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The Glycerol-Dependent Metabolic Persistence of Pseudomonas putida KT2440 Reflects the Regulatory Logic of the GlpR Repressor

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ABSTRACT The growth of the soil bacterium Pseudomonas putida KT2440 on glycerol as the sole carbon source is characterized by a prolonged lag phase, not observed with other carbon substrates. We examined the bacterial growth in glycerol cultures while monitoring the metabolic activity of individual cells. Fluorescence microscopy and flow cytometry, as well as the analysis of the temporal start of growth in single-cell cultures, revealed that adoption of a glycerol-metabolizing regime was not the result of a gradual change in the whole population but rather reflected a time-dependent bimodal switch between metabolically inactive (i.e., nongrowing) and fully active (i.e., growing) bacteria. A transcriptional \( \Phi(\text{glpD-gfp}) \) fusion (a proxy of the glycerol-3-phosphate [G3P] dehydrogenase activity) linked the macroscopic phenotype to the expression of the \( \text{glp} \) genes. Either deleting \( \text{glpR} \) (encoding the G3P-responsive transcriptional repressor that controls the expression of the \( \text{glpFKRD} \) gene cluster) or altering G3P formation (by overexpressing \( \text{glpK} \), encoding glycerol kinase) abolished the bimodal \( \text{glpD} \) expression. These manipulations eliminated the stochastic growth start by shortening the otherwise long lag phase. Provision of \( \text{glpR} \) in \text{trans} restored the phenotypes lost in the \( \Delta\text{glpR} \) mutant. The prolonged nongrowth regime of \( \text{P. putida} \) on glycerol could thus be traced to the regulatory device controlling the transcription of the \( \text{glp} \) genes. Since the physiological agonist of GlpR is G3P, the arrangement of metabolic and regulatory components at this checkpoint merges a positive feedback loop with a nonlinear transcriptional response, a layout fostering the observed time-dependent shift between two alternative physiological states.

IMPORTANCE Phenotypic variation is a widespread attribute of prokaryotes that leads, inter alia, to the emergence of persistent bacteria, i.e., live but nongrowing members within a genetically clonal population. Persistence allows a fraction of cells to avoid the killing caused by conditions or agents that destroy most growing bacteria (e.g., some antibiotics). Known molecular mechanisms underlying the phenomenon include genetic changes, epigenetic variations, and feedback-based multistability. We show that a prolonged nongrowing state of the bacterial population can be brought about by a distinct regulatory architecture of metabolic genes when cells face specific nutrients (e.g., glycerol). Pseudomonas putida may have adopted the resulting carbon source-dependent metabolic bet hedging as an advantageous trait for exploring new chemical and nutritional landscapes. Defeating such naturally occurring adaptive features of environmental bacteria is instrumental in improving the performance of these microorganisms as whole-cell catalysts in a bioreactor setup.

The customary view of prokaryotic metabolism as a homogeneous and cooccurring process in space and time has been increasingly challenged in recent years (1, 2), particularly since the onset of single-cell technologies (3–6). These methodologies revealed a complete repertoire of responses to specific environmental conditions in individual microorganisms (7–12). Diversification of the metabolic regimes in single cells within otherwise clonal populations can be seen as a particular case of phenotypic variation (13, 14), in which different regulatory or epigenetic traits lead to the stochastic manifestation of alternative features in isogenic individuals (15–19). The phenomenon known as persistence, i.e., the occurrence of a live but nongrowing fraction of cells in a bacterial pool (20), is one of the most intriguing cases of phenotypic variation. While the lack of growth may appear negative at a glance, persistence ensures the survival of cells exposed to agents that act on developing bacteria, e.g., some antibiotics (21–23). Once the selective pressure ceases, persistent bacteria can resume growth and fully reconstruct the original population. Regardless of the mechanisms behind this behavior, the standing question is whether persistence is an adaptive trait or just a casual occurrence that happens to be beneficial for antibiotic-sensitive bacteria in the modern era of antimicrobial agents. What we qualify as persistence may just be a particular case of a more common situation in which a starting population stochastically splits between growing and nongrowing cell types when facing a new environmental or physicochemical condition. While persistence re-

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reflects the end of one such scenario (most bacteria grow but a few fail to grow), the opposite extreme (most cells remain static but a few grow) could also occur.

During the course of our studies on the metabolism of the soil bacterium *Pseudomonas putida* KT2440, we noticed that cells cultured on glycerol as the sole carbon source displayed an anomalously long lag period (≥10 h) before initiating any detectable growth. This situation was not observed when the cells were cultured on glucose or succinate under the same conditions. Much of the currently available information on glycerol metabolism in pseudomonads comes from studies of the human pathogen *Pseudomonas aeruginosa* (24, 25), and only recently have the transcriptional and metabolic changes associated with the growth of *P. putida* KT2440 on glycerol been assessed (26). Cells grown on the polyol undergo a complex transcriptional response that includes not only genes involved in central metabolic pathways but also additional ones encoding components of the respiratory chain and others related to stress resistance (27). Yet, the body of data currently available does not provide any clue on the distinct long-lag-phase phenomenon in glycerol. Since this behavior is recurrent upon reincoculation of the cells in fresh medium, we wondered whether the prolonged nongrowing regime of *P. putida* on glycerol was the result of (i) a lengthy, graded, and simultaneous adaptation to the new substrate or (ii) a runaway counterpart of persistence, i.e., the stochastic rise and eventual population takeover of individual substrate-metabolizing cells amid a majority of glycerol-unresponsive bacteria.

In this work, we present a systematic study of the growth and physiology of individual *P. putida* KT2440 cells in cultures with the polyol as the sole carbon source. In particular, we show that exposure to glycerol led to the appearance of two clearly separate subpopulations that differed in their metabolic activity toward the substrate, where the relative proportion of these subpopulations changed with time. Furthermore, such a phenotypic diversification was wired to the dual logic of the GlpR regulator, which represses transcription of the cognate *glp* genes encoding the enzymes needed for glycerol catabolism. Our results suggest that the metabolic heterogeneity observed in glycerol cultures could be an evolutionary trait of *P. putida*, endowing the bacterium with bet-hedging strategies for scounting new substrates-to-be in natural ecological niches.

**RESULTS AND DISCUSSION**

**Glycerol consumption pathways in *P. putida* KT2440 exposed by the genomic organization of the *glp* locus.** Two glycerol dissimilation pathways have been described in bacteria: one begins with a phosphorylation step, followed by phosphorylation, and the other begins with a phosphorylation reaction, followed by dehydrogenation (28, 29). In either case, the terminal metabolic product is dihydroxyacetone-P. In the soil bacterium *P. putida* KT2440, the second pathway is the prevalent (and likely the only) glycerol dissimilatory process (Fig. 1). Uptake of the compound is mediated by the GlpF facilitator, which fosters a diffusion reaction (30). Once inside the cell, glycerol is phosphorylated by an ATP-dependent glycerol kinase (GlpK) to sn-glycerol-3-P (G3P), which cannot diffuse out of the cell. G3P is the substrate for GlpD, a membrane-bound G3P dehydrogenase that yields dihydroxyacetone-P (Fig. 1A). Some microorganisms, such as *Escherichia coli* and *P. aeruginosa*, are able to internalize and utilize G3P or dihydroxyacetone (28), intermediates of the biochemical polyl processing (Fig. 1A). The activities are encoded in *P. putida* by the *glp* gene cluster (26, 31), which includes *glpF, glpK, glpR*, and *glpD* (Fig. 1B). Deep RNA sequencing indicated that *glpR* is constitutively transcribed at a low level in the whole bacterial population irrespective of the carbon source, while the rest of the *glp* genes have a significant expression level only in glycerol cultures (27). Coverage plots demonstrated that there are two transcriptional units within the *glp* gene cluster: one transcriptional unit encompasses *glpF* and *glpK*, and *glpD* is independently transcribed (27). All of these genes are under the transcriptional control of the negative regulator GlpR, a regulatory arrangement previously described in other bacteria (32–34). In *silico* predictions have suggested that the promoters of the *glp* regulon share a regulatory motif in Gram-negative bacteria (35), and on the basis of this feature, we were able to recognize a well-defined GlpR-binding sequence upstream of *glpD* (Fig. 1B). Danilova et al. (35) also hinted at the existence of more GlpR-binding motifs upstream of genes relevant for glycerol uptake and utilization but not up-stream of *glpR*. Against this background, we examined how all
these biochemical and genetic features of *P. putida* KT2440 bring forth a distinctive growth phenotype on glycerol.

Behavior of single *P. putida* KT2440 cells growing on glycerol. When *P. putida* is grown on glycerol as the sole carbon source, the cultures show an unexplainably prolonged lag phase that typically lasts \( \geq 10 \) h (26, 31). The conspicuously long time needed to start growth is independent of the carbon source used to grow the inocula and almost disappears \((1.5 \pm 0.9 \) h\) only when cells are pregrown on the polyol. The lag phase lasted for 17.4 \(\pm\) 3.9 h and 19.8 \(\pm\) 2.4 h when cells were transferred from either succinate or glucose cultures into a glycerol-containing medium, respectively. Interestingly, the addition of substrate mixtures (e.g., glucose and glycerol) to the cultures helps reduce the extension of the lag phase. This situation indicates that a metabolic regulatory phenomenon (e.g., the accumulation of a critical metabolic intermediate) could be involved in the delayed growth on glycerol. Since the growth retardation was even more noticeable when *P. putida* was pregrown in rich LB medium and then passed into M9 minimal medium containing glycerol (the lag phase lasted \(>20 \) h), we adopted this culture approach in all the experiments described in this study. A considerable reorganization of metabolic pathways is expected to occur when cells are shifted from rich to minimal growth conditions (36), including changes in gene expression during the lag phase (37). However, the unexpectedly long lag phase of strain KT2440 on glycerol cannot be easily explained. What are the metabolic and regulatory causes behind such delayed growth of *P. putida* on the polyol?

The first aspect to be addressed was whether (i) the growth retardation applied to all cells of the culture in a more or less synchronous fashion or (ii) what we observed at the population level in reality reflected a stochastic start of growth of individual cells. We monitored the evolution of separate cultures of *P. putida* in microtiter plates inoculated with a dilution of a bacterial suspension that corresponded to an average of 1 cell per well (Fig. 2). Cells previously cultured in LB medium were used as the inoculum, and after washing them with M9 minimal medium without any carbon source (to remove any traces of the complex medium), we transferred the bacteria into M9 minimal medium supplemented with either glucose (i.e., glycolytic substrate), succinate (i.e., gluconeogenic substrate), or glycerol (in which cells adopt a mixed metabolic regime recruiting both gluconeogenic and glycolytic metabolic activities [26]). Inocula were diluted in such a way that each culture (in 96-well microtiter plates) was calculated to receive 1 cell (see Materials and Methods for details on numerical considerations). Growth in each well was periodically monitored as the change in optical density measured at 600 nm \((\text{OD}_{600})\), and results from 1,000 wells were analyzed to ensure a data set large enough to have statistical significance. Figure 2A reveals various informative features. First, the lag phase differed significantly between cultures using different carbon sources (26). Cells started to grow on glucose or succinate in an almost synchronous fashion (data not shown). In contrast, glycerol cultures starting from single cells per well showed a remarkable variation in the extension of the lag phase. Second, once bacterial growth started, the specific growth rate \(\mu_{\text{max}}\) for a given carbon source was virtually identical among all individual wells. Succinate promoted the fastest growth \((\mu_{\text{max}} = 0.43 \pm 0.07 \text{ h}^{-1})\), closely followed by glucose \((\mu_{\text{max}} = 0.39 \pm 0.06 \text{ h}^{-1})\) and then by glycerol \((\mu_{\text{max}} = 0.24 \pm 0.05 \text{ h}^{-1})\). These results qualitatively mirror the behavior of *P. putida* growing in these carbon sources in shaken-flask cultures and using a larger inoculum (26). Regardless of the extension of the lag phase and the individual \(\mu_{\text{max}}\) values, all individual samples reached more or less the same maximal \(\text{OD}_{600}\) values (in the case of glycerol cultures, ca. 0.5 to 0.6 units)

These results made us wonder whether the asynchronous distribution of growth curves on glycerol (Fig. 2A) could be just the spurious consequence of differences in the initial number of cells per well. In other words, we questioned whether the observed phenomenon might be related to the manipulation of the cells rather than arising from physiological or genetic traits. Due to stochastic variations in the experimental procedure (e.g., differences in the inoculum volume transferred into the wells), the number of cells per well is not exactly known and may vary from well to well. The probability \(P\) of inoculating zero, one, two, or three cells per well is shown, indicating that the probability of inoculating a single cell is the most likely outcome.

**FIG. 2.** (A) Growth of *P. putida* KT2440 in single-cell batch cultures. Multwell microtiter plates were inoculated with a highly diluted preculture previously developed in LB medium in order to start each culture from 1 cell per well. Cells were grown at 30°C in 200 \(\mu\)l of M9 minimal medium containing 40 mM glycerol with rotary agitation. The time needed to reach an optical density measured at 600 nm \((\text{OD}_{600})\) of 0.3 (mid-exponential phase of growth) is indicated by a red arrowhead for 50 independent cultures to illustrate the delay in growth initiation among individual wells. This parameter, termed time of metabolic response \((t_{\text{MR}})\), was further used to quantify the response of single-cell cultures. (B) Probability of inoculating a given number of cells in a particular well, as estimated by the Poisson probability distribution. After dilution and inoculation, and due to stochastic variations (e.g., small differences in the volume of the bacterial suspension transferred into each well), the number of cells per well \((n)\) is not exactly known and may vary from well to well. The probability \(P\) of inoculating zero, one, two, or three cells per well is shown, indicating that the probability of inoculating a single cell is the most likely outcome.
theoretical assess the number of cells which can be expected per well (see Materials and Methods). Figure 2B shows that the probability of inoculating zero, one, two, or three cells per well is appreciable, whereas the probability of inoculating more than four cells is negligible. Inoculating a single cell per well is the most likely outcome \( P = 0.343 \), and consequently, the growth pattern observed in each well can be associated with a sole \( P. putida \) cell. The possibility of inoculating more than 1 cell is not to be neglected, but we show that our observations are not produced by the initial variability in the number of cells per well (see below).

Single \( P. putida \) KT2440 cells grown on glycerol undergo different metabolic regimes. The results described above raised the question of how individual cells within a \( P. putida \) KT2440 population react metabolically to exposure to different carbon sources. The gross physiological state of single cells in shaken-flask cultures was examined to study this issue. Since a large number of bacterial oxidases and reductases are involved in central enzymatic functions, the merged redox activity of a cell can be considered a proxy of its overall physiological vitality. On this background, the BacLight RedoxSensor Green vitality stain, 3,8-diamino-5-[3-(diethylmethylammonio)propyl]-6-phenyl diiodide (RSG reagent), was used to diagnose the metabolic state of individual cells. This reagent can be transformed by cell reductases into a highly fluorescent compound, a reliable descriptor of changes in the electron transport chain activity and other vital cellular processes (39, 40). As such, the metabolic heftiness of individual cells could be directly related to the intensity of the RSG signal (17). We stained \( P. putida \) cells harvested during mid-exponential growth from shaken-flask cultures with the same carbon sources described above. Bacteria were precultured in LB medium, washed with M9 minimal medium without any carbon source, and then passed into fresh M9 minimal medium supplemented with either succinate, glucose, or glycerol. Mid-exponential cultures were harvested at an OD\(_{600}\) of 0.5 (note that the duration of the lag phase was different for each carbon source [26]), and cells were stained with the RSG reagent and inspected under the fluorescence microscope. The distribution of RSG-stained cells revealed different patterns depending on the substrate used (Fig. 3). On one hand, the relative intensity of RSG staining increased in the order glycerol < glucose < succinate. On the other hand, the same cell-bound signals appeared to be homogeneously distributed in the population of bacteria grown on glucose and succinate but not so in the glycerol-grown counterpart. No significant signal was observed in cells not treated with the RSG reagent, ensuring that the observed fluorescence stems from redox activity and not from autofluorescence or siderephore production (41). While small cell-to-cell differences in the RSG output of the glucose- and succinate-grown cultures can be considered normal, the clear divide between different metabolic states exposed by the data in Fig. 3 (bottom panel) prompted us to reexamine the case in more detail by means of flow cytometry. This method not only allows for the quantitation of the fluorescent signal in single cells but also enables the recognition of distinct subpopulations (42).

Quantitative image analysis of the fluorescence microscopy pictures of Fig. 3 further exposes the cell-to-cell variability observed in terms of RSG staining in glycerol cultures but not in glucose or succinate cultures (see also Fig. S1 in the supplemental material).

Figure 4 shows the results obtained by flow cytometry of cells from the same shaken-flask cultures described above. In order to validate the positive correlation between RSG staining and metabolic activity of individual cells, samples from each culture condition were either treated with NaN\(_3\) or heated at 85°C for 25 min to inactivate the cells. In either case, \( P. putida \) KT2440 showed no RSG signal in flow cytometry and cells were indistinguishable from the unstained control irrespective of the carbon source (data not shown). Various types of RSG-staining patterns quickly developed in live cells. First, as observed under the fluorescence microscope (Fig. 3), no significant differences were noticed in either the apparent size or the complexity of individual cells irrespective of the carbon source (i.e., forward scatter versus side scatter plots; Fig. 4, left panels). Second, bacteria grown on glucose or succinate were all positive for RSG staining and thus originated a single peak of fluorescence that was clearly displaced from the negative (unstained) control. This situation indicates that all cells were actively metabolizing the substrates on which they were growing. The unimodal distribution of RSG\(^+\) signals and their intensity matched the fluorescence microscopy data of Fig. 3, with \( P. putida \) cells from succinate cultures giving the highest output.

In contrast to cultures developed under entirely glycolytic or gluconeogenic regimes, when cells grew on glycerol a sharp divide of the bacterial...
population into two groups became apparent. The two subpopulations differed in their degree of staining: 62% of the cells were RSG⁺, while the remaining 38% exhibited almost no fluorescence (Fig. 4, bottom panel). One of the subpopulations virtually overlapped with the negative (i.e., unstained) control experiment, meaning that these cells lacked any significant metabolic activity and can be considered metabolically dormant. The divide was consistently observed throughout the mid-exponential phase of growth but tended to gradually merge as a single bacterial pool of metabolically active cells (i.e., most of the cells became RSG⁺) as the growth progressed and the cultures moved into the late exponential phase (see Fig. S2 in the supplemental material).

These results provide a direct link between the time-dependent metabolic bimodality of single cells grown in glycerol and the macroscopically observed lag phase of the corresponding batch cultures. Since the bimodal behavior of RSG⁺ *P. putida* cells was observed only in glycerol cultures and not in the presence of glucose or succinate, we hypothesized that the bifurcation was connected to the metabolism of the polyol itself. However, such plausible connection was by no means evident in the physiological experiments conducted so far, and we therefore set out to explore it with a suite of genetic and biochemical approaches.

**Quantitative analysis of growth patterns.** To analytically study the growth pattern under different culture conditions, we defined a kinetic parameter, termed time of metabolic response (*t*ₘᵣ). This parameter is the time needed for a given culture to attain an optical density measured at 600 nm (OD₆₀₀) of 0.3. By reserving the flow cytometry analysis, the RSG signal was investigated, is indicated in the forward scatter (FSC) versus side scatter (SSC) plots (left column). The histograms in the right column show the distribution of RSG fluorescence in cells harvested from glucose, succinate, and glycerol cultures. The gray rectangle in each histogram plot identifies the region considered negative for the fluorescence signal (as assessed with unstained cells).

![Flow cytometry analysis of the BacLight RedoxSensor Green (RSG) signal in *P. putida* KT2440 growing on different carbon sources. Cells were grown in shaken flasks until they reached the mid-exponential phase of growth and were treated with RSG as indicated in the legend to Fig. 3, and the fluorescence intensity was quantified by flow cytometry. The grayed population, in which the RSG signal was investigated, is indicated in the forward scatter (FSC) versus side scatter (SSC) plots (left column). The histograms in the right column show the distribution of RSG fluorescence in cells harvested from glucose, succinate, and glycerol cultures. The gray rectangle in each histogram plot identifies the region considered negative for the fluorescence signal (as assessed with unstained cells).](image)

![Frequency distribution of the time of metabolic response (*t*ₘᵣ) derived as the time needed for a given culture to attain an optical density measured at 600 nm (OD₆₀₀) of 0.3 in 1,000 single-cell *P. putida* KT2440 cultures carried out in microtiter plates with different carbon sources as described in the legend to Fig. 2. The bars indicate the number of independent wells in which the cultures attained an OD₆₀₀ of 0.3 at a given *t*ₘᵣ. (B) Closeup of the frequency of *t*ₘᵣ values observed in glycerol cultures. The calculated Gaussian distributions are superimposed on the bar graph to identify the possible subpopulations under these conditions (see also Table S2 in the supplemental material). The inset indicates the absolute values of the *t*ₘᵣ derived from the distribution shown in panel A for succinate (S), glucose (G), and glycerol cultures. Note that in the case of glycerol cultures, there are two values corresponding to the subpopulations observed under these conditions. In the inset, each bar represents the mean value of the *t*ₘᵣ parameter ± SD.](image)
whereas a second subpopulation comprised 11% of the entire population and had a \( t_{\text{MR}} \) of 29.8 ± 0.7 h (Fig. 5B). Furthermore, the application of these statistical methodologies to data from succinate and glucose cultures revealed a single population for each condition, as expected from the results in Fig. 5A. In other words, our statistical analysis identified two subpopulations in glycerol cultures, clearly distinguishable by their characteristic \( t_{\text{MR}} \) values (Fig. 5B, inset). These results, in turn, correlate well with the bimodal distribution of cells differing in their metabolic activity (Fig. 4). We simulated a scenario where the only source of stochasticity was the initial number of cells and obtained a unimodal distribution of \( t_{\text{MR}} \) values (see Fig. S3 in the supplemental material). Once the phenomenon of metabolic bimodality on glycerol (which results in a stochastic distribution of growth initiation of single cells and a population-level prolonged lag phase) was well documented, the next question related to the underlying molecular mechanism.

The GlpR repressor imposes a bistable growth pattern of cells grown on glycerol. Inspection of the biochemical and regulatory components underlying glycerol utilization in *P. putida* (Fig. 1A and B) and comparison of the same elements in other bacteria reveal important information about the use of this polyol by Gram-negative bacteria (45). It can be seen that (i) the *glp* structural genes are under the transcriptional control of a single regulatory protein (i.e., GlpR), known to act as a transcriptional repressor (34, 35); (ii) G3P seems to be the key metabolite that relieves the GlpR-dependent repression of the *gln* gene cluster (28); (iii) since G3P is not a substrate of the GlpF facilitator (30, 46), it can neither be taken up by *P. putida* cells nor leaked to the surrounding medium; and (iv) the very driving force for glycerol uptake thus arises from substrate phosphorylation by GlpK. Not surprisingly, this particular arrangement of regulatory elements is reminiscent of other genetic systems known to be submitted to stochastic fluctuations, such as the lac operon in *E. coli* (see below).

A typical bistable behavior at the transcriptional level is imposed by a repressor protein on the transcription of regulated genes (16, 18). The underlying mechanism proposed to explain bistability assumes a specific feedback that acts in combination with a nonlinear response within a transcriptional network (14). The archetype example in this sense is the lac operon of *E. coli*, in which the LacI repressor sets a bimodal expression pattern on the cognate genes (47); the same goes for the structure of the *gln* genes in *P. putida*. Within this framework, it seems likely that the GlpR regulator and the *gln* genes comprise a genetic system prone to exhibiting a bistable regime. To explore this possibility, we eliminated the *glnR* gene in *P. putida* KT2440 (which would determine the constitutive expression of the *gln* genes) and we studied the phenotypic traits discussed in the sections above in the mutant strain.

The first indication of the involvement of GlpR in the bistable growth phenotype of *P. putida* came from shaken-flask culture experiments with the \( \Delta glnR \) mutant using glycerol as the carbon source. Cells used as the inoculum were grown in LB medium. The duration of the lag phase was reduced to 6.5 ± 1.7 h (i.e., approximately 16 h shorter than the lag phase observed for the wild-type strain under the same culture conditions). We then focused on the catabolic enzymes involved in glycerol utilization (Fig. 6A), and we measured the GlpK activity *in vitro* to study the effect of eliminating GlpK in strain KT2440. Figure 6B shows that the GlpK activity is highly responsive to the carbon source. Wild-type cells growing under a glycolytic metabolic regime had the lowest GlpK activity, although the level of enzymatic activity was distinguishable from the background (attaining 7.5 ± 2.5 nmol·min\(^{-1}·mg\) protein\(^{-1}\)). The *in vitro* GlpK activity increased ca. 5-fold when *P. putida* cells were cultured on glycerol, as reported previously (26). However, the specific GlpK activity in the \( \Delta glnR \) strain did not react to the carbon source used to grow the cells. The GlpK activity was detectable even in mutant cells grown in LB medium (data not shown), a culture condition under which the same enzymatic activity was almost completely repressed in the wild-type strain. These results indicate that the elimination of the GlpR repressor protein renders the expression of *glnK* (and therefore the cognate enzymatic activity) constitutive. Moreover, elimination of GlpR completely abolished the bistable growth phenotype, and the entire subpopulation now showed a unimodal behavior (Fig. 6C). This evidence would in principle suffice to explain the observed phenotype of *P. putida* KT2440 on glycerol, and it also provides a biochemical clue to its origin. As an additional control experiment designed to ascribe the bistable growth behavior to the GlpR repressor, we cloned the cognate gene in the low-copy-number pSEVA224 expression vector (giving rise to plasmid pE-pE-glp [Table 1]), and the \( \Delta glnR \) strain was complemented back. The expression of *glnK* in trans restored the stochastic profile of growth (Fig. 6C and D), causing a significant decrease in the *in vitro* GlpK activity (8.7 ± 1.9 nmol·min\(^{-1}·mg\) protein\(^{-1}\), comparable to the activity observed in the wild-type strain under similar growth conditions). The results accredit the role of GlpR as a transcriptional repressor of the cognate *gln* genes and suggest that the delayed growth of part of the bacterial cells is related to a limited activity of the catabolic enzymes encoded therein.

**Time-dependent transcription of the *gln* genes as a descriptor of the macroscopic phenotype on glycerol.** In an attempt to correlate the regulation of the *gln* genes and the phenotype observed in glycerol cultures, we constructed a transcriptional \( \Phi(glpD-gfp) \) fusion based on the green fluorescent protein (GFP) to assess the *gfpD* expression level (Fig. 7A). In this construct, the native promoter and the Shine-Dalgarno sequence of *gfpD* were placed before *gfp* (in the pSEVA637 vector) to drive its expression. In these experiments, *P. putida* KT2440 cells carrying the \( \Phi(glpD-gfp) \) reporter (plasmid pTF-*glpD* [Table 1]) were grown on glycerol in shaken-flask cultures, and broth samples were taken at selected time points to study gene expression by flow cytometry. Note that, besides a negative control (Fig. 7B), the data points at which cells were collected correspond to the lag phase (8 h, Fig. 7C), mid-exponential phase (18 h, Fig. 7D), and stationary phase (36 h, Fig. 7E), reflecting different physiological situations as the bacteria proceed through the growth curve. Figure 7C to E summarizes the evolution of the *gfpD* transcriptional activity in strain KT2440 carrying pTF-*glpD* over time. At 8 h, before cells had started noticeable growth, most of the bacteria were negative for \( \Phi(glpD-gfp) \) transcription, although a small subpopulation of GFP+ cells was consistently detected (accounting for <20% of the total population; Fig. 7C). After 18 h of glycerol-dependent growth, when cells were actively growing, the separation of two subpopulations in terms of GFP content was still evident (Fig. 7D), although the percentage of GFP+ cells under these circumstances increased to 57%. Moreover, the analytical deconvolution of the flow cytometry histogram indicated that the whole signal was the result of two overlapped Gaussian distributions.
Fig. 7D, inset). This analysis indicates that the GFP\(^{-}\)/H11002 and GFP\(^{+}\)/H11001 populations represent 35% and 65% of the total fluorescence signal in the histogram, respectively. When cells attained the early stationary phase (Fig. 7E), all of them were GFP\(^{+}\)/H11001 in a unimodal distribution. Interestingly, the glpD expression pattern qualitatively mirrored the distribution of metabolically active and inactive cells (Fig. 4; see also Fig. S2 in the supplemental material).

The experimental evidence so far indicated that GlpR is ult-

**TABLE 1** Bacterial strains and plasmids used in this work

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<tr>
<th>Bacterial strain or plasmid</th>
<th>Relevant characteristics(^a)</th>
<th>Reference or source</th>
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<td><strong>Strains</strong></td>
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<td>Cloning host; (\Delta(ara-leu)) araD (\Delta lacX174) galE galK phoA thiE1 rpsE rpoB (Rif(^{R})) argE(Am) recA1 Apir lysogen</td>
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<td>HB101</td>
<td>Helper strain; F(^{-}) (\lambda^{+}) hsdS20(t(<em>{R}) m(</em>{R})) recA13 leuB6(Am) araC14 (\Delta(gpt-proA)) 62 lacY1 galK2(Oc) xyl-5 mtl-1 thiE1 rpsL20(Sm(^{r})) glnX44(AS)</td>
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<td>Wild-type strain; mt-2 derivative cured of the TOL plasmid pWW0</td>
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<tr>
<td>pRK600</td>
<td>Helper plasmid used for conjugation; ori(ColE1), RK2(mob(^{+}) tra(^{+})); Cm(^{r})</td>
<td>83</td>
</tr>
<tr>
<td>pSEVA637</td>
<td>Cloning vector; ori(pBBR1), promoter-less gfp; Gm(^{r})</td>
<td>84, 85</td>
</tr>
<tr>
<td>pTF-glpD</td>
<td>pSEVA637 derivative with a DNA fragment corresponding to the 5(^{\prime}) UTR of glpD, including the glpD promoter and Shine-Dalgarno sequence, cloned before the gfp gene [(\Phi(glpD-gfp))]; Gm(^{r})</td>
<td>This work</td>
</tr>
<tr>
<td>pSEVA224</td>
<td>Expression vector; ori(RK2); Km(^{r})</td>
<td>84, 85</td>
</tr>
<tr>
<td>pE-glpR</td>
<td>pSEVA224 derivative carrying gfpR; Km(^{r})</td>
<td>This work</td>
</tr>
<tr>
<td>pE-glpFK</td>
<td>pSEVA224 derivative carrying gfpF and gfpK; Km(^{r})</td>
<td>This work</td>
</tr>
</tbody>
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\(^a\) Antibiotic markers: Cm, chloramphenicol; Gm, gentamicin; Km, kanamycin; Rif, rifampin; Sm, streptomycin; 5\(^{\prime}\) UTR, 5\(^{\prime}\) untranslated region.
mately responsible for the bistable growth phenotype of *P. putida* KT2440 on glycerol. The next question was whether the transcription of the glp genes in a ΔglpR background also reflects the macroscopic effects observed in the mutant strain. We repeated the transcriptional assessment of the Φ(glpD-gfp) expression in *P. putida* ΔglpR and in the same strain complemented in trans with glpR (Fig. 8), using cells transformed with the empty pSEVA637 vector as the negative control (Fig. 8A). The transcription of glpD at 18 h in the mutant strain showed an entirely unimodal distribution (Fig. 8B), indicating that the transcription of Φ(glpD-gfp) becomes constitutive in the absence of the repressor protein in a fashion similar to what was observed in terms of the overall metabolic activity (Fig. 4) and the *in vitro* GlpK activity (Fig. 6B). This behavior was reverted by supplying glpR in an expression plasmid (Fig. 8C), and the resulting bimodal distribution mirrored the one observed for the wild-type strain under the same growth conditions (Fig. 7D). These results accredit the role of GlpR-dependent repression of the glp genes as the main cause of the phenotypic and growth properties of *P. putida* KT2440, drawing a direct correlation between the transcription of the glp genes, the enzymatic activity of the cognate catabolic pathway, and the overall physiology (as exposed by the duration of the lag phase) when cells are grown in mass on glycerol.

**Increasing the intracellular glycerol-3-P content relieves the bimodal growth pattern on glycerol.** The driving force for the uptake of glycerol is the substrate phosphorylation to G3P by GlpK (Fig. 1A and 6A). G3P, in turn, relieves the GlpR-associated repression of the glp genes in a positive feedback loop. We reasoned that this repression could also be lifted by artificially increasing the G3P pool. Is it possible for *P. putida* cells to transport G3P directly from the extracellular environment? A genomic anal-

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**FIG 7** Transcriptional analysis of glpD expression in *P. putida* KT2440. (A) Schematic representation of the transcriptional Φ(glpD-gfp) fusion used to explore the expression of glpD. The 5′ untranslated region preceding glpD (which spans the P<sub>glpD</sub> promoter and a Shine-Dalgarno [SD] sequence [Fig. 1B]) was cloned into the pSEVA637 vector, which carries a promoter-less gfp, generating plasmid pTF-glpd. The elements in this outline are not drawn to scale. (B to E) Time-lapse of GFP fluorescence in *P. putida* KT2440 cells carrying the Φ(glpD-gfp) transcriptional fusion (plasmid pTF-glpd) and grown in shaken-flask cultures in M9 minimal medium containing glycerol as the sole carbon source. The gray rectangle in each plot identifies the region considered negative for the fluorescence signal (as assessed with cells carrying the empty pSEVA637 vector). The inset in panel D identifies the percentages of GFP<sup>+</sup> and GFP<sup>−</sup> cells as identified by analytical deconvolution of the raw flow cytometry data.

**FIG 8** Role of GlpR repressor and of availability of glycerol-3-P in the expression of the glp genes in *P. putida*. (A to C) Transcriptional analysis of the expression of Φ(glpD-gfp) (carried on plasmid pTF-glpd) in the ΔglpR background at 18 h. Cells were grown in shaken-flask cultures using glycerol as the sole carbon source. The gray rectangle in each plot identifies the region considered negative for the fluorescence signal (as assessed with cells carrying the empty vector pSEVA637). (D) *In vitro* quantitation of the specific (Sp) GlpK enzymatic activity in the wild-type strain grown on glycerol and carrying the empty expression vector (pSEVA224) or overexpressing both glpF and glpK from plasmid pE-glphK. Each bar represents the mean value of the corresponding enzymatic activity ± the standard deviation of duplicate measurements from at least three independent experiments. (E) GFP fluorescence in *P. putida* cells carrying the Φ(glpD-gfp) transcriptional fusion in the wild-type strain overexpressing both glpF and glpK after 18 h of growth in shaken-flask cultures using glycerol as the sole carbon source.
ysis of the *P. putida* KT2440 catalogue of glycerol-processing genes suggests that there are not specific transporters for this small molecule (26, 48), and when cells were incubated on M9 minimal medium containing G3P as the sole carbon source, no growth was observed (data not shown). An alternative possibility would be to boost the GlpK activity alone, but in this case, glycerol (the substrate for the kinase) might become limiting if the GlpF transporter (also dependent on GlpR) is not sufficiently active (49).

We attempted to increase the G3P input by overexpressing both *glpF* and *glpK* from strain KT2440 by cloning these genes in an expression plasmid (pSEVA224), giving rise to pE-*glpFK* (Table 1). Upon induction of the P<sub>trc</sub> promoter, which drives the expression of *glpF* and *glpK* with isopropyl-β-d-1-thiogalactopyranoside (IPTG), we measured the specific GlpK activity in vitro in glycerol cultures of the recombinants (Fig. 8D). The overexpression of *glpF* and *glpK* from plasmid pE-*glpFK* resulted in a 3-fold increase in the in vitro GlpK activity compared to a control experiment with *P. putida* KT2440 cells carrying the empty pSEVA224 vector. The activity values obtained for the control strain in this experiment are similar to those reported in Fig. 6B, indicating that the presence of the plasmid does not alter the in vitro GlpK activity. Although we did not measure the actual G3P concentration in these cells, it can be safely assumed that the turnover of this metabolite should be also increased in the strain overexpressing *glpK*, as observed in *Ralstonia eutropha* (49). We then explored the transcription level of the *glpD* gene using the transcriptional fusion described above (Fig. 7A) in glycerol cultures of the wild-type KT2440 strain overexpressing *glpF* and *glpK*. At 18 h, all the cells gave a positive signal for GFP in a unimodal fashion (Fig. 8E), similarly to the distribution observed in the Δ*glpR* mutant under the same growth conditions (Fig. 8B). This result indicates that the GlpR-dependent repression of the *glp* genes is relieved by increasing the intracellular G3P availability—demonstrating that G3P is the key metabolite that mediates the bistable phenotype of *P. putida* KT2440 on glycerol. When the analysis of single-cell batch cultures in microtiter plates was performed with these recombinant cells, the two subpopulations previously seen for the wild-type strain were not observed (data not shown). These results are in line with the notion that if cells are pregrown on glycerol (ensuring a sufficiently high G3P supply), the stochastic growth phenomenon is no longer observed.

**Conclusion.** Phenotypic variation within an otherwise genetically clonal population endows bacterial populations with functionalities that, under given particular circumstances, protect the genetic pool of the group under adverse conditions. The essence of the phenomenon is that either a genetic or a metabolic device generates differential expression of a certain trait, whether structural (e.g., phase variation of pathogens [50]), biochemical, or physiological. Pseudomonads exhibit such phenomena in the natural niches which they inhabit (17, 51). While virtually all prokaryotic promoters are subject to a degree of noise (52), certain regulatory architectures translate such noise into bi-/multimodal or bistable manifestation of the phenotype at stake in single cells. The ON/OFF ratio in these cases can be permanently set by a dedicated molecular device, or it can vary with time or other conditions. Each of these scenarios seems to have been selected to cope with evolutionary challenges and environmental stresses. A dramatic case is that of persister cells (22), which avoid killing by remaining in a static physiological state when most of the population succumbs to the action of antibiotics that target growing bacteria. In other instances, the same population splits into two or more groups that operate different portions of the same metabolic pathway (17).

In the scenario documented in this paper, the stochastic expression of genes needed for glycerol metabolism leads to a prolonged non-growing state of most of the population of *P. putida* KT2440 when cells face this substrate (Fig. 9). Cells will start growing only if the low-probability effector-independent stochastic lifting of the GlpR-mediated repression allows for the expression of *glpF* and *glpK* (the latter being responsible for G3P formation). Once this repression is overcome stochastically, the full expression of the *glp* genes can proceed—finally returning to an OFF state when the substrate is completely depleted. Different levels of metabolic activity are observed in the cells (i.e., their ability to catabolize glycerol) while the transcriptional derepression process is occurring. This situation, in turn, explains the very long lag phase in mass *P. putida* cultures on glycerol. Along this line of reasoning, it has been recently shown that the level of activity of a given metabolic pathway (and the accumulation of critical metabolic intermediates stemming from it) governs the transition from a non-growing to an actively growing state in bacteria and yeast (10–12).

What could be the advantage for bacteria holding such regulatory devices for glycerol metabolism? Glycerol is not an abundant carbon source in the environment, but it does appear, e.g., in root exudates in which pseudomonads are known to thrive (53, 54). As hexoses and organic acids seem to be the preferred carbon sources for *P. putida* (55–57), it could well happen that the prolonged inactivity of the cells when facing glycerol enables carbon source-
dependent metabolic bet hedging (58) to explore new chemical and nutritional landscapes. This concept is reminiscent of foraging in animal ecology (59, 60). In this case, some members of the population (but not the whole population) take risks to broaden the search for alternative food sources (61). Under this scheme, the cost of randomly expressing metabolic genes in *P. putida* is outweighed by the potential benefit of locating (and being prepared to utilize) alternative carbon sources. A similar situation was previously proposed for *E. coli* (62), based on the whole-population transcriptomic fingerprint of cells grown on different carbon sources. The concept is further refined by the results discussed in our contribution, in which the phenotypic cell-to-cell variations are taken into account and integrated into the foraging theory. While metabolic stochasticity may be beneficial in the environment (63), it certainly becomes a hurdle for the use of *P. putida* in a bioreactor setup as a microbial cell factory (64–66). In this sense, the present results suggest genetic strategies to bring about a coordinated consumption of glycerol as the carbon source in an industrial setting, where the polyl is becoming one of the preferred and most readily available substrates for biotechnology (67–69).

**MATERIALS AND METHODS**

**Bacterial strains and growth conditions.** The bacterial strains and the plasmids used in this study are listed in Table 1. *P. putida* and *E. coli* were routinely cultured in 250-ml Erlenmeyer flasks containing 50 ml of the corresponding culture medium with agitation at 170 rpm at either 30°C (*P. putida* strains) or 37°C (*E. coli* strains). For the propagation and construction of vectors, *E. coli* CC118 apr was cultured in LB medium (70) containing the appropriate antibiotics. Physiological experiments were carried out in M9 minimal medium (70, 71), containing 6 g · liter⁻¹ Na₂HPO₄, 3 g · liter⁻¹ KH₂PO₄, 1.4 g · liter⁻¹ (NH₄)₂SO₄, 0.5 g · liter⁻¹ NaCl, 0.2 g · liter⁻¹ MgSO₄·7H₂O, and 2.5 ml · liter⁻¹ of a trace element solution (72). Either 40 mM glyceral, 20 mM glucose, or 30 mM sodium succinate was added to M9 minimal medium as filter-sterilized solutions as the sole carbon source (i.e., the final medium contained 120 mM carbon atoms in all the experiments). Solid medium also contained 15 g · liter⁻¹ agar. Antibiotics were added to medium when needed at the following final concentrations: ampicillin, 150 μg · ml⁻¹ for *E. coli* strains or 500 μg · ml⁻¹ for *P. putida* strains (used during mutant construction); kanamycin, 50 μg · ml⁻¹; and gentamicin, 15 μg · ml⁻¹. IPTG was used at 1.5 mM to induce the activity of the P⁰ promoter in overexpression experiments.

Growth experiment mixtures were inoculated with cells previously grown for 18 h in LB medium, previously spun, and washed twice with M9 minimal medium (previously added with the corresponding carbon source), each well would contain 1 × 10⁹ cells per microliter (73, 74). Kinetic parameters, such as the duration of the lag phase, were calculated as described previously (26), and in all cases, the mean value of the corresponding parameter is given along with the standard deviation calculated from at least five technical replicates in four independent experiments.

**General genetic techniques and mutant construction.** Unless otherwise stated, DNA manipulations followed well-established methods (70) and specific recommendations from manufacturers. Oligonucleotides were purchased from Sigma-Aldrich Co. (St. Louis, MO), and their sequences are listed in Table S1 in the supplemental material. A *P. putida* KT2440 Δ*glpR* mutant was constructed using the method of allelic replacement described by Martínez-García and de Lorenzo (75), with oligonucleotides Δ*glpR*-TS1-F, Δ*glpR*-TS1-R, Δ*glpR*-TS2-F, and Δ*glpR*-TS2-R.

**Cloning of the *glpD* promoter into the *glp* reporter plasmid pSEVA637.** The *glpD* promoter was identified by in silico searching of conserved motifs in the 5′ untranslated region within the *glp* gene cluster (35). The promoter was amplified by PCR from chromosomal DNA of *P. putida* KT2440 using oligonucleotides *glpD*-TF-EcoRI-F and *glpD*-TF-BamHI-R listed in Table S1 in the supplemental material. The 236-bp amplicon was digested and cloned as an EcoRI-BamHI fragment upstream of *gfp* into the multiple cloning site of pSEVA637. After transformation into *E. coli* CC118 apr, plasmids were isolated and the correct transcriptional *Φ*(*glpD*-gfp) fusion was verified by PCR, restriction analysis, and sequencing (Secugen SL, Madrid, Spain). The resulting *Φ*(*glpD*-gfp) reporter vector, termed pTF-glpd, was introduced into *P. putida* KT2440 and its Δ*glpR* mutant by triparental mating (76), using *E. coli* HB101/pRK600 (Table 1) as the helper strain.

**Overexpression of *glpF* and *glpK* as a synthetic operon and construction of a *glpR* complementation plasmid.** A synthetic operon, comprising *glpF* and *glpK*, was constructed by crossover (sewing) PCR (77). A first fragment, comprising *glpF*, was amplified using genomic DNA from *P. putida* KT2440 as the template and oligonucleotides *glpF*-EcoRI-F and *glpF*-EcoRI-R. The *glpK* gene was separately amplified with oligonucleotides *glpK*-F and *glpK*-SacI-R. Finally, selling PCR amplification was used to generate the synthetic operon using a mixture of the amplification products from the first and second PCR steps as the template and the external oligonucleotides (i.e., *glpF*-EcoRI-F and *glpK*-SacI-R). The 2,392-bp amplicon spanned the *glpF* and *glpK* genes as a single transcriptional unit bracketed by EcoRI and SacI restriction sites, with a Shine-Dalgarno motif included before the ATG codon of each gene. This fragment was cloned into the expression vector pSEVA224 (LacI∥Ptrc) digested with the same enzymes, giving rise to plasmid pE-glPFK. A similar procedure was used to construct an expression plasmid for *glpR* complementation. In this case, genomic DNA from *P. putida* KT2440 was used as the template in a PCR amplification using oligonucleotides *glpR*-EcoRI-F and *glpR*-BamHI-R. The 797-bp amplicon was digested with EcoRI and BamHI and cloned into pSEVA224, resulting in plasmid pE-glPR.

**Fluorescence microscopy, image analysis, and flow cytometry.** BacLight RedoSensor Green (RSG) vitality staining was used to diagnose the metabolic state of individual cells. The signal intensity of cells stained with the RSG reagent is altered when cells are treated with reagents that disrupt electron transport (such as NaN₃), and we resorted to this compound to run negative controls in each set of experimental samples. *P. putida* reacted significantly to the addition of 10 mM NaN₃. In each set of experiments, 10⁶ cells suspended in 0.5 ml of phosphate-buffered saline (PBS) buffer (70) were stained with 1 μl of the RSG solution provided by Life Technologies Corp. (Grand Island, NY). Staining procedures followed the instructions of the manufacturer. In some experiments, cells were also stained with propidium iodide to evaluate vitality. Bacteria with damaged cell membranes represented a very small population (<10%) during logarithmic growth in all the carbon sources tested, and excluding this subpopulation by gating on the propidium iodide-negative population did not alter the distribution of RSG subpopulations (data not shown). For immobilization of the cells in microscopy experiments, 2 μl
of the cell suspension was placed onto 0.01% (wt/vol) poly-l-lysin (Sigma-Aldrich Co.)-coated coverslips and dried. Then, each coverslip was assembled with a slide including Prolong (Life Technologies Corp.) to suppress photobleaching and sealed using clear nail polish. Microscopy was performed using an Olympus BX61 microscope equipped with a 100X phase-contrast objective and a DP70 camera (Olympus Corp., Tokyo, Japan). GFP signals were measured using wide-field excitation with an MNIBA2 fluorescence mirror unit. We resorted to our in-house software coded in Python (Python Software Foundation, Delaware) for image analysis read and explored all the pixels within the fluorescence microscope images, reading and quantifying the information from the green channel. We used the Python Imaging Library (PIL) and Matplotlib to plot the results.

Flow cytometry methods followed well-established protocols specifically adapted to P. putida (66, 72, 76, 78), analyzing at least 25,000 cells under each experimental condition. Analysis of flow cytometry data was conducted using Cytlogic 1.2.1 software (CyFlo Ltd., Turku, Finland). Histogram plots have a measure of fluorescence intensity shown on the x axis and the number of bacteria (i.e., events) counted at the specific fluorescence intensity used for RSG/GFP detection shown on the y axis.

In vitro quantitation of the Glpk activity. The Glpk assay mixture contained (in a final volume of 1.0 ml) 2.5 mM ATP, 3 mM glycerol, 0.5 mM NAD+®, 0.8 M hydrazine monohydrate buffer (pH 10.4), 125 mM glycine buffer (pH 8.5), 2 mM MgCl2, 4.5 U α-glycerol-3-P dehydrogenase from rabbit muscle (Sigma-Aldrich Co.), and 80 to 200 μl of the cell extract obtained from cells harvested during exponential growth (OD600 of ca. 0.5). Preparation of cell extracts and other analytical procedures were described in previous publications (26, 72, 76, 78).

SUPPLEMENTAL MATERIAL

Figure S2, PDF file, 0.1 MB.

Figure 15: PDF file, 0.7 MB.

Figure 61: PDF file, 0.1 MB.

Table S1, PDF file, 0.03 MB.

Table S2, PDF file, 0.03 MB.

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The authors declare that there are no conflicts of interest.

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