Tailor-made exopolysaccharides—CRISPR-Cas9 mediated genome editing in *Paenibacillus polymyxa*

Marius Rütering¹,², Brady F. Cress²,³, Martin Schilling⁴, Broder Rühmann¹, Mattheos A. G. Koffas²,³, Volker Sieber¹,⁵,⁶, and Jochen Schmid¹,*

¹Chair of Chemistry of Biogenic Resources, Technical University of Munich, Straubing, Germany, ²Center for Biotechnology and Interdisciplinary Studies, Rensselaer Polytechnic Institute, Troy, NY, USA, ³Department of Chemical and Biological Engineering, Rensselaer Polytechnic Institute, Troy, NY, USA, ⁴Evonik Nutrition and Care GmbH, Kirschenallee, Darmstadt, Germany, ⁵Fraunhofer IGB, Straubing Branch Bio, Electro, and Chemocatalysis BioCat, Straubing, Germany and ⁶Catalysis Research Center, Technical University of Munich, Garching, Germany

*Corresponding author: E-mail: j.schmid@tum.de

Abstract

Application of state-of-the-art genome editing tools like CRISPR-Cas9 drastically increase the number of undomesticated micro-organisms amenable to highly efficient and rapid genetic engineering. Adaptation of these tools to new bacterial families can open up entirely new possibilities for these organisms to accelerate as biotechnologically relevant microbial factories, also making new products economically competitive. Here, we report the implementation of a CRISPR-Cas9 based vector system in *Paenibacillus polymyxa*, enabling fast and reliable genome editing in this host. Homology directed repair allows for highly efficient deletions of single genes and large regions as well as insertions. We used the system to investigate the yet undescribed biosynthesis machinery for exopolysaccharide (EPS) production in *P. polymyxa* DSM 365, enabling assignment of putative roles to several genes involved in EPS biosynthesis. Using this simple gene deletion strategy, we generated EPS variants that differ from the wild-type polymer not only in terms of monomer composition, but also in terms of their rheological behavior. The developed CRISPR-Cas9 mediated engineering approach will significantly contribute to the understanding and utilization of socially and economically relevant *Paenibacillus* species and extend the polymer portfolio.

Key words: exopolysaccharides; CRISPR-Cas9; genome editing; *Paenibacillus polymyxa*

1. Introduction

Value-added compounds synthesized by microorganisms, such as alkaloids, flavonoids, terpenoids, polyketides, lipopeptides, biofuels and exopolysaccharides (EPSs), are of huge interest for a variety of applications in the food, medicine, agriculture and consumer goods industries (1–3). Although advances in synthetic biology and metabolic engineering have significantly contributed to the design of improved microbial factories, robust heterologous expression of complex pathways is often hampered by product toxicity, low yields and the absence or insufficient availability of biosynthetic precursors (4, 5). State-of-the-art genome editing tools like CRISPR-Cas9 rapidly increase the accessibility of undomesticated strains to genetic engineering, and therefore pave the way for taming wild-type (WT) species in order to construct new, biotechnologically relevant production strains (6). Thorough implementation of such tools in a species of interest is of fundamental importance for efficient rewiring of metabolic circuits and optimization or alteration of the produced...
metabolites. Gram-positive bacteria of the phylum firmicutes are promising candidates for this endeavor. Their robustness toward environmental stress in combination with their promiscuity toward different carbon sources, the huge variety in gene clusters dedicated to secondary metabolite production and their well-established cultivability exemplify their potential as customizable microbial factories (7–11). Several genera of this phylum, like Lactobacillus and Clostridium, have been used in biotechnology for centuries, both consciously and unconsciously (12,13). Numerous recent studies demonstrated the successful application of CRISPR-Cas9 in a variety of firmicute families including Bacillaceae, Lactobacillaceae, Clostridiceae and Staphylococcaceae (14–19). However, no reports on CRISPR-Cas9 based genome editing tools for Paenibacillaceae are available yet, and existing genetic engineering approaches are limited due to low efficiencies or the dependence on integration of selectable markers (20,21). Nevertheless, this family comprises several species of economic and social relevance (Figure 1). Paenibacillus larvae, for example, is the causative agent of American Foulbrood, a lethal disease of honeybee larvae (22), posing grave concern for the future of agriculture, and some Paenibacilli are known to be opportunistic human pathogens (23,24). Easily deployable vectors facilitating rapid elucidation of the genetic basis for pathogenicity, immunogenicity and toxicity would hold tremendous scientific value. On the constructive side, numerous studies describe the beneficial use of Paenibacilli in miscellaneous fields. In agriculture, they are critical because of their intrinsic capacity for nitrogen fixation (25) and phosphate solubilization (26), thereby directly promoting plant growth. Furthermore, they produce a variety of insecticides (27) and antimicrobials like polymyxins (28) and fusaricidins (29), which protect plants from phytopathogens and have potential use in medical applications. Their extensive enzymatic capabilities to degrade complex carbohydrates and to produce and tolerate high levels of commercially relevant chemicals like 2,3-butanediol, for example, make Paenibacilli an interesting genus for the fermentative production of this and other platform chemicals from renewable resources (30). Critically, they produce a class of still underappreciated but highly promising EPSs possessing antioxidant activity and outstanding rheological properties, qualifying them for applications in therapeutics or as thickeners (31–35). Grady et al. (36) recently reviewed the potential of Paenibacilli in agriculture and industrial biotechnology in detail.

EPSs are linear or branched, high-molecular weight polymers composed of sugars molecules, which are secreted into the extracellular environment during microbial growth. Due to variations in monomer composition, molecular weight and decoration with functional groups, these polymers exhibit an immense physicochemical versatility making them interesting for various applications (37). This structural variability is highlighted by the existence of over 350 annotated EPSs from prokaryotes (38). EPSs have been primarily used as rheological additives for food, agricultural feed, oil recovery and cosmetic applications (39). Prominent representatives of commercialized EPSs are sphinganes, xanthan, pullulan, dextran and levan (40–42). Although these compounds hold big shares of the markets for bio-based viscosifiers and gelling agents, they are only narrowly suited for specialized applications matching their imparted rheologies. Especially high-value niche applications like tissue engineering, cell encapsulation or drug delivery require explicitly defined physicochemical characteristics, which are not covered by the existing EPSs (43) without further, post-biosynthetic chemical or enzymatic modification.

Using synthetic biology for design and synthesis of tailor-made EPSs is a highly promising approach to fill these gaps. The aforementioned structural diversity of existing EPSs makes their associate biosynthetic pathways ideal templates for generating polymers with tunable properties through the rational engineering of novel structures (44). Typical targets for EPS engineering are functional groups like pyruvyl groups, which contribute to polymer charge density and thereby influence the rheological traits (45). However, the adjustment of substituent patterns only allows for alterations in the degree of superficial decorations while leaving the core, underlying glycan structure and sequence unchanged. Other engineering targets are the glycosyltransferases (GTs), which transfer defined sugar moieties to the nascent, pre-assembled repeating units and thereby determine the composition and linkage pattern of the mature EPS (46,47). Complementation experiments have shown that exchange of GTs with distinct monosaccharide preferences is feasible, indicating that the EPS polymerization and secretion machinery of one organism can potentially be harnessed for the production of various polymers with disparate structures and properties (37). The combination of in-depth characterizations of known and to-be-discovered GTs with protein-engineering will eventually yield a catalog of enzymes, which can be used for the directed incorporation of user-specified sugars imparting desired properties (48). Polymer variants produced in this fashion will successively contribute to the still superficial understanding of EPS structure–function relationships and thereby ultimately allow for the rational design of application-defined properties (43). Modern synthetic biology tools, such as CRISPR-Cas9, will not only facilitate engineering of structural features but will also drastically accelerate strain improvements, spurring the rise of robust and economical production processes for competitive, custom-made EPSs.

In this study, we describe the development of a CRISPR-Cas9 based genome-editing tool for Paenibacillus polymyxa. The single plasmid system was employed for highly efficient, homology directed deletions as well as integrations. The developed CRISPR method was subsequently used to annotate and provide the first experimental evidence of the gene cluster responsible for EPS biosynthesis in P. polymyxa DSM 365. Besides shutting down and significantly attenuating EPS biosynthesis, we were also able to produce structurally altered EPSs exhibiting fundamentally distinct rheological properties. On the basis of these findings, putative substrate specificities of two GTs were assigned.

**Figure 1.** Relevance of Paenibacilli for agriculture, society and industry. The main subject of this study, exopolysaccharides, are highlighted. A detailed review on all aspects can be found in reference [36].
2. Materials and methods

2.1 Plasmids, bacterial strains, primers and growth conditions

The bacterial strains, plasmids and oligonucleotides used in this study are listed in Supplementary Tables S1–S3. Escherichia coli strains were grown in Lysogen-broth (LB; 10 g l⁻¹ sodium chloride, 10 g l⁻¹ peptone and 5 g l⁻¹ yeast extract) at 37 °C. Paenibacillus polymyxa DSM365 (DSMZ, Braunschweig, Germany) was cultured at 30 °C in LB for genetic manipulations and in MM1 P100 for EPS production and phenotype evaluations [30 g l⁻¹ glucose, 1.35 g l⁻¹ magnesium sulfate heptahydrate, 1.67 g l⁻¹ potassium dihydrogen phosphate, 0.05 g l⁻¹ calcium chloride dihydrate, 2 ml l⁻¹ RPMI 1640 vitamins solution (Sigma-Aldrich) and 1 ml l⁻¹ trace elements solution containing 2.5 g l⁻¹ iron (II) sulfate heptahydrate, 2.1 g l⁻¹ sodium tartrate dihydrate, 1.8 g l⁻¹ manganese (II) chloride tetrahydrate, 0.075 g l⁻¹ cobalt (II) chloride hexahydrate, 0.031 g l⁻¹ copper (II) sulfate heptahydrate, 0.258 g l⁻¹ boric acid, 0.023 g l⁻¹ sodium molybdate and 0.021 g l⁻¹ zinc chloride] (32). Antibiotics were added at 50 µg ml⁻¹ for neomycin and 20 µg ml⁻¹ for polymyxin.

2.2 Construction of pCasPP

For construction of the pCasPP plasmid, the Cas9 gene was amplified alongside with the BbsI flanked lacZ cassette from pCRISPRomyces-2, which was a gift from Huimin Zhao (Addgene plasmid # 61737). The neomycin resistance and the repU gene were amplified from pUBoriMCS, which is a PUB110 derivate (49), containing a BbsI inserted origin of replication as well as a multiple cloning site from pUC18. A Bsal site within repU was removed by introducing a silent mutation with the utilized primers. The sgf33 promoter (50) was obtained as artificial gBlock (Integrated DNA Technologies). A cytosine was added by introducing the desired sequence mutations while PCR-amplifying the pCasPP plasmid in two pieces with corresponding primers. To inactivate Cas9, the same procedure was deployed and the two active sites were mutated to D10A and H840A, yielding pdCasPP. All cloning and mutation steps were verified by sequencing (Eurofins, Ebersberg, Germany).

2.3 Bioinformatics

In order to identify genes involved in EPS biosynthesis, the P. polymyxa DSM 365 genome was uploaded to RAST for automated genome annotation (51). Obtained data was screened manually and putatively identified genes involved in EPS-biosynthesis were annotated in detail using NCBI blastx (52) and UniProt blast (53). Intracellular protein localizations were predicted using CELLO v.2.5 (54). Biosynthetic pathways involved in nucleotide sugar production were identified using KEGG (55). Twenty base pair long spacers for Cas9 mediated edits were selected based on on- and off-target scores determined with benchling (http://www.benchling.com) using NGG as PAM motif and the uploaded P. polymyxa DSM 365 genome as reference. In silico cloning and sequence alignments were performed with SnapGene software (GSL Biotech, Chicago, IL, USA).

2.4 Genome editing in Paenibacillus polymyxa DSM 365

CRISPR-Cas9 mediated genome editing in P. polymyxa DSM 365 was performed as follows. Subsequent to spacer selection using benchling (http://www.benchling.com), 24 bp oligos for guide annealing were designed as described in the supplemental material of Cobb et al. (56). Oligonucleotides were phosphorylated, annealed and finally inserted into the pCasPP backbone via Golden Gate assembly. Subsequent to guide cloning, homologous regions were constructed by overlap extension PCR and inserted by restriction and ligation into the SpeI site. A detailed description of the entire cloning procedure can be found in the supplementary information. Constructed plasmids were sequenced and then transferred to E. coli S17-1 for conjugation events. Plasmid transfer from E. coli S17-1 to P. polymyxa DSM 365 was performed as follows. Overnight cultures of recipient and donor strains were subcultured 1:100 in non-selective or selective LB media, respectively, and grown until early exponential phase (4 h). Afterwards, 900 µl of the recipient culture were heat shocked for 15 min at 42 °C and mixed with 300 µl of the donor strain culture. Cells were harvested by centrifugation for 3 min at 8000 × g and the pellet was gently resuspended and dropped on non-selective LB agar. After overnight incubation at 30 °C, cells were scraped from the agar, resuspended in 500 µl 0.9% NaCl and plated on LB agar containing neomycin and polymyxin for counter selection. Paenibacillus polymyxa conjugants were obtained after 48 h incubation at 30 °C and screened for editing events using colony PCR. Sequence verification of genome edits was performed by amplifying edited regions from isolated genomic P. polymyxa DNA using suitable primers. Obtained PCR products were purified, adjusted to a concentration of 10 ng µl⁻¹, mixed with corresponding primers, and sent for sequencing.

2.5 EPS production and purification

For polymer production, baffled 250 mL shake flasks sealed with aluminum caps were filled with 100 mL MM1 P100 media and inoculated with 1 mL of a P. polymyxa overnight culture. Cultures were incubated at 30 °C and 170 rpm for 28 h. To extract EPS from shake-flask experiments, the culture broth was diluted 1:3 with distilled water to decrease viscosity and centrifuged for 30 min at 17 600 × g and 20 °C to separate cells. Subsequently, EPS was precipitated by slowly pouring the supernatant into two volumes of 2-propanol. The precipitated polymer was collected using a spatula and dried overnight at 45 °C in a VDL 53 vacuum drying oven (Binder, Tuttingen, Germany).

2.6 Molecular weight determinations

Gel permeation chromatography was performed using an Agilent 1260 Infinity system (Agilent Technologies, Waldbronn, Germany) equipped with a refractive index detector and a SECurity SLD7000 7-angle static light-scattering detector (PSS Polymer Standards Service GmbH, Mainz, Germany). Samples were analyzed using a Suprema guard column, one Suprema 100 Å (8 × 300 mm) and two
Suprema 10 000 Å (8 × 300 mm) columns (PSS Polymer Standards Service). The eluent, 0.1 M LiNO₃, was pumped at a flow rate of 1 ml min⁻¹ and the column compartment was kept at 50 °C. Samples were injected in 60 min intervals. Qualitative molecular weight results were obtained by comparing sample elution profiles with a 12-point pullulan standard curve.

2.7 Carbohydrate fingerprint

Simultaneous high resolution detection of carbohydrates which can be derivatized with 1-phenyl-3-methyl-5-pyrazolone (PMP) was performed via HT-PMP as described before (57). Briefly, 1 g l⁻¹ solutions of EPS were hydrolyzed in 2 M trifluoroacetic acid (TFA) for 90 min at 121 °C and subsequently neutralized with 3.2% (v/v) ammonium hydroxide. Thereafter, 25 µl neutralized sample was mixed with 75 µl derivatization reagent (0.1 M methanolic-PMP-solution:0.4% ammonium hydroxide solution 2:1) and incubated for 100 min at 70 °C. Finally, 130 µl of 19.23 mM acetic acid were added to 20 µl cooled sample and HPLC separation was performed on a reverse phase column (Gravity C18, 100 mm length, 2 mm i.d.; 1.8 µm particle size; Macherey-Nagel) tempered to 50 °C. For gradient elution, mobile phase A [5 mM ammonium acetate buffer (pH 5.6) with 15% acetonitrile] and mobile phase B (pure acetonitrile) were pumped at a flow rate of 0.6 ml min⁻¹. The HPLC system (Ultimate 3000RS, Dionex) was composed of a degasser (SDR 3400), a pump module (HPG 3400RS) auto sampler (WPS 3000TRS), a column compartment (TCC5000RS), a diode array detector (DAD 3000RS) and an ESL-ion-trap unit (HCT, Bruker). Standards for each sugar (2, 3, 4, 5, 10, 20, 30, 40, 50 and 200 mg l⁻¹) were processed as the samples, starting with the derivatization step. Data were collected and analyzed with BrukerHystar, QuantAnalysis and Dionex Chromleon software.

2.8 Pyruvate assay

To determine the pyruvate content of the polymers, 1 g l⁻¹ EPS solutions were hydrolyzed and neutralized as described for the sugar monomer analyses (58). To start the reaction, 10 µl neutralized sample + 90 µl of ddH₂O or standard were mixed with 100 µl assay mixture [50 µM N-(carboxymethylamino-carbonyl)-4.4'-bis(dimethylamino)-diphenylamine sodium salt (DA-64), 50 µM thiamine pyrophosphate, 100 µM MgCl₂ × 6H₂O, 0.05 U ml⁻¹ pyruvate oxidase, 0.2 U ml⁻¹ horseradish peroxidase, 20 mM K₂HPO₄ buffer pH 6] and incubated at 37 °C, 700 rpm for 30 min in a microplate shaker. The extinction was measured at 727 nm and subtracted with values measured at 540 nm to eliminate background signals. Nine standards in the range from 0.5 to 1000 µM pyruvate were used for calibration.

2.9 Rheological measurements

Rheological measurements were performed with an air-bearing MCR300 controlled-stress rheometer (Anton Paar Germany GmbH) using a cone plate geometry (50 mm diameter, 1° cone angle, 0.05 mm gap) at a constant, Peltier-controlled temperature of 20 °C. Data was collected and analyzed with Rheoplus V.3.61 software (Anton Paar). Viscosity curves were obtained during a logarithmic shear-rate ramp (γ = 0.1–1000 s⁻¹). The linear viscoelastic and further structural features of polymer solutions were determined with a shear stress amplitude sweep from 0.1 to 1000 Pa at a constant frequency of f = 1 Hz. Time dependent flow behavior on non-destructive stress was assessed during frequency sweeps from 0.01 to 100 Hz. All measurements were carried out in triplicates.

3. Results and discussion

3.1 CRISPR-Cas9 vector system

Design and construction of the pCasPP CRISPR-Cas9 expression plasmid (Figure 2) was inspired by a vector system which was efficiently used for genome editing in Streptomyces (56). Streptococcus pyogenes Cas9 (SpCas9) and synthetic guide RNA (sgRNA) region, including the BbsI flanked IncZ selection cassette and origin of transfer (oriT), were amplified from pCRISPyromyces-2 plasmid. The Cas9-encoding gene was set under transcriptional control of the broad-host-range S-layer gene promoter sgsE from Geobacillus stearothermophilus, which was previously shown to be functional in Paenibacillus alvei (21,50). Since the transcription start for guide expression is crucial for guide length and thereby targeting efficiency, the constitutive gapdh promoter for sgRNA expression was not changed. To the best of our knowledge, this is the first report on gapdh promoter functionality in Paenibacilli. Origin of replication (ori), neomycin resistance gene and the repU gene, involved in plasmid replication (59), were amplified from pUBori, a pUB110-derived plasmid which is readily taken up and propagated by P. polymyxa (unpublished data). All DNA parts were assembled using Golden Gate cloning (60). A unique SpeI site was utilized for insertion of homologous arms (1 kb upstream and 1 kb downstream, fused by overlap extension PCR), necessary for homology directed repair (HDR) after successful double strand break by the activity of Cas9.

Insertion of 20 bp long single guide spacers was performed via BbsI based Golden Gate cloning of annealed and phosphorylated primers. For multiplexing, gBlocks comprising two copies of the sgRNA region were utilized as described for the pCRISPyromyces-2 system (56). Plasmid transfer to P. polymyxa DSM 365 was conducted via conjugation using the E. coli S17-1 donor strain and counter-selection on polymyxin-containing media. The functionality of the designed system was first tested on a putative glycosyltransferase (pefF) of the hypothetical EPS cluster. Using this target, different variants of the pCasPP plasmid were assembled, and transformation efficiencies were evaluated (Table 1, Supplementary Figure S1).

Guided Cas9 resulted in high lethality in the absence of a repair template, yielding less than 0.2% colonies compared to the dummy plasmid pCasPP. Although not examined further, it is likely that these colonies resulted from escape mutation, which is probably due to reduced conjugation efficiency of the larger vector, but in noticeably more colonies than the guided variants. Colony PCR confirmed that all tested colonies transformed with the guided plasmid containing a repair template were successful knockout events (eight individual conjugants), whereas tested colonies from all other plasmid transformations showed to be the unedited WT (Supplementary Figure S2). Sequencing of at least three clones from pCasPP, pCasPP-pepFs1 and pCasPP-pepFs1-harms corroborated this finding. These results suggest that the non-homologous end-joining (NHEJ) DNA repair based on the Ku and LigD enzymes is not functional or insufficient to introduce indels in P. polymyxa, although the required genes for NHEJ are encoded within the organism’s genome (62). To more
we found that curing of the plasmid after editing was readily achieved by incubating liquid cultures of the knockout mutant for 72 h at elevated temperatures (37 °C) in absence of antibiotic with a single 1:100 sub-culturing after 36 h. Curing efficiency was assessed for the WT strain and six knockout strains. Specifically, eight colonies obtained from a culture of each knockout strain (by plating dilution series on non-selective agar) were picked and streaked on neomycin plates to test for neomycin sensitivity and to thus determine the curing efficiency. Out of the eight colonies obtained from subculture of each knockout strain, a cured colony sensitive to neomycin was readily found in each case (typically approximately 75% of tested colonies), confirming that the vector can be cured easily (Supplementary Figure S4). To investigate the possibility of performing genome-integration of heterologous DNA with the constructed system, a 300 bp pgrac promoter region was cloned between the 1 kb homologous regions of sacB, a gene encoding for a levansucrase in the P. polymyxa genome, and the editing experiments were performed as described above. All eight tested clones were colony PCR positive and sequencing verified the successful integration of the artificial pgrac promoter sequence, proving the functionality of pCasPP for genome integration (Supplementary Figure S5).

### Table 1. Conjugation efficiency of different pCasPP derivatives

<table>
<thead>
<tr>
<th>Plasmid</th>
<th>Colonies/conjugation</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>pCasPP</td>
<td>&gt;10 000</td>
<td>Non-targeting plasmid; no repair template</td>
</tr>
<tr>
<td>pCasPP-pepFsg1</td>
<td>&lt;20</td>
<td>Targeting pepF at Site 1; no repair template</td>
</tr>
<tr>
<td>pCasPP-pepFsg1-harms</td>
<td>&gt;200</td>
<td>Targeting pepF at Site 1; repair template provided</td>
</tr>
<tr>
<td>pCasPP-pepFsg2-harms</td>
<td>&gt;200</td>
<td>Targeting pepF at Site 2; repair template provided</td>
</tr>
<tr>
<td>pCasPP-harms</td>
<td>&gt;1000</td>
<td>Non-targeting plasmid; repair template provided</td>
</tr>
</tbody>
</table>

thoroughly assess the efficiency of the system, another knockout (ΔugdH1, found to result in phenotypically distinguishable colonies as a consequence of reduced EPS production) was investigated. Due to the ease of screening this knockout by colony morphology, a much larger sample of 50 colonies for each of two distinct spacers within the ugdH1 ORF were analyzed for their phenotypic appearance. Edited conjugants appeared as irregular, flat, opaque colonies with a brittle, dry consistency. WT colonies in contrast were loosely attached, circular and convex with a highly mucoid consistency. All conjugants for both spacers showed the phenotype consistent with an edited strain (Supplementary Figure S3). For all other edits described in this study, at least eight isolated colonies were analyzed using colony PCR, and each successful knockout was verified via sequencing.

In order to perform deletions in series, it is critical that the CRISPR-Cas9 plasmid is curable, so that the vector can be recursively transformed with new sgRNA inserts. Toward this end, we found that curing of the plasmid after editing was readily achieved by incubating liquid cultures of the knockout mutant for 72 h at elevated temperatures (37 °C) in absence of antibiotic with a single 1:100 sub-culturing after 36 h. Curing efficiency was assessed for the WT strain and six knockout strains. Specifically, eight colonies obtained from a culture of each knockout strain (by plating dilution series on non-selective agar) were picked and streaked on neomycin plates to test for neomycin sensitivity and to thus determine the curing efficiency. Out of the eight colonies obtained from subculture of each knockout strain, a cured colony sensitive to neomycin was readily found in each case (typically approximately 75% of tested colonies), confirming that the vector can be cured easily (Supplementary Figure S4). To investigate the possibility of performing genome-integration of heterologous DNA with the constructed system, a 300 bp pgrac promoter region was cloned between the 1 kb homologous regions of sacB, a gene encoding for a levansucrase in the P. polymyxa genome, and the editing experiments were performed as described above. All eight tested clones were colony PCR positive and sequencing verified the successful integration of the artificial pgrac promoter sequence, proving the functionality of pCasPP for genome integration (Supplementary Figure S5).

#### 3.2 Exopolysaccharide biosynthesis in Paenibacillus polymyxa DSM 365

Fundamental mechanisms of polysaccharide synthesis in bacteria have been extensively investigated in organisms such as E. coli, X. campestris and S. pneumoniae (63–65). While research on polysaccharide gene clusters in Gram-negatives started as early as the 1980s, the first studies on EPS genetics in Gram-positive bacteria were published approximately 20 years later (66). The best-characterized clusters of Gram-positive representatives are those of Lactobacillaceae and Streptococccaceae. EPS production by Paenibacillaceae is also described by several reports (34). However, except for the levansucrase catalyzed production of levansucrase in the presence of sucrase as carbon source (67), the genetic basis for the biosynthesis of heteropolysaccharides in this genus is unknown territory. To use the implemented genome editing tool for elucidation and engineering of EPS synthesis, the published P. polymyxa DSM 365 genome (62) was mined, and putative EPS-related genes were annotated thoroughly (accession number: BK010330).

The majority of coding sequences putatively involved in EPS synthesis were found clustered together, as is typical for most heteropolysaccharides (37) (Figure 3, Table 2). The size of the locus is at the high end of known bacterial EPS gene clusters, spanning almost 35 kb and comprising 28 coding sequences that could be assigned to polysaccharide production or hydrolysis. EPS gene clusters from Lactobacilli typically comprise 14 to 18 kb (68,69), whereas clusters from S. thermophilus can be up to 35 kb in size (70). Interestingly, several functional elements within the P. polymyxa DSM 365 EPS-cluster appear to be encoded twice. This could indicate an evolutionary development of the strains towards reliable EPS production. Another explanation could be that the strain is capable of producing two different polymers, with some genes being involved in both pathways (e.g. polymerases and precursors) and some being unique for each polymer (GTs). Some genes associated with precursor supply are encoded one (υgdH1) or two more times (manC, galU) at different loci of the genome. Fcl and gmd are exclusively found within the cluster.

![Figure 2. CRISPR-Cas9 vector system. Promoter sgSE controls SpCas9 expression, gupdh promoter regulates guide RNA levels. Spacer insertion is performed via Golden gate using BbsI. The unique SpeI site allows for insertion of homologous regions for HDR.](https://academic.oup.com/synbio/article-abstract/2/1/ysx007/4772606 by Rensselaer Polytechnic Institute user on 10 September 2018)
The encoded proteins suggest that EPS assembly and secretion in *P. polymyxa* follows the Wzx/Wzy dependent pathway, which is common in Gram-positive and Gram-negative bacteria and is also utilized for EPS formation in *Lactococci* and *Streptococci* (64). Through this pathway, the repeating units of the polymer are assembled on the inner side of the cytoplasmic membrane by GTs, starting with the transfer of the first sugar to a membrane bound lipid undecaprenyl carrier via the so-called priming GT. Thereafter, the nascent polysaccharide repeating unit is elongated through successive addition of distinct sugars by the other GTs. The fully assembled repeating units are then transferred over the membrane by a Wzx flippase and finally polymerized by the Wzy protein in reducing-end growth via a block transfer mechanism. A schematic representation of the hypothetical Wzx/Wzy dependent biosynthesis machinery of *P. polymyxa* is presented in Figure 4.

### 3.3 Exopolysaccharide engineering

The ultimate objective of the presented EPS engineering approach was to alter the chemical structure of the produced EPS in order to influence the physicochemical characteristics of the secreted polymer, yielding EPSs with new or superior material properties.
To achieve this goal, the developed CRISPR-Cas9 system was deployed for gene disruption studies targeting different genes within the identified EPS cluster. Five different genes were deleted individually, three of which are putative GTs (pepF, pepJ, pepE), and two of which are probably involved in precursor synthesis (ugdH1, manC). Additionally, an 18 kb fragment was deleted (clu), to generate an EPS deficient mutant (Figure 5). Details on the deletion sites can be found in Supplementary Table S4. After successful verification of deletions, strains were cured of the plasmid, and EPS of each mutant and WT strain was produced and purified under standardized conditions in biological triplicates (Table 3). To prevent falsification of EPS data by levan production, the utilized fermentation media contained glucose as sole carbon source. In a previous study, we showed that this media composition results in heteropolysaccharide production only upon glucose utilization (31).

All mutant strains produced less EPS than the WT, with ΔpepF and ΔpepJ secreting considerably less and ΔpepC and ΔugdH1 not producing any precipitable polymer. Diminished EPS titers after deletions of GTs are in accordance with studies in Lactobacilli, Xanthomonas and Streptococci reporting similar effects upon inactivation of transferases (71–73). In particular, priming GTs are known to be essential for EPS formation, resulting in EPS deficient mutants if deleted. The importance of enzymes involved in biosynthesis of nucleotide sugars is also described in literature. It has been shown many times that levels of key intermediates for nucleotide sugar biosynthesis directly correlate with obtained EPS titers (74). Li et al. (75) described the crucial role of two UDP-xyllose synthases (uxs) for EPS formation in a related Paenibacillus strain, secreting a xylene containing EPS. Inactivation of the first uxs gene reduced EPS titers by 50% and deletion of the second copy abrogated EPS biosynthesis completely. By disruption of the 18 kb fragment (Δclu) in P. polymyxa, all genes involved in polymerization (pepF, pepG) of the glycan were removed, logically resulting in an EPS deficient mutant. Knockout of manC creates a mutant that is not capable of producing GDP-Mannose and GDP-Fucose (Figure 6). At least one of these nucleotide sugars is probably an essential building block of the repeating unit, because deletion results in non-polymerizable or non-precipitable carbohydrates.

To characterize the EPSs produced by the P. polymyxa mutants and WT strain, molecular weights and monomer compositions of obtained polymers were analyzed. Molecular weights of all polymers were found to be in the same range (Table 3) and comparable to values found for Paenibacillus heteropolysaccharides in literature (76).

Of note, the ΔmanC mutant still showed a small but non-negligible polymer peak in GPC analyses, which was in the same order of magnitude in MW as all other polymers, indicating that still some full-length polymer was formed, but too little to allow for processing (Supplementary Figure S6). A possible explanation for this observation is that basal expression of the two other manC copies encoded in the genome compensates the deletion to a very little extent. Since regulation of cluster expression is presumably decoupled from the regulation of the other manC versions, this compensation is only marginal.

Similar findings were reported for capsule biogenesis in Streptococci (77). No polymer peak was observable by GPC for the Δclu mutant, experimentally proving that EPS biosynthesis was eliminated.
To compare sugar monomer compositions of produced WT and mutant EPSs, dried and ground polymer was re-dissolved, hydrolyzed and analyzed via HT-PMP analysis. The HPLC-MS data shows that all obtained polymers were composed of the same sugar monomers: glucose (Glc), mannose (Man), galactose (Gal), fucose (Fuc) and glucuronic acid (GlcA) (Figure 7). We assume that traces of glucosamine (GlcN) found in the samples are probably impurities from cell debris since its low amount does not suggest a stoichiometrically plausible participation in a conserved repeating unit. Sugar recoveries of about 50% during HPLC-MS (Table 3) are in the expected range for crude EPS batches (78). Since different sugars show different susceptibilities to release and degradation during hydrolysis, some are underrepresented in the final data.

Uronic acids, for example, are prone to degradation and can only be recovered partially (79). In contrast to this, dimers of uronic acids and hexoses are fairly stable, resulting in a reduced release and thereby reduced detection of attached hexose (78). Salts, co-precipitated protein and water, attracted by the hygroscopic EPS powders also contribute to recoveries below 100%. Uronic acids, for example, are prone to degradation and can only be recovered partially (79). Since colony morphology and viscosity of liquid cultures in-...
structure with a pronounced network, delivering high viscosity and stability to mechanical stress. No viscosity plateau at low shear rates can be observed in the viscosity curves of the WT EPS, indicating that polymer strands are linked with each other via inter-molecular forces. The remarkable difference between $G'_0$ and $G''_0$ in amplitude sweeps describes the elastic character of the WT solution. Upon increase of applied strain, the degeneration of the sample network begins with a $G''_0$ overshoot, which probably occurs due to micro crack formation within the structured gel (81). The depicted frequency test points out the stability of the sample over time. Even at low frequencies, simulating long-term stress, the sample character remains dominated by the elastic modulus and does not show any tendencies to start flowing. The WT EPS could potentially be applied as stabilizer for suspensions or as hydrogel for biomedical, cosmetic or food applications. The generated mutant EPS, however, reveals remarkably different attributes. A distinct viscosity plateau can be observed in the flow curves, assigning this sample to the group of entangled polymers without a strong physicochemical network. At low shearing, polymer strands simultaneously ravel and unravel, resulting in constant friction (82). Similar contributions of elasticity and viscosity to the sample structure and the lack of a $G''_0$ overshoot in strain sweeps emphasize the viscous but not gel-like character of the mutant EPS. In frequency

Figure 6. Biosynthetic pathways dedicated to the production of activated nucleotide sugars in *P. polymyxa*. The map was constructed based on a metabolic network model of a *P. polymyxa* strain annotated on KEGG database. All pathways present in the *P. polymyxa* genome are shown in black. Grey routes are not annotated. Enzymes written in purple are located within the EPS cluster, black ones somewhere else in the genome. Nucleotide sugars highlighted with color are constituents of the WT polymer.

Figure 7. Sugar monomer compositions of EPSs from mutants and WT *P. polymyxa*. Pyruvate content was determined with a pyruvate oxidase assay. All other components were analyzed and quantified via HPLC-ESI-MS/MS.
sweeps, this solution behaves like a typical Maxwell material. Sudden deformations, simulated by high frequencies, result in a rather elastic response of the material, whereas long-term stress induces viscous flowing of the solution. This polymer is suited for any application in which a shear-thinning, viscous thickener is needed. Examples are cosmetic lotions, oil drilling fluids or paints and lacquers.

4. Conclusion

In the present study, we designed and adapted a CRISPR-Cas9 based genome-editing tool for *P. polymyxa* for the first time. We proved its functionality in knockout studies of single genes and large regions via sgRNA multiplexing and harnessed it for genome integration experiments. After implementation, we utilized the system to study the yet undescribed EPS biosynthesis machinery of *P. polymyxa*. Results obtained here yield the first insights into basic principles of this Wzx/Wzy pathway, and putative roles of selected key genes were assigned. Furthermore, we exemplified EPS tailoring through genetic recoding. Specifically, we generated mutant EPSs with significantly altered monomer compositions. Rheological characterization revealed that one of these polymer variants exhibits fundamentally different physicochemical properties than the WT, making it suitable for an entirely different set of applications. Future work will focus on the in-depth characterization of all genes involved in biosynthesis of *P. polymyxa* EPS in order to construct a biotechnologically relevant production strain for tailor-made EPS. Thorough chemical structure analysis via NMR will enhance our understanding of structure-function relationships of generated EPS variants. Furthermore, an already constructed, inactivated variant of the pCasPP plasmid (pCasPP) will be used for CRISPRi-mediated repression studies in *P. polymyxa*. We also anticipate that the vector system will expedite research in distinct fields related to *Paenibacillus*, including the production of other value-added products like 2,3-butanediol and health related issues like *P. larvae* pathogenesis in honeybee larvae.

Supplementary data

Supplementary Data are available at SYNBIOL Online.

Acknowledgements

Special thanks go to José Guillermo Ortiz Tena for technical and scientific support in analytical measurements.

Funding

Evonik Industries; DEHEMA (in part).

Conflict of interest statement. None declared.

References


