35), these results indicated that cells carrying the enzymes that are needed to process 2,4-DNT undergo oxidative stress conditions upon exposure to the substrate of the degradation pathway.

**Assessment of effects of 2,4-DNT degradation on the SOS response and recombinogenic activity.** Since the hypothesis underlying this work is that endogenous oxidative stress brought about by the biotransformation of an aromatic substrate could result in genetic novelty, we explored two types of mutagenic effects on the genome of \( P. \) putida. In the first place, and in order to evaluate if the degradation of 2,4-DNT promotes recA-mediated homologous DNA recombination, a reporter strain was designed as follows. An internal region of the \( pyrF \) gene of \( P. \) putida, which results in uracil auxotrophy, was used to disrupt the \( pyrF \) gene of \( P. \) putida, which results in uracil auxotrophy. The frequency of excision of the plasmid from the chromosome of \( P. \) putida EM-DNT-U (i.e., reversion to prototrophy) was adopted as an indication of DNA recombination. (b) DNA recombination frequency upon exposure to 2,4-DNT or norfloroxacin (NOR). Bars represent mean values ± SD \((n = 4)\), and the asterisks identify significant differences at a \( P \) level of <0.05, determined using Student’s \( t \) test. (c) General mechanism of SOS response upon DNA damage. The RecA protein, stimulated by either damaged or single-stranded DNA, triggers the inactivation of LexA (a repressor of the SOS response genes), thereby inducing the response. The LexA degradation-dependent activation of the recA promoter of \( P. \) putida was used as a proxy of the SOS response by constructing a msf-GFP-based biosensor. Elements in the outline are not drawn to scale. (d) Testing of the SOS response biosensor in soft agar experiments. \( P. \) putida EM-DNT was transformed with the reporter plasmid, and two filter paper disks, soaked with either NOR or 2,4-DNT, were applied onto the bacterial lawn. The plates were photographed under blue light after 24 h of incubation. (e) Quantification of the SOS response biosensor activity in cultures of \( P. \) putida EM-DNT in the presence of the additives indicated. Bars represent the mean values of the fold change in msf-GFP fluorescence ± SD \((n = 4)\), and the asterisks identify significant differences at a \( P \) level of <0.05 as determined using Student’s \( t \) test. The baseline in cultures with no additives (Nil) is indicated with a dashed gray line.
0.5 mM, and cells were plated onto M9 minimal medium with or without Ura. The frequency of appearance of *P. putida* prototroph (Ura<sup>−</sup>) clones was quantified (Fig. 3b), and the strain degrading 2,4-DNT showed a 4-fold increase in the frequency of recombination compared to control conditions. Norfloxacin (NOR) was added in cultures used as positive controls. NOR is a fluoroquinolone that interferes with the maintenance of chromosomal topology by targeting DNA gyrase and topoisomerase IV, trapping these enzymes at the DNA cleavage stage and thereby preventing strand rejoining (59). Introduction of double-stranded DNA breaks following topoisomerase inhibition by NOR thus induces the SOS response (38, 60). In the presence of NOR, we detected a 7-fold increase in DNA recombination using our reporter system (Fig. 3b).

Since degradation of 2,4-DNT brings about an increase in the frequency of DNA recombination, we wondered whether this would result in the activation of the SOS response (i.e., DNA damage response), which is stimulated by single- or double-stranded breaks in genomic DNA (61). When DNA is injured, the RecA protein binds to DNA in the damaged region to form a filament. This filament interacts with a dimer of the LexA transcriptional repressor, activating its self-cleavage and causing the dissociation of LexA from its targets and inducing the SOS regulon (Fig. 3c). The promoter of recA is one of the targets of LexA (62), an occurrence that was exploited in this work by constructing a biosensor of RecA activity. The promoter region of recA (PP_1629), including the putative LexA binding site, was cloned in front of a promoterless GFP::luxCDABE dual reporter system (63). The resulting reporter plasmid was introduced into *P. putida* EM·DNT, and the system was tested by exposing the cells to NOR and 2,4-DNT in a qualitative soft agar diffusion test (Fig. 3d). Exposure of strain EM·DNT carrying the RecA reporter to NOR resulted in a distinct halo of growth inhibition, the boundaries of which gave off a strong GFP signal when observed under UV light. 2,4-DNT, in contrast, did not seem to elicit a similar response in the cells carrying the RecA reporter. A similar pattern was observed in GFP-dependent fluorescence when the experiment was repeated in liquid cultures of *P. putida* EM·DNT bearing the SOS response biosensor, as quantified by flow cytometry (Fig. 3d). Addition to NOR at subinhibitory concentrations (250 ng ml<sup>−1</sup>) caused a 4-fold induction of the RecA reporter, whereas neither 2,4-DNT nor H<sub>2</sub>O<sub>2</sub> resulted in any significant activation of the SOS response. Taken together, these results indicate that degradation of the aromatic substrate promotes DNA recombination extent, without significantly affecting the activity of the SOS response. The next question was whether other forms of mutagenesis, such as point mutations in genomic DNA, could be also correlated to 2,4-DNT biotransformation by the engineered *P. putida* strain.

**Biotransformation of 2,4-DNT in *P. putida* barely affects DNA mutagenesis.** Although ROS production stemming from 2,4-DNT metabolism in *P. putida* EM·DNT does not trigger a significant SOS response and only stimulates recombination to a moderate degree, it was still possible that oxidative damage to DNA, generation of 8-oxoG, and general stress-induced relaxation of mismatch repair could increase the overall level of DNA mutagenesis. To inspect this possibility (which was observed in *Burkholderia* sp. R34) the mutagenesis rate was assessed by exposing *P. putida* EM·DNT cells to 2,4-DNT and counting the number of rifampin-resistant (Rifr) colonies after plating on a solid culture medium. The antibacterial effects of Rif are based on its ability to bind the β-subunit of the RNA polymerase, thereby blocking the elongation of the nascent RNA molecule. Rif<sup>+</sup> clones usually harbor RpoB mutations in amino acid residues that make contact with Rif (64), rendering RNA polymerase insensitive to the antibiotic (65, 66). Under our experimental conditions, the background rate (M) of Rif<sup>+</sup> clones (i.e., *P. putida* EM·DNT cells incubated in the presence of dimethyl sulfoxide [DMSO]), was (9.1 ± 0.7) × 10<sup>−9</sup>. Exposure of *P. putida* EM·DNT cells to 2,4-DNT increased this level of mutagenesis by a mere 28% (and the difference from control experiments was not statistically significant). This was an unexpected occurrence, as high levels of endogenous ROS are translated onto a mutagenic state in other bacteria. That this was not the case in the strain constructed in this study indicated that the molecular mechanisms that connect ROS stress with DNA mutagenesis in *P. putida* are
plausibly different from those operating in other bacterial species that other factors could be at play. Yet, what could such key factors be? One metabolic signature that characterizes \textit{P. putida} is its highly reductive redox metabolism. We thus wondered next whether the interplay between ROS production and DNA mutagenesis could be shielded by such a background metabolism that is so characteristic of this species. 

Altering the redox status of \textit{P. putida} increases the mutagenic effect of the 2,4-DNT degradation pathway. \textit{P. putida} maintains an adequate supply of reducing power through the activity of the EDEMP cycle (47), a core metabolic architecture that combines individual biochemical steps from the Entner-Doudoroff, the Embden-Meyerhof-Parnas, and the pentose phosphate pathways (Fig. 4a). By recycling part of the pool of triose-phosphate back to hexose phosphates, this metabolic cycle mediates NAD(P)H formation and enables catabolic overproduction of reduced pyridine nucleotides. This evolutionary-driven metabolic occurrence helps explain the very high resistance to environmental insults (e.g., oxidative stress) displayed by \textit{P. putida} (67–69).

Moreover, antioxidant responses in bacteria exposed to xenobiotics rely on the generation of the reducing power that the cells use to counteract ROS formation. For instance, AhpC, a hydroperoxide-detoxifying enzyme (Fig. 2c), is reduced by peroxiredoxin reductase (AhpF), a process that requires NADH (70). On this basis, and taking into consideration that (i) 2,4-DNT degradation results in the generation of ROS and (ii) the oxidative damage caused by ROS stimulates DNA damage, we set to explore the relationship between redox status and DNA mutagenesis in \textit{P. putida} EM-DNT. To investigate this issue, the redox status of \textit{P. putida} EM-DNT was artificially perturbed by altering the intracellular availability of reduced nicotinamide cofactors through conditional expression of the \textit{nox} gene of \textit{Streptococcus pneumoniae} (71, 72). Nox is a NADH-specific oxidase enzyme from that converts O$_2$ to H$_2$O with negligible formation of H$_2$O$_2$ (Fig. 4b). Cellular energy demands are not affected by Nox, and thus \textit{nox} expression allows for the specific investigation of the impact of NADH oxidation without affecting the overall fitness of the cells (73). A synthetic metabolic module was designed for this purpose, in which the gene encoding NADH oxidase from \textit{S. pneu-
moniae was placed under control of the orthogonal ChnR/Pchne expression system (Fig. 4c), which is inducible by cyclohexanone (74, 75). We tested the effect of such an NADH-burning device on the overall physiology of P. putida EM-DNT by measuring the specific Nox activity in cell-free extracts and evaluating growth rates on succinate cultures (Fig. 4d). Induction of the synthetic device by addition of 0.1 mM cyclohexanone resulted in a specific Nox activity of 3.9 ± 0.6 μmol/mg of protein/min, ca. 40-fold higher than the background oxidase activity in the cell extract of the control strain transformed with the empty vector. Under these conditions, the specific growth rate of P. putida EM-DNT was reduced by ca. 30%. As a direct indication of the role of Nox in mediating a redox imbalance, the reduced nucleotide content was evaluated in cell-free extracts of P. putida EM-DNT. The NADH concentration was 176 ± 25 and 103 ± 38 μM for the strain carrying the empty pSEVA2311 vector or plasmid pS2311·Nox, respectively, demonstrating that the overexpression of nox in P. putida cells decreases the reducing power content. The next step was to evaluate if an alteration in the NADH content affects DNA mutagenesis.

P. putida EM-DNT was transformed with the nox-expressing plasmid or the empty pSEVA2311 vector, and the resulting strains were exposed to different combinations of 2,4-DNT and cyclohexanone. The rate of mutagenesis, M, was explored by assessing the appearance of Rifr clones after each treatment (Fig. 5a). As shown in the figure, the exposure of the cells to 2,4-DNT or activation of Nox individually did not result in a significant increase in DNA mutagenesis. The combination of the two treatments, however, produced a 6-fold increase in M, indicating that the redox imbalance introduced by Nox exacerbated the mutagenic impact of 2,4-DNT. Yet, what is the nature of the mutations that lead to Rifr in such a redox-stressed scenario?

The spectrum of rpoB mutations depends on the intracellular redox status. Mutations in rpoB that confer Rifr are well conserved among prokaryotes, and the amino acid changes that produce this phenotype can be grouped in three clusters in a central region of RpoB (65). We sequenced ca. 100 clones per experimental condition to obtain the spectrum of mutations in each case (Table 1). All the mutations detected in this analysis were located in cluster I of RpoB (between nucleotides 1,528 and 1,626...
of rpoB, as defined by Jatsenko et al. [65]), and they could be grouped into eight categories. Five of the categories were transitions (i.e., A → G and C → T), two of them were transversions (i.e., A → T and C → A), and the rest of the rpoB mutations found in Rif clones were grouped as “other” (a category that included the less abundant rpoB mutations). The two predominant mutations found resulted in Q518L and D521G changes in RpoB, and their frequency of appearance did not suffer any significant change across the experimental treatments tested here. Interestingly, the frequency of C → A transversion that generates an L538I change in RpoB was influenced by both exposure to 2,4-DNT (2-fold increase) and the activation of Nox (4-fold increase). The effect of the two treatments was additive, further linking biotransformation of the substrate and redox status of the cells with DNA mutagenesis. C → A transversions are known to be caused by the presence of 8-oxoG in the DNA (76), which results from the attack of ROS on purine residues. This result thus indicated that oxidative stress caused by 2,4-DNT degradation results in oxidation of purines (as observed in Burkholderia sp. R34), but it also shows that the more reductive redox status of P. putida provides effective protection against this mutagenic effect. Other mutations arising by exposing the cells to 2,4-DNT had a less clear origin, e.g., the activity of the widely distributed Y-family of DNA polymerases (77). Members of this family are characterized not only by their ability to replicate damaged DNA but also by their lack of processivity and low fidelity when copying undamaged template. In P. putida KT2440, this protein category is represented by DinB (PP_1203) and the error-prone DNA polymerase DnaEB (PP_3119) (78). In sum, the spectrum of rpoB mutations detected in our study suggests that ROS-induced mutagenesis triggered by 2,4-DNT metabolism merges direct damage to DNA caused by misincorporation of 8-oxoG with other stress response mechanisms that relax fidelity of DNA replication and/or prevent repair.

**DISCUSSION**

Evolvability of bacteria for conquering novel nutritional and physicochemical landscapes is the result of the interplay between endogenous and exogenous factors. Propagation of new pathways through a bacterial population is fostered by different mechanisms of horizontal gene transfer, e.g., conjugation, transformation, and phage-mediated or ICE (integrative and conjugative elements)-mediated DNA acquisition (79, 80). In this case, the emergence of genetic and functional diversity largely relies on the balance between the effectiveness of the transfer events and the ability of the recipient bacteria to gain an advantage thereof. Not surprisingly, microbial species with a high capacity to receive foreign DNA usually evolve new traits (81)—sometimes as undesirable as antibiotic resistances (6). However, horizontal gene transfer does not account
for the onset of fresh routes for xenobiotic catabolism, which require mutation of preexisting pathways to reshape their substrate range and rewire and connect their biochemical outputs to the existing metabolic network in the host. Since these changes must necessarily occur in single bacteria, one can safely predict that the more highly mutagenic regime they display, the faster the solution space will be explored—and an optimum will be eventually found. Two common triggers of such mutagenic regimes include carbon starvation and endogenous production of ROS (82–85). Given that these mechanisms are virtually universal through all aerobic bacteria, the question arises why most aerobic routes for degradation of xenobiotic aromatics involve the action of Rieske nonheme iron oxygenases borne by pseudomonads and closely related bacteria. The results described in this article provide some rationale for this longstanding observation. The mechanism of this type of oxygenases at their reaction center involves activation of an oxygen molecule by an Fe-S cluster to make the oxygen reactive toward the aromatic substrate (86, 87). Even with a good enzyme-substrate fit, purely stochastic effects at the atomic level may result in coupling failure and release of ROS, the discharge of which will increase if the fitting of the substrate to the enzyme is worse (24–27). We have argued that, because of its mutagenic potential, the release of ROS is a major driver of the evolution of the cognate bacterial population toward a new biodegradative target (35). Yet, why are such biodegradative abilities so conspicuous in pseudomonads? The data above suggest that, while ROS production during 2,4-DNT degradation is unavoidable, DNA mutagenesis is ultimately controlled by the endogenous redox status of the corresponding cells. The highly reductive NAD(P)H regime associated with the distinct central carbon metabolism of \textit{P. putida} thus seems to render this species well suited for hosting pathways involving strong redox transactions (Fig. 5b and 6a). At the same time, a dearth in NAD(P)H availability would result in a high DNA mutagenesis regime. Although data supporting this notion have been generated in only two environmental bacteria (\textit{Pseudomonas putida} and \textit{Burkholderia} spp.), it is likely that they are specific cases of a general evolutionary principle that is reminiscent of the concept of antifragility (35, 88) or hormesis (89, 90), i.e., beneficial effects of a low dose of a toxic input (91, 92), observed in other biological systems. Interestingly, hormetic dose-response mechanisms have been put forward to explain the survival of bacteria exposed to different types of antibiotics (93–95). Under this framework, there seems to be an optimum in the diversity-generating regime caused by stress (e.g., ROS) that is enough to generate sufficient evolutionary bet-hedging for coping with a new nutrient or physicochemical condition while keeping the thereby diversified population—but also damaged population—within viable levels (Fig. 6b). It is within this solution space that bacterial cells equipped with novel catabolic pathways for new substrates are to be found. The data above suggest that such a window of productive innovation is shaped by the background metabolic network, specifically by the redox regime. ROS formation has been observed in other environmental bacteria that employ biodegradation pathways for xenobiotics (96–99), and it is plausible that similar evolutionary processes apply in other instances.

That the redox metabolic status (either inherent or transient) controls evolvability of selectable traits in bacteria raises some corollaries worth follow-up. For instance, ROS may accelerate the emergence of new antibiotic resistances in species subject to endogenous oxidative stress but endowed with a weaker redox metabolism. This opens the possibility of targeting redox biochemistry of virulent strains with drugs that perturb the NAD(P)⁺/NAD(P)H balance. Moreover, our data suggest that some members of a given microbial community could be more innovative than others on the basis of their innate redox metabolism. There could be innovation specialists that nurture genetic diversification and pass the results to less mutable members of a multispecies bacterial consortium through horizontal gene transfer. Finally, ROS-mediated genetic innovation could be repurposed for the sake of generating new pathways through directed evolution or for the optimization of biocatalytic routes for metabolic engineering (100). In any case, the connection between metabolism and evolvability should
not be ignored when addressing the emergence of new traits whether for fundamental understanding of evolutionary processes or for designing new whole-cell biocatalysts.

MATERIALS AND METHODS

Bacterial strains and culture conditions. Bacterial strains are listed in Table S1 in the supplemental material. *Burkholderia* sp. R34 has been previously described as a 2,4-DNT-degrading specimen (33). *P. putida* EM173 was used as the host of the dnt pathway (51). *E. coli* strains DH5α, CC118, CC118λpir, and HB101 were employed for DNA cloning procedures as indicated in Text S1 in the supplemental material. Bacteria were grown in rich LB medium or in M9 minimal medium with sodium succinate added at 0.4% (wt/vol) with rotary agitation at 170 rpm (101). *P. putida* was grown at 30°C, while *E. coli* was grown at 37°C. When required, 2,4-DNT was added at 0.25 or 0.5 mM from a 0.5 M stock solution freshly prepared in DMSO. Control experiments included an equivalent volume of DMSO. Overnight-grown *P. putida* in M9 minimal medium and succinate was used as inoculum by diluting the bacterial suspension to a starting OD<sub>600</sub> of 0.05. Cells were grown in the same culture medium until the OD<sub>600</sub> reached 0.5, at which point the cell suspension was split: one culture served as a control, and the other one was challenged with 2,4-DNT. Other additives were as indicated, and samples were periodically taken as specified. Additives on the mutation rate

Determination of mutation frequencies and detection of rpoB mutations. *P. putida* cultures were grown overnight with the appropriate antibiotics, diluted 1:10,000 into 50 ml of fresh culture medium (initial OD<sub>600</sub> of 0.05) in 250-ml Erlenmeyer flasks, and incubated for 4 h. These cultures were diluted 1:4 into fresh medium (1 ml in 10-ml test tubes) and incubated for 16 h in the presence of relevant additives. Appropriate dilutions were spread on LB medium plates containing 50 μg ml<sup>-1</sup> Rif. Colonies were counted after 48 h at 30°C. Luria-DelBrück fluctuation analysis was performed to study the impact of additives on the mutation rate. *M. M* values were computed using the Ma-Sandri-Sarkar maximum likelihood estimator (ratio between the number of Rif<sup>R</sup> colonies and the total viable count, corrected by plating efficiency) from at least six independent experiments per condition (104, 105). The statistical significance of mean *M* values across experiments was evaluated with Student’s *t* test. To identify the nature and frequency of point mutations in rpoB, ca. 100 Rif<sup>R</sup> colonies were randomly collected from at least four independent experiments, and the locus was sequenced with primers indicated in the supplemental material. The BioEdit software (BitSize Bio) was used for sequence comparison.

Analytic procedures. The ROS-sensitive green fluorescent dye H<sub>2</sub>DCF-DA (Sigma-Aldrich) was used to quantitate ROS formation. Flow cytometry, preparation of cell-free extracts, metabolite extraction, and
enzymatic assays followed protocols from our laboratory (106–109), and further details can be found in Text S1 in the supplemental material.

**Statistical analysis.** All reported experiments were independently repeated at least three times (as indicated in the figure legends), and mean values of the corresponding parameter and standard deviation (SD) are presented. The significance of differences when comparing results was evaluated via use of Student’s t test.

**SUPPLEMENTAL MATERIAL**

Supplemental material for this article may be found at https://doi.org/10.1128/mBio.01512-18.

**TEXT S1**, PDF file, 0.2 MB.

**TABLE S1**, PDF file, 0.2 MB.

**ACKNOWLEDGMENTS**

V.D.L., P.I.N., and D.R.P.P. designed the project and the experimental layout. Ö.A. performed experiments related to strain construction, oxidative stress, substrate degradation, and DNA mutagenesis. B.C. constructed strains and plasmids used in the study. P.I.N. and D.R.P.P. analyzed the data. P.I.N. and V.D.L. wrote the manuscript with further contributions from all authors to data analysis and interpretation of the results. We thank J. Blázquez (Spain), M. H. H. Narholm (Denmark), A. Smania (Argentina), and L. Elts (Canada) for inspiring discussions. Ö.A. was supported by TUBITAK-BIDEP through the International Postdoctoral Research Scholarship Programme 2219.

We declare that there are no competing interests associated with the contents of our article.

This study was supported by HELIOS Project of the Spanish Ministry of Economy and Competitiveness BIO2015-66960-C3-2-R (MINECO/FEDER), and by ARISYS (ERC-2012-ADG-322797), EmPowerPutida (EUH2020-BIOTEC-2014-2015-6335536), and MADONNA (H2020-FET-OPEN-RIA-2017-1-766975) contracts of the European Union to V.D.L. The study was also supported by the Novo Nordisk Foundation (grant NNF10CC1016517) and the Danish Council for Independent Research (SWEET, DFF Research Project 8021-00039B) to P.I.N.

**REFERENCES**

1. Copley SD. 2009. Evolution of efficient pathways for degradation of anthropogenic chemicals. Nat Chem Biol 5:559–566. https://doi.org/10.1038/nchembio.197.
2. Wackett LP. 2004. Evolution of enzymes for the metabolism of new chemical inputs into the environment. J Biol Chem 279:41259–41262. https://doi.org/10.1074/jbc.R400014200.
3. Furusawa C, Horinouchi T, Maeda T. 2018. Toward prediction and control of antibiotic-resistance evolution. Curr Opin Biotechnol 54:45–49. https://doi.org/10.1016/j.copbio.2018.01.026.
4. Durão P, Balbontín R, Gordo I. 2018. Evolutionary mechanisms shaping the maintenance of antibiotic resistance. Trends Microbiol 26:677–691. https://doi.org/10.1016/j.tim.2018.01.005.
5. Palmer AC, Kishony R. 2013. Understanding, predicting and manipulating the genotypic evolution of antibiotic resistance. Nat Rev Genet 14:243–248. https://doi.org/10.1038/nrg3351.
6. Baquero F, Martínez JL. 2017. Interventions on metabolism: making antibiotic-susceptible bacteria. mBio 8:e01950-17. https://doi.org/10.1128/mBio.01950-17.
7. Seffernick JL, Wackett LP. 2016. Ancient evolution and recent evolution converge for the biodegradation of cyanuric acid and related triazines. Appl Environ Microbiol 82:1638–1645. https://doi.org/10.1128/AEM.03594-15.
8. Nishino SF, Paoli GC, Spain JC. 2000. Aerobic degradation of dinitrotoluene and pathway for bacterial degradation of 2,6-dinitrotoluene. Appl Environ Microbiol 66:2139–2147. https://doi.org/10.1128/aem.66.7.2139-2147.2000.
9. Spain JC. 1995. Biodegradation of nitroaromatic compounds. Annu Rev Microbiol 49:523–555. https://doi.org/10.1146/annurev.mi.49.100.195.002515.
10. Kurt Z, Minoia M, Spain JC. 2018. Resveratrol as a growth substrate for bacteria from the rhizosphere. Appl Environ Microbiol 84:e00104-18. https://doi.org/10.1128/AEM.00104-18.
11. Nikel PI, Pérez-Pantoja D, de Lorenzo V. 2013. Why are chlorinated pollutants so difficult to degrade aerobically? Redox stress limits 1,3-dichloroprop-1-ene metabolism by *Pseudomonas putida*. Philos Trans R Soc Lond B Biol Sci 368:20120377. https://doi.org/10.1098/rstb.2012.0377.
12. van der Meer JR. 1997. Evolution of novel metabolic pathways for the degradation of chloroaromatic compounds. Antonie Van Leeuwenhoek 71:159–178. https://doi.org/10.1023/A:1000164009336.
13. Kivisaa M. 2011. Evolution of catabolic pathways and their regulatory systems in synthetic nitroaromatic compounds degrading bacteria. Mol Microbiol 82:265–268. https://doi.org/10.1111/j.1365-2958.2011.07824.x.
14. Johnson GR, Spain JC. 2003. Evolution of catabolic pathways for synthetic compounds: bacterial pathways for degradation of 2,4-dinitrotoluene and nitrobenzene. Appl Microbiol Biotechnol 62:110–123. https://doi.org/10.1007/s00253-003-1341-4.
15. Pérez-Pantoja D, Donoso R, Junca H, González B, Pieper DH. 2017. Phylogenomics of aerobic bacterial degradation of aromatics, p 1–48. In Rojo F (ed), Aerobic utilization of hydrocarbons, oils and lipids. Springer, Cambridge, England.
16. Pérez-Pantoja D, Donoso R, Agulló L, Córdova M, Seeger M, Pieper DH, González B. 2012. Genomic analysis of the potential for aromatic compounds biodegradation in Burkholderiales. Environ Microbiol 14:1091–1117. https://doi.org/10.1111/j.1462-2980.2011.02613.x.
17. Nikel PI, Martínez-García E, de Lorenzo V. 2014. Biotechnological domestication of pseudomonads using synthetic biology. Nat Rev Microbiol 12:368–379. https://doi.org/10.1038/nrmicro3253.
18. Nikel PI, Silva-Rocha R, Benedetti I, de Lorenzo V. 2014. The private life of environmental bacteria: pollutant biodegradation at the single cell
Akkaya et al.

35. 19. Beldá E, van Heek RGA, José Lopez-Sanchez MJ, Cruveiller S, Barbe V, Fraser C, Klenk HP, Petersen J, Morgat A, Nikel PI, Vallenten D, Rouy Z, Sekowska A, Martins dos Santos VAP, de Lorenzo V, Danchin A, Méquige C. 2016. The revisited genome of *Pseudomonas putida* KT2440 enlightens its value as a robust metabolic chassis. Environ Microbiol 18:3403–3424. https://doi.org/10.1111/1462-2920.12320.

24. 22. Dvorák P, Nikel PI, Damborský J, de Lorenzo V. 2017. Bioremediation 3.0: engineering pollutant-removing bacteria in the times of systemic biology. Biotechnol Adv 35:845–866. https://doi.org/10.1016/j.biotechadv.2017.08.001.

27. 21. Patrauchan MA, Florizone C, Eapen S, Gómez-Gil L, Sethuraman B, Dvorák P, Nikel PI, Damborský J, de Lorenzo V. 2017. Characterization of biphenyl dioxygenase of *Pandoraea pnomenusa* bacteria: mutational processes in pseudomonads. FEMS Microbiol Lett 241:151–156. https://doi.org/10.1016/j.femsle.2004.10.013.

37. 34. Parales RE, Resnick SM. 2004. Aromatic hydrocarbon dioxygenases, p 39–70. In Chen J, Gottesman S (ed), Advances in applied biotechnology. InTech, Rijeka, Croatia.

38. 55. Bolten JL, Trush MA, Penning TM, Dryhurst G, Monks TJ. 2000. Role of *E. coli* RpoS-mediated repression of *Rhodococcus jostii* RHA1. J Bacteriol 190:37–47. https://doi.org/10.1128/JB.190.1.0037-047.2000.

39. 40. Bolton JL, Trush MA, Penning TM, Dryhurst G, Monks TJ. 2000. Role of *E. coli* RpoS-mediated repression of *Rhodococcus jostii* RHA1. J Bacteriol 190:37–47. https://doi.org/10.1128/JB.190.1.0037-047.2000.

41. 42. Kim J, Park W. 2014. Oxidative stress response in *Pseudomonas putida* sp. DNT with the substrate-blind regulator DntR via *via* RpoS-mediated repression of *Rhodococcus jostii* RHA1. J Bacteriol 190:37–47. https://doi.org/10.1128/JB.190.1.0037-047.2000.

43. 44. Kim J, Park W. 2014. Oxidative stress response in *Pseudomonas putida* sp. DNT with the substrate-blind regulator DntR via *via* RpoS-mediated repression of *Rhodococcus jostii* RHA1. J Bacteriol 190:37–47. https://doi.org/10.1128/JB.190.1.0037-047.2000.

45. 46. Kim J, Park W. 2014. Oxidative stress response in *Pseudomonas putida* sp. DNT with the substrate-blind regulator DntR via *via* RpoS-mediated repression of *Rhodococcus jostii* RHA1. J Bacteriol 190:37–47. https://doi.org/10.1128/JB.190.1.0037-047.2000.

47. 48. Kim J, Park W. 2014. Oxidative stress response in *Pseudomonas putida* sp. DNT with the substrate-blind regulator DntR via *via* RpoS-mediated repression of *Rhodococcus jostii* RHA1. J Bacteriol 190:37–47. https://doi.org/10.1128/JB.190.1.0037-047.2000.

49. 50. Kim J, Park W. 2014. Oxidative stress response in *Pseudomonas putida* sp. DNT with the substrate-blind regulator DntR via *via* RpoS-mediated repression of *Rhodococcus jostii* RHA1. J Bacteriol 190:37–47. https://doi.org/10.1128/JB.190.1.0037-047.2000.

51. 52. Kim J, Park W. 2014. Oxidative stress response in *Pseudomonas putida* sp. DNT with the substrate-blind regulator DntR via *via* RpoS-mediated repression of *Rhodococcus jostii* RHA1. J Bacteriol 190:37–47. https://doi.org/10.1128/JB.190.1.0037-047.2000.

53. 54. Kim J, Park W. 2014. Oxidative stress response in *Pseudomonas putida* sp. DNT with the substrate-blind regulator DntR via *via* RpoS-mediated repression of *Rhodococcus jostii* RHA1. J Bacteriol 190:37–47. https://doi.org/10.1128/JB.190.1.0037-047.2000.

55. 56. Kim J, Park W. 2014. Oxidative stress response in *Pseudomonas putida* sp. DNT with the substrate-blind regulator DntR via *via* RpoS-mediated repression of *Rhodococcus jostii* RHA1. J Bacteriol 190:37–47. https://doi.org/10.1128/JB.190.1.0037-047.2000.
NADPH Quenches Endogenous Stress

58. Galvão TC, de Lorenzo V. 2005. Adaptation of the yeast URA3 selection system to Gram-negative bacteria and generation of a ΔbetCDE Pseudomonas putida strain. Appl Environ Microbiol 71:883–892. https://doi.org/10.1128/AEM.71.2.883-892.2005.

59. Martínez-García E, Nikel PI, Aparicio T, de Lorenzo V. 2014. Pseudomonas 2.0: genetic upgrading of P. putida KT2440 as an enhanced host for heterologous gene expression. Microb Cell Fact 13:159. https://doi.org/10.1186/s12934-014-0159-3.

60. Drlica K, Zhao X. 1997. DNA gyrase, topoisomerase IV, and the 4-quinolones. Microbiol Mol Biol Rev 61:377–392.

61. Harms A, Maissonneuve E, Gerdes K. 2016. Mechanisms of bacterial persistence during stress and antibiotic exposure. Science 354:aa4268. https://doi.org/10.1126/science.aa4268.

62. Baharoglu Z, Mazel D. 2014. SOS, the formidable strategy of bacteria against infections. FEMS Microbiol Rev 38:1126–1145. https://doi.org/10.1111/1574-6967.12077.

63. Abella M, Campoy S, Erill I, Rojo F, Barbé J. 2007. Cohabitation of two different lexA regulons in Pseudomonas putida. J Bacteriol 189: 8855–8862. https://doi.org/10.1128/JB.01213-07.

64. Benedetti IM, de Lorenzo V, Silva-Rocha R. 2012. Quantitative, non-disruptive monitoring of transcription in single cells with a broad-host range GFP-luxCDABE dual reporter system. PLoS One 7:e52000. https://doi.org/10.1371/journal.pone.0052000.

65. Campbell EA, Korzheva N, Mustaev A, Murakami K, Nair S, Goldfarb A, Darst SA. 2001. Structural mechanism for rifampicin inhibition of bacterial RNA polymerase. Cell 104:901–912. https://doi.org/10.1016/S0092-8674(01)00286-0.

66. Jatsenko T, Tover A, Tegova R, Kivisaar M. 2010. Molecular characterization of RifR mutations in Pseudomonas aeruginosa and Pseudomonas putida Mutat Res 683:106–114. https://doi.org/10.1016/j.mrfmmm.2009.10.015.

67. Artisovitch O, Vassylyeva MN, Svetlov D, Svetlov P, Perederina A, Igarashi N, Matsugaki N, Wakatsuki S, Tahirov TH, Vassylyev DG. 2005. Allosteric modulation of the RNA polymerase catalytic reaction is an essential component of transcription control by rnas. Cell 122: 351–363. https://doi.org/10.1016/j.cell.2005.07.014.

68. Nikel PI, Chinn J, Martínez-García E, Taylor AC, de Lorenzo V. 2013. Accumulation of inorganic polyphosphate enables stress endurance and catalytic vigour in Pseudomonas putida KT2440. Microb Cell Fact 12:50. https://doi.org/10.1186/1475-2859-12-50.

69. Nikel PI, Romero-Campero FJ, Zeidman JA, Gorhi-Moreno A, de Lorenzo V. 2015. The glycerol-dependent metabolic persistence of Pseudomonas putida KT2440 reflects the regulatory logic of the Gpr repressor. mBio 6:e00340-15. https://doi.org/10.1128/mBio.00340-15.

70. Tarassova K, Tegova R, Teras R, Tark M, Saumaa S, Kivisaar M. 2009. Elevated mutation frequency in surviving populations of carbon-starved pspD-deficient Pseudomonas putida is caused by reduced expression of superoxide dismutase and catalase. J Bacteriol 191: 3604–3614. https://doi.org/10.1128/JB.01803-09.

71. Teras R, Mikkel K, Berkovitch N, Tegova R, Tider A, Teras R, Siddronesk J, Tegova R, Tower A, Dame RT, Kivisaar M. 2012. Homologous recombination is facilitated in starving populations of Pseudomonas putida by phenol stress and affected by chromosomal location of the recombination target. Mutat Res 737:12–24. https://doi.org/10.1016/j.mrfmmm.2012.07.004.

72. Bowman LAM, McLean S, Poole RK, Fukuto JM. 2011. The diversity of microbial responses to nitric oxide and agents of nitrosative stress: close cousins but not identical twins. Adv Microb Physiol 59:135–219. https://doi.org/10.1016/S0065-2911(09)05588-1.

73. Cadet J, Douki T, Garaschutto D, Ravant JL. 2003. Oxidative damage to DNA: formation, measurement and biochemical features. Mutat Res 531:5–23. https://doi.org/10.1016/j.mrfmmm.2003.09.001.

74. Ferraro DJ, Gahar K, Ramaswamy S. 2005. Rieske business: structure-function of Rieske non-heme oxygenases. Biochem Biophys Res Commun 338:175–190. https://doi.org/10.1016/j.bbrc.2005.08.222.

75. Chávez FP, Lünsdorf H, Jérez CA. 2004. Growth of polychlorinated-biphenyl-degrading bacteria in the presence of biphenyl and chlorobiphenyls generates oxidative stress and massive accumulation of compounds—from one strategy to four. Nat Rev Microbiol 9:803–816. https://doi.org/10.1038/nrrmicro2652.

76. Danchin A, Binder PM, Noraia S. 2011. Antifragility and tinkering in biology (and in business) flexibility provides an efficient epigenetic way to manage risk. Genes 2:998–1016. https://doi.org/10.3390/genes40900999.

77. Mathieu A, Fleurier S, Frénoy A, Dairou J, Bredeche MF, Sanchez-Vizuete P, Song X, Matic I. 2016. Discovery and function of a general core horometic stress response in E. coli induced by sublethal concentrations of antibiotics. Cell Rep 17:46–57. https://doi.org/10.1016/j.celrep.2016.09.004.

78. Chávez FP, Lünsdorf H, Jérez CA. 2004. Growth of polychlorinated-biphenyl-degrading bacteria in the presence of biphenyl and chlorobiphenyls generates oxidative stress and massive accumulation of...
inorganic polyphosphate. Appl Environ Microbiol 70:3064–3072. https://doi.org/10.1128/AEM.70.5.3064-3072.2004.
97. Czechowska K, van der Meer JR. 2012. Reversible and irreversible pollutant-induced bacterial cellular stress effects measured by ethidium bromide uptake and efflux. Environ Sci Technol 46:1201–1208. https://doi.org/10.1021/es203352y.
98. Kanaly RA, Harayama S. 2010. Advances in the field of high-molecular-weight polycyclic aromatic hydrocarbon biodegradation by bacteria. Microb Biotechnol 3:136–164. https://doi.org/10.1111/j.1751-7915.2009.00130.x.
99. Kang YS, Lee Y, Jung H, Jeon CO, Madsen EL, Park W. 2007. Overexpressing antioxidant enzymes enhances naphthalene biodegradation in Pseudomonas sp. strain As1. Microbiology 153:3246–3254. https://doi.org/10.1099/mic.0.2007/008896-0.
100. Nikel PI, de Lorenzo V. 2018. Pseudomonas putida as a functional chassis for industrial biocatalysis: from native biochemistry to trans-metabolism. Metab Eng https://doi.org/10.1016/j.ymben.2018.05.005.
101. Sambrook J, Russell DW. 2001. Molecular cloning: a laboratory manual, 3rd ed. Cold Spring Harbor Laboratory, Cold Spring Harbor.
102. Martínez-García E, de Lorenzo V. 2012. Transposon-based and plasmid-based genetic tools for editing genomes of Gram-negative bacteria. Methods Mol Biol 813:267–283. https://doi.org/10.1007/978-1-61779-412-4_16.
103. Nikel PI, Kim J, de Lorenzo V. 2014. Metabolic and regulatory rearrangements underlying glycerol metabolism in Pseudomonas putida KT2440. Environ Microbiol 16:239–254. https://doi.org/10.1111/1462-2920.12224.
104. Foster PL. 2005. Stress responses and genetic variation in bacteria. Mutat Res 569:3–11. https://doi.org/10.1016/j.mrfmmm.2004.07.017.
105. Rosche WA, Foster PL. 2000. Determining mutation rates in bacterial populations. Methods 20:4–17. https://doi.org/10.1006/meth.1999.0901.
106. Martínez-García E, Aparicio T, de Lorenzo V, Nikel PI. 2014. New transposon tools tailored for metabolic engineering of Gram-negative microbial cell factories. Front Bioeng Biotechnol 2:46. https://doi.org/10.3389/fbioe.2014.00046.
107. Nikel PI, de Lorenzo V. 2013. Implantation of unmarked regulatory and metabolic modules in Gram-negative bacteria with specialised mini-transposon delivery vectors. J Biotechnol 163:143–154. https://doi.org/10.1016/j.jbiotec.2012.05.002.
108. Nikel PI, de Lorenzo V. 2013. Engineering an anaerobic metabolic regime in Pseudomonas putida KT2440 for the anoxic biodegradation of 1,3-dichloroprop-1-ene. Metab Eng 15:98–112. https://doi.org/10.1016/j.ymben.2012.09.006.
109. Dvořák P, Chríst L, Nikel PI, Fedr R, Soucek K, Sedlacková M, Chaloupková R, de Lorenzo V, Prokop Z, Damborský J. 2015. Exacerbation of substrate toxicity by IPTG in Escherichia coli BL21(DE3) carrying a synthetic metabolic pathway. Microb Cell Fact 14:201. https://doi.org/10.1186/s12934-015-0393-3.
110. Couce A, Blázquez J. 2009. Side effects of antibiotics on genetic variability. FEMS Microbiol Rev 33:531–538. https://doi.org/10.1111/j.1574-6976.2009.00165.x.