Metabolic fluxes in the central carbon metabolism of *Dinoroseobacter shibae* and *Phaeobacter gallaeciensis*, two members of the marine Roseobacter clade

Tobias Fürch¹, Matthias Preusse¹, Jürgen Tomasch², Hajo Zech³, Irene Wagner-Döbler², Ralf Rabus³ and Christoph Wittmann*¹

Address: ¹Institute of Biochemical Engineering, Technische Universität Braunschweig, Gaußstraße 17, D-38106 Braunschweig, Germany, ²Helmholtz Centre for Infection Research, Research Group Microbial Communication, D-38124 Braunschweig, Germany and ³Institute for Chemistry and Biology of the Marine Environment (ICBM), University of Oldenburg, D-26111 Oldenburg, Germany

Email: Tobias Fürch - t.fuerch@tu-bs.de; Matthias Preusse - m.preusse@tu-bs.de; Jürgen Tomasch - jtomasch@gmx.de; Hajo Zech - hajo.zech@icbm.de; Irene Wagner-Döbler - Irene.Wagner-Doebler@helmholtz-hzi.de; Ralf Rabus - rabus@mpi-bremen.de; Christoph Wittmann* - c.wittmann@tu-bs.de

* Corresponding author

Abstract

**Background:** In the present work the central carbon metabolism of *Dinoroseobacter shibae* and *Phaeobacter gallaeciensis* was studied at the level of metabolic fluxes. These two strains belong to the marine Roseobacter clade, a dominant bacterial group in various marine habitats, and represent surface-associated, biofilm-forming growth (*P. gallaeciensis*) and symbiotic growth with eukaryotic algae (*D. shibae*). Based on information from recently sequenced genomes, a rich repertoire of pathways has been identified in the carbon core metabolism of these organisms, but little is known about the actual contribution of the various reactions in vivo.

**Results:** Using ¹³C labelling techniques in specifically designed experiments, it could be shown that glucose-grown cells of *D. shibae* catabolise the carbon source exclusively via the Entner-Doudoroff pathway, whereas alternative routes of glycolysis and the pentose phosphate pathway are obviously utilised for anabolic purposes only. Enzyme assays confirmed this flux pattern and link the lack of glycolytic flux to the absence of phosphofructokinase activity. The previously suggested formation of phosphoenolpyruvate from pyruvate during mixotrophic CO₂ assimilation was found to be inactive under the conditions studied. Moreover, it could be shown that pyruvate carboxylase is involved in CO₂ assimilation and that the cyclic respiratory mode of the TCA cycle is utilised. Interestingly, the use of intracellular pathways was highly similar for *P. gallaeciensis*.

**Conclusion:** The present study reveals the first insight into pathway utilisation within the Roseobacter group. Fluxes through major intracellular pathways of the central carbon metabolism, which are closely linked to the various important traits found for the Roseobacter clade, could be determined. The close similarity of fluxes between the two physiologically rather different species might provide the first indication of more general key properties among members of the Roseobacter clade which may explain their enormous success in the marine realm.
Background
The Roseobacter lineage, representing a group of Alphaproteobacteria [1], is found in various marine habitats where it is present in high abundance, comprising up to 25% of the total bacterial community [2]. Overall, the diverse metabolic properties of the Roseobacter clade and its ubiquitous occurrence in marine ecosystems suggest that members of this clade play an important role in global biogeochemical processes such as cycling of carbon or sulphur [3]. Members of the Roseobacter clade participate in DMSP demethylation [4], the oxidation of carbon monoxide [5] and degradation of aromatic compounds [6,7]. Typically, they use external organic substrates as carbon sources [8]. Of outstanding interest is the fact that they are able to generate energy from light (aerobic anoxygenic phototrophy) [9] and thus contribute significantly to phototrophic energy generation [10,11]. All these important traits are linked to the core part of central carbon metabolism involved in the breakdown of nutrients and the supply of metabolites and energy for various cellular requirements. Recent efforts in genome sequencing and annotation of Roseobacter members have provided a first insight into the repertoire of underlying metabolic reactions available (Figure 1) and have led to different suggestions for possible pathways that might be involved in important physiological functions [12]. As an example, a mixotrophic CO₂ assimilation pathway has been proposed for R. denitrificans, in which CO₂ is fixed either (i) via the combined action of pyruvate-orthophosphate dikinase and phosphoenolpyruvate carboxylase or (ii) via pyruvate carboxylase [13]. For glucose catabolism, up to three alternative routes are encoded in the genome: glycolysis, the pentose phosphate pathway and the Entner-Doudoroff pathway. At this point, it seems highly relevant to study the contribution of these potential pathways to the metabolism of bacteria in the Roseobacter clade to improve our understanding of their physiology. Our current knowledge of the in vivo fluxes through intracellular pathways among the Roseobacter lineage is still very limited.

To address this issue, we applied metabolic flux analysis using ¹³C labelled isotopes to gain a first insight into the central catabolic pathways of Dinoroseobacter shibae DFL12 [1] and Phaeobacter gallaeciensis DSM 17395 [14]. These species represent two prominent members of the Roseobacter clade. P. gallaeciensis has received strong interest due to its ability to produce the antibiotic tropodithietic acid. D. shibae was isolated as a novel species from marine dinoflagellates and lives in a symbiotic relationship with eukaryotic algae [15]. Metabolic flux analysis using ¹³C labelled isotopes has proven a key technology in the unravelling of metabolic pathways and has recently been used to study different microorganisms mainly linked to biotechnological production processes [16-19]. No such study has yet been performed for members of the Roseobacter clade.

Results and Discussion
Cultivation profile
The cultivation profile of D. shibae on defined medium with glucose as the sole carbon source is displayed in Figure 2. After an initial adaptation phase, cells grew exponentially with a constant specific growth rate of 0.11 h⁻¹.
Pathways for glucose catabolism

The carbon core metabolism of *D. shibae* and *P. gallaeciensis* consists of three potential routes for glucose catabolism. Glucose can be alternatively catabolised via glycolysis (EMP), the pentose phosphate pathway (PPP) and the Entner-Doudoroff pathway (EDP). The use of [1-13C] glucose by each individual pathway leads to a different labelling pattern in specific fragments of alanine and serine, which can be taken as a clear differentiation of flux (Figure 3). For *D. shibae* the corresponding [M-57] fragment of serine did not show any enrichment of 13C but rather reflected the pattern resulting from the natural abundance of 13C only (Table 1). Any contribution of glycolysis to formation of this metabolite and its precursor 3-phosphoglycerate can therefore be excluded as this would lead to enrichment of 13C at the C4 position, yielding a higher fraction of M+1 labelled molecules of Ser. Thus glycolytic flux obviously was not present. The two remaining possibilities, the PPP and the ED pathway, can be differentiated by the labelling pattern of alanine, which represents the pyruvate pool in the cell. The high enrichment of 13C label in the [M-57] fragment of alanine indicates a large contribution of the ED pathway, since formation via the PPP would lead to non-labelled alanine. The [M-85] fragment of alanine, comprising only the carbon atoms C2 and C3 of pyruvate, was not enriched in 13C, showing that pyruvate was labelled only at its C4 position. This perfectly matches the isotope pattern expected for the ED pathway, whereas glycolytic flux would have resulted in label enrichment at the C3 position, and further confirms the flux distribution.

Interestingly, *P. gallaeciensis* showed almost identical characteristics and obviously also uses mainly the ED pathway during growth on glucose. The quantification of relative flux (Eqs. 2 and 3) revealed that the use of the ED pathway amounts to >99%, whereas glycolysis and PPP contribute only <1% (Table 2). Compared to other microorganisms such as *E. coli* [20], *B. subtilis* [21], *B. megaterium* [18] or *C. glutamicum* [22] grown on glucose, this is a rather unusual flux pattern. Most organisms use glycolysis and the pentose phosphate pathway concomitantly but at varying ratios (Table 2). Exclusive utilisation of the ED pathway, as found here, has been previously observed in selected species of *Pseudomonas* or *Arthrobacter* where this behaviour was attributed to a lack of phosphofructokinase [23,24]. Among the two microorganisms studied, *D. shibae* does contain a gene encoding for this enzyme, whereas *P. gallaeciensis* does not. For both *Roseobacter* species, in contrast to *E. coli* as positive control, phosphofructokinase activity could not be detected, clearly explaining the lack of glycolytic flux (Figure 4B). While this matches with the genomic repertoire of *P. gallaeciensis*, we conclude at this stage that the phosphofructokinase in *D. shibae* is either not expressed, might have another function or even is a non-functional protein. The flux pattern for both organisms is supported by enzymatic assays showing high in vitro activity of 6-phosphogluconate dehydrogenase and 2-dehydro-3-deoxyporphogluconate aldolase, the two key enzymes in the Entner-Doudoroff pathway (Figure 4A).

Pathways for PEP synthesis - contribution of pyruvate-orthophosphate dikinase and phosphoenolpyruvate carboxykinase

Based on the labelling data given above, the formation of PEP from pyruvate by pyruvate-orthophosphate dikinase or via pyruvate carboxylase and phosphoenolpyruvate carboxykinase would result in the presence of PEP with 13C enrichment at position C1. However, the [B02] fragments of Phe and Tyr, each corresponding to the carbon atoms C3 and C2 of PEP, do not show significant enrichment of 13C (Table 1). The same holds for the [M-57] fragment, which corresponds to the entire carbon skeleton of Phe and Tyr and thus all precursors, that is, PEP and E4P. Flux quantification using Equations 4 and 5 confirms that PEP is solely synthesised by the reactions of lower glycolysis (Table 2). This is an interesting finding with respect to the recently suggested mixotrophic CO2 assimilation pathway for some members of the *Roseobacter* clade, which also involves the potential contribution of pyruvate orthophosphate dikinase (PDPK) [13]. Despite the putative gene for this protein also being annotated for the species investigated here, we could clearly
The theoretical labelling pattern of the C\textsubscript{3} pool (GAP and PYR) derived from 99\% [1-\textsuperscript{13}C] glucose depending on activities in the carbon core metabolism. The three major routes for glucose catabolism are presented: glycolysis (solid lines), the pentose phosphate pathway (dashed lines) and the Entner-Doudoroff pathway (dotted lines). White circles represent unlabelled (\textsuperscript{12}C) carbon whereas black circles indicate labelled (\textsuperscript{13}C) carbon. The numbers reflect the position of the carbon atom within the molecule. Ala: alanine; G6P: glucose 6-phosphate; 6PG: 6-phosphogluconate; KDPG: 2-keto-3-deoxy-6-phosphogluconate; F6P: fructose 6-phosphate; R5P: ribose 5-phosphate; GAP: glyceraldehyde 3-phosphate; PYR: pyruvate.
demonstrate that the formation of PEP from PYR is not active in vivo under the conditions studied.

Pathways for oxaloacetate synthesis - contribution of CO₂ assimilation and oxidative TCA cycle

Oxaloacetate as a central metabolite can be formed by two major pathways, that is, carboxylation involving pyruvate carboxylase or via pyruvate dehydrogenase and the energy-generating reactions of the TCA cycle. The following data clearly suggest that both pathways are active simultaneously in the two Roseobacters. For the experimental setup chosen and carbon transfer in the underlying metabolic reactions, the carboxylation of pyruvate is the only reaction that leads to ¹³C labelled oxaloacetate (Figure 5). The label can be present in carbon positions C₁ or C₄, whereby single- or double-labelled molecules can be formed, depending on the incorporation of ¹²CO₂ or ¹³CO₂. In contrast, the alternative route via the cyclic respiratory mode of the TCA cycle yields exclusively non-labelled oxaloacetate. In all possible cases the labelled carbon atoms from either pyruvate or oxaloacetate are released in the decarboxylation steps of the TCA cycle as ¹³CO₂. Inspection of the labelling pattern of aspartate, corresponding to the oxaloacetate backbone, immediately

Table 1: Selected TBDMS-α-amino acid fragments used in the study derived from D. shibae and P. gallaeciensis grown on 99% [¹³C]glucose.

<table>
<thead>
<tr>
<th>Fragment</th>
<th>C-Atoms</th>
<th>Mass isotopomer distribution (% of total pool)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>D. shibae</td>
</tr>
<tr>
<td></td>
<td></td>
<td>M+0</td>
</tr>
<tr>
<td>Ala M-57</td>
<td>1-3</td>
<td>50.0 ± 0.2</td>
</tr>
<tr>
<td>Ala M-85</td>
<td>2-3</td>
<td>96.8 ± 0.1</td>
</tr>
<tr>
<td>f302 M-1-2</td>
<td>51.2 ± 0.1</td>
<td>48.2 ± 0.1</td>
</tr>
<tr>
<td>Asp M-57</td>
<td>1-4</td>
<td>72.4 ± 0.7</td>
</tr>
<tr>
<td>Asp M-85</td>
<td>2-4</td>
<td>83.3 ± 0.6</td>
</tr>
<tr>
<td>f302 M-1-2</td>
<td>82.1 ± 0.3</td>
<td>17.6 ± 0.3</td>
</tr>
<tr>
<td>Glu M-57</td>
<td>1-5</td>
<td>80.7 ± 0.3</td>
</tr>
<tr>
<td>Glu M-85</td>
<td>2-5</td>
<td>92.1 ± 0.2</td>
</tr>
<tr>
<td>f302 M-1-2</td>
<td>83.4 ± 0.2</td>
<td>16.2 ± 0.2</td>
</tr>
<tr>
<td>Gly M-57</td>
<td>1-2</td>
<td>96.1 ± 0.0</td>
</tr>
<tr>
<td>Gly M-85</td>
<td>2</td>
<td>98.8 ± 0.1</td>
</tr>
<tr>
<td>Phe M-57</td>
<td>1-9</td>
<td>85.7 ± 0.6</td>
</tr>
<tr>
<td>f302 M-1-2</td>
<td>95.9 ± 0.3</td>
<td>4.1 ± 0.3</td>
</tr>
<tr>
<td>Ser M-57</td>
<td>1-3</td>
<td>95.3 ± 0.3</td>
</tr>
<tr>
<td>Ser M-85</td>
<td>2-3</td>
<td>97.7 ± 0.1</td>
</tr>
<tr>
<td>f302 M-1-2</td>
<td>95.6 ± 0.0</td>
<td>3.9 ± 0.0</td>
</tr>
<tr>
<td>Tyr M-57</td>
<td>1-9</td>
<td>86.2 ± 0.7</td>
</tr>
<tr>
<td>f302 M-1-2</td>
<td>96.1 ± 0.2</td>
<td>3.9 ± 0.2</td>
</tr>
</tbody>
</table>

* tert-butyldimethylsilyl; fragmentation patterns are described elsewhere [27]

Table 2: Comparison of catabolic pathway activity and origins of metabolic intermediates in central carbon metabolism of D. shibae, P. gallaeciensis and other bacteria derived from carbon labelling experiments.

<table>
<thead>
<tr>
<th>Pathway activity/Fractional pool composition [%]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glycolysis</td>
</tr>
<tr>
<td>PPP</td>
</tr>
<tr>
<td>ED pathway</td>
</tr>
<tr>
<td>PEP from PYR</td>
</tr>
<tr>
<td>PEP from OAA</td>
</tr>
</tbody>
</table>

* this study
n.a. = not available in the organism
shows that single- and double-labelled mass isotopomers are present in significant amounts for *D. shibae* and *P. galaeiensis*, indicating *in vivo* activity of pyruvate carboxylase in both strains (Table 1). However, the relative fractions of these $^{13}$C-enriched mass isotopomers are relatively small, excluding sole contribution of this reaction to oxaloacetate synthesis. The dominant fraction consists of non-labelled molecules, obviously derived via the oxidative TCA cycle. We thus conclude that the cyclic respiratory mode of the TCA cycle is active *in vivo* in both strains. For *D. shibae*, which possesses a photosystem for energy generation, this mode might display an important strategy to derive energy under conditions where the photosystem is not active, for example, during the night or in deeper water regions.

**Conclusion**

Being one of the first metabolic studies of members of the Roseobacter clade using the $^{13}$C labelling experimental approach, a deeper insight into the activity of the important metabolic routes of *D. shibae* and *P. galaeiensis* was achieved. Interestingly, the use of intracellular pathways is highly similar in the studied species *D. shibae* and *P. galaeiensis*. This stands in surprising contrast to the overall differences in phenotypic behaviour exhibited by these two strains, since *D. shibae* is an algal-associated microorganism whereas *P. galaeiensis* is free-living in marine habitats. However, this may be a first indication of more general key properties among members of the Roseobacter clade that explain their enormous success in the marine realm.

**Methods**

**Strains, medium and growth conditions**

The strains used in this study are the genome sequenced strains *Dinoroseobacter shibae* DFL12 [1] and *Phaeobacter galaeiensis* DSM 17395 [14]. For cultivation of both strains a defined, synthetic seawater medium (minimal medium) was used [25], containing the following compo-
ments per litre of medium: 4.0 g NaSO₄, 0.2 g KH₂PO₄, 0.25 g NH₄Cl, 20.0 g NaCl, 3.0 g MgCl₂·6 H₂O, 0.5 g KCl and 0.15 g CaCl₂·2 H₂O, 0.19 g NaHCO₃. 1 ml trace element solution and 10 ml vitamin solution. The final glucose concentration in the medium was in the range of 0.4 to 0.9 g l⁻¹. The trace element solution contained 2.1 g Fe(SO₄)·7 H₂O, 13 ml 25% (v/v) HCl, 5.2 g Na₂EDTA·2 H₂O, 30 mg H₂BO₃, 0.1 g MnCl₂·4 H₂O, 0.19 g CoCl₂·6 H₂O, 2 mg CuCl₂·2 H₂O, 0.144 g ZnSO₄·7 H₂O and 36 mg Na₂MoO₄·2 H₂O per litre. The vitamin solution for _D. shibae_ contained the following components per litre: 0.2 g biotin, 2.0 g nicotinic acid and 0.8 g 4-aminobenzoic acid. All solutions were sterilised separately and mixed at room temperature prior to inoculation. For carbon labelling experiments 99% [1-¹⁳C] glucose (Euriso-Top, Saint-Aubin, France) was used as substrate. The cultivations were carried out on orbital shakers at 200 rpm in 500 ml shaken flasks with a culture volume of 50 ml at 37°C ( _D. shibae_ ) and 28°C ( _P. gallaeciensis_ ). To ensure comparable conditions between the two microorganisms and avoid any potential influencing effects of phototrophy in _D. shibae_, both organisms were cultivated in the light. Under these conditions, no bacteriochlorophyll is synthesised and therefore no active photosystem is present that might affect energy metabolism and thus induce changes in the fluxes through the main metabolic pathways.

**Analytics**

Cell concentration was monitored by measuring the optical density (OD) at 600 nm or by gravimetry [26]. The ¹³C labelling pattern of the amino acids contained in the cell protein was determined as follows [27]. Cells were harvested during exponential growth phase at half-maximal optical density including a washing step in 0.9% NaCl solution, followed by lyophilisation. Subsequently, 4 mg of lyophilised cells was resuspended in 200 µl of 6 M HCl and incubated at 110°C for 24 h. The obtained hydrolysate was neutralised by addition of 6 M NaOH and cleared of insoluble matter (0.2 µm centrifugal filter device Ultrafree MC, Millipore, Bedford, MA, USA). Subsequently, 50 µl of the hydrolysate was transferred to a 2 ml sample vial, lyophilised and derivatised at 80°C for 60 min with 50 µl N, N-dimethylformamide (Carl Roth, Karlsruhe, Germany) containing 0.1% (v/v) pyridine and 50 µl N-methyl-( tert-butyl(dimethyl)silyl)-trifluoroacetamide (MBDSTFA, Macherey-Nagel, USA). GC/MS measurements were carried out as described earlier [27]. Subsequent MS data processing was carried out according to Fürch et al. [18] and Lee et al. [28,29].

**Preparation of cell extracts for enzyme activity measurements**

Cells were harvested by centrifugation at 10,000 g for 10 min, washed twice with 100 mM Tris-HCl (pH 7.0) containing 20 mM KCl, 5 mM MnSO₄, 2 mM DTT and 0.1 mM EDTA, and then resuspended in the same buffer. Afterwards the cells were disrupted by sonication for 1 min using an ultrasonic disrupter (Sonifier W250 D, Branson, Danbury, USA) with an amplitude of 30%. Cell debris was removed by centrifugation. The resulting crude cell extract was immediately used to determine specific enzyme activity. All operations were carried out on ice.

**Enzyme assays**

Enzyme activities in crude cell extract were measured spectrophotometrically. All compounds of the reaction mixture were pipetted into a cuvette with a 1 cm light path and reactions were initiated by adding the cell extract or substrate respectively. The total protein concentration of the crude cell extract was determined using RotiQuant (Carl Roth GmbH, Karlsruhe, Germany). The overall activity of 6-phosphogluconate dehydratase (EDD) and 2-dehydro-3-deoxyphosphogluconate aldolase (EDA) was measured using a two-step reaction [30]. For this purpose 0.8 µmol 6-phosphogluconate, 1 µmol MgCl₂, 5 µmol Tris-HCl buffer (pH 7.65) and 100 µl of extract were incubated in a total volume of 0.5 ml for 5 min at room temperature. The reaction was stopped by dilution with 2 ml of the same buffer and then by heating in a boiling water bath for 2 min. After centrifugation, the supernatant solution was assayed for pyruvate with NADH and lactate dehydrogenase according to Peng and Shimizu [31]. The activity of 6-phosphofructokinase (PFK) in the crude cell extract was assayed as described by Gancedo and Gancedo [32]. The reaction mix contained 50 mM imidazole HCl (pH 7.0), 0.05 mM ATP, 5 mM MgCl₂, 1 mM EDTA, 0.25 mM NADH, 0.25 mM fructose 6-phosphate (F6P), 0.5 U aldolase, 0.5 U glycerolphosphate dehydrogenase and 0.5 U triosephosphate isomerase.

**Metabolic flux calculations**

Metabolic flux calculations were performed as described previously [18]. Briefly, metabolic flux ratio analysis was used to gain information about the flux distribution at important branch points within the network. As several alternative pathways may lead to a particular product, the fractional contribution (metabolic flux ratio) of each pathway was determined based on the molecular mass distributions of the reactants and the product according to Fischer and Sauer [33]. For the performed calculations, corrected mass spectra of selected fragments of serine, glycine, alanine, phenylalanine, tyrosine, aspartate and glutamate were used in this study (see Table 1). As the amino acids are synthesised from precursor metabolites of the central carbon metabolism with a known and well conserved carbon transition, their labelling pattern can be used to conclude the corresponding labelling pattern of their precursors [34]. To gain important information about the position of the labelling within the molecule, different fragments were considered simultaneously. In general, TBDMS-derivatised amino acids yield characteris-
tic fragments by electron impact ionisation. The \([\text{M}-57]\) fragment of each amino acid contains the complete carbon backbone, whereas the \([\text{M}-85]\) fragment lacks the carbon at the \(C_1\) position that corresponds to the carbon atom of the carboxyl group of the amino acid. The third fragment considered - \([f_{302}]\) - always contains the \(C_1\) and \(C_2\) carbon of the corresponding amino acid.

In the case of alternative pathways yielding a specific product, the fractional contribution of each pathway can be determined concerning the mass distributions of the reactants and the product according to Eq. (1) [33].

\[
\begin{bmatrix}
  f_{n-1} \\
  f_{n-2} \\
  \vdots \\
  f_1 \\
  f_n = 1 - f_{n-1} - f_{n-2} - \cdots - f_1
\end{bmatrix}
= \begin{bmatrix}
  \text{MDV}_1 - \text{MDV}_n \\
  \text{MDV}_{n-1} - \text{MDV}_n \\
  \vdots \\
  \text{MDV}_2 - \text{MDV}_n \\
  \text{MDV}_1 - \text{MDV}_n
\end{bmatrix}
\]

In Eq. (1) index \(X\) indicates the product molecule whereas the consecutive numbers 1 through \(n\) represent reactant molecules of alternative pathways contributing to the mass distribution of the product pool. The corresponding fractional amount of each pathway \(f\) can then be calculated by considering two additional constraints: (i) all fractions must have a positive value and (ii) their sum has to equal 1. A more detailed description will be given in the following respective sections.

**Theoretical framework for flux estimation**

To carry out metabolic flux calculations for \(D.\ shibae\) and \(P.\ gallaeciensis\), a metabolic network was constructed based on genome data (GenBank accession numbers NC_009952 \([D.\ shibae]\) and NZ_ABIF00000000 \([P.\ gallaeciensis]\]). As we focused on the central carbon metabolism, the major catabolic routes for glucose as well as the reactions linking the \(C_3\) and \(C_4\) pools were considered. In terms of glucose catabolism, the annotated genome revealed the presence of the genes encoding for glycolytic enzymes, enzymes of reactions in both the PPP and the ED pathway and TCA cycle. For \(D.\ shibae\), pyruvate carbox-
ylase (PYRCx) and phosphoenolpyruvate carboxykinase (PEPCK) were found to be the interconnecting reactions between the C$_3$ and the C$_4$ pool. Furthermore, a gene encoding for pyruvate orthophosphate dikinase (PPDK) is annotated, indicating a potential exchange flux between the PYR and PEP pool. A summary of all reactions considered is presented in Figure 1. To resolve the metabolic fluxes through catabolic pathways and around important branch points within the metabolic network, appropriate approaches involving the mass patterns of different amino acid fragments were developed.

**Strategy for the estimation of glucose catabolic fluxes**

In Figure 3 the theoretical labelling patterns of the C$_3$ pool directly follow from the mass pattern of the [f302] fragments of the precursors 3-phosphoglycerate (serine), oxaloacetate (aspartate), pyruvate (alanine) and the product PEP (phenylalanine). The corresponding isotopomer of each molecule is illustrated next to the experimental data (mass spectra). White circles represent $^{12}$C whereas black circles indicate labelled $^{13}$C. The numbers given reflect the position of the carbon atom within the molecule. PEPCK: phosphoenolpyruvate carboxykinase; PPDK: pyruvate-orthophosphate dikinase.

**Strategy for estimating fluxes around the PEP pool**

The metabolic reaction network around the PEP node is presented in Figure 7. It contains all reactions for which the corresponding genes have been annotated in the KEGG database. The pathways through lower glycolysis and the reactions catalysed by phosphoenolpyruvate carboxykinase (PEPCK) and pyruvate orthophosphate dikinase (PPDK) yielding PEP from either OAA or PYR are considered. Fluxes into the PEP pool were resolved using the mass distribution patterns of the [f302] fragments (carbon atoms at position C$_1$ and C$_2$) of the amino acids directly connected to the PEP pool according to Equations 4 and 5.

$$f_{\text{EMP}} = \frac{\text{PEP}[\text{f302}]_{\text{exp}} - \text{Asp}[\text{f302}]_{\text{exp}}}{\text{Ser}[\text{f302}]_{\text{exp}} - \text{Asp}[\text{f302}]_{\text{exp}}}$$

(4)

$$f_{\text{PEP}} = \frac{\text{PYR}[\text{f302}]_{\text{exp}} - \text{Asp}[\text{f302}]_{\text{exp}}}{\text{PEP}[\text{f302}]_{\text{exp}} - \text{Asp}[\text{f302}]_{\text{exp}}}$$

(5)

**Abbreviations**

[1-$^{13}$C] glucose: Glucose labelled at C1-position; 3PG: 3-phosphoglycerate; 6PG: 6-phosphogluconate; AcCoA: Acetyl-Coenzyme A; Ala: Alanine; CLE: labelling experiment; EDA: 2-keto-3-deoxy-6-phosphogluconate; ald-
lase; EDD: 6-phosphogluconate dehydrogenase; EDP: Entner-Doudoroff pathway; EMP: Embden-Meyerhof-Par- 
nas; F6P: Fructose-6-phosphate; FUM: Fumarate; G6P: Glucose 6-phosphate; GAP: Glycerate-3-phosphate; GC/MS: Gas chromatography/mass spectrometry; KDPG: 2-keto-3-deoxy-6-phosphogluconate; MAL: Malate; MDV: Mass distribution vector; OAA: Oxaloacetate; OD: Optical density; OGA: 2-oxoglutarate; PEP: Phosphoenolpyru- 
vate; PEPCk, pCa: Phosphoenolpyruvate carboxykinase; PFK: 6-phosphofructokinase; Phe: Phenylalanine; PPP: Pentose phosphate pathway; PPDK: ppdK Pyruvate ortho- 
phosphate dikinase; PVR: Pyruvate; PYRcx, pyCa: Pyru-
vate carboxylase; R5P: Ribose 5-phosphate; Ser: Serine; SUC: Succinate; TCA: Tricarboxylic acid; Tyr: Tyrosine

Authors’ contributions
TF carried out the labelling analytics and data processing, performed the flux calculations and drafted the manu-
script together with CW. MP performed the cultivation experiments for D. shibae. HZ performed the cultivation experiments for P. gallaeciensis. JT assisted in method set-
up for cultivation and analytics. IWD helped to draft the manuscript. RR helped to draft the manuscript. CW con-
ceived, designed and coordinated the study and drafted the manuscript together with TF. All authors read and 
approved the final manuscript.

Acknowledgements
JT and IWD gratefully acknowledge the support of the Volkswagen Foun-
dation under the grant VW-Vorab (ZN 2182, “Comparative functional genome analysis of representative members of the Roseobacter Clade”). We are grateful to Renate Gahl-Janssen (Oldenburg) for technical assistance. HZ and RR acknowledge support from the Volkswagen Foundation under the grant VW-Vorab (ZN2235, “Comparative functional genome analysis of representative members of the Roseobacter clade”) and the Marine Microbiology Initiative of the Moore Foundation (USA).

References
6. Buchanan A, Collier LS, Neidle EL, Moran MA: Key aromatic-ring-
15. Alavi MR: Predator/prey interaction between Pfiesteria pisci-
cida and Rhodomonas mediated by a marine alpha proteo-
16. Christensen B, Nielsen J: Metabolic network analysis of Penicil-
lium chrysogenum using (13)C-labeled glucose. Biotechnol Bio-
18. Fuch T, Hollmann R, Witzmann C, Wang W, Deckwer WD: Compari-
tive study on central metabolic fluxes of Bacillus megute-
19. Wittmann C, Hans M, van Winden WA, Ras C, Heijnen JJ: Dynamics of intracellular metabolites of glycolysis and TCA cycle during cell-cycle-related oscillation in Saccharomyces cerevi-
26. Kiefer P, Heinze E, Zälder O, Wittmann C: Comparative meta-


