Creating metabolic demand as an engineering strategy in *Pseudomonas putida* – Rhamnolipid synthesis as an example

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A B S T R A C T
Metabolic engineering of microbial cell factories for the production of heterologous secondary metabolites implicitly relies on the intensification of intracellular flux directed toward the product of choice. Apart from reactions following peripheral pathways, enzymes of the central carbon metabolism are usually targeted for the enhancement of precursor supply. In *Pseudomonas putida*, a Gram-negative soil bacterium, central carbon metabolism, i.e., the reactions required for the synthesis of all 12 biomass precursors, was shown to be regulated at the metabolic level and not at the transcriptional level. The bacterium’s central carbon metabolism appears to be driven by demand to react rapidly to ever-changing environmental conditions. In contrast, peripheral pathways that are only required for growth under certain conditions are regulated transcriptionally. In this work, we show that this regulation regime can be exploited for metabolic engineering.

We tested this driven-by-demand metabolic engineering strategy using rhamnolipid production as an example. Rhamnolipid synthesis relies on two pathways, i.e., fatty acid *de novo* synthesis and the rhamnose pathway, providing the required precursors hydroxylalkanoxyloxy-alkanoic acid (HAA) and activated (dTDP-)rhamnose, respectively. In contrast to single-pathway molecules, rhamnolipid synthesis causes demand for two central carbon metabolism intermediates, i.e., acetyl-CoA for HAA and glucose-6-phosphate for rhamnose synthesis.

Following the above-outlined strategy of driven by demand, a synthetic promoter library was developed to identify the optimal expression of the two essential genes (*rhlA*B) for rhamnolipid synthesis. The best rhamnolipid-synthesizing strain had a yield of 40% rhamnolipids on sugar [CmolRL/CmolGlc], which is approximately 55% of the theoretical yield. The rate of rhamnolipid synthesis of this strain was also high. Compared to an exponentially growing wild type, the rhamnose pathway increased its flux by 300%, whereas the flux through *de novo* fatty acid synthesis increased by 50%.

We show that the central carbon metabolism of *P. putida* is capable of meeting the metabolic demand generated by engineering transcription in peripheral pathways, thereby enabling a significant rerouting of carbon flux toward the product of interest, in this case, rhamnolipids of industrial interest.

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Metabolic engineering of secondary metabolite producers implicitly relies on high flux through central carbon metabolism (CCM). This high flux caused by the demand for carbon and energy for the synthesis of the molecule of interest, however, is rarely matched, requiring substantial improvements in CCM operation. The reactions of the CCM are providing the twelve precursors for biomass, i.e., for proteins, nucleic acids, polysaccharides, and lipids (Noor et al., 2010).

An excellent example of rational strain engineering by optimizing the flux distribution and channeling it to the product of choice was reported by Becker et al. (2011). The authors introduced 12 genome-based changes, including the overexpression of five genes encoding for enzymes fueling precursor-synthesizing pathways. In addition, the deletion or down-regulation of two genes encoding enzymes catalyzing competing reactions were introduced, yielding an L-lysine-overproducing strain of Corynebacterium glutamicum introduced 12 genome-based changes, including the overexpression of five genes encoding for enzymes fueling precursor-synthesizing pathways. In addition, the deletion or down-regulation of two genes encoding enzymes catalyzing competing reactions were introduced, yielding an L-lysine-overproducing strain of Corynebacterium glutamicum.

In contrast with this example, we here propose an approach that does not require substantial modifications of the host strain but, rather, relies on the capacity of the organism to reroute flux driven by a given (in our case, engineered) demand.

The strategy used here exploits the findings of Koebmann et al., (2002), who observed a strong increase in flux through glycolysis in Escherichia coli after introducing an ATP sink, thus discovering the concept of “driven by demand.”

It was recently shown that the CCM in P. putida is not transcriptionally regulated but, rather, is metabolically regulated (Sudarsan et al., 2014). Despite substantial rerouting of flux during growth on glucose, fructose, and benzoate, the transcription levels of the genes for CCM remain constant. Notably, the carbon substrate degradation pathways beta-ketoadipate and Entner-Doudoroff are transcriptionally regulated (Koebmann et al., 2002). Indeed, there is also evidence from extreme growth conditions (e.g., growth in the presence of a second phase of octanol) that P. putida can match metabolic demand by tripling the glucose uptake rate without producing any side products; thus, only biomass and CO2 are formed by this bacterium (Blank et al., 2008).

An example of this strategy is an engineered P. putida that hyper-produces polyhydroxyalkanoate (PHA) (Poblete-Castro et al., 2013). The authors deleted one gene (gcd, encoding for glucose dehydrogenase), which led to the rerouting of fluxes to the desired product. They were able to increase PHA accumulation by 100%.

In contrast to these results, peripheral pathways, which are not part of the CCM, are not as easily modified. Establishing a high production rate of aromatics from sugars in P. putida again entails substantial modifications to reroute intracellular flux resulting from the regulation of the synthesis pathways of aromatics. This regulation relies on the biosynthesis pathways of specific amino acids, which are regulated allosterically (Wierckx et al., 2005).

To verify the engineering-by-demand approach, we chose rhamnolipid synthesis as an example. It was earlier shown that P. putida is able to produce rhamnolipids after introducing two genes of the rhamnolipid synthesis pathway from P. aeruginosa, encoding RhlA and RhlB (Wittgens et al., 2011; Ochsner et al., 1994). The demand for precursors (i.e., increased flux through the rhamnose activation pathway and de novo lipid synthesis) varied according to the different promoter strengths of the rhlAB operon. The flux redistribution is estimated and the results are discussed in the context of the implications of the driven-by-demand concept for constructing superior production strains based on P. putida.

2. Materials and methods

2.1. Bacterial strains and culture conditions

The bacterial strains used, Pseudomonas putida KT2440 (Nelson et al., 2002) and Escherichia coli DH5α (Hanahan, 1983), were routinely cultivated in lysogeny broth (LB) medium (10 g/L tryptone, 5 g/L yeast extract, 10 g/L NaCl) (Bertani, 1951) at 30 °C for P. putida and at 37 °C for E. coli. Cells containing the vector pVT131 (de Lorenzo et al., 1993) and its derivatives were selected by adding tetracycline at concentrations of 10 μg/mL for recombinant E. coli and 20 μg/mL for P. putida. For the selection of pBRRIMCS-3 and derivatives, 20 μg/mL tetracycline was added. Rhamnolipid production by P. putida was conducted in LB medium containing 10 g/L glucose and 20 μg/mL tetracycline. The cells were cultivated in a 500 mL shake flask without baffles filled with 50 mL of the cultivation medium and using a MultiTron shaker (INFORS HT Bottmingen, Switzerland) at 250 rpm, with a throw of 25 mm and humidity of 80%.

2.1.1. Isotope-labeling experiments

Rhamnolipid-producing P. putida KT2440 pPS05 was grown under the conditions stated above. As media, LB medium and M9 minimal medium (Na2HPO4 × 2H2O 12.8 g/L, KH2PO4 3 g/L, NaCl 0.5 g/L, NH4Cl 1.0 g/L, 2 μM MgSO4 and 2 μL/L US trace elements solution (37% fuming HCl 82.81 mL/L, FeSO4 × 7H2O 4.87 g/L, CaCl2 × 2H2O 4.12 g/L, MnCl2 × 4H2O 1.50 g/L, ZnSO4 × 7H2O 1.87 g/L, H3BO3 0.30 g/L, Na2MoO4 × 2H2O 0.25 g/L, CuCl2 × 2H2O 0.15 g/L, Na2EDTA × 2H2O 0.84 g/L)) (Sambrook et al., 1989) were used. As a carbon source, 10 g/L regular glucose or uniformly isotope-labeled U-13C6 glucose (Cambridge Isotope Laboratories, Inc., Tewksbury, MA, USA) was applied. Tetracycline was added to a final concentration of 20 μg/mL. Samples were obtained twice: once after 8 h and the second after 24 h.

2.2. Expression vectors

For the promoter libraries, degenerated primers (Table 1) were applied carrying the consensus sequences at the −35 and the −10 position of a σ54-promoter (according to Jensen and Hammer (1998)). In this approach (already described in (Beuker et al., 2016)), the two regions of the promoter are conserved, while the remaining nucleotides are randomized. The here used promoter contains 40 nucleotides of which 12 are conserved in the RNA polymerase binding sites, hence 28 nucleotides are degenerated resulting in an high number of possible promoters (428).

The first fragment carrying the mono-rhamnolipid production genes rhlAB and the synthetic promoter were amplified from the P. aeruginosa genome using primers Fw-Psyn-thlAB and Rev-Psyn-

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**Table 1**

PCR primers used in this work. Underlined letters in the nucleotide sequences mark the restriction sites.

| Name       | Direction | Used for       | Template                        | Contains                        | Sequence |
|------------|-----------|----------------|---------------------------------|---------------------------------|----------|
| Fw-Psyn-thlAB | fward    | pSynProXX      | pBBR1                           | SglI, SynPro                    | AABBAGGCSCGCCGCCCC(N)5TTGACA(N)17TATAAT(N)6 |
| Rev-Psyn-thlAB | rev      | pSynProXX      | pBBR1                           |                                  | TCAACCAAGGCTTCTTCCACATCATCCCTTCCACATCATCAGCATCC |
The 5’ and 3’ regions were complementary to the vector backbone and were used for insertion. Between these regions, the new terminator sequence T_O (underlined) was introduced. This sequence was exchangeable through the two restriction sites: 5’ BglII AGATCT and 3’ KpnI GTTACC (restriction sites are marked with bold letters). Other exchangeable elements were the promoter (small dots) with 5’ KpnI and 3’ BsiWI CGTACG. The promoter was a synthetic promoter originating from the above-mentioned promoter library. In addition, we introduced a ribosome binding site (large dots), which is exchangeable with the restriction enzymes 5’ BsiWI CGTACG and 3’ XbaI TCTAGA. The transcription start was at the 3’ end (broken line).

Between the promoter and the transcription start, the ribosomal binding site sequence AGGGG was introduced because this sequence was determined by thermodynamic calculations to lead to particularly strong translation (Rühl, 2012). The distance to the start codon was chosen to be eight base pairs and included a restriction site. All of the elements were exchangeable via single cutter restriction sites. The plasmid was digested with PciI and BglII, and Gibson Assembly was used for combining of the synthetic DNA fragment with the vector backbone. The new modified plasmid was called pPS05 (Fig. 1) and was transformed via electroporation into E. coli DH5α. The new plasmid mixture was again transformed, this time into P. putida KT2440. A first screening was carried out using selective blood agar plates, which were examined regarding hemolytic activity. The construct that showed the highest titer was named pSynPro8 (Wittgens, 2013).

2.3. Analysis of rhamnolipids

The rhamnolipid concentration was determined by reversed-phase high-performance liquid chromatography (RP-HPLC). As an analytical column, a NUCLEODUR C18 Gravity (Macherey-Nagel GmbH & Co. KG, Düren, Germany) was used (dimensions: 150 x 4.6 mm; particle size: 3 μm). The HPLC system Ultimate 3000 was connected to a Corona-charged aerosol detector (CAD) (all Thermofisher Scientific Inc., Waltham, MA, USA). The flow rate was set to 1 mL/min, and the column oven temperature was maintained at 40 °C. Five microliters of the sample were injected. A binary gradient of acetonitrile and ultra-pure water supplied with 0.2% (v/v) formic acid was used. The acetonitrile concentration was increased linearly from 70% to 100% between 1 min and 9 min, and it was decreased linearly from 100% to 70% between 11 min and 12 min. One measurement was terminated after 15 min. The rhamnolipid concentration of P. putida cultures was measured after 3 days of cultivation. One milliliter of the supernatant was centrifuged for 5 min at 17,000 g. Five hundred microliters of the supernatant were subsequently mixed with 500 μL of acetonitrile and stored at 4 °C overnight. Subsequently, the sample was centrifuged for 5 min at 17,000 g. One hundred fifty microliters of the supernatant were filtered using Phenex-RC syringe filters (diameter 4 mm, pore size 0.2 μm) (Phenomenex,

T4 Ligase (Thermo Fisher Scientific, Waltham, MA, USA), the Gibson Assembly Cloning Kit (NEB, Ipswich, MA, USA), and restriction enzymes (Thermo Fisher Scientific, Waltham, MA, USA) were used routinely.

![Fig. 1. Vector pPS05. Depicted are all of the important elements, including the genes rhIA and rhIB, the tetracycline resistance cassette (TetR), the terminators (T0 and T1), the origin of replication (ori), the promoter (SynPro), and the ribosomal binding site (RBS). The plasmid also carries a mobilization gene (mob).](image-url)
Aschaffenburg, Germany) and pipetted into HPLC vials for subsequent HPLC analysis.

2.4. Flux estimation in supply pathways

The following assumptions were made to estimate the flux through the activated dTDP-rhamnose synthesis pathway. First, the content of lipopolysaccharides (LPS) per CDW in *P. putida* is similar to that in *P. aeruginosa*. Second, the number of rhamnose molecules incorporated into LPS is similar in both organisms. Darveau et al. determined the LPS of *P. aeruginosa* to account for 7.3% of the total CDW (Darveau and Hancock, 1983), whereas de Weger et al. reported that the LPS of *P. aeruginosa* consisted of 3.3% of rhamnose (de Weger et al., 1987). Hence, 0.24% of the CDW of *P. aeruginosa* and *P. putida* consists of rhamnose, amounting to 0.015 mmol rhamnose per g_{CDW}.

The flux through fatty acid *de novo* synthesis can be calculated accordingly. Approximately 10% of the CDW is composed of fatty acids (Stephanopoulos et al., 1998). A major constituent of the cell membrane in *P. putida* KT2440 is hexadecanoic acid (Rühl et al., 2012), with a molecular weight of 256 g/mol. *De novo* fatty acid synthesis hence requires 8C₂ units, which results in 3.12 mmol fatty acid formed per g_{CDW}. Multiplied by the growth rate, the rate for fatty acid biosynthesis can be estimated.

2.5. Measurement of 13C-labeled rhamnolipids

Sample preparation was performed according to Behrens et al. (2016). Rhamnolipid characterization was carried out on an Alliance 2695 separations module coupled to a micromass Quattro micro triple quadrupole mass spectrometer (both Waters Corporation, Milford, MA, USA). For separation, reversed-phase chromatography on a NUCLEODUR Sphinx RP (3 μm) was performed with a 2 mm x 150 mm column (Merck, Darmstadt, Germany) at 40 °C with a flow rate of 0.3 mL/min. A binary gradient of acetonitrile (for HPLC, Gradient Grade; VWR International, Fontenay-sous-Bois, France) and 5 mM ammonium formate buffer (LC-MS Ultra, ≥ 99.0%; Sigma-Aldrich Chemie GmbH, Steinheim, Germany) at pH 3.3 were used as eluents. Mass spectrometric detection was performed by electrospray ionization in negative mode.

2.6. Measurement of carbon dioxide production

The produced carbon dioxide was measured using a BCP-CO₂ sensor with a PA-plastics housing and recorded using the software BacVis (all from BlueSens Gas Sensor GmbH, Herten, Germany). The sensor was screwed onto a special 1 L shake flask without baffles. The flask was filled with 100 mL of medium and shaken at 200 rpm with a throw of 50 mm at 30 °C.

3. Results and discussion

3.1. Synthetic promoter library

The expression vector used previously was pVLT31 (de Lorenzo et al., 1993) featuring the ori RSF1010 and the inducible tac promoter (de Boer et al., 1983). This vector was used here as a reference. The newly constructed plasmid is based on pBBR1-MS3 (Kovach et al., 1995), containing the pBBR1 ori. The native RBS of the genes were kept. With *P. putida* carrying the synthesis pathway under the control of a synthetic promoter library indeed high rhamnolipid producing mutants were identified. From the first screening of a couple of thousand mutants using the blood agar assay, only 75 transformants were found to produce rhamnolipids (these results have partly been published in Beuker et al. (2016)). 23 of these were characterized in more detail. Shake flask experiments were carried out with these strains to determine physiological data on rhamnolipid production. Surprisingly nine strains did not produce detectable amounts of rhamnolipids. The remaining 14 transformants covered a rhamnolipid titer range from 0.05 g/L to 2.8 g/L (Fig. 2). While the transformant with the reference expression system, featuring a tac promoter, only produced 0.31 g/L rhamnolipids (Wittgens et al., 2011), the best producing transformant carrying the variant no. eight of the synthetic promoter library produced up to 2.8 g/L and was called *P. putida* KT2440 pSynPro8. This microbial cell factory, with its constitutive expression system features simplified handling, as induction is no longer required.

Seven promoters were chosen for sequencing, covering the whole range of rhamnolipid titers. The data shows, that indeed the −35 and −10 box remained unchanged, while the randomized regions featured great variations in nucleotides. As all these promoters showed rhamnolipid production during the first screening, one might conclude that only intact promoters lead to the production of rhamnolipids.

**SynPro8**: AGCTC TTGACA AGGTCGAAAATTGAAG TATAAT ATCAGT  
**SynPro5**: TTTCG TTGACA AGCCTAGTTTCGCCATT TATAAT GACTCG  
**SynPro16**: GTGTA TTGACA AAGCGCTTACCTCTTC TATAAT ATAGAG  
**SynPro1**: GGTGCG TTGACA TTGGCATTACAACGTAT TATAAT TTACCG

![Fig. 2.](image-url) Final rhamnolipid titers of *P. putida* with different synthetic promoters. The previously used tac promoter based system served as reference. The error bars represent the deviation from the mean of two biological replicates.
growth frees more resources for rhamnolipid synthesis. Indeed an increase in the rhamnolipid production rate, indicated by a deviation of the experimental data above the fitted line can be inferred.

### 3.3. Meeting metabolic demands

We observed in the above experiments an increased glucose uptake rate of 1.2 mmol/(gCDW h) for pPS05 and 3.2 mmol/(gCDW h) for pSynPro8, which we here investigate conceptually. Can the metabolic network operation in *P. putida* be hijacked by creating demand for central carbon metabolism (CCM) intermediates? We previously showed that energetic demand could be met by *P. putida* by increasing the glucose uptake rate dramatically, for example, in response to a second phase of octanol (Blank et al., 2008) or the uncoupler 2,4-dinitrophenol (DNP) (Ebert et al., 2011).

To evaluate whether the metabolism of the *P. putida* used here, grown on a complex medium (e.g., amino acids available) and complemented with glucose, responds comparably, we added alternating DNP concentrations. When adding 700 mg/L DNP to *P. putida* KT2440 growing in LB medium supplemented with 10 g/L glucose, the strain had a glucose uptake rate of 2.0 mmol/(gCDW h) instead of 1.1 mmol/(gCDW h) without DNP. At a concentration of 1,100 mg/L DNP, the glucose uptake rate reached 2.2 mmol/(gCDW h). This increase in glucose uptake rate did not affect the growth rates, which were 0.86 h⁻¹ (no DNP), 0.88 h⁻¹ (700 mg/L DNP), and 0.79 h⁻¹ (1,100 mg/L DNP) (Fig. 4). Importantly, no byproduct formation was observed, consistent with Blank et al. (2008) and Ebert Ebert et al. (2011). Hence, creating a sink for NADH by the addition of an uncoupler resulted in a doubling of the glucose uptake rate. This finding provides strong support that the metabolic network operation of *P. putida* can be substantially altered by the here-exploited concept of driven by demand.

### 3.4. Hijacking central carbon metabolism operation

Can the observed increase in glucose uptake rate be exploited for the production of valuable chemicals? We previously reported that the CCM in *P. putida* is, under many conditions, not regulated transcriptionally but, rather, at the metabolic (and potentially posttranslational) level. In contrast, peripheral anabolic and catabolic pathways are transcriptionally regulated (Sudarsan et al., 2014). This organization of the metabolic network allows for rapid flux rerouting to adapt to ever-changing growth conditions. We here argue again that these flux changes are the result of changing metabolic demands (Table 2).

Consequently, we examined whether this structural

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[Fig. 3. Rhamnolipid production with *P. putida* KT2440 pSynPro8. Development of biomass (rectangles, •), rhamnolipid (triangles, ▲), and glucose (diamonds, ◆) concentrations. The experimental data are depicted as symbols, and the lines present the fitted courses. The CDW trend was determined using a logistic growth model. A multivariable least squares fit was used to illustrate the development of all three fermentation parameters depending on each other, according to Wittgens et al. (2011). The error bars represent the deviation from the mean of two biological replicates. CDW, cell dry weight.](#)

[Fig. 4. 2,4-Dinitrophenol (DNP) addition leads to enhanced glucose uptake rates. The specific glucose uptake rate is shown in white, and the growth rate is depicted by the black-filled columns. *P. putida* KT2440 pPS05 was cultivated without DNP as a reference and with 700 and 1,100 mg/L DNP, according to Ebert et al. (2011). The error bars represent the deviation from the mean of two biological replicates.](#)
organizational of the metabolic network explained the high performance of the rhamnolipid-synthesizing strain (Fig. 3). We introduced the genes for mono-rhamnolipid synthesis into P. putida (rhlA and rhlB), thereby creating additional demand in the CCM for glucose-6-phosphate (for rhamnose synthesis) and acetyl-CoA (for hydroxy-fatty acid production via de novo lipid synthesis) (Fig. 5).

To examine different demands, i.e., different rhamnolipid production rates, we used the two plasmids pSynPro8 and pPS05 and two strains, namely, the reference P. putida KT2440 and the derivative P. putida KT40CZC. In P. putida KT40CZC, the PHA synthesis operon is deleted (Escapa et al., 2012). PHA synthesis starts from activated hydroxy-fatty acids competing directly with rhamnolipid synthesis for the common precursor. Quantitative physiology experiments allowed for the estimation of flux via the precursor-supplying pathways, the glucose uptake rate, and the growth rate. For comparison, the physiology of the reference strain P. putida KT2440 not producing rhamnolipids was examined.

It was discovered that P. putida KT40CZC was better suited for rhamnolipid synthesis with the new expression plasmid pPS05 than the wild-type strain (Table 3). This recombinant strain has a higher specific production rate and also reaches a higher titer.

With an increasing rhamnolipid production rate, the glucose uptake increased while the growth rate remained constant (Fig. 6). In the best producing strain with a rhamnolipid production rate of 0.1 mmol/(gCDW h), the glucose uptake increased by 24% to 1.3 mmol/(gCDW h) compared to the non-producing strain. P. putida KT2440 pPS05 produced rhamnolipids with a glucose uptake rate of 1.2 mmol/(gCDW h), which is 12% more than the wild-type strain. These findings are a strong indication that flux through the CCM is indeed increased. The additional glucose is used to satisfy the increased demand created by the production of the rhamnolipids.

To assess whether the extra glucose, which is taken up is directed to rhamnolipid synthesis or just burned and oxidized to CO2, CO2 production was measured in a separate experiment. As can be seen, carbon dioxide production of the reference P. putida KT2440 is higher compared to the rhamnolipid producer (Fig. 7). The reduction in CO2 formation might be a response of P. putida to the availability of an additional redox sink, namely rhamnolipid synthesis. The reference strain oxidizes glucose to CO2 via the tricarboxylic acid (TCA) cycle. To regenerate the reduced redox cofactors and thereby balance the electrons, the electron transport chain is employed by P. putida using oxygen as terminal electron acceptor. The rhamnolipid-producing strain can use the rhamnolipids as electron acceptors as the carbons in the fatty acid moieties are higher reduced than glucose. Thus, we assume that this strain under the given conditions does not need the electron transport chain to the same extent as the reference strain, with the result of lower CO2 formation.

As less glucose is wasted for the generation of CO2, the rhamnolipid production strain should be able to allocate more carbon to the synthesis of rhamnolipids. These considerations again hint to the conclusion that via reorganization of metabolic fluxes the cell is able to meet the opposed demand.

Based on the physiological data, we estimated the flux through the pathways providing the precursors for rhamnolipid production, i.e., activated dTDP-rhamnose, and the hydroxy-fatty acids for HAA synthesis (Fig. 8). The P. putida wild type not producing rhamnolipids still synthesizes rhamnose as part of the lipopolysaccharides on the outer cell wall. A constant synthesis rate was estimated based on the growth rate and on the approximate rhamnose content in the lipopolysaccharides. For de novo synthesis, basic activity was calculated via the fatty acid share in the biomass. When introducing the enzymes for rhamnolipid production, the particular flux via the activated rhamnose pathway was dramatically enhanced. Moreover, de novo fatty acid synthesis increased by 50% in the best producing strain.

Again, these enhancements of flux in the metabolic network
ponents were utilized for cell growth. The numbers in parentheses show the percentages of the theoretical maximum.

The deviation shown in light gray originates from the mean of two biological replicates. The black-filled columns represent the rate via the de novo reaction network of a given pathway to distinguish the enzymatic steps that control the overall production rate. This specific reaction rate can then be enhanced, which should lead to a higher production rate toward the product of choice. This process is continued iteratively until no further enhancement can be achieved (Stephanopoulos et al., 1998).

Fig. 7. Carbon dioxide produced during cultivation by P. putida KT2440 (gray line) and P. putida KT2440 pPS05, which is capable of rhamnolipid synthesis (black line). The deviation shown in light gray originates from the mean of two biological replicates.

Table 3
Growth and production parameters of rhamnolipid synthesis. For comparison, wild-type P. aeruginosa PA01 is listed in line 1.

| Organism       | Medium | Substrates | Cell Dry Weight [g(cw)/L] | Maximal Titer [g/L] | Carbon Yield[^a] [Cmol RL/Cmol subs] | Process Time [h] | Space-Time Yield [mg RL/(L h)] | Specific RL-Prod. Rate[^b] [g RL/(g cw h)] | Reference         |
|----------------|--------|------------|---------------------------|--------------------|--------------------------------------|------------------|---------------------------------|--------------------------------------------|------------------|
| P. aeruginosa PA01 | MS      | Sunflower Oil 250 | 16.3                      | 39.0               | 0.07 (8%)                            | 90               | 433.3                           | 0.027                                      | Müller et al. (2010) |
| P. putida KT2440 pSynPro8 | LB      | Glucose 12 | 3.1                       | 3.2                | 0.40 (55%)                           | 22               | 146.4                           | 0.047                                      | Wittgens (2013)    |
| P. putida KT40CZC pPS05 | LB      | Glucose 11 | 3.4                       | 2.4                | 0.35 (48%)                           | 23               | 104.3                           | 0.031                                      | This study         |
| P. putida KT2440 pPS05 | LB      | Glucose 11 | 4.5                       | 2.2                | 0.32 (44%)                           | 22               | 100.0                           | 0.022                                      | This study         |

[^a]: For the calculation of yields during production on complex media, it was assumed that rhamnolipids were synthesized from glucose, whereas other medium components were utilized for cell growth. The numbers in parentheses show the percentages of the theoretical maximum.

[^b]: The specific rhamnolipid production rate was calculated as the average over the entire fermentation time using the maximal rhamnolipid titer and the maximal cell dry weight determined.

Figure 6. Performance parameters of rhamnolipid producers compared to the wild-type strain. The black-filled columns represent the growth rate, and the unfilled columns indicate the glucose uptake rate. On the second y-axis, specific rhamnolipid production is plotted. The respective columns are filled in gray. For better comprehension, the second y-axis is also drawn in gray. The error bars represent the deviation from the mean of two biological replicates.

Figure 8. Estimated intracellular fluxes in P. putida strains with different activity levels of rhamnolipid production. The black-filled columns represent the rate via the rhamnose pathway, and the gray-filled columns indicate the de novo fatty acid synthesis rate (second y-axis). For better comprehension, the second y-axis is also shown in gray. The error bars represent the deviation from the mean of two biological replicates.

CCM is clearly altered to meet the RL synthesis demand.

Metabolic engineering strategies often involve the increase of flux in precursor-providing pathways. To identify valuable targets for overexpression, metabolic control analysis is frequently performed. Metabolic control analysis examines the underlying reaction network of a given pathway to distinguish the enzymatic step that controls the overall production rate. This specific reaction rate can then be enhanced, which should lead to a higher production rate toward the product of choice. This process is continued iteratively until no further enhancement can be achieved (Stephanopoulos et al., 1998).

To increase rhamnolipid production, it would thus be advisable to overexpress the entire rml operon harboring the genes responsible for the synthesis of the activated dTDP-rhamnose from glucose-1-phosphate or the pyruvate dehydrogenase, which is the first step from pyruvate toward de novo lipid synthesis (Scriba and Holzer, 1961). A substantial enhancement of flux in glucose-6-phosphate- and pyruvate-supplying pathways via CCM is required to sustain an unchanged rate of growth and to generate the resources for the heterologous production of rhamnolipids (Fig. 5).

We hence conclude that CCM operation in P. putida is driven by the demand created. Again, the mechanistic explanation might be that the CCM in this organism is regulated at the metabolic level (Sudarsan et al., 2014). Hence, generating additional demand by
putida is its superior glucose uptake system. In the past, we overexpressed two enzymes producing rhamnolipids, which burdened CCM at glucose-6-phosphate and acetyl-CoA, resulting in flux increases by 300% and 50%, respectively.

There is however one major difference to the experiment carried out by Sudarsan et al. (2014). While they were working in a chemostat environment at constant growth rate, the experiments described here were carried out in batch cultures, featuring different growth phases. In the previous study, the putative impact of growth rate on transcriptional regulation was thus excluded by the experimental setup, while this has not been done in the experiments carried out in this study. Differences in growth could thus be partly responsible for the observed phenomena.

The conditions under which an organism responds to being driven by demand is not generally known. Pseudomonas, as a soil bacterium, lives in environments where nutrients are scarce and must compete for these resources. Other industrial microorganisms, such as E. coli, Lactobacillus, and Saccharomyces cerevisiae, have been isolated from nutrient-rich environments, such as the lower intestines of mammals, milk, and crops, respectively. With many resources in abundance, these organisms might have been selected for rate rather than for efficiency (yield) (Novak et al., 2006; Jessup and Bohannan, 2008).

A good example for the high efficiency of P. putida is its glucose uptake system. Whereas E. coli and other microorganisms absorb glucose at high rate, P. putida converts part of the glucose into gluconate and further into ketoglucanate (del Castillo et al., 2007). This process has two advantages in a competitive environment. First, while growing on a preferred carbon source, the bacterium renders this carbon source useless for competitors not able to absorb gluconate or ketoglucanate. Second, gluconate and ketoglucanate are organic acids and thus greatly lower the pH. A low pH not only stops the growth of some organisms but also frees some of the scarce resources, such as iron and phosphate (Rodriguez and Fraga, 1999).

Another interesting feature of the glucose uptake system of P. putida is its superior glucose affinity. Glucose uptake in P. putida into the periplasmic space occurs through outer membrane porins encoded by oprB1 and oprB2. The importation into the cytosol is mediated through an ABC uptake system encoded by the open reading frames PP1015 to PP1018 (del Castillo et al., 2007). ABC transporters in Gram-negative bacteria often feature a binding protein that is located in the periplasm. This binding protein has high affinity for the specific substrate (Roos et al., 1996). The transport system of Thermococcus litoralis, featuring a similar binding protein, for example, has a K_m of approximately 20 nM, whereas the K_m of the glucose uptake system of E. coli exhibits 1 μM (Xavier et al., 1996). After binding the substrate, the actual transport is driven by the hydrolysis of ATP molecules (Davidson and Chen, 2004). This biochemical investment allows for glucose uptake when concentrations are very low.

These two examples show the adaptation of P. putida to environments where nutrients are scarce. Under such conditions, changes in carbon sources are frequent. Whenever a change in the environment occurs, P. putida is able to reconfigure its CCM rapidly without relying on transcriptional regulation. In the context of heterologous rhamnolipid production, which is an engineered demand, P. putida increases carbon uptake while maintaining a high growth rate. The metabolism simply reacts to the “theft” of precursors needed for growth by increasing the flux through the essential precursor pathways.  

3.5. Glucose satisfies the enhanced carbon demand

As described above, the rate of glucose uptake increases when an additional demand is created. Is this glucose directed toward rhamnolipid synthesis and hence to the created demand, or is it simply distributed throughout the bacterium’s metabolism? Via stable isotope-labeling experiments and the analysis of the resulting labeling patterns in the rhamnolipid molecules, the carbon source used for rhamnolipid synthesis was determined.

For the corresponding experiments, on the one hand, minimal medium supplemented with unlabeled glucose and, on the other hand, LB medium with uniformly 13C-labeled glucose were used. Two samples were obtained: one in the exponential phase and the other in the stationary phase. At the latter time point, the glucose was completely consumed. The synthesized rhamnolipids were analyzed by HPLC coupled to mass spectrometric detection (LC-MS). The produced biomass was investigated by gas chromatography coupled to mass spectrometry (GC-MS) measurement. Sections of the LC-MS mass spectra at the retention time of Rha-C_{10–10} (11.8 min) are depicted in Fig. 9.

The molecular weight of the Rha-C_{10–10} rhamnolipid molecule is 504 u. It is detected as a deprotonated molecule (mass to charge ratio [m/z] of 503) during LC-MS analysis with electro spray ionization. In the mass spectra shown in Fig. 9, the measured signals are higher than m/z 503 caused by 13C-incorporation from the labeled glucose precursor. The examined rhamnolipid Rha-C_{10–10} contains 26 carbon atoms; thus, signals from m/z 503 to m/z 529 can occur. A fully labeled rhamnolipid would give rise to m/z 529.

Mass spectra a and d show the rhamnolipid Rha-C_{10–10} synthesized from unlabeled glucose in different growth phases as a reference. The highest peak is at m/z 503. Mass spectrum b, obtained from cultivation in minimal medium with fully labeled glucose, should show only the m/z 529 signal. However, as shown in Fig. 9b, the measured rhamnolipids show a distribution with its maximum at m/z 517. The explanation for this distribution in partially labeled rhamnolipid is the presence of unlabeled biomass from the preculture and the percentage of unlabeled glucose, which is contained in the 13C-labeled glucose. The enrichment of 13C-labeled glucose is 99% for each carbon atom, resulting in 94% (0.99%) completely labeled glucose molecules. Interestingly, the mass spectra for the rhamnolipid derived from the cultivation with 13C-labeled glucose with minimal medium (Fig. 9b) and LB medium (Fig. 9c) are very similar, indicating that during the exponential phase, the main source for rhamnolipid production is glucose, whereas the components of the LB medium are consumed for cell growth. Spectrum e shows the expected distribution: almost exclusively fully labeled rhamnolipid Rha-C_{10–10}. The distribution in spectrum f is also shifted toward higher masses, again showing that the main carbon source for rhamnolipid synthesis is glucose. During the exponential phase (Fig. 9c), this leads to a spectrum similar to the spectrum of the rhamnolipid when using isotopically labeled glucose as sole carbon source (Fig. 9b). Later, when the unlabeled compounds are further attenuated by the depletion of media components and the comparatively higher influx of labeled carbon atoms, the labeling pattern of the rhamnolipid shifts to higher masses (Fig. 9f).

An interesting observation is that, in the mass spectra, the m/z values with even numbers also appear, despite de novo fatty acid synthesis assembling fatty acids using C\textsubscript{2} molecules as basic modules. In theory, the labeling of the rhamnolipid molecules should thus increase in steps of two. The unlabeled carbon that is incorporated into the rhamnolipid molecule, however, also contains natural 13C-isotopes. The natural abundance of 13C is 1.08. In the rhamnolipid molecule, which contains 26 carbon atoms, this results in almost 29% of the molecules having one 13C-isotope and approximately 4% of the total rhamnolipid molecules carrying two 13C-isotopes. Furthermore, isotope ratios of O (18O=0.00%; 17O=0.04%; 16O=0.21%) and H (4H=100%; 2H=0.1%) contribute to the signals. This naturally occurring isotope distribution is
reflected in mass spectra a and d.

We also performed the reverse test, measuring the labeling in the proteinogenic amino acids. The fractional labeling of the examined amino acids showed that most amino acids were absorbed from the medium, whereas only a minor fraction was synthesized de novo from glucose (Fig. 10). Whereas some amino acids showed a fractional labeling of more than 0.2, at least half of the analyzed amino acids featured a fractional labeling less than 0.1; hence, more than 90% of these amino acids were absorbed from the medium.

To conclude, we showed that the synthesized rhamnolipids mainly originated from the $^{13}$C-labeled glucose, whereas the biomass predominantly originated from unlabeled medium compounds. Hence, the increased glucose uptake caused by the created demand was most likely used by the cell to satisfy this demand. The sugars were directed by the metabolism of *P. putida* toward the reactions providing precursors for rhamnolipid synthesis. Only a minor fraction of the sugar was used for biomass production. In addition, the carbon dioxide production experiments point to the conclusion that rhamnolipid synthesis offers a possibility for the cell to use the offered carbon source more efficiently. Thus, the production strain is able to invest more glucose for the synthesis of rhamnolipids.

The yield of product on substrate is an important parameter in biocatalysis. One challenge is the reduction of yield caused by the growth of the whole-cell catalyst. Hence, production under reduced- or non-growth conditions is an engineering goal. We reported previously that engineered *P. putida* KT2440, under reference conditions, produced rhamnolipids independently of growth (Wittgens et al., 2011). The carbon source used (LB or glucose) was inferred from CO$_2$ evolution. Here, we report the direct measurements of the $^{12}$C/$^{13}$C-labeling of rhamnolipids and the produced biomass.

The detected labeling patterns supported the hypothesis that rhamnolipid synthesis occurred independently of growth. When supplying LB medium and $^{13}$C-labeled glucose, the majority of the rhamnolipid molecules contained labeled $^{13}$C-carbon atoms emanating from the labeled glucose. The fractional labeling in the exponential growth phase of the C$_{10}$-C$_{10}$ rhamnolipids was 0.63. Only a small part (approximately 15%) of the rhamnolipid molecules carried ten or fewer labeled carbon atoms. Hence, the compounds from the LB medium contributed only a small fraction to rhamnolipid synthesis. In contrast, only 16% of the carbon atoms in the biogenic amino acids were labeled. The majority of the biomass is thus derived from unlabeled compounds of the medium and not from glucose. The assumption that medium compounds are used for growth while glucose is utilized in rhamnolipid synthesis occurred independently of growth. When supplying LB medium and $^{13}$C-labeled glucose, the majority of the rhamnolipid molecules contained labeled $^{13}$C-carbon atoms emanating from the labeled glucose. The fractional labeling in the exponential growth phase of the C$_{10}$-C$_{10}$ rhamnolipids was 0.63. Only a small part (approximately 15%) of the rhamnolipid molecules carried ten or fewer labeled carbon atoms. Hence, the compounds from the LB medium contributed only a small fraction to rhamnolipid synthesis. In contrast, only 16% of the carbon atoms in the biogenic amino acids were labeled. The majority of the biomass is thus derived from unlabeled compounds of the medium and not from glucose. The assumption that medium compounds are used for growth while glucose is utilized in rhamnolipid synthesis occurred independently of growth.
synthesis is thus strengthened. A possible explanation for the uncoupling of growth and rhamnolipid production might be found in the lipidic precursor, i.e., the activated hydroxy-fatty acids. These molecules are also used for the synthesis of the internal storage polymer PHA, which P. putida produces during secondary substrate-limiting conditions when the carbon source is still abundant (Prieto et al., 2007).

4. Conclusion

In this work, we showed that the concept of “driven by demand” described for E. coli by Koebmann et al. (2002) could also be exploited for flux rerouting in P. putida. To rapidly implement heterologous pathways, tools for creating the required demand are becoming important (Zobel et al., 2015; Nikel et al., 2014). Interestingly, in the engineered microorganism, the production of the product of choice is uncoupled from growth, which might be caused by overcoming the host’s intrinsic regulation cascades by introducing the target genes under the control of a synthetic promotor. The results suggest that P. putida is a suitable host for bio-technological applications. Its versatile metabolism, which responds to the demand created by heterologous pathways, is well suited for metabolic engineering. Metabolic engineering of bacteria yielding a production strain, which is competitive on an industrial scale, is a complex task. Even for scientific groups with both the equipment and the required expertise, this process requires considerable time and effort, as demonstrated by the excellent extant examples (Becker et al., 2011; Becker and Wittmann, 2012a, 2012b; Buschke et al., 2013). P. putida might provide a good starting point for establishing novel synthesis pathways via classic metabolic engineering strategies.

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