Dynamic Metabolic Rewiring Enables Efficient Acetyl Coenzyme A Assimilation in *Paracoccus denitrificans*

Katharina Kremer,* Muriel C. F. van Teeseling, Lennart Schada von Borzyskowski, Iria Bernhardsgrütter,* Rob J. M. van Spanning,* Andrew J. Gates, Mitja N. P. Remus-Emsermann, Martin Thanbichler,* Tobias J. Erb*<sup>a,b,g</sup>

*Max Planck Institute for Terrestrial Microbiology, Marburg, Germany
†Faculty of Biology, Philipps-Universität, Marburg, Germany
‡Department of Molecular Cell Biology, Vrije Universiteit, HV, Amsterdam, The Netherlands
§School of Biological Sciences, University of East Anglia, Norwich, United Kingdom
∥School of Biological Sciences, University of Canterbury, Christchurch, New Zealand
¶Biomolecular Interaction Centre, University of Canterbury, Christchurch, New Zealand
#LOEWE Center for Synthetic Microbiology, Marburg, Germany

**ABSTRACT** During growth, microorganisms have to balance metabolic flux between energy and biosynthesis. One of the key intermediates in central carbon metabolism is acetyl coenzyme A (acetyl-CoA), which can be either oxidized in the citric acid cycle or assimilated into biomass through dedicated pathways. Two acetyl-CoA assimilation strategies in bacteria have been described so far, the ethylmalonyl-CoA pathway (EMCP) and the glyoxylate cycle (GC). Here, we show that *Paracoccus denitrificans* uses both strategies for acetyl-CoA assimilation during different growth stages, revealing an unexpected metabolic complexity in the organism’s central carbon metabolism. The EMCP is constitutively expressed on various substrates and leads to high biomass yields on substrates requiring acetyl-CoA assimilation, such as acetate, while the GC is specifically induced on these substrates, enabling high growth rates. Even though each acetyl-CoA assimilation strategy alone confers a distinct growth advantage, *P. denitrificans* recruits both to adapt to changing environmental conditions, such as a switch from succinate to acetate. Time-resolved single-cell experiments show that during this switch, expression of the EMCP and GC is highly coordinated, indicating fine-tuned genetic programming. The dynamic metabolic rewiring of acetyl-CoA assimilation is an evolutionary innovation by *P. denitrificans* that allows this organism to respond in a highly flexible manner to changes in the nature and availability of the carbon source to meet the physiological needs of the cell, representing a new phenomenon in central carbon metabolism.

**IMPORTANCE** Central carbon metabolism provides organisms with energy and cellular building blocks during growth and is considered the invariable “operating system” of the cell. Here, we describe a new phenomenon in bacterial central carbon metabolism. In contrast to many other bacteria that employ only one pathway for the conversion of the central metabolite acetyl-CoA, *Paracoccus denitrificans* possesses two different acetyl-CoA assimilation pathways. These two pathways are dynamically recruited during different stages of growth, which allows *P. denitrificans* to achieve both high biomass yield and high growth rates under changing environmental conditions. Overall, this dynamic rewiring of central carbon metabolism in *P. denitrificans* represents a new strategy compared to those of other organisms employing only one acetyl-CoA assimilation pathway.

**KEYWORDS** *Paracoccus denitrificans*, acetate, carbon metabolism, ecophysiology, metabolic regulation

Citation Kremer K, van Teeseling MCF, Schada von Borzyskowski L, Bernhardsgrütter I, van Spanning RJM, Gates AJ, Remus-Emsermann MNF, Thanbichler M, Erb TJ. 2019. Dynamic metabolic rewiring enables efficient acetyl coenzyme A assimilation in *Paracoccus denitrificans*. mBio 10:e00805-19. https://doi.org/10.1128/mBio.00805-19.

Editor Caroline S. Harwood, University of Washington

Copyright © 2019 Kremer et al. This is an open-access article distributed under the terms of the Creative Commons Attribution 4.0 International license.

Address correspondence to Tobias J. Erb, toerb@mpi-marburg.mpg.de.

Received 28 March 2019
Accepted 7 June 2019
Published 9 July 2019
During growth, bacteria have to distribute metabolic flux between catabolism and anabolism. An important central metabolic pathway is the tricarboxylic acid (TCA) cycle, which interfaces anabolism and catabolism. In the tricarboxylic acid (TCA) cycle, the C2-unit acetyl coenzyme A (acetyl-CoA) is catabolized into CO2, generating reducing equivalents and ATP for the cell. At the same time, the TCA cycle produces several intermediates that are committed to biosynthesis. Note that the TCA cycle poses a special challenge for many compounds that are exclusively metabolized via acetyl-CoA, such as acetate, alcohols, short- and long-chain fatty acids, esters, and waxes. Because all carbon of acetyl-CoA is lost to CO2, this allows energy conservation but not carbon assimilation through the TCA cycle. Consequently, dedicated pathways for the assimilation of acetyl-CoA are required to allow growth on these ubiquitous substrates.

Two different acetyl-CoA assimilation pathways in bacteria have been described: the glyoxylate cycle (GC) (1, 2) and the ethylmalonyl CoA pathway (EMCP) (3, 4). The GC uses two enzymes in addition to the enzymes of the TCA cycle. The first enzyme of the GC, isocitrate lyase (Icl), cleaves isocitrate into succinate and glyoxylate. This step is followed by the condensation of glyoxylate with a second molecule of acetyl-CoA to form malate and free CoA in a reaction catalyzed by the second enzyme of the GC, malate synthase (1, 2).

The EMCP also forms malate and succinate. However, unlike the GC, the EMCP is a linear pathway that employs 13 different enzymes that collectively convert a total of three acetyl-CoA and two CO2 molecules into the TCA cycle intermediates malate and succinate. The key enzyme of the EMCP is crotonyl-CoA carboxylase/reductase (Ccr), which catalyzes the reductive carboxylation of crotonyl-CoA to ethylmalonyl-CoA (3–5).

Overall, the GC and the EMCP differ substantially in terms of cofactor and carbon requirements. The GC requires four ATPs and generates reducing equivalents at the level of two nicotinamides as well as two quinols per four acetates converted into malate (estimated change in reaction Gibbs free energy \( \Delta_rG^\circ = -129 \text{ kJ/mol} \)). To generate two malates from acetate in the EMCP, three molecules of acetate and two CO2 molecules are converted at the expense of three ATPs and two reduced nicotinamides, while two quinols are generated (estimated \( \Delta_rG^\circ = -142 \text{ kJ/mol} \)). When acetate is used to generate the CO2 and fuel the cycle, this changes the stoichiometry to three consumed ATPs and one reduced nicotinamide, as well as three quinols, generated (estimated \( \Delta_rG^\circ = -190 \text{ kJ/mol} \)). While the EMCP requires more enzymatic steps (and likely more cellular resources), it enables the fixation of inorganic carbon into biomass, as well as the coassimilation of different C-5 and C-3 carboxic acids compared to the GC, indicating that the two pathways provide different physiological advantages. In most bacteria, only one of the two acetyl-CoA assimilation pathways is present. Importantly, the alphaproteobacterium Paracoccus denitrificans encodes the enzymes for both acetyl-CoA assimilation pathways (Fig. 1), raising the question of which role each individual pathway plays in the cell.

In this work, we show that P. denitrificans uses both the GC and the EMCP. The EMCP serves as the default acetyl-CoA assimilation pathway, which is always present with low activity in the cell during growth on various substrates, including carbon sources that do not require acetyl-CoA assimilation. In contrast, the GC is specifically induced after an organism switches to carbon sources that depend on acetyl-CoA assimilation, such as acetate or crotonate. We further demonstrate that the two acetyl-CoA assimilation pathways confer distinct physiological advantages. Growth with the EMCP results in increased biomass yield in P. denitrificans, while the GC allows for higher growth rates, suggestive for a rate yield tradeoff between the two pathways. Phylogenetic analysis indicates that P. denitrificans and several other alphaproteobacteria acquired the GC through lateral gene transfer, consisting with a specific adaptation of the central carbon metabolism of these organisms during evolution. Collectively, our experiments show a surprising flexibility in central carbon metabolism of alphaproteobacteria that allows the complete rewiring of metabolic flux depending on the type and availability of the carbon and free-energy source according to the physiological needs of the cell.
RESULTS

*P. denitrificans* uses the EMCP and the GC for acetyl-CoA assimilation. Earlier studies suggested that *Paracoccus* might use the EMCP as well as the GC for acetyl-CoA assimilation (4, 6, 7). We therefore searched the genome of the fully sequenced strain *P. denitrificans* Pd1222 for homologs of genes involved in the EMCP and the GC from *Rhodobacter sphaeroides* 2.4.1 and *Escherichia coli* K-12, respectively. Our analysis verified that *P. denitrificans* has the genetic potential for both the GC and the EMCP (Fig. 1; Table 1) (4).

To test whether both the EMCP and the GC are functional and thus operate in *P. denitrificans*, we grew bacterial cells in batch culture on minimal medium supplemented with different carbon sources and quantified the activity of Ccr, the key enzyme of the EMCP, as well as Icl, the key enzyme of the GC, in cell extracts obtained from the cultures at mid-log phase (Fig. 2).

The activity of Ccr was almost undetectable in cells grown on succinate and at a very low basal level (≤100 mU/mg) in extracts of cells grown on glucose and glycolate. In contrast, the activity of Ccr was significantly higher in extracts of cells grown on crotonate, acetate, and ethanol. In these extracts, Icl activity was also present, while no Icl activity was detected in extracts of cells grown on glucose, succinate, or glycolate. Together, our data suggest that both the EMCP and the GC are active in *P. denitrificans*. However, while the EMCP is always active at low levels and upregulated in response to growth on substrates requiring acetyl-CoA assimilation, the GC is specifically activated only during the growth of *P. denitrificans* on these substrates.
The EMCP is sufficient to sustain the growth of *P. denitrificans* on acetate. Next, we tested whether both pathways are essential for acetyl-CoA assimilation in *P. denitrificans*. To that end, we aimed to selectively block the GC or EMCP using external inhibitors. Several inhibitors of Icl are known (8). We screened three of them to identify 3-nitropropionate (3-NPA) as a potential candidate for a selective GC inhibitor of *P. denitrificans* in vivo.

3-NPA inhibited purified *P. denitrificans* Icl *in vitro* with an apparent 50% inhibitory concentration (IC50) of 34 μM at concentrations of 100 μM D,L-isocitrate (Fig. 3A). Moreover, 3-NPA did not affect the growth of *P. denitrificans* on carbon substrates that do not require Icl activity, such as succinate (Fig. 2B; see also Fig. S1E in the supplemental material). Notably, 3-NPA also did not affect the growth of *Methylobacterium extorquens* AM1, an EMCP-positive organism that is closely related to *P. denitrificans* on acetate (Fig. S1B and E). This suggested to us that 3-NPA acts as a GC inhibitor that shows negligible off-target effects in the cell, although the possibility of additional effects of 3-NPA could not be completely excluded.

**TABLE 1** Genes coding for enzymes of the EMCP and the GC in *Rhodobacter sphaeroides* 2.4.1 and *Escherichia coli* K-12, respectively, and their homologues identified in *P. denitrificans* Pd1222*

| Pathway or cycle | Enzyme | Gene name | Gene ID in: | Pathway or cycle | Enzyme | Gene name | Gene ID in: |
|------------------|--------|-----------|-------------|------------------|--------|-----------|-------------|
| **EMCP**         | β-Ketothiolase | phaA | RSP_0745 | - | **P. denitrificans** | Pd1222 | - |
|                  | Acetoacetyl-CoA reductase | phaB | RSP_0747 | - |
| **Crotonyl-CoA carboxylase/reductase** | ccr | RSP_0960 | - | **E. coli** | K-12 | - |
| **Ethylmalonyl-CoA/methylmalonyl-CoA epimerase** | epi | RSP_0812 | - | **P. denitrificans** | Pd1222 | - |
| **Ethylmalonyl-CoA mutase** | ecm | RSP_0961 | - | - |
| **(2S)-Methylsuccinyl-CoA dehydrogenase** | mcd | RSP_1679 | - | - |
| **Mesaconyl-CoA hydratase** | mch | RSP_0973 | - | - |
| **(2R)-Methylmalonyl-CoA/thymine-CoA lyase** | mcl-1 | RSP_1771 | - | - |
| **(S)-Malyl-CoA thioesterase** | mcl-2 | RSP_0970 | - | - |
| **Propionyl-CoA carboxylase** | pccA | RSP_2191/2189 | - | - |
| **Ethylmalonyl-CoA/methylmalonyl-CoA epimerase** | epi | RSP_0812 | - | - |
| **Propionyl-CoA carboxylase** | pccA | RSP_2191/2189 | - | - |
| **(2R)-Methylmalonyl-CoA mutase** | ecm | RSP_2912 | - | - |
| **GC** | Isocitrate lyase | aceA | - | CSY66_09370 | **P. denitrificans** | Pd1222 | - |
| Malate synthase | aceB | - | CSY66_09365 | Pd1222 | - |

*Key enzymes of the two pathways are in bold. Missing enzyme homologues are indicated by dashes.*

**FIG 2** Ccr and Icl enzyme activities in cell extracts of *P. denitrificans* grown on different carbon sources. Ccr activity is given as a combined activity of reduction and reductive carboxylation of crotonyl-CoA, quantified in the absence (light red) and presence (dark red) of NaHCO₃. Error bars indicate standard deviations.
When we added increasing concentrations of 3-NPA to *P. denitrificans* growing on acetate, the growth rate was successively decreased from $0.26 \pm 0.043$ h$^{-1}$ (0 M 3-NPA) to $0.13 \pm 0.005$ h$^{-1}$ (600 µM 3-NPA) but not completely abolished (Fig. 3B; Fig. S1D). This indicated that the GC is not essential in *P. denitrificans* and that the EMCP is sufficient for acetyl-CoA assimilation.

The use of the GC increases the growth rate of *P. denitrificans* on acetate. To estimate the contribution of each pathway to acetyl-CoA assimilation, we individually deleted the genes coding for Ccr and Icl from the genome of *P. denitrificans*, yielding Δccr and Δicl strains, respectively (Fig. 4; Fig. S2). During growth on succinate, both deletion strains grew similarly to the wild type. However, a shift from succinate to acetate caused pronounced and different strain-specific growth defects.

The Δccr strain grew with wild type-like growth rates but displayed an extended lag phase (Fig. 4). Notably, this lag phase ranged from 15 to 50 h between different experiments (Fig. S3). However, when the Δccr strain was subcultured on acetate, the variation in the lag phase disappeared (Fig. 4B, solid blue line), and the growth rate corresponded to that of the wild type (Fig. 4B and C), indicating that the mutant adapted to acetate.

The Δicl strain, in contrast, did not exhibit a prolonged lag phase but showed a decreased growth rate on acetate, which is consistent with the results of the 3-NPA inhibition experiments. Together, these results suggest that the GC enables fast growth of *P. denitrificans* on acetate, but unlike with the EMCP, some time is required for the strain’s growth to be induced after a switch to acetate.

The EMCP and GC show a complex expression pattern during switches from succinate to acetate. To follow the expression of the EMCP and the GC in vivo, the native ccr and icl genes of *P. denitrificans* were replaced with constructs encoding translational fusions of Ccr to the red fluorescent protein mCherry and of Icl to the cyan fluorescent protein Cerulean, respectively. The resulting strain (*P. denitrificans* ccr::ccr-mCherry icl::icl-Cerulean) was continuously monitored by determining the total fluorescence of the population during cultivation in 96-well plates (Fig. 5).

During growth on succinate, cells showed very low levels of mCherry fluorescence and even lower levels of Cerulean fluorescence, which is consistent with our earlier finding that basal Ccr, but no Icl, activity could be detected in cell extracts of *P. denitrificans* grown on succinate (Fig. 2). When shifted from succinate to acetate, the reporter strain showed biphasic ccr-mCherry and icl-Cerulean expression patterns. Ccr-mCherry fluorescence increased within the first 20 h and then decreased gradually at the onset of exponential phase until it again reached a low basal level. In contrast, the production of Icl-Cerulean started only after 20 h but continued to increase until
shortly before the cells entered stationary phase. Subsequently, Icl-Cerulean fluorescence dropped transiently before continuing to rise again in the late stationary phase. Very similar growth-linked expression patterns were observed in reporter strains of *P. denitrificans* carrying only a single Ccr-mCherry or Icl-mCherry fusion (Fig. S5). In summary, our results suggest that acetyl-CoA assimilation in *P. denitrificans* follows a complex regulatory pattern in which the GC is used for fast and efficient acetyl-CoA assimilation during exponential phase, while the EMCP is used to bypass the time needed for activation of the GC.

**Population-wide switch responses monitored by single-cell microscopy.** To understand whether the transition from EMCP- to GC-driven acetyl-CoA assimilation during the switch to acetate occurred in the whole population or in only a subset of
cells, we followed the production of Ccr-mCherry and Icl-Cerulean in *P. denitrificans* at the single-cell level using time-lapse fluorescence microscopy. Here, we imaged cells growing on succinate (Fig. 6A and B) and acetate (Fig. 6C and D) at 2-h intervals and subjected the images to automated analysis (10) to determine the average fluorescence per cell (Fig. 6A and C). While repeated handling of the cultures for sample collection led to slightly decreased growth rates, the overall expression patterns of *ccr*-mCherry and *icl*-Cerulean matched those measured during continuous cultivation in the 96-well plates (Fig. 5).

While no fluorescence was detected for the wild type (negative control) (Fig. S4), the reporter strain showed different production patterns for Ccr-mCherry and Icl-Cerulean. On succinate, Ccr was produced at low levels during exponential phase, with increased expression in the stationary phase. In contrast, Icl was detected only at the end of the exponential phase and only in a small fraction of cells, which appeared strongly fluorescent. This implies that Icl expression is heterogeneous in the late exponential phase on succinate and thus may suggest a bet-hedging-like behavior under these conditions (Fig. 6A and B; Fig. S6).

Upon the switch to acetate, Ccr was produced first, whereas Icl was induced only at the onset of the early exponential phase. While some cells switched earlier to Icl production than others, all cells produced Icl in the exponential phase, indicating that the whole population and not only a subset of cells shifted from using the EMCP to using the GC (Fig. 6C and D; Fig. S6).

To follow the expression dynamics of *ccr* and *icl* continuously in individual cells, we trapped succinate-grown *P. denitrificans* *ccr:ccr*-mCherry *icl:icl*-Cerulean in a microfluidic device. We then provided acetate as the sole source of carbon and tracked cell growth and division as well as reporter production by microscopy over the course of time (Movie S1). This analysis confirmed that essentially all cells switched from the EMCP to the GC, demonstrating that faster acetyl-CoA assimilation is switched on in the whole population and not by individual cells that outgrow the population (Fig. 7).

**FIG 5** Expression of Ccr and Icl in *P. denitrificans* during growth on succinate and acetate. The optical density (OD$_{600}$) and fluorescence intensity (FI) of the *P. denitrificans* wild type (A) and *ccr:ccr*-mCherry *icl:icl*-Cerulean strain (B) were monitored during growth on succinate and acetate using a 96-well plate reader. Fluorescence was normalized to the OD and corrected by the background signal of the wild type. The Ccr-mCherry level is shown in red, and the Icl-Cerulean level is shown in cyan. Growth curves and fluorescent measurements from triplicates are shown, and in some cases the lines overlap.

**DISCUSSION**

Here, we describe an unprecedented (and unexpected) metabolic redundancy in bacterial central carbon metabolism. The chromosome of *P. denitrificans* carries genes for two fundamentally different acetyl-CoA assimilation pathways, the EMCP and the
Fluorescence of *P. denitrificans* ccr:ccr-mCherry ict:ict-Cerulean during growth on succinate (A, B) and acetate (C, D). At the indicated time points, the cultures were sampled and analyzed microscopically. Images were subjected to automated analysis using BacStalk (10). (A and C) Distribution of cellular fluorescence intensities (n = 450 cells per time point) shown as box plots. The values were corrected for the background fluorescence of the wild type. The margins of the boxes encompass the 25th to 75th percentiles; the lines indicate the median, and the whiskers extend to the 5th and 95th percentiles, respectively. The colored lines connect the medians. (B and D) Representative microscopy images. Ph3, phase contrast; RFP, red fluorescent protein, mCherry channel (overlaid with Ph3); CFP, cyan fluorescent protein; Cerulean channel (overlaid with Ph3).
Both pathways are functional and regulated dynamically depending on the growth stage of the cells. The EMCP serves a default function and is expressed at basal levels at all times and on all carbon and free-energy sources tested. Upon a switch to acetate, the EMCP is strongly induced at the early stage of growth but then decreases in activity again. The GC, in contrast, is exclusively induced on acetate and reaches peak activity at the late stage of growth, indicating a surprising rewiring of central carbon metabolism in *P. denitrificans* at the onset of the exponential phase.

What might be the reasons for the rewiring of central carbon metabolism in *P. denitrificans*? Individual knockout strains of the EMCP and GC show that the two pathways confer distinct advantages. The EMCP increases the growth yield, while the GC allows faster growth on acetate. How can these differences be explained, and what are their consequences?

Importantly, the EMCP is able to fix inorganic carbon to increase biomass yield according to the following equation: $3 \text{acetate} + 2 \text{CO}_2 + 2 \text{NADPH} + 2 \text{quinones} \rightarrow 2 \text{malate} + 2 \text{NADP}^+ + 2 \text{quinols}$. This stoichiometry allows organisms using the EMCP to gain extra carbon from the environment if additional reducing equivalents are available to the cell, e.g., through internal storage compounds, such as polyhydroxybutyrate (PHB), or through growth substrates that are comparatively more reduced than average cellular carbon. This is supported by recent calculations that predict an approximately 3%-higher yield for *M. extorquens* on methanol when the EMCP is used than when the GC is used (11). Another important feature of the EMCP is that it not only enables acetyl-CoA assimilation but also is directly linked to PHB metabolism and can also function in the assimilation of propionate, in addition to several dicarboxylic acids. This versatility makes the EMCP an all-purpose pathway that might allow *P. denitrificans* to assimilate several different carbon and free-energy sources in parallel, which may explain the constant expression of the EMCP on diverse growth substrates.

The GC, on the other hand, is a very specialized route that requires only two additional enzymes. The GC requires fewer protein resources, and the thermodynamics of the cycle (i.e., the free energy of the overall process) might become more favorable with the upper limit of the free Gibbs energy dissipation rate in *P. denitrificans* (12). Rerouting acetyl-CoA assimilation to the GC thus might allow a higher metabolic flux and consequently faster growth, providing a potential explanation as to why *P. denitrificans* switches to the GC when high concentrations of acetate are available.

Overall, a picture emerges in which metabolic rewiring is a highly coordinated strategy in *P. denitrificans* that allows optimal growth of this species in a changing environment.
environment. When the growth substrate changes from succinate to acetate, it is conceivable that the constant expression of the EMCP facilitates the immediate assimilation of acetate. This might help \textit{P. denitrificans} during the lag phase until the GC is fully induced and thus provide an advantage over other microorganisms that rely on the GC only. Once induced, the GC then may enable increased acetate assimilation rates, which might enable \textit{P. denitrificans} to outcompete microorganisms that possess solely the EMCP.

Out of 48 \textit{Paracoccus} genomes analyzed, 34 possess only a \textit{ccr} homolog, while 9 additionally contain a homolog of \textit{icl} (Fig. 8; see also Fig. S7 and S8 in the supplemental material). Only five strains seem to exclusively possess an \textit{icl} homolog. Notably, \textit{ccr}

\textbf{FIG 8} Maximum-likelihood phylogenetic tree of \textit{Paracoccus} strains. The phylogenetic tree is based on the concatenated alignments of 12 protein sequences from 48 \textit{Paracoccus} strains. Bootstrap values of \(\geq 0.5\) are given on the respective nodes; calculated branch lengths of the tree are ignored for the sake of easier visualization. The presence of \textit{Ccr} is marked with a red dot, the presence of \textit{Icl} is marked with a blue dot, and the presence of both enzymes is marked with a purple dot. The 48 strains are clustered in three distinct groups.
phylogeny correspond largely to overall strain phylogeny, suggesting that the EMCP is the ancient acetyl-CoA assimilation strategy in the genus *Paracoccus*. This hypothesis is in line with our experimental observation that the EMCP serves a default function in *P. denitrificans* and is further supported by the fact that other, closely related alphaproteobacteria possess only the EMCP (4).

Notably, the phenomenon of metabolic rewiring is presumably not restricted to the *Paracoccus* clade alone. Several alphaproteobacteria show the genetic potential to express multiple acetyl-CoA assimilation pathways (4), suggesting that dynamic rewiring of the central carbon metabolism is a more widespread strategy in nature. This discovery extends recent findings that metabolic degeneracy plays an important role in alphaproteobacterial metabolism (13). Future experiments will focus on understanding the molecular mechanisms that underlie the coordination of the two acetyl-CoA pathways and their regulation through transcriptional, posttranslational, or allosteric regulatory mechanisms (14–17) and on clarifying the evolutionary and ecological significance of dynamic metabolic rewiring as a new principle in microbial central carbon metabolism.

**MATERIALS AND METHODS**

**Strains, media, and growth conditions.** All experiments were performed with *Paracoccus denitrificans* strains DSM413 and Pd1222 (Table 2). The strain preferentially used for all experiments was DSM413, which is the original wild-type strain isolated by Beijerinck and Minkman in 1908 (18). In cases where genetic manipulation of this strain was not successful, its genetically more tractable derivative Pd1222 (19) was used alternatively. *P. denitrificans* was grown at 30°C in mineral salt medium (trace elements, TE3-Zn) (20) supplemented with defined carbon sources and adjusted to a total carbon concentration of 120 mM. The density of cultures was determined photometrically at a wavelength of 600 nm. Bacterial growth was monitored over time using Tecan Infinite 200 Pro plate reader systems (Tecan Trading AG, Switzerland) with Nunclon Delta Surface 96-well plates (Thermo Scientific, USA). *Escherichia coli* was grown at 37°C in Luria-Bertani broth or M9 minimal medium (337 mM NaH2PO4, 220 mM KH2PO4, 85.5 mM NaCl, 18.7 mM NH4Cl, 1 mM MgSO4, 0.3 mM CaCl2, 0.13 mM Na2EDTA, 0.03 mM FeSO4, 1 μM thiamin, 1 μM biotin) supplemented with 60 mM acetate. Methylcobalamin derivatives AM1 were grown as described previously (21) at 30°C in mineral medium supplemented with 10 mM acetate. Antibiotic concentrations were used as follows: kanamycin at 25 or 50 μg ml⁻¹, spectinomycin at 50 μg ml⁻¹, and tetracycline at 10 μg ml⁻¹.

**Chemicals.** All chemicals were obtained from Sigma-Aldrich (Steinheim, Germany) and Carl Roth (Karlsruhe, Germany) and were of the highest purity available.

**Construction of plasmids.** Oligonucleotides and synthetic genes were purchased from Eurofins (Brussels, Belgium) and GenScript (Piscataway, USA). The plasmid DNA was purified using the QIAprep Spin Miniprep Kit (Qiagen, Germany) and the GenElute Plasmid Miniprep Kit (Sigma-Aldrich, Germany).

**Construction of strains.** The deletion of the ccr gene in the wild-type strain was achieved by using the suicide plasmid pSYNccrflanks (9). The ccr gene was replaced with mCherry or Cerulean to create strains of *P. denitrificans* expressing the respective ccr-mCherry or ccr-Cerulean expression cassette (Table 2). After transformation of *P. denitrificans*, the suicide plasmids were removed by selecting for sucrose suicide plasmid resistance (4). For the construction of the Δccr strain, the suicide plasmid was replaced with the artificial chromosome pTE1604 (22) to achieve markerless integration of the ccr gene.

**Molecular biology.** All oligonucleotide synthesis, cloning, and sequencing reactions were performed using standard methods (23). Restriction enzymes were purchased from Fermentas (Berlin, Germany) and New England Biolabs (Ipswich, USA). All chemicals were of the highest purity available. E. coli DH5α was grown as described previously (24).

**ACKNOWLEDGMENTS.** We thank the German National Genome Research Network (NGFN) for financial support. We thank Karl-Georg Butzke for his contributions to the design of the manuscript. We thank Dominik Krämer and Timo Lenk for their help in the laboratory.

**TABLE 2** Strains used in this study

| Strain |
|--------|
| E. coli TOP 10 | Cloning strain | Invitrogen, USA |
| E. coli DH5α | Cloning strain | ThermoFisher Scientific, USA |
| E. coli BL21 | Protein expression | ThermoFisher Scientific, USA |
| E. coli ST18 | Mating strain | 28 |
| M. extorquens AM1 | Wild type | 39 |
| *P. denitrificans* DSM413 | Wild type | 18 |
| *P. denitrificans* Pd1222 | Increased conjugation frequency | 19 |
| *P. denitrificans* DSM413 Δccr (TJE-KK5) | Δccr | This work |
| *P. denitrificans* DSM413 Δicl (TJE-KK6) | Δicl | This work |
| *P. denitrificans* Pd1222 ccr::ccr-mCherry icl::icl-Cerulean (TJE-KK14) | ccr::ccr-mCherry icl::icl-Cerulean | This work |
| *P. denitrificans* Pd1222 cccr::ccr-mCherry (TJE-KK13) | cccr::ccr-mCherry | This work |
| *P. denitrificans* Pd1222 icl::icl-mCherry (TJE-KK10) | icl::icl-mCherry | This work |
regions, and subsequent ligation into PstI/EcoRI-digested pK18mobsacB resulted in plasmids pK18mobsacB_tccflanks (pTE1606) and pK18mobsacB_ticflanks (pTE1615).

For the introduction of fluorescent reporter fusions, ccr and icl were ordered as synthetic genes fused to evoglowPp1 (24) and mCherry (25), respectively, each preceded by a 30-bp linker and followed by the downstream region of the respective genes, yielding plasmids pEX4_ccrevoglowPp1ccrds (pTE1601) and pEX4_iclmCherryiclds (pTE1603).

pTE1603 was digested with SalI and BamHI. Subsequent ligation into SalI/BamHI-cut pK18mobsacB yielded pK18mobsacB_ticlmCherryiclds (pTE1616).

pTE1601 was amplified with targeted omission of the evoglowPp1 gene via inverted PCR using primers NdeI_pKK21-f and pKK21-r. The mCherry gene was amplified from pTE1603 using primers mCherry-f and NdeI_mCherry-r. Restriction of both fragments with AvrII and NdeI and subsequent ligation of the individual fragments to each other resulted in the plasmid pEX4_ccrmCherryccrds.

pEX4-ccrmCherryccrds was digested with SalI and BamHI. Subsequent ligation into SalI/BamHI-cut pK18mobsacB yielded pK18mobsacB_tccrmCherryccrds (pTE1624).

pTE1603 was amplified with targeted omission of the mCherry gene via inverted PCR using primers NdeI_pSYNiclXiclds-f and HindIII_pSYNccrXccrds-r. The Cerulean gene (26) was amplified from plasmid pVCERC-6 (27) using primers HindIII_cerulean-f and NdeI_cerulean-r. Restriction of both fragments with HindIII and NdeI and subsequent ligation of the individual fragments to each other resulted in the plasmid pEX4_iclceruleaniclds. Restriction of pEX4_iclceruleaniclds with SalI and BamHI and ligation into SalI/BamHI-cut pK18mobsacB yielded pK18mobsacB_ticlceruleaniclds (pTE1625).

For the heterologous expression of icl, icl was amplified from pTE1603 using primers NdeI_icl-for and HindIII_icl. The resulting product was digested with NdeI and HindIII and ligated into NdeI/HindIII-digested pET16b, yielding pET16b_icl (pTE1614).

**Genetic manipulation of *P. denitrificans***. The transfer of plasmids into *P. denitrificans* was achieved by biparental mating with the donor strain *E. coli* ST18 (28). Selection of the first integration was performed on LB or methanol mineral medium plates supplemented with 25 μg/ml kanamycin. Colonies were picked and restreaked on LB with kanamycin and LB with 3% sucrose in parallel. Colonies that were kanamycin resistant and sucrose sensitive were grown in plain LB for 2 days. Subsequently, cells were plated on methanol mineral medium plates supplemented with 6% sucrose in serial dilution. Colonies were restreaked on LB with kanamycin and LB with 3% sucrose in parallel. Kanamycin-sensitive, sucrose-resistant clones were screened for the successful deletion or integration of genes by colony PCR.

**Preparation of cell extracts.** *P. denitrificans* cultures were grown at 30°C in mineral medium supplemented with various carbon sources. Cells were harvested at mid-exponential phase, resuspended

---

**TABLE 3 Oligonucleotides used for plasmid construction and mutant verification in this study**

| Oligonucleotide       | Sequence                                      | Purpose                       |
|-----------------------|-----------------------------------------------|-------------------------------|
| NdeI_pKK21-f          | TATATCATATATGCCCCGGCCCGCCCGCATTG             | Generation of pTE1624         |
| pKK21-r               | TATATCTAGGGCGCCCGCCCGCCCGACTTACG             |                               |
| mCherry-f             | TATATCTAGGGCGCCCGCCCGCCCGACTTACG             |                               |
| NdeI_mCherry-r        | TATATCATATATGCCCCGGCGCCCGCCCGACTTACG         |                               |
| NdeI_pSYNiclXiclds-f  | TATATCATATGCCCCGGCGCCCGCCCGACTTACG           | Generation of pTE1625         |
| HindIII_pSYNicrXccrds-r | TATATCATATGCCCCGGCGCCCGCCCGACTTACG            |                               |
| HindIII_cerulean-f    | TATATCATATATGCCCCGGCGCCCGCCCGACTTACG         |                               |
| NdeI_icl-f            | TATATCATATGCCCCGGCGCCCGCCCGACTTACG           | Generation of pTE1614         |
| HindIII_icl-r         | TATATCATATGCCCCGGCGCCCGCCCGACTTACG           |                               |
| Pden_3873_ds-f        | GTCAGGCGCTTGTATTGCGCCGAAATG                 | Verification of TJE-KK5       |
| Pden_3873_ups-r       | GCGAAAGCGCCGACCTGTGCGGAAATATG               |                               |
| Pden_1363_ds-f        | CATATCTAGGGCGCCGACCTGTGCGGAAATG             | Verification of TJE-KK6       |
| Pden_1363_ups-r       | GCTGAGCCATATCTGTACGCTATGACGGAGTACTAC        |                               |
| Pden_1363_ds-f        | CATATCTAGGGCGCCGACCTGTGCGGAAATG             | Verification of TJE-KK10      |
| iclmCherryiclds_seq-f | CATATCTAGGGCGCCGACCTGTGCGGAAATG             |                               |
| Pden_1363_ds-f        | GACGAGCGCCGACCTGTGCGGAAATG                 |                               |
| Pden_1363_ups-r       | GCTGAGCCATATCTGTACGCTATGACGGAGTACTAC        |                               |
| NdeI_mCherry-r        | TATATCATATGCCCCGGCGCCCGCCCGACTTACG           | Verification of TJE-KK13      |
| Pden_3873_ups-r       | GCGAAAGCGCCGACCTGTGCGGAAATATG               |                               |
| 3’-ccr-f              | CAGGAGCGCCGACCTGTGCGGAAATG                 |                               |
| Pden_3873_ds-f        | GTCAGGCGCTTGTATTGCGCCGAAATG                 |                               |
| Pden_1363_ds-f        | CATATCTAGGGCGCCGACCTGTGCGGAAATG             |                               |
| Pden_1363_ups-r       | GCTGAGCCATATCTGTACGCTATGACGGAGTACTAC        |                               |
| HindIII_cerulean-f    | TATATCATATGCCCCGGCGCCCGCCCGACTTACG           |                               |

For the introduction of fluorescent reporter fusions, ccr and icl were ordered as synthetic genes fused to evoglowPp1 (24) and mCherry (25), respectively, each preceded by a 30-bp linker and followed by the downstream region of the respective genes, yielding plasmids pEX4_ccrevoglowPp1ccrds (pTE1601) and pEX4_iclceruleaniclds (pTE1603).

pTE1603 was digested with SalI and BamHI. Subsequent ligation into SalI/BamHI-cut pK18mobsacB yielded pK18mobsacB_ticlmCherryiclds (pTE1616).

pTE1601 was amplified with targeted omission of the evoglowPp1 gene via inverted PCR using primers NdeI_pKK21-f and pKK21-r. The mCherry gene was amplified from pTE1603 using primers mCherry-f and NdeI_mCherry-r. Restriction of both fragments with AvrII and NdeI and subsequent ligation of the individual fragments to each other resulted in the plasmid pEX4_ccrmCherryccrds.

pEX4_ccrmCherryccrds was digested with SalI and BamHI. Subsequent ligation into SalI/BamHI-cut pK18mobsacB yielded pK18mobsacB_tccrmCherryccrds (pTE1624).

pTE1603 was amplified with targeted omission of the mCherry gene via inverted PCR using primers NdeI_pSYNiclXiclds-f and HindIII_pSYNicrXccrds-r. The Cerulean gene (26) was amplified from plasmid pVCERC-6 (27) using primers HindIII_cerulean-f and NdeI_cerulean-r. Restriction of both fragments with HindIII and NdeI and subsequent ligation of the individual fragments to each other resulted in the plasmid pEX4_iclceruleaniclds. Restriction of pEX4_iclceruleaniclds with SalI and BamHI and ligation into SalI/BamHI-cut pK18mobsacB yielded pK18mobsacB_ticlceruleaniclds (pTE1625).

For the heterologous expression of icl, icl was amplified from pTE1603 using primers NdeI_icl-for and HindIII_icl. The resulting product was digested with NdeI and HindIII and ligated into NdeI/HindIII-digested pET16b, yielding pET16b_icl (pTE1614).

**Genetic manipulation of *P. denitrificans***. The transfer of plasmids into *P. denitrificans* was achieved by biparental mating with the donor strain *E. coli* ST18 (28). Selection of the first integration was performed on LB or methanol mineral medium plates supplemented with 25 μg/ml kanamycin. Colonies were picked and restreaked on LB with kanamycin and LB with 3% sucrose in parallel. Colonies that were kanamycin resistant and sucrose sensitive were grown in plain LB for 2 days. Subsequently, cells were plated on methanol mineral medium plates supplemented with 6% sucrose in serial dilution. Colonies were restreaked on LB with kanamycin and LB with 3% sucrose in parallel. Kanamycin-sensitive, sucrose-resistant clones were screened for the successful deletion or integration of genes by colony PCR.

**Preparation of cell extracts.** *P. denitrificans* cultures were grown at 30°C in mineral medium supplemented with various carbon sources. Cells were harvested at mid-exponential phase, resuspended...
in ice-cold morpholinepropanesulfonic acid (MOPS)-KOH buffer (100 mM, pH 7.2), and lysed by sonication
(MS 72 microtip, 40%, 15 pulses three times). Cell debris was removed by centrifugation at 35,000 × g and 4°C for 1 h. The total protein concentration of the cell extracts was determined with the Bradford assay (29) using bovine serum albumin (BSA) as the standard. The catalytic activities of crotonyl-CoA carboxylase/reductase and isocitrate lyase were measured spectrophotometrically as described previously (3, 30).

**Synthesis of crotonyl-CoA.** Crotonyl coenzyme A was synthesized from its anhydride as described before (31).

**Live-cell imaging.** To monitor gene expression in vivo, cells were immobilized on 1% agarose pads and analyzed microscopically using an Axio Observer.Z1 (Zeiss) microscope equipped with a Plan Apochromat 100×/1.4 oil Ph3 phase-contrast objective, an ET-mCherry filter set (Chroma, USA), and a pco-edge sCMOS camera (PCO AG, Germany). Images were recorded with VisiView 3.3.0.6 (Visitron Systems, Puchheim, Germany) and processed with Fiji (32) and Adobe IllustratorCS5 (Adobe Systems, USA). For time-lapse imaging, cells were transferred to a B04 CellASIC ONIX2 microfluidic plate (Merck, Germany) and cultivated at 30°C under continuous medium flow (1 lb/in²) and imaged at regular intervals using the microscope setup described above. For automated data analysis, images were processed with Fiji (32) and BacStalk (10).

**Heterologous production and purification of 6×His-Icl.** Competent *E. coli* BL21 cells were transformed with the expression plasmid pET16b_Icl and grown at 37°C in terrific broth (23.6 g/liter yeast extract, 11.8 g/liter tryptone, 9.4 g/liter K2HPO4, 2.2 g/liter KH2PO4, and 4% [vol/vol] glycerol) supplemented with ampicillin. Gene expression was induced at an optical density at 600 nm (OD600) of 0.8 by addition of 0.5 mM isopropyl thiogalactopyranoside (IPTG). After additional growth at 18°C for 12 h, cells were harvested, resuspended in 3 volumes of buffer A (500 mM NaCl, 50 mM Tris-HCl, pH 7.9), followed by sonication. Lysates were cleared by centrifugation at 42,000 × g and 4°C for 45 min. The supernatant was applied onto a preequilibrated 1-ml Ni-Sepharose fast-flow column (HiTrap, FF; GE Life Sciences, United Kingdom) and washed with buffer A. Proteins were eluted from the column by the addition of increasing concentrations of buffer B (500 mM NaCl, 50 mM Tris-HCl, 500 mM imidazole, pH 7.9), followed by buffer exchange to storage buffer (buffer A, 10% glycerol). Protein concentration was determined using a NanoDrop 2000 spectrometer (Thermo Scientific, USA), and purity was analyzed by SDS-PAGE according to the procedures in reference 33.

**Measurement of enzyme activities in cell extracts and purified protein. (i) Ccr.** The enzyme activity of crotonyl-CoA carboxylase/reductase was measured by monitoring the consumption of NADPH spectrophotometrically as described previously (3).

(ii) Icl. The enzyme activity of isocitrate lyase was measured by monitoring the production of a phenylhydrazine-glyoxylate complex as described previously (30). For the Icl inhibition assay with 3-nitropropionate (3-NPA), the compound was preequilibrated in 100 mM MOPS-KOH, pH 7.2, overnight. Purified isocitrate lyase was incubated in the reaction mixture containing 3-NPA at various concentrations at 30°C for 10 min before the assay was started by the addition of 2 mM D,L-isocitric acid.

**Phylogenetic analysis.** Sequences of 12 proteins (GapA, GyrA, RecA, RpoA, RpoB, TrpB, 30S ribosomal protein S2, 30S ribosomal protein S3, 30S ribosomal protein S12, 50S ribosomal protein L2, 50S ribosomal protein L3, 50S ribosomal protein L11) from the core proteome of 48 strains of *Paracoccus* were downloaded from the IMG database (34) and aligned using MUSCLE 3.8.31 (35). The alignments were concatenated and used to calculate a phylogenetic tree with MEGA 7.0.14 (36) using the maximum-likelihood method with the Le-Gascuel substitution model and 100 bootstrap replicates. The resulting tree was visualized using iTOL (37).

Sequences of Ccr and Icl from 43 and 14 strains of *Paracoccus*, respectively, were downloaded from the IMG database and used to calculate maximum-likelihood phylogenetic trees as described above.

**SUPPLEMENTAL MATERIAL**

Supplemental material for this article may be found at https://doi.org/10.1128/mBio.00805-19.

**FIG S1,** TIF file, 2.2 MB.

**FIG S2,** TIF file, 1 MB.

**FIG S3,** TIF file, 1.4 MB.

**FIG S4,** TIF file, 2 MB.

**FIG S5,** TIF file, 2.6 MB.

**FIG S6,** TIF file, 2.1 MB.

**FIG S7,** TIF file, 2.6 MB.

**FIG S8,** TIF file, 1.8 MB.

**MOVIE S1,** AVI file, 3.4 MB.

**ACKNOWLEDGMENTS**

This work was supported by funds from the Germany Research Foundation (project 192445154-SFB 987 to T.J.E. and M.T.). M.C.F.V.T. acknowledges support from the
European Molecular Biology Organization (EMBO long-term fellowship ALTF 1396-2015).

K.K., I.B., L.S.V.B., and T.J.E. conceptualized the project. R.J.M.V.S. and A.J.G. contributed by giving fundamental advice in molecular biology and training in genetic manipulation of Paracoccus denitrificans. K.K. and T.J.E. designed the study. K.K. performed all genetic and biochemical experiments and analyses. K.K., M.C.F.V.T., and M.T. planned and performed all microscopic analyses. K.K. and M.N.P.-E. analyzed the microscopic images of the time-lapse experiments. K.K. and L.S.V.B. performed homology and phylogenetic analyses. K.K. and T.J.E. wrote the manuscript.

We declare no competing interests.

REFERENCES

1. Kornberg HL, Madsen NB. 1957. Synthesis of C4-dicarboxylic acids from acetate by a glyoxylate bypass of the tricarboxylic acid cycle. Biochim Biophys Acta 24:651–653. https://doi.org/10.1016/0006-3002(57)90268-8.

2. Kornberg HL, Krebs HA. 1957. Synthesis of cell constituents from C2-units by a modified tricarboxylic acid cycle. Nature 179:988–991. https://doi.org/10.1038/179988a0.

3. Ebright RJ, Berg JA, Brecht V, Muller M, Fuchs G, Alber BE. 2007. Synthesis of C5-dicarboxylic acids from C2-units involving crotonyl-CoA carboxylase/reductase: the ethylmalonyl-CoA pathway. Proc Natl Acad Sci U S A 104:10631–10636. https://doi.org/10.1073/pnas.0702791104.

4. Ebright RJ, Fuchs G, Alber BE. 2009. (2S)-Methylsuccinyl-CoA dehydrogenase closes the ethylmalonyl-CoA pathway for acetyl-CoA assimilation. Mol Microbiol 73:992–1008. https://doi.org/10.1111/j.1365-2958.2009.06837.x.

5. Ebright RJ, Brecht V, Fuchs G, Muller M, Alber BE. 2009. Carboxylation mechanism and stereochemistry of crotonyl-CoA carboxylase/reductase, a carboxylating enoyl-thioester reductase. Proc Natl Acad Sci U S A 106:8871–8876. https://doi.org/10.1073/pnas.0903931096.

6. Claassen PAM, van den Heuvel MHMJ, Zehnder AJB. 1987. Enzyme profiles of Thiobacillus-versutus after aerobic and denitrifying growth—regulation of isocitrate lyase. Arch Microbiol 147:30–36. https://doi.org/10.1007/BF00492901.

7. Claassen PAM, Zehnder AJB. 1986. Isocitrate lyase activity in Thiobacillus-versutus grown anaerobically on acetate and nitrate. J Gen Microbiol 132:3179–3185. https://doi.org/10.1099/00221287-132-131-3179.

8. Lee YW, Wahab HA, Choong YS. 2015. Potential inhibitors for isocitrate lyase of Mycobacterium tuberculosis and non-M. tuberculosis: a summary. Biomed Res Int 2015:895453. https://doi.org/10.1155/2015/895453.

9. Schloss JV, Cieland WW. 1992. Inhibition of isocitrate lyase by 3-nitrocinnamaldehyde: a reaction-intermediate analogue. Biochemistry 21:4420–4427. https://doi.org/10.1021/bi00261a035.

10. Hartmann R, van Teesseling MCF, Thanbichler M, Drescher K. 2018. BacStalk: a comprehensive and interactive image analysis software tool for bacterial cell biology. bioRxiv https://doi.org/10.1101/360230.

11. Peyraud R, Kiefer P, Christen P, Massou S, Portais JC, Vorholt JA. 2009. Demonstration of the ethylmalonyl-CoA pathway by using 13C metabolomics. Proc Natl Acad Sci U S A 106:4846–4851. https://doi.org/10.1073/pnas.0810932106.

12. Sambrook J, Russel DW. 2001. Molecular cloning: a laboratory manual. Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY.

13. Schauer A, Tauch A, Jager W, Kalinowski J, Thierbach G, Puhler A. 1994. Small mobilizable multi-purpose cloning vectors derived from the alkaliphilic bacterium Sporosarcina ureae. Gene 145:69–73. https://doi.org/10.1016/0378-1119(94)90324-7.

14. Drepper T, Eggert T, Circolone F, Heck A, Krauss U, Guter JL, Wendorff M, Losi A, Gartner W, Jaeger KE. 2007. Reporter proteins for in vivo fluorescence without oxygen. Nat Biotechnol 25:443–445. https://doi.org/10.1038/nbt1293.

15. Sharer NC, Campbell RE, Steinhbach PA, Giepmans BN, Palmer AE, Tsien RY. 2004. Improved monomeric red, orange and yellow fluorescent proteins derived from Discosoma sp red fluorescent protein. Nat Biotechnol 22:1567–1572. https://doi.org/10.1038/nbt1037.

16. Rizzo MA, Springer GH, Granada B, Piston DW. 2004. An improved cyan fluorescent protein variant useful for FRET. Nat Biotechnol 22:445–449. https://doi.org/10.1038/nbt945.

17. Thanbichler M, Iniesta AA, Shapiro L. 2007. A comprehensive set of proteins derived from Discosoma sp red fluorescent protein. Nat Biotechnol 25:443–445. https://doi.org/10.1038/nbt1293.

18. Balasubramanian S, Sauer U, Pethe K, McKinney JD. 2016. A rheostat mechanism governs the bifurcation of carbon flux in mycobacteria. Nat Commun 7:12527. https://doi.org/10.1038/ncomms12527.

19. Beijerinck M, Minkman D. 1910. Bildung und Verbrauch von Stickoxydul durch Bakterien. Zentralbl Bakteriol Parasitenk Infektionskr Hyg Abt II 7:125–132.

20. Thanbichler M, Iniesta AA, Shapiro L. 2007. A comprehensive set of proteins derived from Discosoma sp red fluorescent protein. Nat Biotechnol 25:443–445. https://doi.org/10.1038/nbt1293.

21. Thanbichler M, Iniesta AA, Shapiro L. 2007. A comprehensive set of proteins derived from Discosoma sp red fluorescent protein. Nat Biotechnol 25:443–445. https://doi.org/10.1038/nbt1293.

22. Thanbichler M, Iniesta AA, Shapiro L. 2007. A comprehensive set of proteins derived from Discosoma sp red fluorescent protein. Nat Biotechnol 25:443–445. https://doi.org/10.1038/nbt1293.

23. Schafer A, Tauch A, Jager W, Kalinowski J, Thierbach G, Puhler A. 1994. Small mobilizable multi-purpose cloning vectors derived from the alkaliphilic bacterium Sporosarcina ureae. Gene 145:69–73. https://doi.org/10.1016/0378-1119(94)90324-7.

24. Drepper T, Eggert T, Circolone F, Heck A, Krauss U, Guter JL, Wendorff M, Losi A, Gartner W, Jaeger KE. 2007. Reporter proteins for in vivo fluorescence without oxygen. Nat Biotechnol 25:443–445. https://doi.org/10.1038/nbt1293.

25. Balasubramanian S, Sauer U, Pethe K, McKinney JD. 2016. A rheostat mechanism governs the bifurcation of carbon flux in mycobacteria. Nat Commun 7:12527. https://doi.org/10.1038/ncomms12527.

26. Beijerinck M, Minkman D. 1910. Bildung und Verbrauch von Stickoxydul durch Bakterien. Zentralbl Bakteriol Parasitenk Infektionskr Hyg Abt II 7:125–132.

27. Thanbichler M, Iniesta AA, Shapiro L. 2007. A comprehensive set of proteins derived from Discosoma sp red fluorescent protein. Nat Biotechnol 25:443–445. https://doi.org/10.1038/nbt1293.

28. Thoma S, Schobert M. 2009. An improved Escherichia coli donor strain for diparental mating. FEMS Microbiol Lett 294:127–132. https://doi.org/10.1111/j.1574-6968.2009.01556.x.

29. Thanbichler M, Iniesta AA, Shapiro L. 2007. A comprehensive set of proteins derived from Discosoma sp red fluorescent protein. Nat Biotechnol 25:443–445. https://doi.org/10.1038/nbt1293.

30. Thanbichler M, Iniesta AA, Shapiro L. 2007. A comprehensive set of proteins derived from Discosoma sp red fluorescent protein. Nat Biotechnol 25:443–445. https://doi.org/10.1038/nbt1293.

31. Thanbichler M, Iniesta AA, Shapiro L. 2007. A comprehensive set of proteins derived from Discosoma sp red fluorescent protein. Nat Biotechnol 25:443–445. https://doi.org/10.1038/nbt1293.
map to the synthesis of CoA esters. Molecules 21:517. https://doi.org/10.3390/molecules21040517.

32. Schindelin J, Arganda-Carreras I, Frise E, Kaynig V, Longair M, Pietzsch T, Preibisch S, Rueden C, Saalfeld S, Schmid B, Tinevez JY, White DJ, Hartenstein V, Elceiri K, Tomancak P, Cardona A. 2012. Fiji: an open-source platform for biological-image analysis. Nat Methods 9:676–682. https://doi.org/10.1038/nmeth.2019.

33. Laemmli UK. 1970. Cleavage of structural proteins during the assembly of the head of bacteriophage T4. Nature 227:680–685. https://doi.org/10.1038/227680a0.

34. Markowitz VM, Chen IMA, Palaniappan K, Chu K, Szeto E, Grechkin Y, Ratner A, Jacob B, Huang JH, Williams P, Huntemann M, Anderson I, Mavromatis K, Ivanova NN, Kyrpides NC. 2012. IMG: the integrated microbial genomes database and comparative analysis system. Nucleic Acids Res 40:D115–D122. https://doi.org/10.1093/nar/gkr1044.

35. Edgar RC. 2004. MUSCLE: multiple sequence alignment with high accuracy and high throughput. Nucleic Acids Res 32:1792–1797. https://doi.org/10.1093/nar/gkh340.

36. Kumar S, Stecher G, Tamura K. 2016. MEGA7: Molecular Evolutionary Genetics Analysis version 7.0 for bigger datasets. Mol Biol Evol 33:1870–1874. https://doi.org/10.1093/molbev/msw054.

37. Letunic I, Bork P. 2016. Interactive tree of life (iTOL) v3: an online tool for the display and annotation of phylogenetic and other trees. Nucleic Acids Res 44:D115–D122. https://doi.org/10.1093/nar/gkw190.

38. Remus-Emsermann MN, Gisler P, Drissner D. 2016. MiniTn7-transposon delivery vectors for inducible or constitutive fluorescent protein expression in Enterobacteriaceae. FEMS Microbiol Lett 363:fnw178. https://doi.org/10.1093/femsle/fnw178.

39. Peel D, Quayle JR. 1961. Microbial growth on C1 compounds. I. Isolation and characterization of Pseudomonas AM 1. Biochem J 81:465–469. https://doi.org/10.1042/bj0810465.