Efficient Cas9-based genome editing of *Rhodobacter sphaeroides* for metabolic engineering

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**Abstract**

**Background:** *Rhodobacter sphaeroides* is a metabolically versatile bacterium that serves as a model for analysis of photosynthesis, hydrogen production and terpene biosynthesis. The elimination of by-products formation, such as poly-β-hydroxybutyrate (PHB), has been an important metabolic engineering target for *R. sphaeroides*. However, the lack of efficient markerless genome editing tools for *R. sphaeroides* is a bottleneck for fundamental studies and biotechnological exploitation. The Cas9 RNA-guided DNA-endonuclease from the type II CRISPR-Cas system of *Streptococcus pyogenes* (SpCas9) has been extensively employed for the development of genome engineering tools for prokaryotes and eukaryotes, but not for *R. sphaeroides*.

**Results:** Here we describe the development of a highly efficient SpCas9-based genomic DNA targeting system for *R. sphaeroides*, which we combine with plasmid-borne homologous recombination (HR) templates developing a Cas9-based markerless and time-effective genome editing tool. We further employ the tool for knocking-out the uracil phosphoribosyltransferase (*upp*) gene from the genome of *R. sphaeroides*, as well as knocking it back in while altering its start codon. These proof-of-principle processes resulted in editing efficiencies of up to 100% for the knock-out yet less than 15% for the knock-in. We subsequently employed the developed genome editing tool for the consecutive deletion of the two predicted acetoacetyl-CoA reductase genes *phaB* and *phbB* in the genome of *R. sphaeroides*. The culturing of the constructed knock-out strains under PHB producing conditions showed that PHB biosynthesis is supported only by PhaB, while the growth of the *R. sphaeroides ΔphbB* strains under the same conditions is only slightly affected.

**Conclusions:** In this study, we combine the SpCas9 targeting activity with the native homologous recombination (HR) mechanism of *R. sphaeroides* for the development of a genome editing tool. We further employ the developed tool for the elucidation of the PHB production pathway of *R. sphaeroides*. We anticipate that the presented work will accelerate molecular research with *R. sphaeroides*.

**Keywords:** *Rhodobacter sphaeroides*, Cas9, Genome editing, PHB
Introduction

The purple non-sulphur bacterium Rhodobacter sphaeroides is a microorganism with an extremely adaptable metabolism [1]. It is a facultative phototroph that can grow on many different carbon substrates and can respire aerobically and anaerobically with different electron acceptors [2, 3]. The metabolic versatility of Rhodobacter sphaeroides has raised curiosity for white biotechnological applications. In particular, this microorganism has been largely studied for photoheterotrophic hydrogen production and chemoheterotrophic terpene biosynthesis. In both processes, approaches for improving the metabolic performances of the microorganism were investigated [4–8]. Reducing formation of by-products like the polymer poly-β-hydroxybutyrate (PHB) holds potential for eliminating competition for carbon and reducing power, as already proven to work for H₂ biosynthesis [5, 9]. Nonetheless, a lot of work is still required to further increase our knowledge on R. sphaeroides metabolism and exploit its biotechnological potential.

The high-throughput exploration of R. sphaeroides metabolism requires the development of highly effective and time-efficient genome engineering tools [4]. Currently, the introduction of genomic modifications in R. sphaeroides is based on suicide-plasmid driven homologous recombination (HR) systems [10, 11]. These systems depend on the incorporation of the non-replicating vector into the R. sphaeroides genome via an initial single crossover (SCO) event and consequent excision of the vector via a second SCO event upon induction of counter-selection pressure i.e. employing a levansucrase gene (sacB) carrying vector [11–17]. The second SCO can either lead to the reconstitution of a wild type genomic background or to the desired genetic modification [11]. The lack of a strict counter-selection method targeting specifically the genomic region of interest usually leads to high rates of wild-type revertants, especially for essential genes. Therefore, screening of mutants can be time consuming and is frequently unsuccessful.

A wide range of CRISPR-Cas (Clustered Regularly Interspaced Short Palindromic Repeats-CRISPR associated proteins) bacterial and archaeal adaptive immune systems have been repurposed as tools for eukaryotic, prokaryotic and archaeal genome engineering [18–26]. Most of these tools exploit the RNA-guided DNA endonuclease from the type II CRISPR-Cas system of Streptococcus pyogenes, denoted as SpCas9. As any Cas9 orthologue, SpCas9 can be easily programmed to precisely introduce double stranded DNA breaks (DSB) to a selected DNA sequence, denoted as protospacer. A single, customizable guide RNA (sgRNA) molecule directs Cas9 to the protospacer via complementarity between its exchangeable 5’-end sequence, denoted as spacer, and the protospacer sequence. The only additional requirement for Cas9-based targeting is the presence of a specific, 2–8 nt long motif right after the 3’-end of the selected protospacer, denoted as protospacer adjacent motif (PAM) [27]. All in all, the simplicity in the design and construction of a Cas9-based DNA targeting system has made it popular as the basis for numerous genome manipulation applications.

A plethora of Cas9-based tools have been developed the last 5 years for prokaryotes [24, 28–33]. Cas9 has been extensively used as an efficient HR induction and counter selection tool, when combined with plasmid-based homologous recombination (HR) or recombineering; alternative DSB repairing mechanisms (like the template independent non-homologous end joining, NHEJ) are lacking in most prokaryotes [34]. Therefore, Cas9-based DSBs after HR are lethal if introduced to the wild type genomes: unmodified cells will be eliminated from the treated population, allowing survival only for the recombined ones [34].

In this study, we develop a highly efficient Cas9-based targeting system for R. sphaeroides. We further combine it with HR templates, developing an HR-Cas9 counter selection tool. We then employ the tool for the proof-of-principle efficient generation of uracil-phosphoribosyltransferase (up) gene knock-out and knock-in strains, the latter combined with single nucleotide substitution. The developed process simplifies and accelerates R. sphaeroides genome editing, as it requires only 3 days from conjugation to screening of clean mutants. Additionally, we use the developed tool for the construction of acetoacetyl-CoA reductase deletion mutant strains, elucidating the dominant metabolic pathway of R. sphaeroides towards PHB production.

Materials and methods

Bacterial strains, media and growth conditions

The strains used in this study are listed in Table 1. E. coli DH5α was used for cloning and routine amplification. E. coli S17-1 was used as vector donor for R. sphaeroides in diparental conjugation.

Culturing of R. sphaeroides was performed in RÄ minimal or LB (Luria–Bertani) medium at 250 rpm, 30 °C. Growth on solid medium was performed on RÄ supplemented with 15% w/v agar. Culturing of E. coli strains was performed in LB medium at 250 rpm, 37 °C. Growth on solid medium was performed on LB supplemented with 15% w/v agar. When required, kanamycin (50 µg/mL) was added for all the mentioned growing conditions.
E. coli TOP10 cells for storage and amplification. Subsequently, the sgRNA construct was ligated into the pUC19 vector and transformed into DH5α (Thermo Fisher Scientific), the sgRNA construct was validated using the Galaxy/Codon harmonizer online tool [35]. The codon-harmonized cas9 was synthetically constructed by Baseclear (the Netherlands) (Additional file 1: Table S4) with a 6×His-tag fused at its C-terminus, and delivered inserted in a pUC57 vector backbone. The cas9 gene was PCR amplified using the pUC57hSpCas9 vector as template and primers BG10937/BG10938. The pBBR-sgRNA vector was linearized using primers BG10939/BG10941. The construction of the non-targeting plasmid pBBR_Cas9_NT vector (Additional file 1: Figure S1) was done via HiFi assembly of the two aforementioned PCR amplicons and the cas9 ampiclon was inserted downstream of the lac promoter site of the pBBR-sgRNA. The lack of lacI repressor gene or its homolog in R. sphaeroides genome allowed constitutive transcription of the harmonized cas9 gene.

### Generation of the upp targeting plasmids

The spacers introduced in the sgRNA construct of the pBBR_Cas9 vectors are listed in the Additional file 1: Table S5. The spacers were selected employing the chopchop web tool (https://chopchop.cbu.uib.no/). The spacer- selection parameters were set to avoid self-complementarity of the sgRNA, have efficiency more than 50 and no predicted off-targets. Using the plasmid pBBR_Cas9_NT as template, it was possible to substitute the non-targeting spacer contained in the sgRNA construct with three different targeting spacers for the gene uracil phosphoribosyltransferase upp (sp1–sp3). The pBBR_Cas9_NT was used as template for PCR amplification of the backbone, the start codon and part of the cas9 gene, employing primers BG11415/BG11416 in combination with primers BG11411, which also contained the spacer 1, 2 and 3 sequences, respectively as overhangs. 3 fragments assemblies were performed and the reaction mixtures were transformed in DH5α E. coli competent cells (NEB) via heat shock, for storage and plasmid amplification. Upon sequence verification, the pBBR_Cas9_NT plasmid was transformed to E. coli S17-1 cells.
Generation of vectors for gene knock-out (KO) and gene knock-in (KI)

Homologous recombination (HR) followed by Cas9-based counter selection was used for the KO of the upp gene, as well as the KI of the upp gene back to its original place in the R. sphaeroides genome combined with single nucleotide substitution.

Plasmids for upp KO

We started by constructing the HR control, non-targeting plasmids pBBR_Cas9_Δupp500HR_NT and pBBR_Cas9_Δupp1000HR_NT. The pBBR_Cas9_NT backbone was PCR linearized into two fragments using the primers sets BG11866/BG11182 and BG11887/BG11888. The 500 bp or 1 kb long upstream and downstream flanking sites of the upp gene were PCR amplified from the R. sphaeroides genome employing the primer sets BG11866/BG11867 and BG11869/BG11871 for the 500 bp sizes and the primer sets BG11866/BG11868 and BG11870/BG11871 for the 1 kb sizes. HiFi assembly and transformation to E. coli DH5α cells was followed by sequence verification of the obtained pBBR_Cas9_Δupp500HR_NT and pBBR_Cas9_Δupp1000HR_NT plasmids. The sequence verified plasmids were transformed to E. coli S17-1 cells.

We proceeded with the construction of the HR-editing, upp targeting plasmids pBBR_Cas9_Δupp500HR_sp2 and pBBR_Cas9_Δupp1000HR_sp2. The previously mentioned couples of primer sets were used for the generation of the 500 bp and 1 kb flanking sites. The pBBR_Cas9_NT vector was employed as template for the PCR amplification of the conserved backbone region with the BG11887/BG11888 primer set. In addition, the spacer sp2 was included in the corresponding position employing the primer sets BG11182/BG11487 and BG11489/BG11886 for the amplification of the cas9 gene and the sgRNA module, respectively. HiFi assembly and transformation to E. coli DH5α cells was followed by sequence verification of the obtained pBBR_Cas9_Δupp500HR_sp2 and pBBR_Cas9_Δupp1000HR_sp2 plasmids. The sequence verified plasmids were transformed to E. coli S17-1 cells.

Plasmids for upp KI

We proceeded with the construction of the upp knock-in (KI) control pBBR_Cas9_Klupp1000HR_NT plasmid and the upp KI plasmids pBBR_Cas9_Klupp1000HR_sp4 and pBBR_Cas9_Klupp1000HR_sp5. The pBBR_Cas9_NT plasmid was used as template for the amplification of the fragments containing the backbone (primer set BG11187/BG11188), the cas9 gene and the targeting sgRNAs. The cas9- and the sgRNA-containing fragments were generated via PCR amplification with primer sets BG11182/BG12907 and BG11186/BG12908 for spacer sp4 and primer sets BG11182/BG12909 and BG11186/BG12910 for spacer 5. Additionally, the HR template for KI of the upp gene back to its native position were comprised of the upp gene sequence flanked by the 1000 bp genomic regions upstream and downstream of the upp gene. The genomic amplification for the 1000-bp long flanking sites was performed using the primer pairs BG12347/BG11868 and BG11870/BG12348. The primers BG12347 and BG12348 introduce a single point mutation in the upp start codon (ATG to GTG). The construction of the two editing plasmids, namely pBBR_Cas9_Klupp1000HR_sp4 and pBBR_Cas9_Klupp1000HR_sp5, was performed via 5-fragment assemblies. The assembly mixtures were then transformed to E. coli DH5α cells. Upon sequence verification, the plasmids were transformed to E. coli S17.

Plasmids for pha8 KO

We employed the previously described cloning strategy for generating three phaB KO plasmids (pBBR_Cas9_ΔphaBHR_sp1, pBBR_Cas9_ΔphaBHR_sp2 and pBBR_Cas9_ΔphaBHR_sp3). The pBBR_Cas9_NT plasmid was used as template for the amplification of the fragments containing the backbone (primer set P302/P303), the cas9 and the targeting sgRNAs. Each plasmid carried a different spacer designed to target a protospacer within the phab sequence. For spacer sp1, the cas9- and the sgRNA-containing fragments were generated via PCR amplification with primer sets P301/P383 and P304/P384, respectively. For spacer sp2 the cas9- and the sgRNA-containing fragments were generated via PCR amplification with primer sets P301/P385 and P304/P386, respectively. For spacer sp3 the cas9- and the sgRNA-containing fragments were generated via PCR amplification with primer sets P379/P380 and P381/P382, respectively. Each of the three phaB editing plasmids contained the same HR template, consisted of the genomic regions 1 kb upstream of the phaB start codon and 1 kb downstream of the phaB stop codon. The regions were generated via PCR amplification with primer sets P379/P380 and P381/P382, respectively. The construction of the three editing plasmids was performed via 5-fragment assemblies. The assembly mixtures were then transformed to E. coli S17 cells and the constructed plasmids were sequence verified.

We employed the previously described cloning strategy for generating three phbB KO plasmids (pBBR_Cas9_ΔphbBHR_sp1 and pBBR_Cas9_ΔphbBHR_sp2). The pBBR_Cas9_NT plasmid was used as template for the amplification of the fragments containing the backbone (primer set P302/P303), the cas9 and the targeting
sgRNAs. Each plasmid carried a different spacer designed to target a protospacer within the removed phbB sequence. For spacer sp1, the cas9- and the sgRNA-containing fragments were generated via PCR amplification with primer sets P301/P418 and P304/P417, respectively. For spacer sp2 the SpCas9- and the sgRNA-containing fragments were generated via PCR amplification with primer sets P301/P420 and P304/P419, respectively. Each of the two phbB editing plasmids contained the same HR template, consisted of a sequence of four stop codons (TAG) and an EcoRI restriction site, flanked on the one side by the genomic region 500 bp upstream and 546 bp downstream the phbB start codon and on the other side by the genomic region 145 bp upstream and 850 bp downstream the phbB stop codon. The regions were generated via PCR amplification with primer sets P415/P416 and P413/P414, respectively, using the R. sphaeroides genome as template. Moreover, each plasmid carried a different spacer designed to target a protospacer within the 59 nt long substituted phbB region. The construction of the two editing plasmids was performed via 5-fragment assemblies. The assembly mixtures were then transformed to E. coli S17 cells and the constructed plasmids were sequence verified.

**Diparental conjugation and plasmid curing**

Rhodobacter sphaeroides was inoculated from glycerol stock in 10 mL RÄ medium and incubated for 2 days at 30 °C and 200 rpm prior to conjugation. After 48 h, 50 μL of culture were transferred to fresh 10 mL RÄ medium, incubated for 24 h under the same conditions, and harvested when the OD at 600 nm was approximately 3. E. coli S17-1 harbouring the desired plasmid was grown overnight in LB supplemented with kanamycin at 37 °C and 200 RPM. Then, the overnight culture was transferred to fresh 2xYT media with antibiotic and grown for 2 h until the OD at 600 nm was approximately 1. The cell suspension was harvested and washed twice with 1 mL of RÄ medium, mixed with R. sphaeroides culture at a 1:1 ratio (1 mL each) and centrifuged for 1 min (maximum speed). The pellet was concentrated by suspension in 100 μL of RÄ medium, which was transferred to a sterile 0.22 μm 47 mm diameter nitrocellulose filter disc (Sigma-Aldrich) on a PY agar plate and incubated for 6 h at 28 °C. The conjugation mixture was harvested and resuspended in 2 mL of RÄ medium. 100 μL of diluted culture was spread on RÄ agar plates containing kanamycin and incubated at 30 °C for 3 days until colonies appeared. Purification of R. sphaeroides conjugants from E. coli S17-1 donors was performed by subsequent transfer of R. sphaeroides colonies to LB agar medium supplemented with kanamycin until no E. coli colonies appeared. Finally, a colony PCR was performed using primer set P3/P4 to verify that E. coli was not present in the culture.

Plasmid curing from positive R. sphaeroides mutants was carried out by cultivating the mutants in 5 mL of RÄ medium without antibiotic for 24 h, twice. The cells from the second cultivation were then diluted and spread on RÄ agar plates. The individual colonies were streaked onto LB agar with and without kanamycin. Those cells that grew on LB without antibiotic but were unable to grow in the presence of antibiotic were considered to have lost the plasmid. Final verification of plasmid curing was conducted by performing PCR with primers BG10937/BG10938, which are specific for the Cas9 gene.

**RNA extraction and reverse transcriptase PCR (RT-PCR)**

To check expression of the cas9 gene under the lac promoter, total RNA of R. sphaeroides conjugated with the pBBR_Cas9_NT plasmid was extracted employing the RNeasy mini kit (Qiagen, Germany) according to the manufacturer’s protocol and treated with DNAse I (NEB) to remove genomic DNA contamination in the sample. The SuperScript III Reverse Transcriptase kit (Invitrogen) was used for RT-PCR. When the first strand of cDNA was synthesized, the primers BG11112 and BG11115 were used to verify transcription activity of Cas9 gene using standard PCR.

**5-FU screening**

For the 5-fluorouracil (5-FU) screening process we prepared RÄ agar plates supplemented with 5-FU up to 100 μg/mL final concentration, using a 50 mg/mL stock solution prepared in dimethyl sulfoxide (DMSO). Colonies obtained from conjugation were picked and streaked on RÄ-5-FU plates with kanamycin. Survival colonies were isolated and their upp locus was then PCR amplified and sequenced.

**Analytics**

**Carbon and nitrogen concentrations**

To determine active biomass levels and C/N ratios of the biomass, analysis was done using a TOC-L analyzer (Shimadzu). Biomass samples were diluted 3–5 times in a total volume of 15 mL using MilliQ and a stirring bar was added. The TOC-L analysed both non-purgeable organic carbon (NPOC) as well as total nitrogen (TN) content. The TN content was used to calculate the active biomass concentration using the elemental composition of CH$_{1.99}$O$_{0.5}$N$_{0.19}$ of R. sphaeroides described previously [36]. The C/N ratio was calculated by dividing molar concentrations resulting from NPOC and TN measurements.
**Polyhydroxybutyrate (PHB) concentrations**

Volumes ranging from 2 to 5 mL of biomass were spun down and resuspended in MilliQ water. This was spun down and pellets were dried in an oven at 100 °C. The dried biomass was resuspended in 1 mL of concentrated H₂SO₄ and heated to 90 °C for 40 min. The samples were diluted by addition of 9 mL of 0.09 mM H₂SO₄. Analogous to the organic acid analysis, samples were mixed 1:1 with propionic acid. Separation of compounds was done using the same flow and temperature as the organic acids. For quantification, standards with known concentrations of commercially available PHB (Sigma-Aldrich) were treated the same as the samples.

**Results and discussion**

**SpCas9 targeting in *R. sphaeroides***

The first aim of this study was to assess potential toxic effects of SpCas9 expression in *R. sphaeroides* cells, as previously reported for other microbial species [37–39]. For this purpose, we constructed the pBBR_Cas9_NT control vector by cloning the *cas9* gene, codon optimized for *R. sphaeroides*, and a sgRNA expressing module with a non-targeting (NT) spacer in the *E. coli–Rhodobacter* shuttle vector pBBR1MCS2. The expression of the *cas9* gene was under control of the Plac promoter that due to the absence of the lacI repressor gene in the *R. sphaeroides* genome, has constitutive transcription activity. Moreover the sgRNA expressing module was under the control of the synthetic constitutive P_BbaJ95023 promoter [40] and the NT spacer sequence was designed in such way that any candidate protospacer within the genome of *R. sphaeroides* would contain at least 6 mismatches in the non-seed region (the 12 PAM-distal nucleotides) and at least 2 extra miss-matches in its seed region (the 8 PAM-proximal nucleotides). The pBBR_Cas9_NT vector in parallel with the pBBR1MCS2 control vector were conjugated in *R. sphaeroides* cells (Additional file 1: Figure S2). Even though the conjugation efficiency for the pBBR_Cas9_NT vector was reduced compared to the control pBBR1MC2 vector (Fig. 1a), it remained high and the size of the pBBR_Cas9_NT containing colonies was comparable to the size of the pBBR1MC2 containing colonies. We then randomly selected 5 of the colonies conjugated with the pBBR_Cas9_NT vector for cultivating and plasmid isolation. The sequence integrity of the *cas9* gene, the sgRNA expressing locus and the corresponding promoters was verified for all the 5 isolated plasmids, leading us to the conclusion that the drop in the conjugation efficiency of the pBBR_Cas9_NT can most likely be attributed to the almost double size of the vector (9566 bp) compared to the control vector (5144 bp) and, at least under the tested conditions, not to toxicity of SpCas9. Moreover, we confirmed the transcription of the *cas9* gene in *R. sphaeroides* by RT-PCR (Fig. 1b).

We further developed a SpCas9-based system for efficient introduction of lethal double stranded DNA breaks (DSBs) in *R. sphaeroides*, genome, selecting the *upp* (uracil phosphoribosyltransferase) gene as the proof of principal target. Knock-out mutations of the *upp* gene in other bacteria resulted in mutant strains resistant to 5-fluorouracil [41, 42]. For *R. sphaeroides* we determined that 5-Fluorouracil is toxic at concentrations as low as 1 μg/mL in RÄ medium and its use could facilitate future screening steps. The pBBR_Cas9_NT vector was employed for the construction of three targeting plasmids (pBBR_Cas9_sp1-3) each containing a unique spacer that corresponds to a different target sequence (protospacer) within the *upp* gene. If the constructed SpCas9 system is efficiently expressed in *R. sphaeroides*, the number of obtained colonies upon conjugation with the targeting plasmids is going to be substantially lower compared to the number of obtained colonies upon conjugation with the control plasmid.

The pBBR_Cas9 series of plasmids was conjugated in *R. sphaeroides* (Fig. 1c). The conjugation efficiency for the pBBR_Cas9_sp1 and pBBR_Cas9_sp2 constructs dropped more than 3 orders of magnitude compared to the pBBR_Cas9_NT control, while for the pBBR_Cas9_sp3 the conjugation efficiency dropped only by 50% (Fig. 1c). This result shows that the constructed SpCas9 targeting system is highly active in *R. sphaeroides*. To further study the observed fluctuation of targeting efficiency of different spacers, we randomly selected 5 surviving colonies conjugated with the pBBR_Cas9_sp3 plasmid, isolated the plasmids. The plasmid sequencing results did not reveal any indel mutations or single nucleotide substitutions in the *cas9* gene and the sgRNA module that could lead to the deactivation of the SpCas9 targeting system. Previous studies have shown that CRISPR-Cas based chromosomal targeting can induce genomic island excision events, resulting to the removal of the chromosomal target in the surviving population [43, 44]. We PCR amplified the *upp* genomic regions from the previously selected 5 surviving colonies conjugated with the pBBR_Cas9_sp3 plasmid; all the amplicons showed the expected wild-type size (Additional file 1: Figure S3) and were also sequence verified. This result excluded a genomic island excision event. It was previously reported that the targeting efficiency of a SpCas9 system relies heavily on the selected spacer and can substantially differentiate amongst different targets, even within the same gene [45]. Hence, the most possible explanation for the low targeting efficiency of the SpCas9-based system employing spacer 3 is the comparatively higher efficiency of the native RecA-based repair mechanism that employs
untargeted chromosomal copies as repairing templates [45].

**Homologous recombination-**SpCas9** counter selection for gene deletions**

There are multiple studies on bacterial genome editing employing (i) homologous recombination (HR) via plasmid-borne templates that carry the desired modifications, and (ii) Cas9-induced DSBs for the induction of the cellular HR machinery and as a counter-selection system for the unedited cells [24, 34]. Previous studies reported HR activity in *R. sphaeroides* [11] and here we developed a SpCas9 targeting system. Hence, we set out to develop a HR-SpCas9 counter selection system for efficient genome editing in *R. sphaeroides*.

As a proof of principle, we programmed the designed HR-SpCas9 system to knock out the *upp* gene of *R. sphaeroides*, excluding the start codon and the last 12 nucleotides of the gene in order to avoid potential polar effects for neighboring genes of the operon. Two editing plasmids were constructed and tested, both containing the previously described spacer 2 for efficient *upp* targeting and HR templates consisted of either 500 bp (pBBR_Cas9_Δ*upp*500HR_sp2) or 1 kb (pBBR_Cas9_Δ*upp*1000HR_sp2) upstream and downstream genomic regions flanking the selected for deletion *upp* fragment. Two control plasmids (pBBR_Cas9_Δ*upp*500HR_NT and pBBR_Cas9_Δ*upp*1000HR_NT), containing the same HR templates as the editing plasmids but a non-targeting spacer, were also taken along for assessing the contribution of SpCas9 to the efficiency of the tool.

Upon conjugation of the above described constructs in *R. sphaeroides*, a drop was observed 1 order of magnitude in the number of surviving colonies upon conjugation.
Homologous recombination-SpCas9 counter selection for gene insertions and single nucleotide substitutions

To further expand the genome editing toolbox of *R. sphaeroides*, we proceeded with the development of a plasmid-based HR-SpCas9 counter selection knock-in (KI) system. As a proof of principle, we designed a system for reinsertion of the *upp* gene in the genome of the previously constructed Δ*upp* mutant strain. We selected two spacers (sp4 and sp5, Additional file 1: Table S5) for the editing plasmids, aiming to target the remaining fraction of the *upp* gene in the genome of the Δ*upp* mutant (Fig. 3a). Moreover, we constructed HR templates containing the *upp* gene - with the “ATG” start codon substituted by the “GTG” as sequence verification marker-flanked by 1 kb upstream and downstream genomic regions (pBBR_Cas9_Klupp1000HR_sp4 and pBBR_Cas9_Klupp1000HR_sp5). Upon reinsertion of the *upp* gene into the Δ*upp* genome the corresponding protospacers would be disrupted, providing resistance from the SpCas9 targeting.

The two editing plasmids and the pBBR_Cas9_NT control plasmid were conjugated in the plasmid-cured *R. sphaeroides* Δ*upp* mutant and the experiment was performed in 3 biological replicates. As expected, a decrease in the number of surviving conjugants was observed for both editing plasmids compared to the non-targeting control plasmid (Fig. 3b). The *upp* genomic region from 38 to 40 colonies per construct and per conjugation was amplified by colony PCR with genome specific primers.

The editing constructs resulted in low editing efficiencies; 16/118 screened colonies for spacer 4 and only 2/120 screened colonies for spacer 5 were KI mutants (Additional file 1: Figure S5A, B). All the mutants were subsequently sequence verified for the existence of the “GTG” start codon (Fig. 3c). This result shows that, unlike the *upp* deletion process, the efficiency of the *upp* knock-in process is much lower, at least for the selected target.

Eliminating PHB accumulation in *R. sphaeroides*

The intracellular accumulation of PHB in *R. sphaeroides* under nitrogen-limited culturing conditions acts as a carbon drain and a NADP⁺ regeneration mechanism, using acetoacetyl-CoA (AA-CoA) as precursor [6, 7, 46] (Fig. 4a). Hence, engineering a *R. sphaeroides* strain with reduced PHB production capacity could facilitate the increment of the intracellular NADPH and AA-CoA pools, and the channelling of the *R. sphaeroides* metabolism towards the production of alternative biotechnologically interesting molecules, such as hydrogen and terpenes.

AA-CoA reductases catalyse the NADPH-dependent conversion of AA-CoA into (R)3-hydroxybutyrate-CoA ((R)3HB-CoA) which is the precursor of PHB. There are two predicted AA-CoA reductase genes in the genome of *R. sphaeroides*, namely *phaB* (RSP_0747) and *phbB* (RSP_3963), and the contribution of each corresponding enzyme to the AA-CoA reduction into (R)3HB-CoA was the focus of this study. For this purpose, the *phaB* and *phbB* genes were separately and jointly knocked-out of the *R. sphaeroides* genome employing the previously described tool. More specifically, three editing plasmids were constructed and employed for the deletion of the *phaB* gene (namely pBBR_Cas9_AphaBHR_sp1, pBBR_Cas9_AphaBHR_sp2 and pBBR_Cas9_AphaBHR_sp3) and two for the knockout of the *phbB* gene (namely pBBR_Cas9_AphbBHR_sp1 and pBBR_Cas9_AphbBHR_sp2) via the introduction of premature stop codons in the sequence of the gene. We reasoned to not completely delete the *phbB* gene as it is located within an operon in the *R. sphaeroides* genome, unlike *phaB* which is located at the end of an operon (Additional file 1: Figure S6). The
efficiency of the phaB deletion processes was 100% for constructs harbouring spacer sp1 (20/20 PCR screened colonies were clean phaB deletion mutants) and spacer sp2 (10/10 PCR screen colonies were clean phaB deletion mutants), but 0% efficiency for constructs harbouring spacer sp3 (0/13 PCR screened colonies were clean phaB deletion mutants) (Additional file 1: Figure S7). The efficiency of the phbB knockout processes was 100% when
the pBBR_Cas9_ΔphbHR_sp1 construct was employed (7/7 PCR screened colonies were clean phbB deletion mutants) and 0% when the pBBR_Cas9_ΔphbHR_sp2 construct was employed (0/8 PCR screened colonies were clean phbB deletion mutants) (Additional file 1: Figure S8), confirming that the efficiency of the developed Cas9-based is spacer-dependent.

The C/N ratio of microbial biomass increases when PHB is produced. The C/N ratio of the wild type R. sphaeroides strain and the generated ΔphaB, ΔphbB and ΔphaB_ΔphbB strains was tested for PHB production after 48 h of cultivation under nitrogen excess and nitrogen-limited conditions. Analysis by TOC-L revealed no statistically relevant difference between the C/N ratios of the strains under nitrogen excess conditions (Fig. 4b, left panel). Under nitrogen limitation conditions, a substantial decrease was noticed in the C/N ratio of the ΔphaB strain as compared to the wild-type strain (Fig. 4b, Additional file 1: Table S6). This decrease indicated reduced carbon accumulation in the ΔphaB and ΔphaB_ΔphbB strains in comparison to the wild-type strain, which could be attributed to reduced PHB production.

In order to confirm the reduction in the PHB production, HPLC analysis was performed to quantify the intracellular PHB content of the wild-type, ΔphaB, ΔphbB and ΔphaB_ΔphbB strains. The analysis showed that the PHB content of the wild-type and ΔphbB strains can fluctuate between 40 and 70% (w/w) of the dry cellular weight (DCW), which is consistent with previously reported results [7, 47]. The PHB concentrations in the ΔphaB and ΔphaB_ΔphbB strains were comparable and more than 99% lower compared to the PHB concentrations of the wild type and ΔphbB strains. The active biomass of wild-type and mutant strains was calculated based on the TOC-L data (Fig. 4c, Additional file 1: Table S6). Only a minor reduction in the active biomass of the ΔphaB and ΔphaB_ΔphbB strains, as compared to the active biomass of the wild-type strain, was observed.
Conclusions

In our work we developed a highly active SpCas9-based DNA targeting system for the biotechnologically interesting bacterium *R. sphaeroides*. We further combined the SpCas9 DNA targeting activity with the native homologous recombination (HR) mechanism of *R. sphaeroides* for the development of an efficient HR-SpCas9 counter selection-based genome editing toolbox for gene knock-outs, knock-ins and single nucleotide substitutions. Hence, the developed toolbox can substantially accelerate fundamental and metabolic engineering studies on *R. sphaeroides*, e.g. on membrane and photosynthetic reaction centre proteins, as well as metabolic engineering studies for improved hydrogen production and terpene synthesis, as already have been achieved for other microorganisms with developed SpCas9 toolboxes. Finally, we employed the developed toolbox to elucidate the genetic background of PHB production in *R. sphaeroides* and we constructed mutant strains with dramatically reduced PHB production capacity and almost unaffected growth. Hence, the here constructed *R. sphaeroides* mutant strains can be the basis for further engineering towards the enhanced production of

![Fig. 4 a](image) The PHB biosynthetic pathway of *Rhodobacter sphaeroides*. The annotation is based on KEGG. Enzymatic conversions are indicated by their EC numbers. Double arrowed reactions describe reversible reactions ($\Delta G^\circ > -30$ kJ/mol [48]). Ac-CoA acetyl-CoA; AcAc-CoA acetoacetyl-CoA; (R)-3-HB-CoA (R)-3-hydroxybutanoyl-CoA; (S)-3-HB-CoA (S)-3-hydroxybutanoyl-CoA, Crot-CoA crotonyl-CoA; PHB poly-β-hydroxybutyrate. The interrupted square indicates the metabolic step that is hypothesized to be catalysed by *phaB* and/or *phbB*. b Effect of the *phaB* (RSP_0747) and the *phbB* (RSP_3963) knockout, as well as of the combined knockout, on the C/N ratios of the generated mutants in nitrogen excess and limiting conditions on a defined medium. Lower C/N ratios indicate less PHB accumulation (Student’s *t*-test, *P < 0.05). c Effect of the *phaB* (RSP_0747) and the *phbB* (RSP_3963) knockout, as well as of the combined knockout, on the acids production and the PHB accumulation in *Rhodobacter sphaeroides* (Student’s *t*-test, ***P < 0.001). Concentrations of active biomass, PHB, pyruvate and oxo-glutarate were measured after 24 h cultivation in Sistrom’s minimal medium supplied with 1.0 g/L or 0.25 g/L of NH₄Cl (Nitrogen excess and nitrogen limited conditions, respectively). Error bars indicate standard deviations from at least two replicates.
alternative biotechnologically interesting molecules, such as hydrogen and terpenes.

Supplementary information

Supplementary information accompanies this paper at https://doi.org/10.1186/s12934-019-1255-1.

Additional file 1. Additional figures and tables.

Abbreviations

PHB: poly-β-hydroxybutyrate; HR: homologous recombination; SCO: single crossover; sacB: levansucrase gene; CRISPR-Cas: Clustered Regularly Interspaced Short Palindromic Repeats-CRISPR associated proteins; DSB: double stranded DNA breaks; sgRNA: single guide RNA; PAM: protospacer adjacent motif; upp: uracil-phosphoribosyltransferase gene; 5-FU: 5-fluorouracil; AA-CoA: acetyl-CoA; phbB: AA-CoA reductase; phbA: AA-CoA reductase.

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Authors’ contributions

IM, EO, SK, RW and JvO designed the work. IM, EO, MRG, AM, WP and BAP conducted, analyzed, and interpreted the experiments. IM and EO drafted and wrote the manuscript. All authors read and approved the final manuscript.

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Availability of data and materials

The datasets supporting the conclusions of this article are included within the article (and its additional file).

Ethics approval and consent to participate

Not applicable.

Consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing financial interests.

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