

Polar accumulation of pyoverdin and exit from stationary phase

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Abstract

Pyoverdin is a water-soluble metal-chelator synthesized by members of the genus *Pseudomonas* and used for the acquisition of insoluble ferric iron. Although freely diffusible in aqueous environments, preferential dissemination of pyoverdin among adjacent cells, fine-tuning of intracellular siderophore concentrations, and fitness advantages to pyoverdin-producing versus nonproducing cells, indicate control of location and release. Here, using time-lapse fluorescence microscopy to track single cells in growing microcolonies of *Pseudomonas fluorescens* SBW25, we show accumulation of pyoverdin at cell poles. Accumulation occurs on cessation of cell growth, is achieved by cross-feeding in pyoverdin-nonproducing mutants and is reversible. Moreover, accumulation coincides with localization of a fluorescent periplasmic reporter, suggesting that pyoverdin accumulation at cell poles is part of the general cellular response to starvation. Compatible with this conclusion is absence of non-accumulating phenotypes in a range of pyoverdin mutants. Analysis of the performance of pyoverdin-producing and nonproducing cells under conditions promoting polar accumulation shows an advantage to accumulation on resumption of growth after stress. Examination of pyoverdin polar accumulation in a multispecies community and in a range of laboratory and natural species of *Pseudomonas*, including *P. aeruginosa* PAO1 and *P. putida* KT2440, confirms that the phenotype is characteristic of *Pseudomonas*.

Introduction

Extracellular secreted products perform important functions in microbial populations and communities. They provide structure and protection [1], enable coordinated action [2] [3] and also allow acquisition of recalcitrant nutrients, such as polymers that are too large to be internalized [4], or are otherwise unavailable. An example of the latter is pyoverdin.

Pyoverdin is a naturally fluorescent iron-scavenging chelator (siderophore) produced by members of the genus *Pseudomonas* [5]. Iron is an essential micronutrient that exists in an insoluble state (ferric) in aerobic environments [6]. In response to intracellular iron scarcity, pyoverdin biosynthesis begins in the cytoplasm via nonribosomal peptide synthesis and undergoes maturation in the periplasm [7]. After secretion, it binds Fe³⁺ with high affinity ($K_a = 10^{32} \text{ M}^{-1}$ for PVDI produced by *P. aeruginosa* PAO1 [8]). Once bound to ferric iron the ferripyoverdin complex loses its fluorescent properties, but is recognized by a specific receptor (FpvA) and imported back into the periplasm, where iron is extracted. The pyoverdin molecule is then recycled and can undergo further cycles of export and import [9].

Because pyoverdin is a soluble extracellular product, it has been widely assumed to be equally available to all members of a community [10, 11, 12]. However, recent work shows that its distribution is subject to cell-level control. In one study pyoverdin producers retained an environment-dependent fitness advantage in conditions where invasion by nonproducers was

expected. In light of these experimental results, the possibility of personalization was raised [13]. Other studies have further supported this notion. For example, in growing microcolonies, pyoverdin diffuses primarily through adjacent cells, reducing loss into the environment [14]. Furthermore, *Pseudomonas aeruginosa* cells tune periplasmic concentrations of pyoverdin in order to protect against oxidative stress [15].

Here we use time-lapse fluorescence microscopy to study the relationship between *P. fluorescens* SBW25 cells and pyoverdin. Recognizing the importance of spatial structure and contributions therefrom to μm -scale features of microbial assemblages [16, 17, 18, 19], cells were grown on thin layers of agarose set on top of microscope slides. In actively dividing cells, naturally fluorescent apo-pyoverdin is evenly distributed in the periplasm, however, on cessation of growth we observed pyoverdin to accumulate at cell poles. This surprising discovery motivated quantitative analysis, revealing the process of accumulation to be dynamic, reversible, and ecologically relevant.

Results

P. fluorescens SBW25 (hereafter SBW25) is a model bacterium [20] known to produce pyoverdin [21] and other secreted products [22, 23]. In a previous study Zhang & Rainey [13] provided evidence of pyoverdin personalization that was inferred following contrasting outcomes of fitness assays

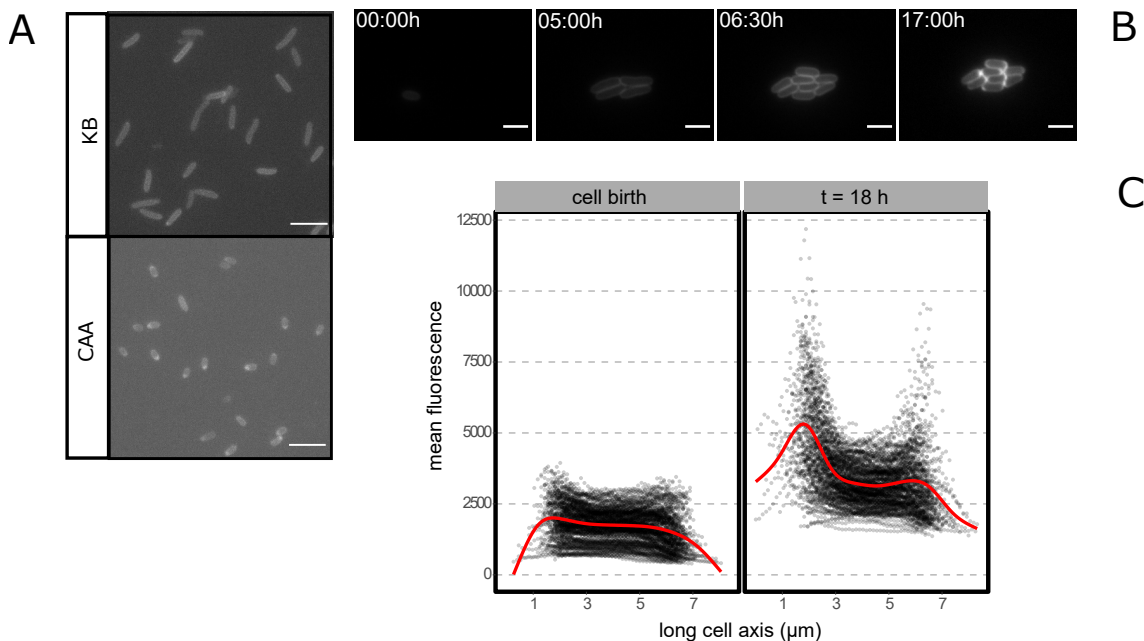


Fig. 1. Accumulation of pyoverdinin in *P. fluorescens* SBW25 at the cell pole. A) Snapshots of experiments described in [13] where fitness assays of pyoverdinin producing SBW25 and a nonproducing *pvdS* defective mutant yield contrasting results depending on the culture medium. In both cases the environment is unstructured and ancestral SBW25 producer cells are rare, inoculated at 1%. A fitness advantage to nonproducing cells in KB was previously reported, but the reverse in CAA [13]. In KB (top) pyoverdinin nonproducing cells rarely showed evidence of accumulation of pyoverdinin, whereas (bottom) this was common in CAA cultured cells where accumulation is visible at the pole. Images were obtained from 3 μ l samples of these experiments imaged under fluorescence light to visualize the distribution of pyoverdinin. All scale bars correspond to 10 μ m. B) Fluorescence time-lapse images of a growing microcolony of SBW25 in a SMM agarose pad. Images represent selected time points including, respectively: the initial inoculum, exponential growth, end of exponential growth (i.e., the final number of cells in the colony) and end of time-lapse acquisition (18 h total) C) Mean fluorescence intensity along the long axis of cells in a growing microcolony, when the last generation of cells is born (left) and at the end of acquisition ($t = 18$ h, right). Black dotted line represents the fluorescence profile of individual cells, the red line represents a smoothed mean of all the cells. Between 0 h and 18 h accumulation of pyoverdinin is evident, especially at the pole.

performed under different conditions. We reproduced these assays, but rather than examining frequencies of producers and nonproducers by plating, aliquots were observed by fluorescence microscopy. In casamino acids medium (CAA), the medium where an unexpected advantage for pyoverdinin producers had been described, cells exhibited internal pyoverdinin accumulation more evident at the pole where fluorescent foci (Fig. 1 A) appeared. This initial observation provoked further analysis.

To accurately characterize subcellular patterns of pyoverdinin, time-lapse fluorescence images of ancestral SBW25 were obtained in defined succinate minimal medium (SMM), where succinate acts both as a carbon source and a weak iron chelator. During exponential phase and early stationary phase, SBW25 cells showed a phenotype typical of fluorescent *Pseudomonas* with pyoverdinin being homogeneously distributed in the periplasm, while in late stationary phase fluorescent pyoverdinin foci appeared at the cell pole (Fig. 1 B). Superposing cell fluorescence profiles obtained by image segmentation confirmed that both polar localization and accumulation of the siderophore occurred at this later stage (Fig. 1 C).

By examining cell division throughout the time-lapse series it is possible to explore the relationship between time and population growth. The corresponding frequency of polarized cells (Fig. 2 A) was tracked by classifying segmented cells automatically as "polarized" (accumulated) or "homogeneous" (non-accumulated) using a machine learning algorithm (Supplementary Materials and Methods). After

inoculating the microscope slide, cells undergo a period of acclimation to the medium without division. We observed no accumulation events during this phase. As division begins, age of cells in microcolonies decreases until it reaches a minimum that marks exponential phase. Pyoverdinin continued to be non-polarized. Note that the small frequency of cells identified as "polarized" in the plot falls within the range of classification error. Finally, the population enters stationary phase and cells continue ageing without division. Accumulation of pyoverdinin at the pole increased within the first few hours, encompassing most cells within the population by 18h (Fig. S1). As stationary phase progresses, cells become classified as "polarized" at different rates while possessing varying amounts of intracellular pyoverdinin, suggesting that the quantity of internal pyoverdinin does not define polar localization per se (Fig S2 A). No significant difference was observed in the total amount of internal pyoverdinin between cells classified as "polarized" or "homogeneous" (Fig S2B), despite the fact that the majority of cells accumulate pyoverdinin over time (Fig 1C).

While the time-averaged state of microcolonies exposes population-level dynamics of polarization – namely, that accumulation happens in stationary phase – it obscures the behavior of individual cells. To specifically analyze single cells, cells were grouped according to division status over three generations: F0 for initial inoculum (cells where birth could not be identified but division was observed), F1 for cells in exponential phase (born from the division of F0 cells

and underwent division later) and F2 for the daughter cells of F1 that entered stationary phase and remained constant for the final hours of the experiment. Growth measurements from F1 cells, and pyoverdinin accumulation measurements from F2 cells, corroborate – despite some variability in the onset of accumulation – that accumulation is incompatible with active cell division. Cells either elongate or accumulate fluorescence (pyoverdinin) at the cell pole. This result holds for both the old and new pole (Fig. 2 B). Curiously, pyoverdinin accumulates preferentially at the new pole, but not exclusively, and sometimes distinct foci are present at both (Fig. S3)

Stationary phase marks cessation of cell division due to nutrient depletion. On an agarose pad, entry into stationary phase is highly variable, with access of cells to nutrients and oxygen depending on position and proximity to neighbouring cells. To study pyoverdinin accumulation under controlled conditions cells were exposed to two stressors that abruptly arrest cell division, but by different mechanisms: 1, an iron chelating agent (2,2'-dipyridil (DP)) and; 2, a protein synthesis-disrupting antibiotic (tetracycline)(Fig. S5 A). Both stressors induced accumulation at the pole (Fig. 2 C).

Precisely because polarization appears when cells are starved and/or stressed, processes associated with cell death, such as cell wall damage, are potential elicitors. We thus asked whether pyoverdinin accumulation was connected to cell viability. Bacteria were treated with DP, which reliably induces polarization (Fig. 2 C) and then transferred to a fresh agarose pad. Time-lapse imaging revealed that polarization is reversible and that depolarization precedes exit from lag phase (Fig. 2 D). Polarized cells consistently re-established a homogeneous pyoverdinin distribution within the first few hours after inoculation, with elongation and division resuming at a later time point (Fig. S5 B). Note that both the initial frequency of polarized cells and the time to depolarization were counted from the start of image acquisition. During the previous experimental manipulation, some individuals might have altered their polarization status. Despite this, the trend is clear, with depolarization occurring before resumption of growth. Polarization is thus not a consequence of cell death but rather a reversible, dynamic process tied to arrest of cell division.

Pyoverdinin polarization is clearly connected to the physiological state of cells, but how this is achieved is unclear. A recent study showed that the cytoplasm of *E.coli* shrinks on starvation leading to enlargement of periplasmic space at cell poles [24]. We hypothesized that this might also occur in SBW25 cells, with the space created by shrinkage allowing accumulation of pyoverdinin. To this end, a red fluorescent protein (mScarlet) was translationally fused to the peptide signal from *dsbA* thus targeting mScarlet to the periplasm [24]. Having confirmed periplasmic localization, cells were grown on agarose pads and the distribution of mScarlet was tracked by microscopy. At inoculation, mScarlet was homogeneously distributed as observed for pyoverdinin, however, on entry into stationary phase, the fluorescent marker accumulated at the poles of the cells (Fig. 3 A). As with pyoverdinin, mScarlet accumulation was biased toward new poles (Fig. 3 B). As expected, unlike pyoverdinin, mScarlet showed no evidence of secretion to the external medium (Fig. S8)

While overlap in patterns of pyoverdinin and periplasm-targeted mScarlet accumulation points towards the fact that pyoverdinin polarization is coupled to general cellular phenomena, possibilities remain for the involvement of genetic determinants of pyoverdinin biosynthesis, regulation and / or

transport. To test this hypothesis, patterns of pyoverdinin accumulation were monitored in a range of mutants.

First, polar accumulation of pyoverdinin is not a consequence of defective periplasmic maturation [25]: SBW25 producer and nonproducer types (the latter with an mCherry fluorescent marker) were co-cultured. Pyoverdinin was localized in nonproducing cells (Fig. 3.1) demonstrating that accumulated pyoverdinin is functional, since any siderophore internalized by Pvd⁻ necessarily comes from the external medium after export by SBW25. Polarization thus involves pyoverdinin that is actively used among cells of the population.

Next, recognizing that mature pyoverdinin interacts with membrane proteins via OpmQ [26, 27], *opmQ* was deleted and the pattern of pyoverdinin accumulation determined. SBW25 $\Delta opmQ$ showed no change in capacity to accumulate pyoverdinin at cell poles (Fig. 3.2), however, the mutant showed increased intracellular levels of pyoverdinin; secretion was not abolished [28](Fig. S6 A).

Pyoverdinin synthesis is subject to positive feedback control via the pyoverdinin receptor FpvA making this protein a candidate for involvement in pyoverdinin accumulation [7]. Production of pyoverdinin by SBW25 $\Delta fpvA$ was significantly reduced compared to the ancestral type (Fig. S6 B), however, where pyoverdinin was visible (thus accumulated), it was typically polarized as in ancestral SBW25 (Fig. 3.3).

While the genes involved in extraction of ferric iron and translocation into the cytoplasm have not been characterized in *P. fluorescens*, a recent study in *P. aeruginosa* offered potential candidates for polarization including a siderophore-binding periplasmic protein [9]. Deletion of the homologs of these genes in SBW25 PFLU_2048, PFLU_2041, and PFLU_2043 did not disrupt accumulation of pyoverdinin (Fig. S7). Thus, neither aggregation of pyoverdinin at trafficking points across the periplasm, nor accumulation via the ferripyoverdinin-complex recycling, are involved in polarization.

Additional possibilities for polar accumulation are via connections to extracellular polymer synthesis and cell morphology. SBW25 secretes cellulose when constructing bacterial mats at the air-liquid interface [23, 29] which could trap pyoverdinin at the perimeter of cells. However, pyoverdinin localization is not altered in SBW25 $\Delta wsp \Delta awa \Delta mws$ [30] that is unable to produce cellulose (Fig. 3.4). Finally, the rod-cell-shape characteristic of *Pseudomonas* was considered a possible contributory factor. To test this, pyoverdinin accumulation was observed in a spherical $\Delta mreB$ mutant [31]. Despite the aberrant cell shape, this mutant displays fluorescence foci of pyoverdinin after extended culture (Fig. 3.5). Curiously, pyoverdinin accumulation at the pole is also independent of cell size in the ancestral genotype (Fig S9).

Polar accumulation of pyoverdinin is linked to cell physiology, but is unperturbed by known genetic determinants of pyoverdinin biosynthesis, regulation, or transport. This leads to the possibility that pyoverdinin accumulation under conditions of cellular stress is an accidental consequence of the biophysics of cell biology and lacks ecological relevance. The alternate hypothesis is one of ecological relevance: a plausible explanation being that release of accumulated pyoverdinin allows cells to reduce the time required to exit stationary phase.

An ideal test would be an experiment in which time to resumption of growth of an ancestral type that accumulates pyoverdinin is compared to a mutant unable to accumulate pyoverdinin, under conditions where pyoverdinin accumulation both occurs, and doesn't occur. If accumulation of pyoverdinin confers a fitness advantage, then ancestral types are expected to

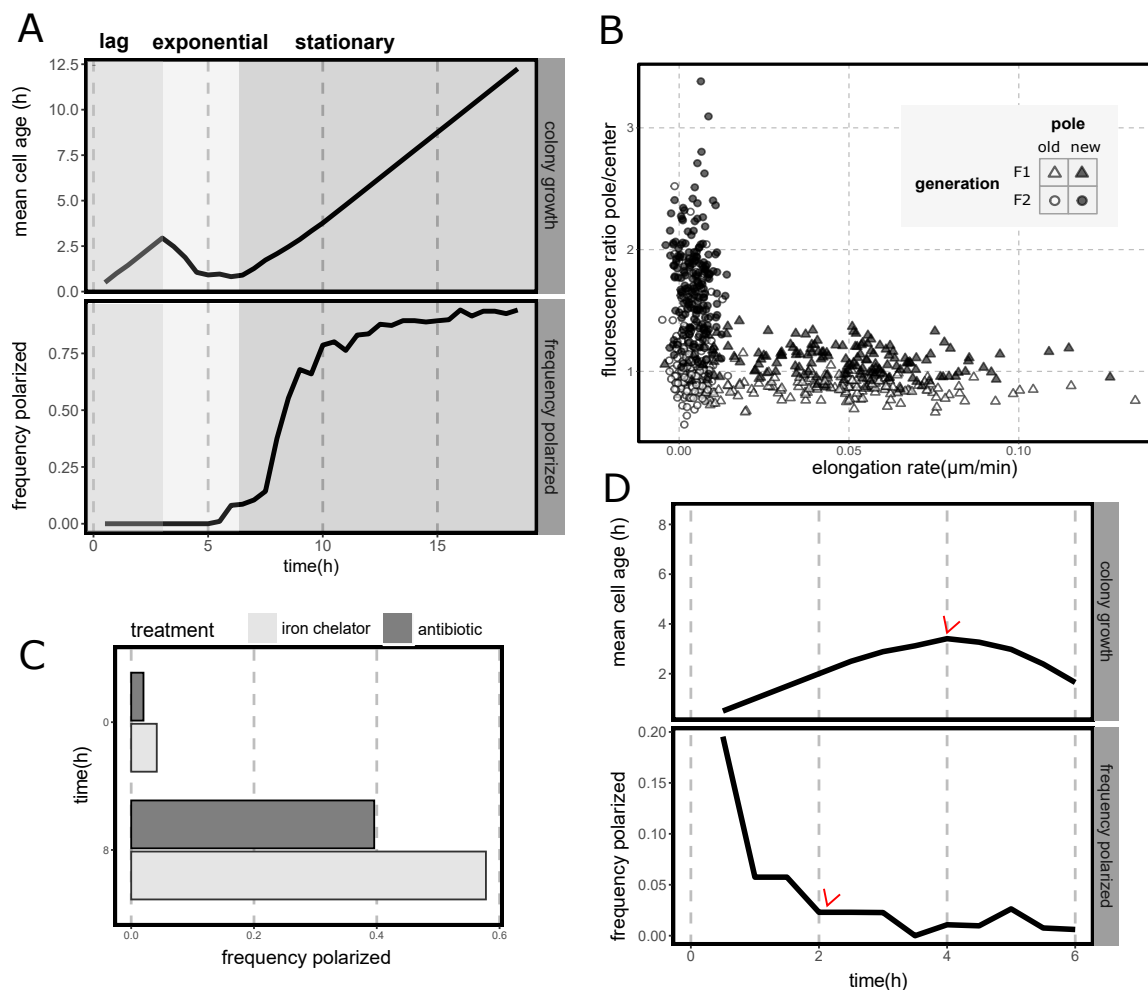


Fig. 2. Polar accumulation is a reversible phenotype associated with arrest of cell division. A) Polarization in different growth stages of a microcolony. Mean age of the cells in a growing microcolony of SBW25 (black line, top) and the corresponding frequency of polarized cells for each time point (black line, bottom). Colored panels represent the growth stages of the microcolony, from left to right: lag phase (generation F0), exponential phase (generation F1), and stationary phase (generation F2). $N = 35, 159, 187$ respectively. In all cases data has been filtered to exclude cells with segmentation errors or other artefacts that preclude proper analysis. Note that as microcolonies begin to form in exponential phase and cells are no longer isolated, overlap between adjacent cells creates regions of high fluorescence that could lead to classification errors. Nevertheless, visual inspection reveals that cells remain in a homogeneous state during exponential growth, with polarization onset being clearly associated to stationary phase. To maintain a consistent physiological response, in this study cells are manipulated from a starting point corresponding to early stationary phase (Fig. S4). B) Elongation rate and accumulation of fluorescence at the cell pole of individual cells in a growing microcolony. Data extracted from A). Markers represent individual cells in different growth phases of the colony (F1, exponential, triangle markers; F2, stationary, circle markers) and the old (white markers) and new (black markers) cell pole. Elongation rate is represented by the average over the lifetime of a cell. Accumulation of fluorescence at the pole is represented by the maximum ratio over the lifetime of a cell of the sum of the pixels in the pole region and central region of a cell. These regions are defined by segmenting the cell and dividing it in 3 portions over the long axis, where the external 1/3 represent each pole and the remaining 1/3 represents the center. C) Polarization in response to chemical stresses related and unrelated to iron metabolism. Bars represent the frequency of polarized cells at the start of treatment and after 8h of treatment with either 100 $\mu\text{g}/\text{ml}$ 2,2'-dipyridil (DP) (light bars) or 5 $\mu\text{g}/\text{ml}$ tetracycline (dark bars). No cell division was observed during treatment. $N = 99$ ($t=0\text{h}$), 94 ($t = 8\text{h}$) and $N = 94$ ($t = 0\text{h}$), 79 ($t = 8\text{h}$) for DP and tetracycline treatments respectively. D) Depolarization and subsequent growth of cells pre-treated with an iron chelator. Plot represents colony growth and polarization as in A). Cells were treated with 100 $\mu\text{g}/\text{ml}$ DP during 4h, washed and inoculated on a fresh SMM agarose pad. Data corresponds to five technical replicates i.e., five positions on the agarose pad. Red arrows indicate the time point where the colony overall starts growing (top) and where the majority of the cells are depolarized (bottom). Total initial number of cells $N=87$.

resume growth more rapidly than non-accumulating mutants, but only after cells have first experienced conditions that promote accumulation of pyoverdinin.

Unfortunately, as evident above, no such pyoverdinin-accumulating mutant exists, nonetheless, progress is possible via an experiment that exploits the fact that a PvdS mutant cannot accumulate pyoverdinin because it is unable to produce it. To this end, SBW25 and a PvdS mutant (SBW25

pvdSG229A(D77N) [13] (hereafter termed Pvd⁻) were grown separately in SMM for sufficient time to ensure cells entered stationary phase (24 h). A sample of cells from both populations was then transferred to independent SMM agarose pads and the time to first cell division determined. While a slight delay in median time for resumption of growth was observed in Pvd⁻ compared to SBW25 the distribution of data points shows significant overlap. The findings were not affected by transfer of

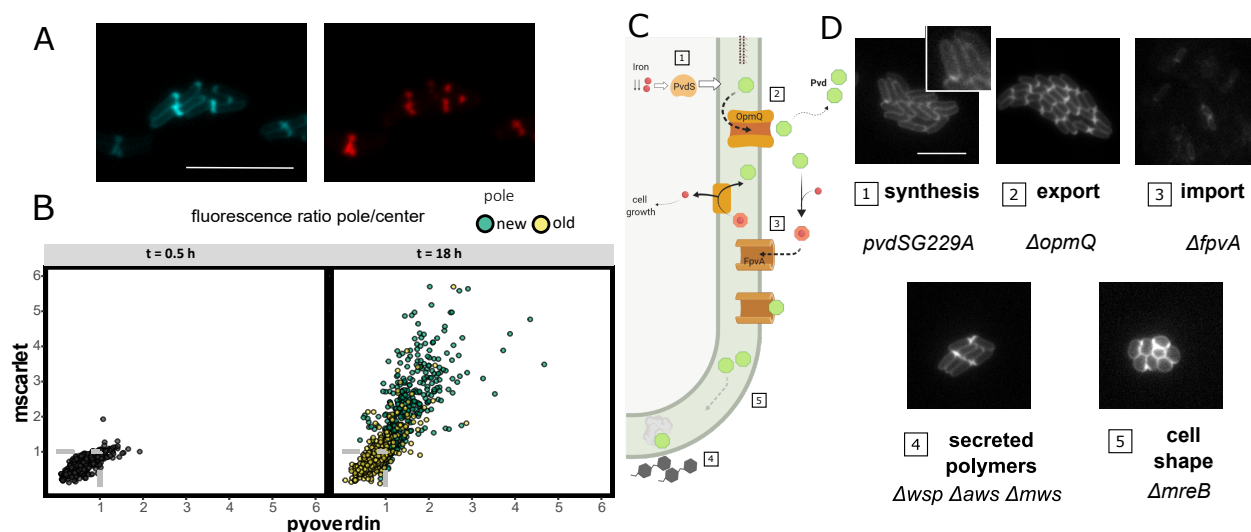


Fig. 3. Untangling the mechanism of polarization. A) A recent study [24] showed that upon starvation the cytoplasm of *E. coli* cells shrinks creating extra space in the periplasm at one of the cell poles. To check if a similar phenomenon happens in *P. fluorescens* SBW25 we expressed a red fluorescent mScarlet protein fused to the localization signal peptide of dsbA, a periplasmic protein. The image obtained at $t = 18 \text{ h}$ shows co-localization of mScarlet (bottom, red) and pyoverdinin fluorescence (top, blue, scale bar represents 10 μm). B) Quantitative analysis of time-resolved images at 30 min and 18 h. Cells were segmented and intensity of fluorescence was measured along the long axis. Fluorescence at the poles was compared with the signal produced at the center of each cell; this was done for both pyoverdinin (x-axis) and mScarlet (y-axis). Dots represent the intensity ratio for both fluorescent molecules. For measurements at 18 h the values for both the new pole (green dots) and old pole (yellow dots) are represented. Polar information is not available for freshly inoculated cells. The gray dashed line highlights the values where the center of the cell and the poles are not substantially different, i.e., their fluorescence ratio is close to 1. Deviation from this value indicates polar accumulation of the molecules. C) Cartoon depicting a simplified version of the pyoverdinin pathway in which candidate genes for polarization are shown. 1. Pyoverdinin (Pvd) synthesis starts in the periplasm in response to iron scarcity mediated by the transcription factor PvdS. Pyoverdinin is then secreted to the bacterial periplasm, where it matures and becomes fluorescent. 2. Periplasmic pyoverdinin is exported into the external medium by a complex that includes the transporter OpmQ. There, it chelates insoluble iron (Fe^{3+}). 3. Ferripyoverdinin complexes (no longer fluorescent) are then imported back into the periplasm after binding to the receptor FpvA. This receptor is known to also bind free pyoverdinin [7]. In the periplasm, iron is extracted and pyoverdinin is again recycled into the external medium by OpmQ. Polarization might also be determined by genes unrelated to the pyoverdinin pathway: 4. SBW25 is known to secrete polymers such as cellulose that might trap pyoverdinin [23]. 5. Pyoverdinin could accumulate at the cell poles due to the rod shape of SBW25. D) Mutants associated to the main processes depicted in A) and their phenotype with regards to pyoverdinin polarization. Mutants were grown on an agarose pad as described and fluorescence images displaying pyoverdinin were taken at the time points indicated in the photo. The pyoverdinin nonproducing mutant *pvdSG229A(D77N)* was co-inoculated with SBW25 to enable access of the mutant to pyoverdinin. Mutants were tagged with a red fluorescent protein, one mutant colony is displayed in the image.

cells from overnight SMM culture to agarose pads additionally supplemented with excess iron (Fig. 4A, left). This indicates that even the Pvd^- mutant was not deficient in iron despite having entered stationary phase.

Results were significantly different when the 24 h pre-culture period in SMM included a final 4 h period during which populations of both SBW25 and Pvd^- experienced iron-stress arising from supplementation of SMM with the iron chelating agent DP. As shown above, a 4 h treatment with DP not only caused cessation of growth, but also triggered polarization and accumulation of pyoverdinin. The effect of this treatment was to greatly extend the time to resumption of growth in the non-pyoverdinin-accumulating mutant. This difference was eliminated when cells, having experienced iron-deprivation, were provided with excess iron (Fig. 4A, right). These results demonstrate a fitness advantage to accumulation of pyoverdinin that stems from a reduction in time to exit stationary phase.

This conclusion is further supported by the results of an additional experiment in which SBW25 and the Pvd^- mutant were grown as previously, treated with DP and then mixed in a 1:1 ratio and then transferred to a fresh SMM agarose pad. Measurement of the time to resume growth showed little difference among the competing genotypes (Fig. 4B). This is consistent with the expectation that SBW25

releases accumulated pyoverdinin into the (fresh) medium to the benefit of both producer and non-producer alike. Notable though is the fact that producer cells reap greatest benefit. When nonproducers are instead inoculated in the presence of rare ancestor cells ($\sim 1\%$) their lag time again increases significantly, supporting the conclusion that newly released pyoverdinin underpins the faster division of producers (Fig. S10). Furthermore, the amount of pyoverdinin accumulated during starvation seems to be sufficient to support at least twice as many cells, as the lag time of SBW25 producer cells was not affected by the presence of nonproducers.

We further tested pyoverdinin polarization in conditions closer to the natural milieu of SBW25, where species interdependencies are common and resources invariably limiting. To implement a minimal bacterial community, SBW25 was co-cultured with a cellulose-degrading *Bacillus* isolated from a compost heap [32]. Because SBW25 is unable to degrade this polymer, it must rely on the *Bacillus* species to obtain carbon for growth. These two strains were grown together in minimal medium with cellulose paper as the sole carbon source and periodically imaged to assess polarization status. Pyoverdinin was readily observed to be polarized (Fig. 5 A). Pyoverdinin accumulation at the cell pole is thus likely to be part of the natural phenotypic repertoire of SBW25, and given that in

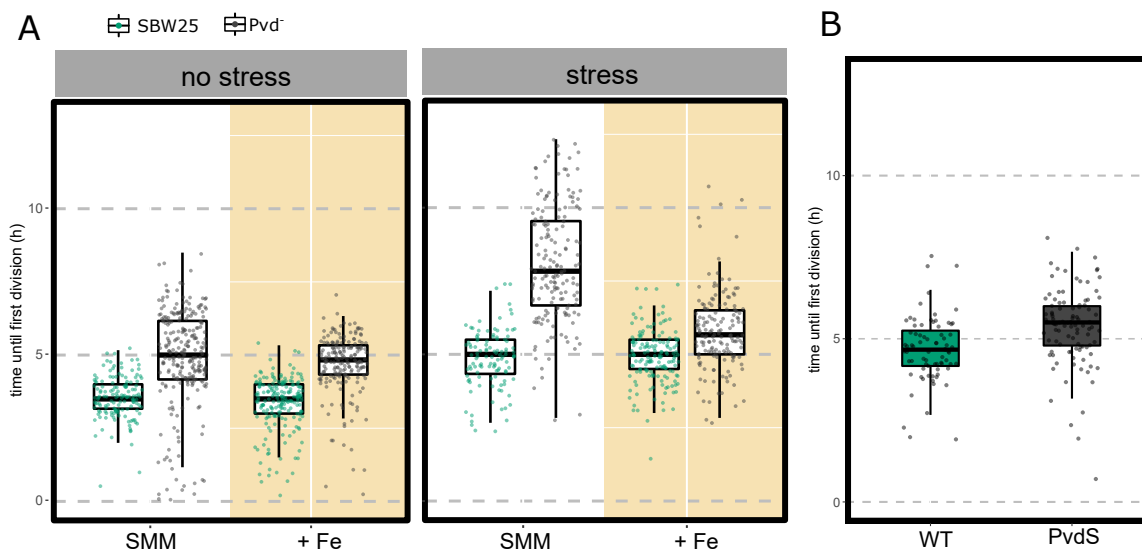


Fig. 4. Pyoverdinin accumulation facilitates recovery of growth after stress. A) Time until first division (lag time) of SBW25 (green) and pyoverdinin defective mutant *pvdS229*(D77N), termed Pvd⁻ (gray) under different treatments and conditions. Prior to inoculation cells were either grown in the usual culture medium SMM (no stress on left panel) or treated with DP for 4h (stress on right panel). Cells were then inoculated on a fresh agarose pad, supplemented with 0.45 mM Fe₂[SO₄]₃; these are labeled "+ Fe" (yellow background); unsupplemented SMM treatments are labeled "SMM" (white background). Dots represent individual cell values, box plots represent the associated distribution (median, 25th and 75th percentiles) N = 145, 261, 199, 193, 111, 168, 156, 156, respectively, from left to right. B) Time until first division of SBW25 (green) and Pvd⁻ mutants co-inoculated in a fresh agarose pad after separate stress treatment (4h in DP). Pvd⁻ mutants were labeled with a red fluorescent protein to allow identification of individual cells. Dots and box plots as in A). N = 175 total cells.

natural environments bacteria likely find themselves in or close to stationary phase most of the time, polar accumulation of pyoverdinin stands to be the normal state.

Since pyoverdins are produced by many species of the genus *Pseudomonas*, investigating subcellular distribution in related strains may provide an evolutionary context for polar accumulation. We selected 11 strains belonging to the genus *Pseudomonas*, comprising both common laboratory strains (such as *P. aeruginosa* PAO1, in which pyoverdinin is usually studied) and natural isolates from different European locations. The ability of these strains to polarize pyoverdinin was classified qualitatively, that is, presence or absence of polarized cells (Fig. 5 B.) All tested strains localized pyoverdinin at the cell poles in conditions similar to those described here for SBW25. Interestingly, different strains polarized pyoverdinin with treatments of varying stringency. In some cases extended culture in SMM was enough to observe the phenotype (e.g., *P. putida* KT2440). In others, amendment with an iron chelator was necessary to observe the effect (e.g., *P. aureofaciens* U149), at the same dosage that induced polarization in SBW25, and in the specific case of *P. aeruginosa* PAO1, very high doses of DP were required. This range of responses could reflect the secretion of secondary siderophores by some strains [34] or differences in the regulation of pyoverdinin production [12], and suggest that polarization is an ecologically relevant trait that varies depending on the evolutionary history of the lineage.

Discussion

Previous work showing that the population-level distribution of pyoverdinin changes depending on nutrient status [13], contact with neighboring cells [14], and environmental stress [15] motivated our investigation. With focus on *P.*

fluorescens SBW25, and using time-resolved microscopy, we have shown that pyoverdinin transiently accumulates within cells (preferentially at cell poles), that localization is a reversible process associated with arrest of cell division, and is affected by factors such as entry into stationary phase and deprivation of specific nutrients (Fig. 2). Particularly significant is demonstration that accumulation of pyoverdinin has ecological relevance (Fig. 4).

Investigations at the level of individual cells allow cell-level behaviors to be linked to the dynamics of populations [17, 18]. For example, Gore et al (2009) [35] showed that the cellular location of invertase responsible for degradation of sucrose in *Saccharomyces cerevisiae* creates diffusion gradients of degradation products that deliver benefit to producing cells, despite costs to producers of synthesizing the enzyme [35]. Similarly, the siderophore enterochelin (a catecholate secreted by *E. coli* and other *Enterobacteriaceae*) can remain associated with the outer membrane under conditions of low cell density, delivering preferential benefit to enterochelin-producing cells [36]. Some localized products display dynamic behaviour [37], with, in some instances, proteins oscillating between poles [38]. In other instances, proteins accumulate via formation of nuclear occlusions in the cytoplasm [39]. In the context of pyoverdinin, complexes termed "siderosomes" localize enzymes for synthesis of pyoverdinin at old cell poles during exponential growth by association of the cytoplasmic membrane with L-ornithine N5-oxygenase (encoded by *pvdA*) [40, 41].

While pyoverdinin in SBW25 was most often observed at new cell poles, it was also evident at old poles; on occasion it was found at both poles. Moreover, although all cells accumulate pyoverdinin (Fig 1C), a minority show no evidence of polar localization. Measurement of the amount of pyoverdinin in cells that localize the iron chelator showed that it is the same as that

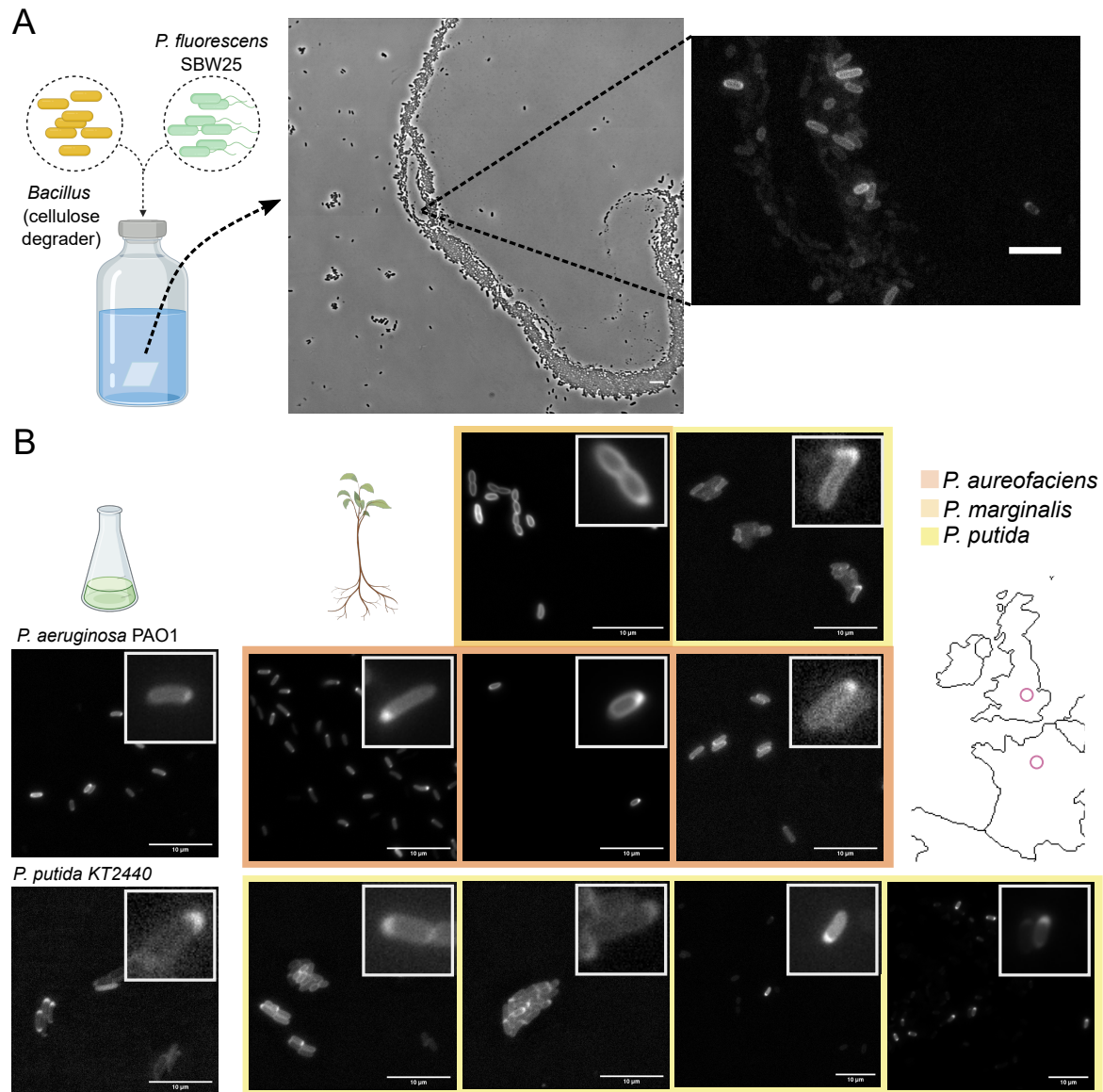


Fig. 5. Accumulation visible by polarization is evident in SBW25 in a multispecies community and is a common phenotype in related species of the genus *Pseudomonas*. A) Polarized SBW25 cells in a multispecies community with interdependencies. SBW25 and a cellulose-degrading *Bacillus* strain isolated from a compost heap in Paris, France [32] were co-cultured in glass vials with minimal medium and cellulose paper as the only carbon source. The community was periodically sampled to assess the polarization state of SBW25. A representative image obtained after 28 days of growth is displayed, where both strains are visible (left, phase contrast image) and in a magnified region where pyoverdinin distribution in SBW25 is visualized (right, fluorescence image). B) Qualitative accumulation assessment in species of *Pseudomonas* other than SBW25. A collection of laboratory (left) and natural (right) strains were tested for accumulation as evident by polarization. Accumulation by polarization in commonly used laboratory strains: *P. aeruginosa* PAO1 tested in SMM agarose pad with 1000 $\mu\text{g/ml}$ DP for 7:30 h; *P. putida* KT2440 tested in SMM agarose pad for 24h. Polarization test in natural isolates, left to right and top to bottom: U106 (liquid KB medium for 24 h); U177 (SMM agarose pad for 24 h); U149 (liquid SMM with DP 100 $\mu\text{g/ml}$ for 24 h); U180 (liquid SMM with DP 100 $\mu\text{g/ml}$ for 24 h); U181 (SMM agarose pad for 16 h); T24 V1a (SMM agarose pad for 24 h); T24 V9b (SMM agarose pad for 24 h); T24 H1b (SMM agarose pad for 24 h); T24 H9b (liquid SMM for 24 h). Natural isolates were collected in Oxford, UK [33] (first and second row) and in Paris, France [32] (Bottom row) (locations are roughly marked with a pink circle on the map). Color represents the identified *Pseudomonas* species. Insets highlight cells with clearly accumulated pyoverdinin.

contained within cells displaying a homogeneous distribution of pyoverdinin (Fig S2).

Taken together, ambiguity surrounding accumulation and polar localization suggested a general connection to biophysical aspects of cell biology. Interestingly, a recent study of *E. coli* under starvation conditions showed polar accumulation of fluorescent markers, including mCherry [24]. Accumulation was connected to shrinkage of the cytoplasm, with shrinkage

creating an expansion of space in the polar region of the periplasm. It thus seemed plausible that accumulation of pyoverdinin in SBW25 in stationary phase may be a consequence of nonspecific cellular behavior under growth arrest. Such a possibility was bolstered by the fact that a PvdS mutant deficient in synthesis of pyoverdinin (and thus lacking siderosomes) nonetheless accumulated pyoverdinin, that deletion of genetic determinants of uptake and recycling had

no observable effects on accumulation, and that pyoverdinin accumulation also occurred at discrete locations in spherical $\Delta mreB$ cells (Fig. 3D.5).

To investigate the hypothesis that pyoverdinin accumulation is connected to cytoplasmic shrinkage, red fluorescent mScarlet was translationally fused to the localization signal peptide of DsbA. As cells entered stationary phase accumulation of the fluorescent marker protein was observed at cell poles in a manner analogous to that observed for pyoverdinin. In fact, single images captured using filters specific for the reporter protein and pyoverdinin showed an almost perfect overlap (3A,B). This lead us to conclude that polar accumulation of pyoverdinin is part of a general cellular response to starvation. Thus observations made in *E. coli* with a chimeric reporter [24] appear to hold for SBW25, but with our observations connecting cytoplasmic shrinkage to accumulation of a biologically relevant molecule. Additionally we demonstrate that accumulation delivers beneficial effects on fitness.

Data in Fig. 4A show that cells that have accumulated pyoverdinin – most of which is localized to cell poles – resume growth more rapidly compared to cells that have not accumulated pyoverdinin. A time-to-first cell division advantage was not evident when growth-arrested cells were transferred to iron-replete conditions. This demonstrates that liberation of the stock of pyoverdinin accumulated during growth cessation speeds the time to growth resumption after growth arrest, presumably through provision of available iron to growing cells 4B.

Of additional interest is evidence from cell-level observations that rare pyoverdinin producers are largely unaffected by the presence of Pvd⁻ mutants (Fig. S10). Such mutants have often been referred to as “cheats” and thus expected to negatively impact the fitness of pyoverdinin producers (so called “cooperators”). Lack of detrimental impact is further evidence that pyoverdinin production preferentially benefits producer cells. This finding further calls into question the fit between social evolution theory and production of extracellular products by microbes [13, 12]. It also gels with work showing that populations of siderophore-producing *Pseudomonas* can be grown in the presence of an excess of nonproducing mutants without significant reduction in overall yield [42].

Recent work suggests important physiological differences between acute (and unexpected) interruptions of cell division, and gradual growth arrest that determines entry into stationary phase [?]. In the former case, sudden arrest induces a disrupted state that results in large variability in cell-level duration of lag phase. In the latter case, cells implement genetic programs that buffer the effects of impending starvation creating cohesive population-level responses [?]. The seemingly adaptive nature of pyoverdinin accumulation under stress is reminiscent of – and perhaps even connected to – the capacity of SBW25 to enter a semi-quiescent capsulated state upon starvation. During starvation, SBW25 cells produce an excess of ribosomes that allow rapid exit from stationary phase once growth-permissive conditions are encountered. Cells unable to provision ribosomes are quickly out-competed by those that do [43]. It is possible that localization of pyoverdinin, followed by fast release under growth permissive conditions, has evolved as a strategy precisely because it maximizes competitive performance upon resumption of growth. Rapid resumption of growth is likely to deliver significant fitness benefits in environments punctuated by periods of nutrient abundance and scarcity.

Mounting evidence indicates that pyoverdinin – in its many variant forms [34] – has multiple impacts on composition and function of microbial communities [44, 45, 46, 47, 48,

49]. The fact that pyoverdinin is accumulated and localized under growth limiting conditions in a diverse range of *Pseudomonas*, combined with evidence of the same behavior in SBW25 cells grown for weeks under nutrient restrictive conditions with reliance on cellulose degrading *Bacillus* (Fig. 5), suggests that polarization has relevance to conditions beyond those experienced by cells in standard, nutritionally-rich, exponential-phase, laboratory culture.

Given that pyoverdinin accumulation appears connected to cytoplasmic shrinkage, it seems reasonable to assume that other molecules may similarly accumulate in the periplasm in response to division arrest signals, which both change size and shape of cellular compartments. Indeed, accumulation of products related to nutrient stress is one possible function of the bacterial periplasm, with roles analogous to vacuoles in eukaryotes [50].

Materials and methods

Strains, culture media, and reagents

Ancestral *Pseudomonas fluorescens* SBW25 originally isolated on beet roots at the University of Oxford farm (Wytham, Oxford, U.K.) [20] and a collection of relevant mutants were used: the pyoverdinin nonproducer *pvdS*G229A(D77N) named Pvd⁻ in the main text (construction described in [13]), the corresponding strain with mCherry fluorescence tagging under IPTG induction, $\Delta mreB$, PBR716 ($\Delta wsp\Delta aws\Delta mws$, described in [30]). Pyoverdinin import and export defective mutants $\Delta PFLU3979$ (OpmQ) and $\Delta PFLU2545$ (FpvA) were created by two-step allelic exchange [51]. The periplasm reporter strain was created by fusing the red fluorescent protein mScarlet to the signal peptide of periplasmic protein, DsbA (see Supplementary Methods). A neutrally marked SBW25 strain [52] was used to replicate the fitness assays from [13]. *Escherichia coli* DH5 α λ_{pir} and pRK2013 were used for cloning. *Bacillus* 002.IH from Steven Quistad’s compost heap collection [32] was used for the community experiment. For the phylogenetic comparison, an assortment of species of the genus *Pseudomonas* were selected from the lab collection. Most experiments were performed in succinate minimal medium (SMM, described in [14]). Overnight cultures were done in Luria-Bertani (LB) broth. Replication of fitness assays was performed in CAA and KB media as described in the original work [13]. Community experiment was done in M9 minimal medium with cellulose as the only carbon source (Whatman). Where indicated media was supplemented with 2,2'-dipyridil (Sigma), tetracycline (Duchefa, France), Fe₂[SO₄]₃(III) (Sigma), IPTG (Melford). For strain construction tetracycline, nitrofurantoin (Sigma), X-gal (Melford), D-cycloserine (Duchefa) were used.

Agarose pad

To prepare the agarose pad, 220 μ L of agarose (Melford) dissolved in SMM (2% w/v) were poured onto a microscope slide fitted with a sticky frame (Gene Frame, Fisher Scientific), pressed with a clean slide and allowed to dry for \sim 2 min. A small \sim 3 mm section of the pad and frame was cut across the slide to provide air for the growing cells. 1.5 μ L of washed culture was inoculated on the agarose pad and sealed with a coverslip.

Microscopy

Inoculated agarose pads were monitored by taking snapshots every 30 min for a typical total time of 18h using the microscope Axio Observer.Z1 (Zeiss, Germany). Cells were imaged under phase contrast (exposure: 100 ms) and fluorescence corresponding to Pvd using the fluorescence Source X-Cite 120 LED and the following filters: 390/40 BrightLine HC, Beamsplitter T 425 LPXR, 475/50 BrightLine HC (exposure: 30 ms, 12 % intensity). Images were taken with 63x and optovar 1.6x magnification.

Image analysis

Image processing was performed using the image analysis software Image J. Segmentation and analysis was performed using the package for Matlab SuperSegger from the Wiggins Lab [53]. Further analysis and data visualization was carried out with the programming software R. After segmentation cells were sorted by polarization status using a classifier obtained with the Statistics and Machine Learning Toolbox for Matlab (More information in Supplementary Methods).

Community experiment

SBW25 and *Bacillus* 002.IH (from [32]) were grown on 20 ml M9 minimal medium and cellulose as the only carbon source (1 cm x 1 cm cellulose paper). After overnight growth cultures were washed and co-inoculated into 20 ml of M9 medium with cellulose paper in a 60 ml vial. These vials were incubated at room temperature with unscrewed caps to allow air exchange, and periodically sampled for imaging.

Phylogenetic comparison

A collection of laboratory and natural strains belonging to the genus *Pseudomonas* were subjected to a binary qualitative polarization test, i.e. the test was considered positive if polarized cells were observed under fluorescence microscopy in conditions similar to SBW25 but no dynamics were assessed. Commonly used laboratory strains *P. aeruginosa* PAO1 and *P. putida* KT2440 were tested. Natural isolates belong to collections from two different locations. Paris, France [32]: T24 V1a, T24 V9b, T24 H1b, T24H9b; all classified as *P. putida*. Oxford, UK [33]: *P. marginalis* U106, *P. putida* U177, *P. aureofaciens* U149, U180, and U181.

Author contributions

PBR, CMF and MA designed research. CMF performed research. CMF and MA analyzed data. PBR, CMF and MA wrote the paper.

Conflicts of interest

The authors declare no competing interests.

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Supplementary Appendix

Supplementary Methods: Image processing and phenotypic classification of cells

Images were pre-processed for segmentation using the image analysis software ImageJ [54]. Time-lapses of growing microcolonies were segmented using the SuperSegger Toolbox [53] for Matlab [55]. This toolbox has been designed specifically for the analysis of bacterial microcolonies imaged under phase contrast and fluorescence microscopy and allows the user to adapt segmentation variables to suit a particular experimental system using neural network training. Individual images of cells and their lifetime properties (including time of birth and death, lineage, polar identity, and others) are the output of the software. This enabled tracking of cells throughout three generations mentioned in the main text: F0 or inoculum cells, F1 or exponentially growing cells resulting from the division of F0, and F2 cells or stationary phase cells resulting from the division of F1. Pyoverdine polarization is thus observed in F2 cells. To obtain a means to automatically classify cells by their pyoverdine polarization status the Statistics and Machine Learning Toolbox for Matlab was used [56]. 646 segmented cell images were collected as the training set, originating from 4 different experiments. The training set included images obtained from microcolonies growing on an agarose pad and from cells treated with an iron chelator in liquid medium, to account for differences between experimental setups that could impact on image analysis (e.g., overlapping cells in microcolonies, changing levels of relative cell fluorescence in different environments, etc.) These cells were visually classified into "polarized" and "homogeneous". Next, image features were extracted to develop a classification algorithm. To extract unidimensional features from 2-dimensional images, these were processed as follows: first, mean fluorescence along the long cell axis was obtained (also named kymograph). Cells were then divided into three equally sized regions representing the center of the cell and both cell poles and different variables were obtained comparing these regions. Many variables were tested and optimum results were obtained when using the following procedure: kymographs were normalized by either subtracting or dividing by the median fluorescence of the cell center. From these, standard deviation, and 75th percentile of either polar region were calculated. These 6 features were fed into the Classification Learner App in Matlab with 5-fold cross-validation. The model with best results was a linear SVM (Support Vector Machine) model, with accuracy >90%. It is estimated that the model has a false negative rate of ~5% for "homogeneous" cells and ~10% for "polarized" cells i.e. it has a slight bias towards under-identification of polarization which we found more desirable than over-identification. This model was used for all subsequent classification mentioned in the main text and SI. To further analyze and visualize data the programming software R was used [57].

Supplementary Methods: Construction of strain with periplasmic red fluorescent marker

The IPTG-inducible periplasmic mScarlet reporter was created by first introducing the mScarlet-I CDS from plasmid pMRE-Tn7-155 (amplified using primers mScarlet_fwd and mScarlet_rev) into plasmid pUC18-mini-Tn7T-LAC (linearized

using primers Tn7LAC_fwd and Tn7LA_rev) using NEBuilder HIFI DNA Assembly Master Mix (New England BioLabs)[58, 59]. The DsbA signal sequence from SBW25 (the first 81 nucleotides of PFLU0083) was then introduced to the start of mScarlet-I in the resulting plasmid by FastCloning [60] using primers dsbAmScarlet_fwd and dsbAmScarlet_rev. The forward primer included the LEGPAGL amino acid linker used by Uehara *et al* (2009) [61] to construct dsbA_{ss}-mCherry in *E. coli*. This final plasmid was introduced into SBW25 by electroporation (mediated by the transposition helper plasmid pUX-BF13 [62]) and integration of the transposable element containing Plac-dsbA_{ss}-mScarlet-I into the attTn7 site downstream of *glmS* was selected by plating to gentamicin. Correct integration of the Tn7 cassette was confirmed by Sanger sequencing. All cloning was performed in *E. coli* OneShot TOP10 chemically competent cells (ThermoFisher Scientific) and PCRs were conducted using either Phusion High-Fidelity PCR Master Mix (ThermoFisher Scientific) or Q5 High-Fidelity 2X Master Mix (New England BioLabs). pMRE-Tn7-155 was a gift from Mitja Remus-Emsermann (Addgene plasmid # 118569 ; <http://n2t.net/addgene:118569>; RRID:Addgene_118569). pUC18-mini-Tn7T-LAC was a gift from Herbert Schweizer (Addgene plasmid # 64965 ; <http://n2t.net/addgene:64965>; RRID:Addgene_64965).

Primers (overlaps for assembly or primer extensions in lower case, annealing region in uppercase, dsbAss in italics, LEGPAGL linker in bold)

Table 1.

Primer	sequence
mScarlet_fwd	cggataacaaGTGAGCAAGGGCGAGGCA
mScarlet_rev	ctgtgtgaaaTTACTTACTTGTACAGCTCGTCCATGC
Tn7LAC_fwd	aagtaagtaaTTTACACAGGAAACAGAATTC
Tn7LAC_rev	cctgctcacTTGTTATCCGCTCACAAATTC
dsbAmScarlet_fwd	<i>ctgccagcctctcgggcatgaccgcacaagctgccgacgtgccgttctcggagggtccggctggtctg</i> atgGTGAGCAAGGGCGAGGC
dsbAmScarlet_rev	<i>ccgaagaggctgagcagtcgagagcggcctgaggatcagattaccgatggtctgttctctgtgtgaaa</i> TTGTTATCCGCTCACAAATTCACAC

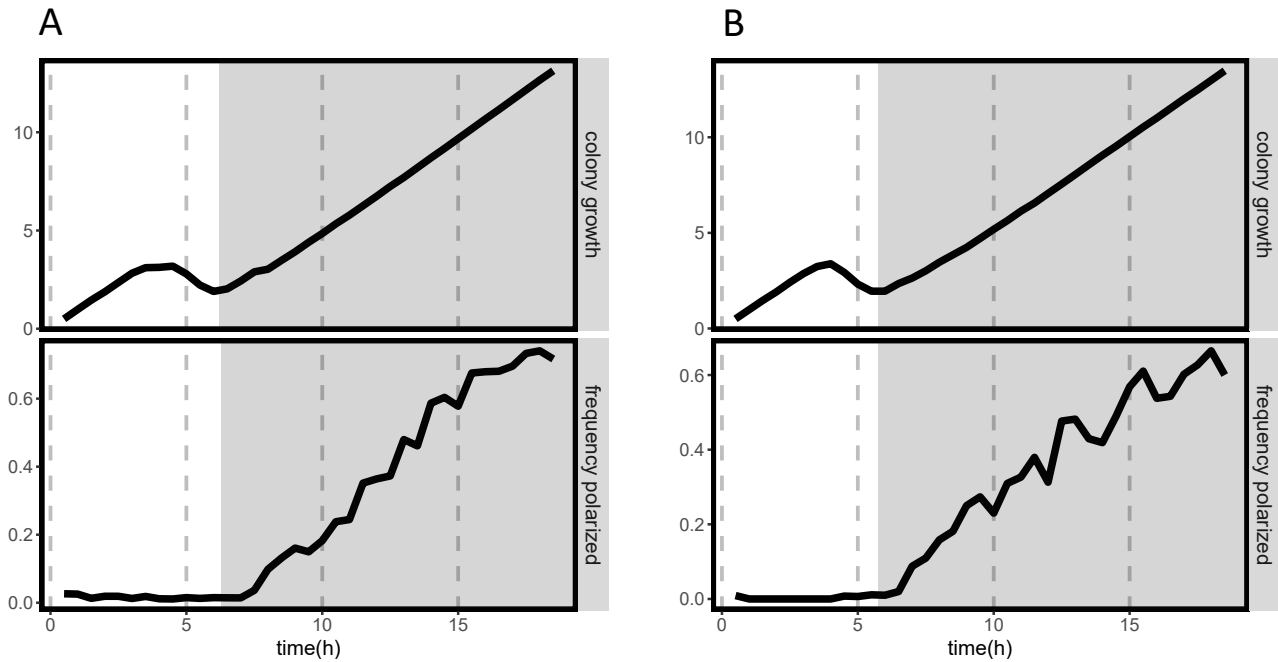


Fig. S1. SBW25 consistently polarizes pyoverdinin upon entry into stationary phase. Polarization in different growth stages of a microcolony. Both plots represent mean age of the cells in a growing microcolony of SBW25 (black line, top) and the corresponding frequency of polarized cells for each time point (black line, bottom). Bacterial colonies undergo different growth stages, from left to right: lag phase (generation F0), exponential phase (generation F1), and stationary phase (generation F2) Colored panels highlight stationary phase of bacterial colony growth, where pyoverdinin is observed to accumulate at cell poles. In all cases data has been filtered to exclude cells with segmentation errors or other artifacts that preclude proper analysis. Each plot depicts a biological replicate of the experiment described in Fig. 2A. A) 5 technical replicates, N = 152 initial cells. B) 4 technical replicates, N = 109 initial cells

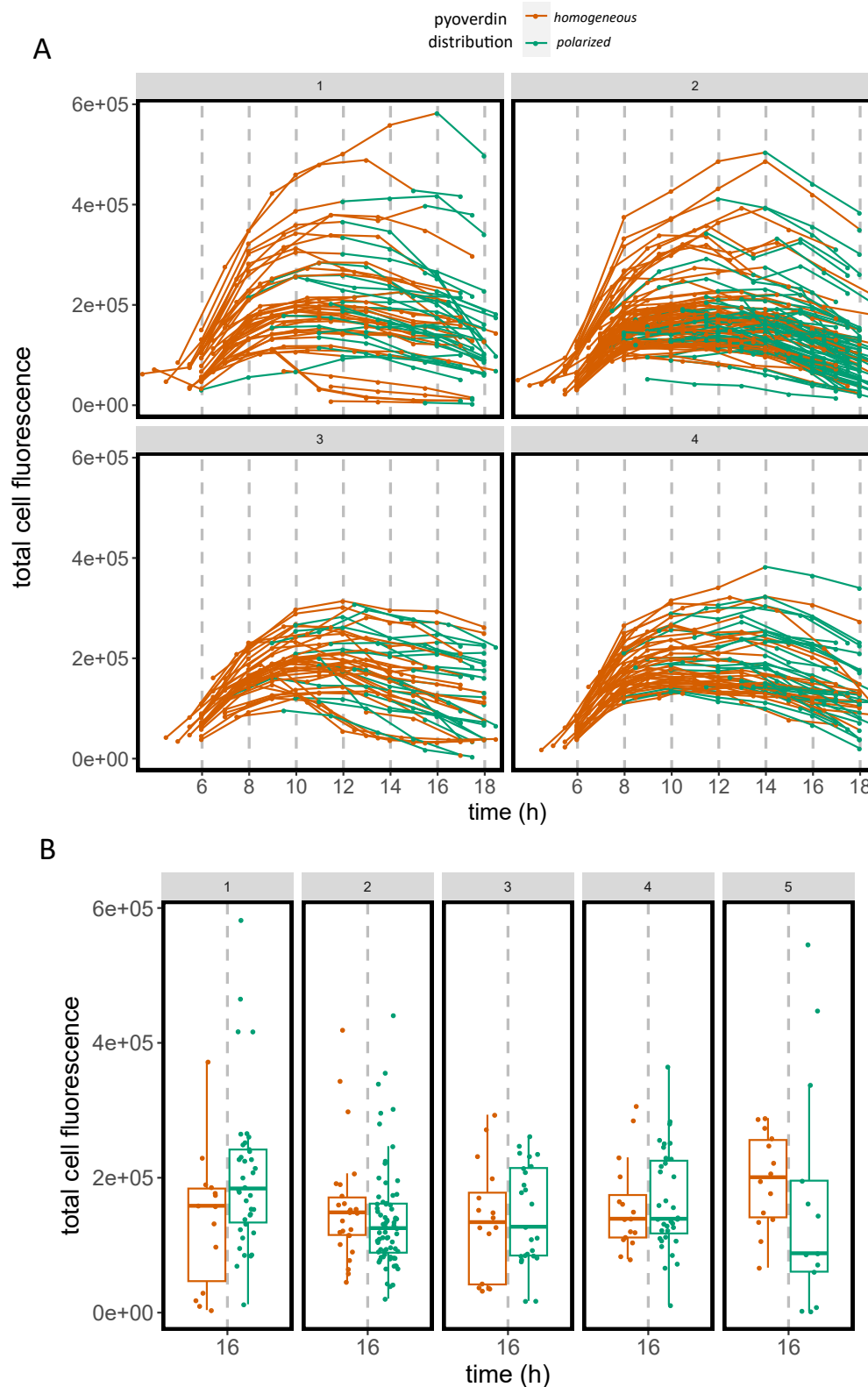


Fig. S2. Total cell fluorescence over time in cells that accumulate pyoverdinin at the pole during stationary phase A) Each line represents an individual cell that is generated by cell division but does not subsequently divide over the course of the time-lapse, i.e. generation F2 as defined in the main text. Values are the sum of the fluorescence intensity of all pixels corresponding to a segmented cell. Lines are colored according to the pyoverdinin distribution inside the cell at each time point, classified by a machine learning algorithm in "homogeneous" (red lines) or "polarized" (green line). Each panel labelled 1-4 represents a technical replicate, i.e. a different position in the microscope slide during image acquisition. B) Distribution of total cell fluorescence in cells that are classified as "homogeneous" (red) or "polarized" (green) at a time point corresponding to mid to late stationary phase, $t = 16$ h. Box plots represent the mean and associated statistical parameters. Dots represent individual cell values. Panels represent technical replicates as in A.

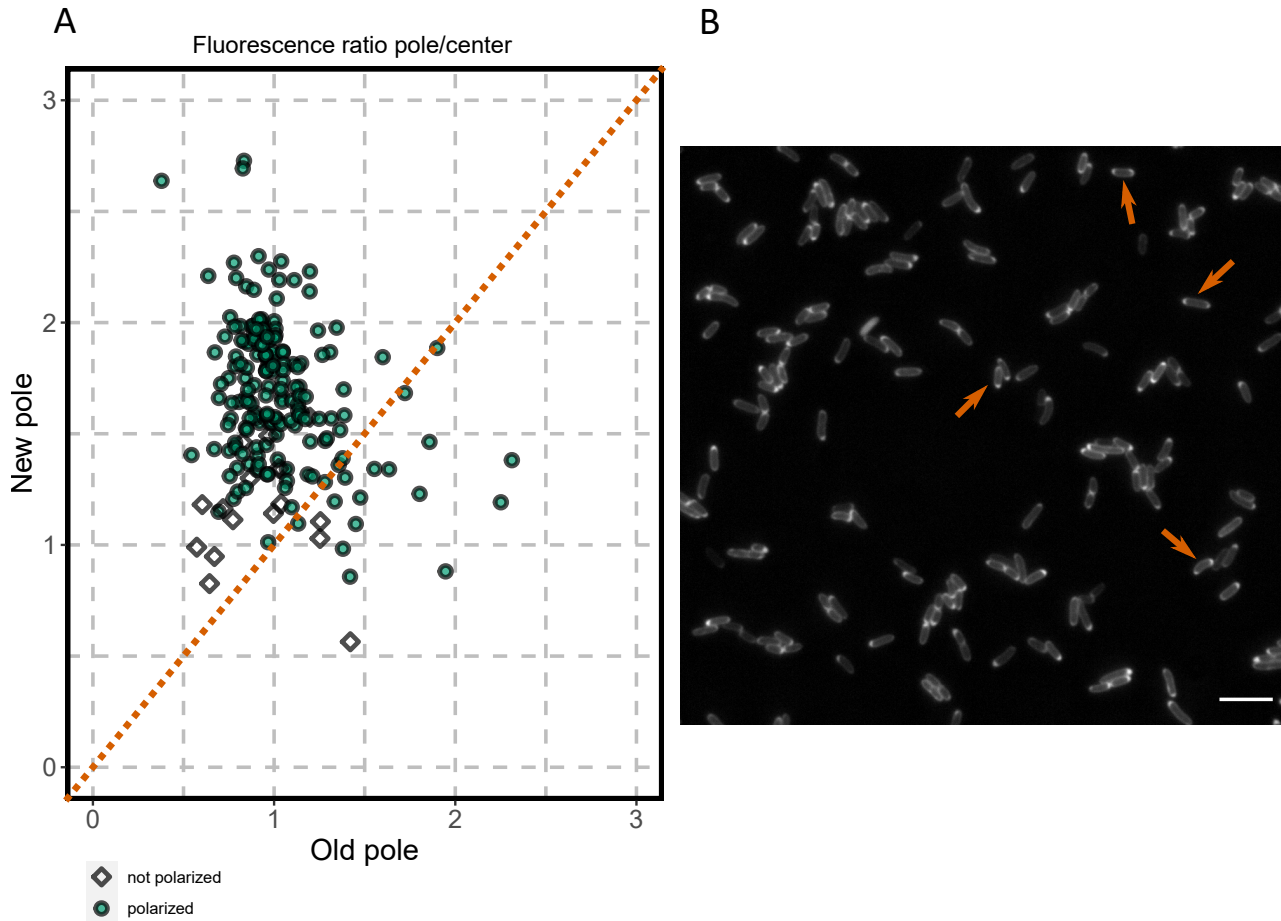


Fig. S3. Accumulation of pyoverdinin occurs predominantly at the new cell pole, but not exclusively and occasionally at both cell poles. A) Ratio of fluorescence between the central region of cells and the region of the new cell pole or old cell pole. Fluorescence values are obtained by dividing segmented cells into 3 proportional regions over long cell axis and averaging fluorescence along both cellular dimensions. Data corresponds to individual F2 cells at time = 18 h, N = 187. Green circles represent cells classified as "polarized", white diamonds represent cells classified as "not polarized". Red line highlights 1:1 ratio of fluorescence in cell centers to cell poles for reference. B) Image of SBW25 cells in conditions that promote polarization, in this case, high cellular density leading to rapid entry into stationary phase (t = 18 h) where accumulation of pyoverdinin at both cell poles can be observed in few individuals (red arrows). Scale bar = 10 μm .

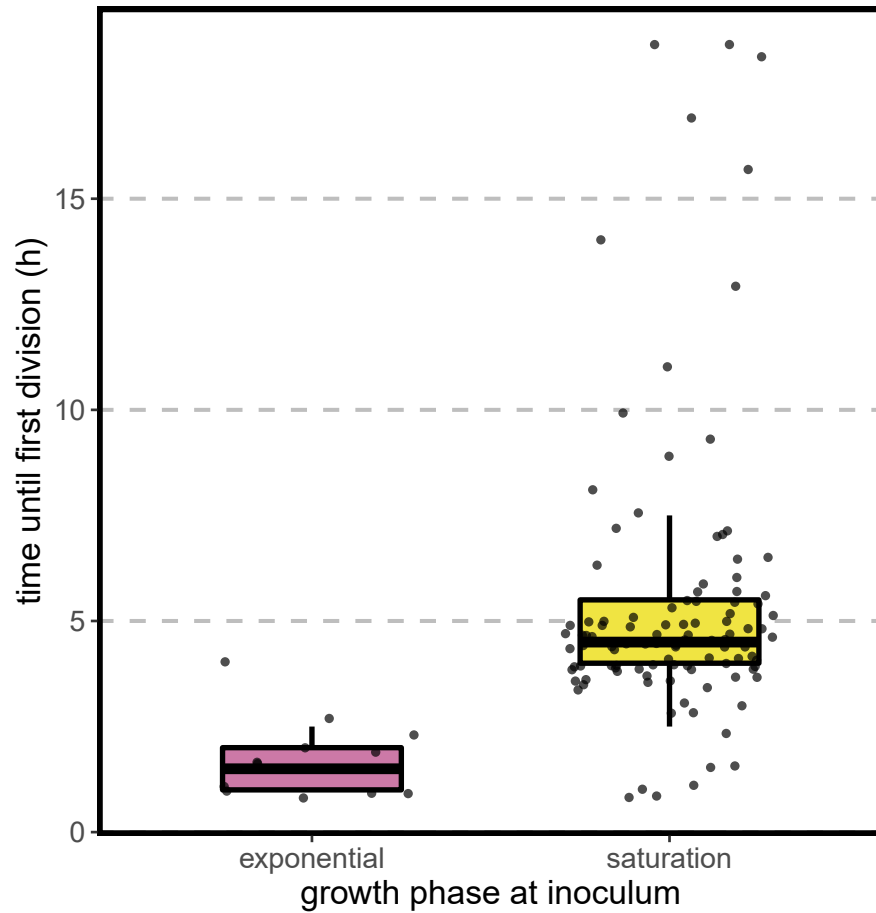


Fig. S4. Lag time difference between cells in stationary and exponential phase To assess the physiological state of cells after 24 h in SMM (whether they are still in exponential phase due to the minimality of the medium or rather they are entering stationary phase) we compared the time until first division of cells previously resuspended in fresh liquid SMM culture for 4h with data from cells inoculated in a fresh agarose pad directly following the usual protocol described in the main text. In the former case the cells are in exponential phase as their lag time is comparable to division time in this medium. (N = 13, 109 respectively)

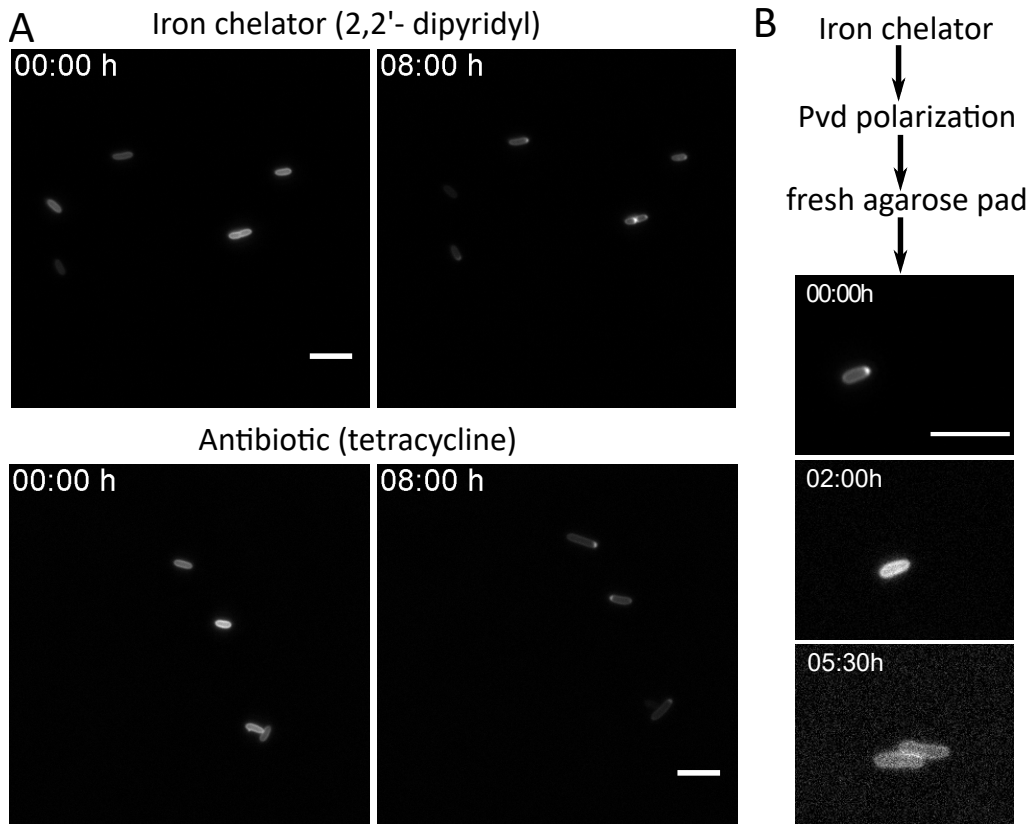


Fig. S5. Exposure to chemicals that prevent cell growth induces reversible accumulation of pyoverdinin at the cell pole. A) Treatment with iron chelator (top) or antibiotic (bottom) triggers accumulation of pyoverdinin at the cell pole without detectable cell growth. SBW25 cells were inoculated on an agarose pad containing either 100 $\mu\text{g}/\text{ml}$ of 2,2'-dipyridyl (top) or 5 $\mu\text{g}/\text{ml}$ tetracycline (bottom). Images were taken at $t = 0$ h and $t = 8$ h at the same position on the agarose pad. Note then that the photos depict the same individual cells at both time points, as no cell division was observed. B) Pyoverdinin polarization is reversible and precedes recovery of growth after stress. Time-lapse images of SBW25 pre-treated with an iron chelator to induce pyoverdinin polarization, then washed and inoculated on a fresh agarose pad. Images follow one individual cell as it recovers homogeneous pyoverdinin distribution in the periplasm, followed by cell division.

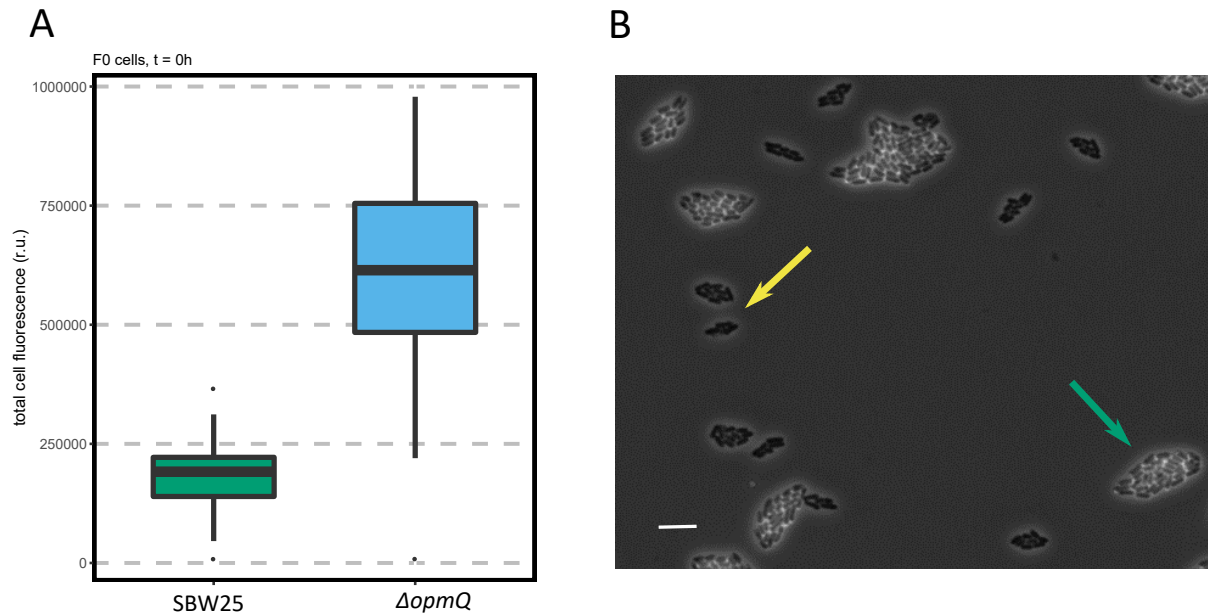


Fig. S6. OpmQ and FpvA deletion mutants are impacted on their ability to recycle, import, and synthesize pyoverdinin. A) Deletion of the export pump OpmQ leads to increased intracellular levels of pyoverdinin compared to ancestral SBW25. Box plots represent the aggregated fluorescence of individual cells of SBW25 (green) and $\Delta opmQ$ (light blue) at $t = 0$ h of inoculation on the agarose pad, $N = 35, 29$, respectively. B) Deletion of the receptor FpvA reduces pyoverdinin synthesis to basal levels, and impedes its import by crossfeeding. Image depicts co-culture of ancestral SBW25 (tagged with mCherry for identification) and $\Delta fpvA$, where phase contrast and fluorescent imaging of pyoverdinin have been overlayed to highlight the differences in pyoverdinin production between both strains. Scale bar = $10 \mu\text{m}$. Strains were mixed in equal proportions and grown on an agarose pad for 18 h. Green arrows point to microcolonies of SBW25, yellow arrows point to $\Delta fpvA$ microcolonies, as representative examples of the two phenotypes.

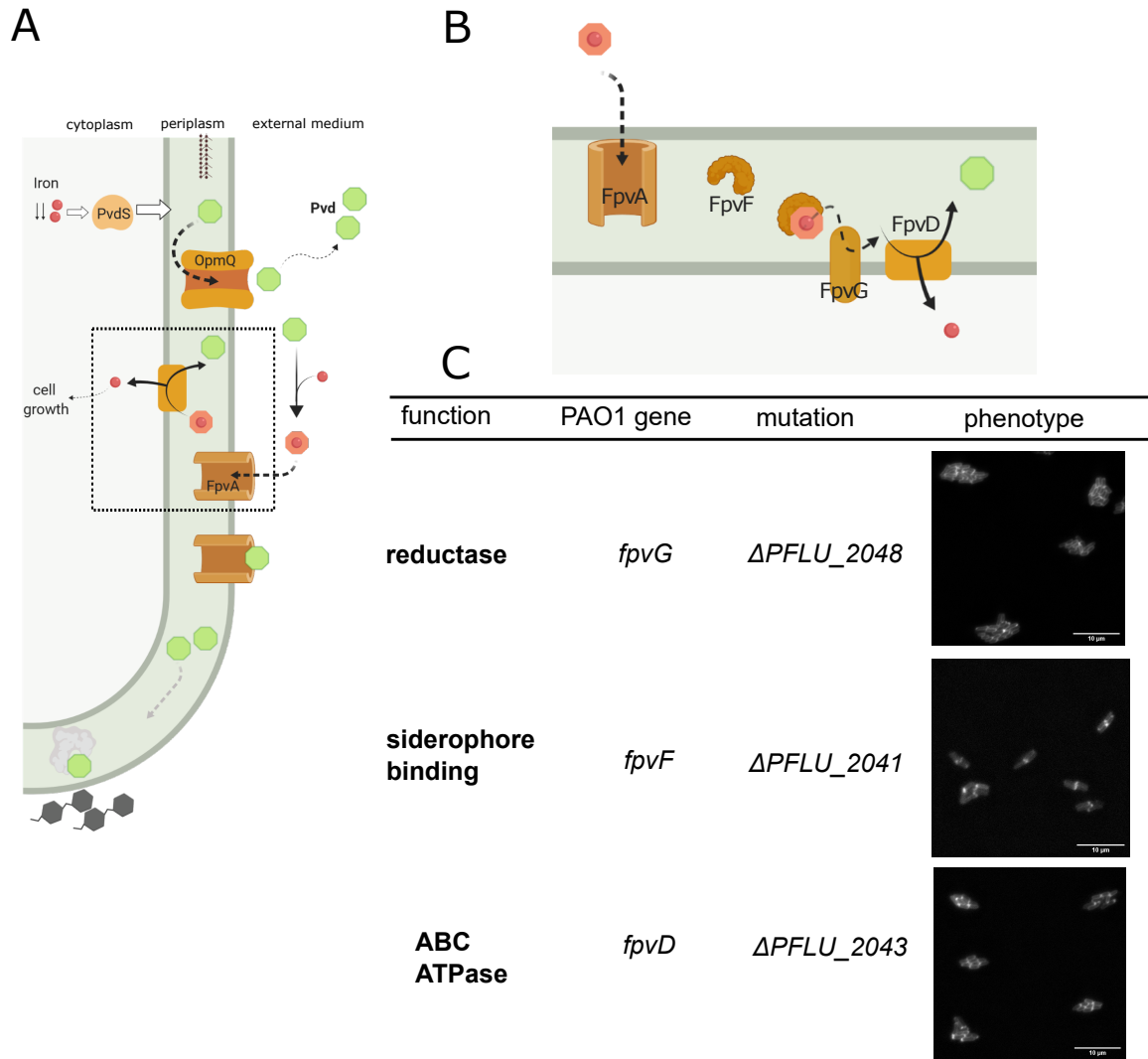


Fig. S7. title A) Cartoon depicting the pyoverdinin pathway, see main text for details. B) Cartoon detailing the mechanism of iron extraction from the ferripyoverdinin complex and recycling of pyoverdinin molecules. C) Pyoverdinin polarization phenotype of a collection of deletion mutants of the processes depicted in B). Genes were selected based on recent work on *P. aeruginosa* PAO1 identifying the function of previously unknown proteins. Orthologs of these genes were identified in *P. fluorescens* SBW25 in the database pseudomonas.com. Genes were deleted by two-step allelic exchange and cells were cultured and imaged as previously described.

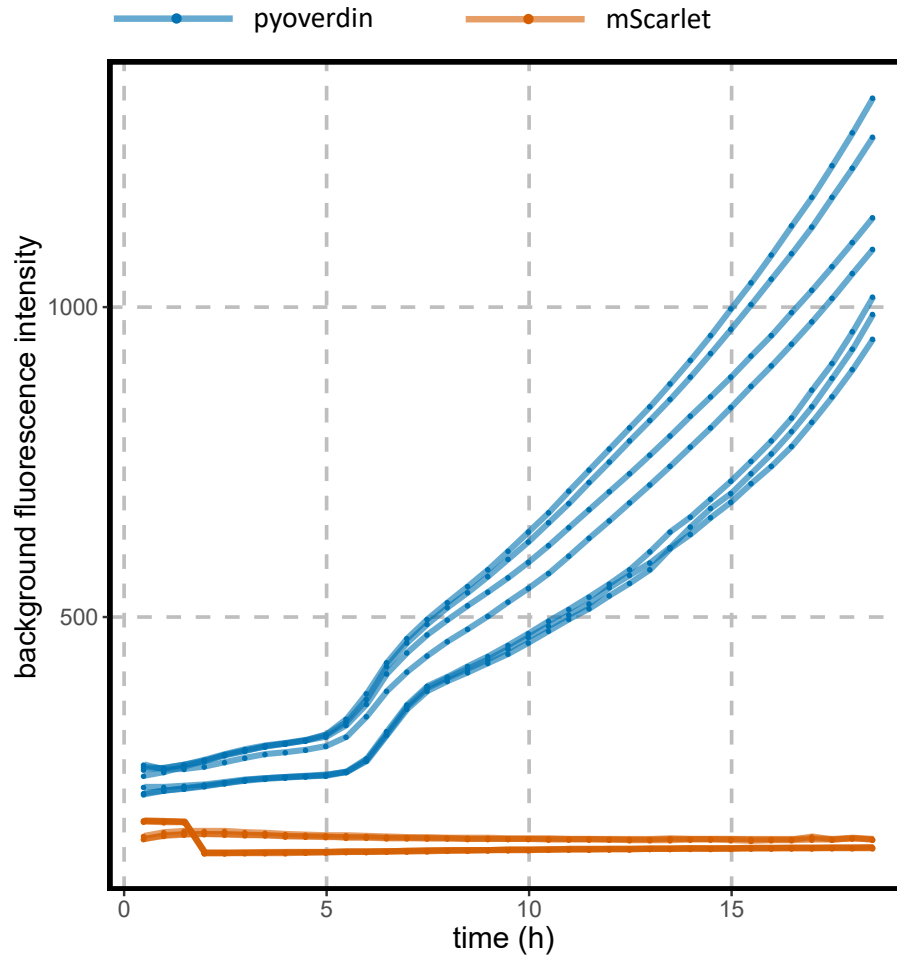


Fig. S8. Fluorescence background measured when imaging cells that produce the naturally-occurring molecule pyoverdinin and an engineered periplasmic mCherry protein Dots and lines represent the median intensity value of a 100x100 px square in the microscope field where cells are not present. Red lines correspond to the red fluorescent protein mCherry while blue lines correspond to pyoverdinin (which exhibits a fluorescence emission peak at 450nm). Each line represents a technical replicate.

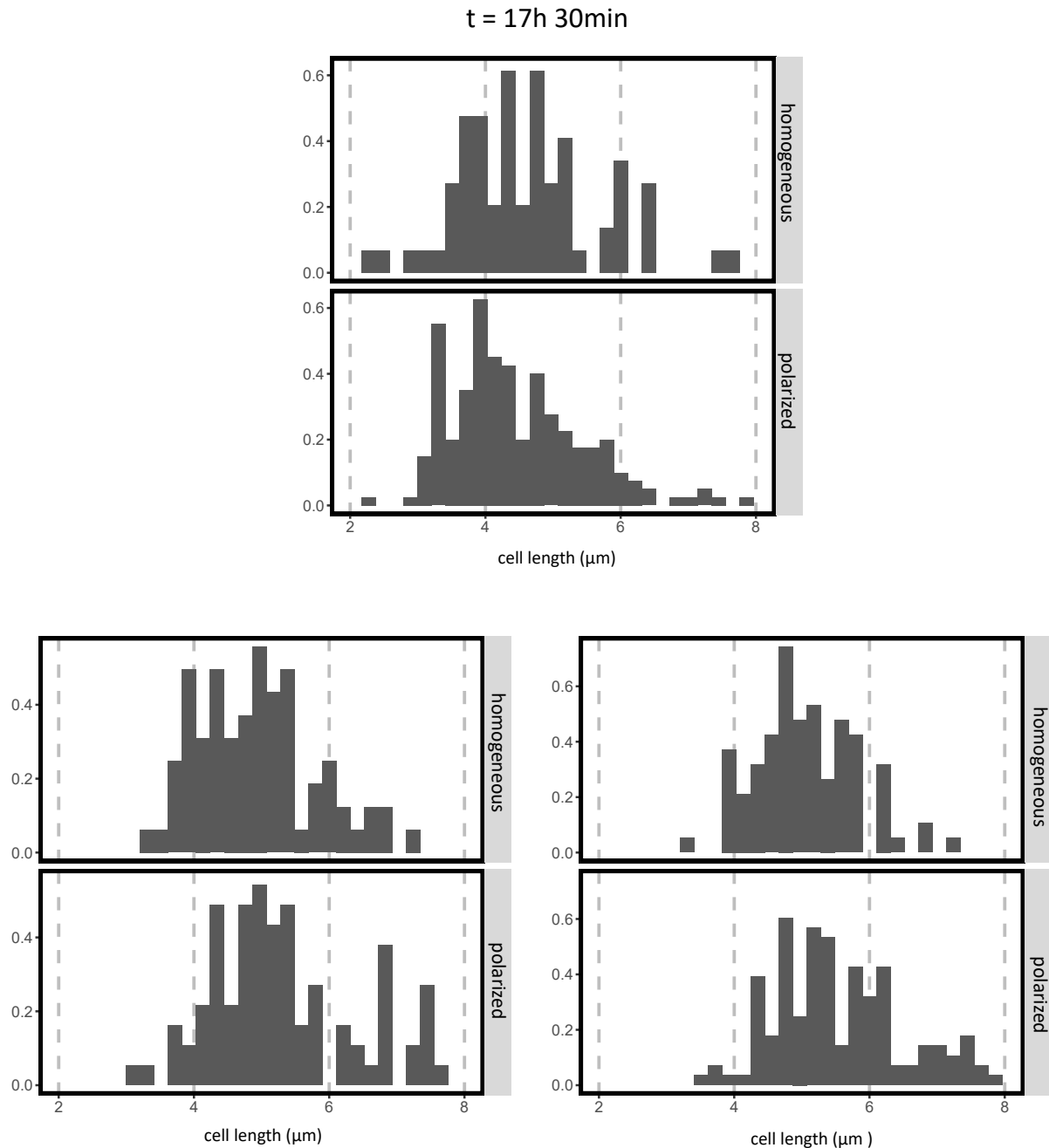


Fig. S9. Accumulation of pyoverdinin as a function of cell size. Distribution of cell size after 17h 30 min of growth in the agarose pad. The population of cells is classified according to the localization of pyoverdinin in "accumulated" and "non-accumulated". Cell size is measured as length on the long axis, since elongation occurs in this direction and thus will determine final cell area. Bars represent the distribution of cell lengths in both groups of cells. Each plot represents an independent experiment. Strains used: top: SBW25, bottom: MPB25340 (SBW25 with periplasmic red fluorescent marker), left and right panels represent 2 biological replicates. N = 261, 167, 235 respectively.

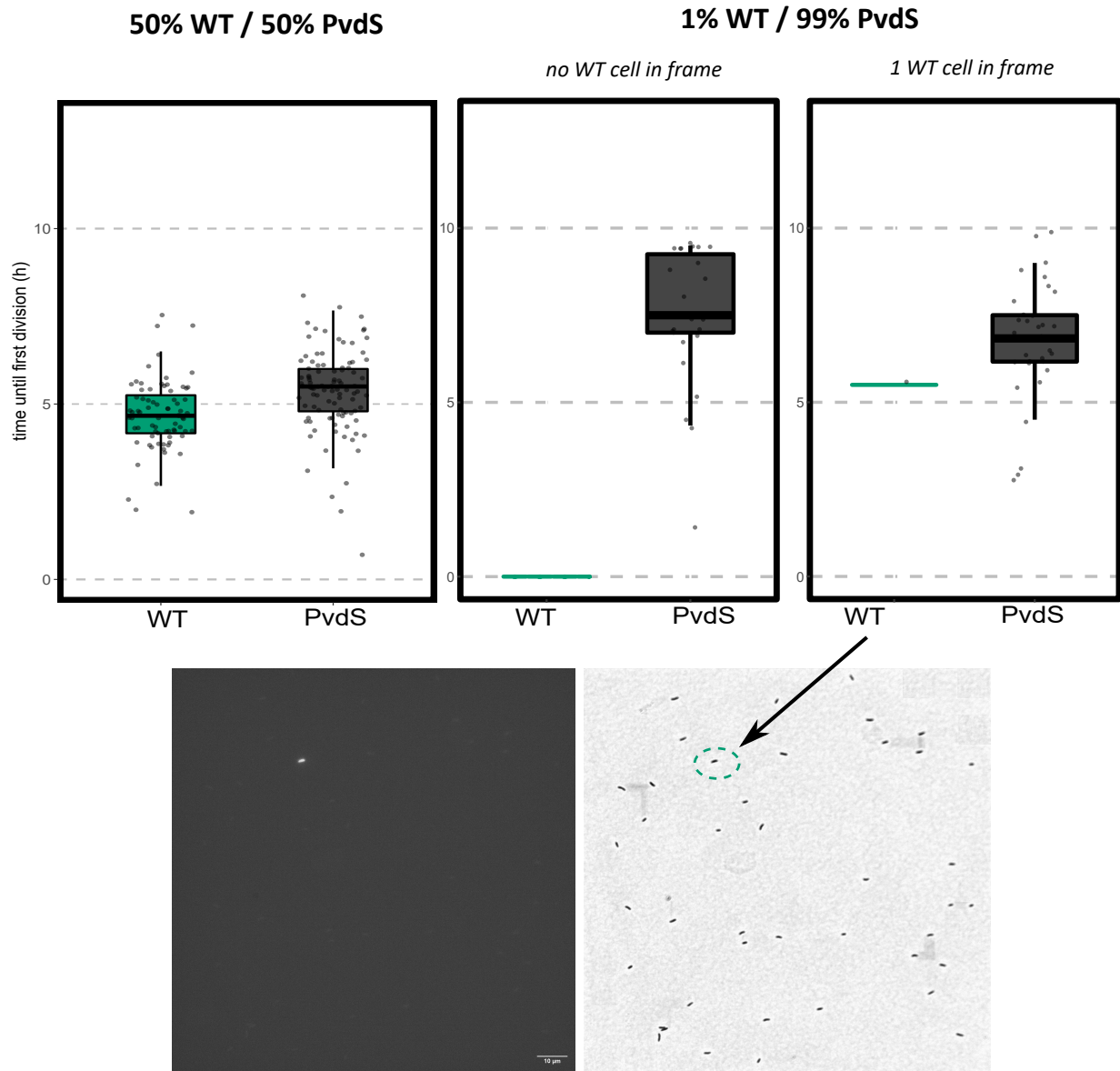


Fig. S10. Lag time of ancestor and nonproducing mutant in co-culture where pyoverdinin producer is rare Time after first division of ancestor and mutant cells pre-treated with DP upon co-inoculation on a fresh agarose pad. Prior to inoculation cells were mixed at 1:1 proportions (left panel, figure is the same as fig 4 B in main text and is included for reference) or ancestor cells at 1%. Center panel depicts data from a technical replicate where no ancestor cells can be identified in the microscope field of acquisition, right panel depicts data from a technical replicate where a single ancestor cell can be identified. Images below correspond to this latter dataset, where left panel represents fluorescent pyoverdinin imaging and right panel represents phase contrast imaging. Size of the field of view is 132.06 x 132.06 μm and scale bar represents 10 μm . The single pyoverdinin producer is highlighted in green dashed line. (N = 27, 35 respectively)