Incorporation of alternative amino acids into cyanophycin by different cyanophycin synthetases heterologously expressed in *Corynebacterium glutamicum*

Ramona Wördemann1, Lars Wiefel1, Volker F. Wendisch2 and Alexander Steinbüchel1,3*

Abstract

Cyanophycin (multi-l-arginyll-poly-l-aspartic acid; also known as cyanophycin grana peptide [CGP]) is a biopolymer that could be used in various fields, for example, as a potential precursor for the synthesis of polyaspartic acid or for the production of CGP-derived dipeptides. To extend the applications of this polymer, it is therefore of interest to synthesize CGP with different compositions. A recent re-evaluation of the CGP synthesis in *C. glutamicum* has shown that *C. glutamicum* is a potentially interesting microorganism for CGP synthesis with a high content of alternative amino acids. This study shows that the amount of alternative amino acids can be increased by using mutants of *C. glutamicum* with altered amino acid biosynthesis. With the DM1729 mutant, the lysine content in the polymer could be increased up to 33.5 mol%. Furthermore, an ornithine content of up to 12.6 mol% was achieved with ORN21(*Pgdh*4). How much water-soluble or insoluble CGP is synthesized is strongly related to the used cyanophycin synthetase. CphADh synthesizes soluble CGP exclusively. However, soluble CGP could also be isolated from cells expressing CphA6308Δ1 or CphA6308Δ1_C59SS in addition to insoluble CGP in all examined strains. The point mutation in CphA6308Δ1_C59SS partially resulted in a higher lysine content. In addition, the CGP content could be increased to 36% of the cell dry weight under optimizing growth conditions in *C. glutamicum* ATCC13032. All known alternative major amino acids for CGP synthesis (lysine, ornithine, citrulline, and glutamic acid) could be incorporated into CGP in *C. glutamicum*.

Keywords: Alternative amino acids, *Corynebacterium glutamicum*, Cyanophycin, Lysine, Ornithine

Introduction

The polyamide cyanophycin is typically composed of poly(aspartic acid) as a backbone to which arginine residues are attached; it is also referred to as multi-l-arginyll-poly-l-aspartic acid or as cyanophycin grana peptide (CGP). CGP is non-ribosomal polypeptide synthesised by cyanophycin synthetases (CphA, EC 6.3.2.29/EC 6.3.2.30). During heterologous production of CGP, other alternative amino acids can also be incorporated into the polymer in addition to aspartic acid and arginine. Instead of arginine, larger amounts of lysine, citrulline or ornithine and several others were also incorporated (Berg et al. 2000; Steinle et al. 2009; Wiefel et al. 2011). Glutamic acid can also be incorporated into the backbone as an alternative to aspartic acid (Wiefel et al. 2019a and references cited therein). In addition, biochemically incorporated constituents can be modified enzymatically or chemically (Frommeyer et al. 2014; Wiefel and Steinbüchel 2016, Wiefel 2019b). These variations of the polymer can change the solubility behavior of the polymer (Wiefel 2019b).
and Steinbüchel 2014). Variations in CGP are especially interesting to extend the possible field of applications of CGP. For example, CGP-derived dipeptides can be used as food supplements or animal feed additives or may have medical and cosmetic applications (Sallam and Steinbüchel 2010). The dipeptides are produced by enzymatic digestion of CGP employing cyanophycinase (intracellular CphB or extracellular CphE; EC 3.4.15.6), usually producing Asp-Arg dipeptides and related oligopeptides (Sallam et al. 2009).

Corynebacterium glutamicum in particular could be an interesting candidate for the production of CGP with a high content of alternative amino acids, as this bacterium is traditionally used for the production of amino acids on an industrial scale (Wendisch 2020). C. glutamicum is best known for the production of glutamate and lysine, both of which can be used for the synthesis of CGP. However, the other amino acids, which can be used for the synthesis of CGP, can also be produced in larger quantities by C. glutamicum. On the other hand, the thick and rigid cell wall of C. glutamicum is disadvantageous for isolating CGP from the cells in comparison to other microorganisms (Wiefel et al. 2019a).

In the early 2000s, CGP synthesis in C. glutamicum was studied for the first time. At that time only a CGP yield of less than 3% of the dry cell mass was obtained from C. glutamicum cells (Aboulmagd et al. 2001b). At that time the existence of water-soluble CGP, which usually contains a high proportion of lysine, was not known, yet. For this reason, research on this topic was recently resumed. During this re-evaluation by Wiefel et al. (2019a), different vectors, CphAs and media were examined. It was shown that the strains can accumulate up to 17% of their cells dry mass CGP. It was also shown for the first time that glutamic acid can replace aspartic acid in the backbone of CGP (Wiefel et al. 2019a). The aim of this study was to improve the CGP synthesis in C. glutamicum with regard to the incorporation of alternative amino acids into the polymer. For this, optimized cultivation conditions and modified strains of C. glutamicum were used to try to increase the proportion of alternative amino acids and to further increase the CGP yield. Two variants of CphA from Synechocystis sp. PCC6308 and CphA from D. hafniense DSM 10664 were used for the investigations. Besides the wild type, mutants of C. glutamicum were used, which produce increased amounts of lysine, ornithine or cadaverine.

### Materials and methods

#### Bacterial strains, media, and growth conditions

All bacterial stains and plasmids used in this study are listed in Table 1. E. coli strains were used for plasmid maintenance and propagation and were cultivated at 30 °C and a stirring rate of 130 rpm in lysogeny broth (LB) medium (Sambrook et al. 1989). Usually, 10 ml medium in 100 ml Erlenmeyer flask were used. For C. glutamicum strains BHIS (Tauch et al. 2002), CASO + G (Persicke et al. 2011) and CGXII (Keilhauer et al. 1993) medium were used. Strains harbouring plasmids were grown in presence of 25 μg kanamycin per ml. For the arginine auxotrophic ORN2(P _gdi Δ) strain (Jensen et al. 2015), 15 mM arginine was added to the CGXII medium. To monitor cell growth, the optical density was measured at 600 nm (OD _600 ). For cyanophycin synthesis in C. glutamicum two precultures with 10 and 50 ml medium in 100 and 250 ml Erlenmeyer flasks, respectively, were

### Table 1 Strains and plasmids used in this study

| Strain or plasmid | Relevant characteristics | Reference or source |
|-------------------|--------------------------|---------------------|
| **Strains**       |                          |                     |
| E. coli TOP10     | F- mcrA Δ(mrr-hsd RMS-mcrBC) Q80lacZΔM15 ΔlacX74 recA1 araD139 Δ(larleu)7697 galU galA K12 (Str^-) endA1 supG | Invitrogen, Carlsbad, USA |
| C. glutamicum ATCC 13,032 | Wild type, Biotin-auxotroph | DSMZ, Braunschweig, Germany |
| C. glutamicum DM1729 | ATCC 13032 pyc K258, hom V89A, lysC T111 | Georgi et al. (2005) |
| C. glutamicum ORN2(P _gdi Δ) | ATCC 13032 with in-frame deletion of prophages CGP1 (cg1507-cg1524), CGP2 (cg1746-cg1752), and CGP3 (cg1890-cg2071), ΔargFRG, F _gdi Δ - 35: TTGCCA - 10: TATAAT | Jensen et al. (2015) |
| **Plasmids**      |                          |                     |
| pVWEx1            | F  _recA, lacI 8, Km^R    | Peters-Wendisch et al. (2001) |
| pVWEx1: cphA3308 Δ1 | pVWEx1 with cphA3308 Δ1 (Pst/Vxbal) | Wiefel et al. (2019a) |
| pVWEx1: cphA3308 Δ1 _CS595S | pVWEx1 with cphA3308 Δ1 _CS595S (Pst/Vxbal) | This study |
| pVWEx1: cphA3001 | pVWEx1 with cphA3001 (SauIII/Vxbal) | Wiefel et al. (2019a) |
| pEC-XT99A: ldcC    | pVWEx1 with cphA3001, ldcC from E. coli K12, Tet^R | Sgobba et al. (2018) |

Km^R kanamycin resistance, Tet^R tetracycline resistance.
used to inoculate the main culture containing 100 ml in a 500 ml Erlenmeyer flask with two baffles. The first preculture with CASO + G medium was inoculated with a cryoculture and was cultivated for about 8 h. Before inoculating the second preculture, the cells were washed with washing buffer (50 mM TRIS, 50 mM NaCl, pH 6.3). For the second preculture, CGXII medium was used as the main culture. The main culture was inoculated by a 16-h second preculture to an initial OD$_{600}$ of 0.2 and induced after 4 h with 1 mM IPTG. In contrast to the previous studies of Wiefel et al. (2019a) the strains were not cultivated under biotin limiting conditions.

**Cultivation of *C. glutamicum* strain DM1729**

pVWEx1::cphA$_{6308}$Δ1_C595S in a stirred tank reactor

A Biostat UD30 stainless steel reactor (B. Braun Biotech International, Melsungen, Germany) was used for batch cultivation of *C. glutamicum* strain DM1729 harboring pVWEx1::cphA$_{6308}$Δ1_C595S in 20-liter scale. Two fermentations were performed. In the first fermentation normal CGXII medium according to Keilhauer 1993 was used, only without addition of MOPS buffer. For the second fermentation, 5 g/l CASO broth (Carl Roth, Karlsruhe, Germany) was added to the medium instead of urea. The fermentations were inoculated with an OD$_{600}$ between 0.7 and 0.8. An IPTG concentration of 0.1 mM was used for induction. Cultivation was carried out at 30 °C and a pO$_2$ saturation of 30% in the medium, which was controlled by stirring rates between 100 and 800 rpm and aeration rates between 0.5 and 2.0 vvm (volume per volume × minute). The pH was kept constant at a value of 7 with 4 M NaOH or 4 M HCl. As antifoam agent Struktol® J 673 A (Schill + Seilacher, Hamburg, Germany) was used. Samples with a volume of 100 ml were taken to determine OD$_{600}$, cells dry masses and CGP content of the cells.

**Molecular biology techniques**

Plasmid DNA from *E. coli* strains was isolated using the GenJET plasmid miniprep kit (Thermo Scientific, Waltham, USA). The plasmids contain the genes for the cyanophycin synthetases CphA$_{6308}$Δ1, CphA$_{6308}$Δ1_C595S or CphADh (see Table 2). Upstream of the genes a ribosome binding site (RBS) is located which was optimized for expression in *C. glutamicum* (GAA AGGAGGCCCTTCAG, Siebert and Wendisch 2015). For isolation of plasmids from *C. glutamicum* strains, 15 mg lysozyme were added per ml to the resuspension buffer, and the resuspended cells were incubated for 2 h at 37 °C before following the protocol of the manufacturer (Thermo Scientific, Waltham, USA).

Ampicillin was added to the medium during the production of electrocompetent *C. glutamicum* cells (Jensen et al. 2015). Transformation of *C. glutamicum* was performed by electroporation and a heat-shock at 46 °C for 6 min (van der Rest et al. 1999). Restriction analyses were carried out for control purposes. The restriction enzymes were purchased from Thermo Scientific.

**Isolation of cyanophycin from *C. glutamicum***

For isolation of CGP from *C. glutamicum*, cells were harvested by centrifugation (20 min, 5000×g, 4 °C), washed with 0.9% (wt/vol) NaCl, and freeze-dried. After determining the cells dry mass, the cell disintegration of the cells was carried out using an ultrasonic disintegrator. For this purpose, the freeze-dried cells were crushed and resuspended in H$_2$O (20 ml/g). The suspension was then sonicated for 1 min/ml with the UP200S ultrasonic processor and shaken overnight at 4 °C. Insoluble material was sedimented by centrifugation (30 min, 5000×g), and the water-soluble CGP, which was contained in the supernatant, was precipitated using 1.5 vol ice-cold ethanol. After centrifugation, the CGP pellet was washed with acetone and dried at 70 °C. The CGP was dissolved in H$_2$O and precipitated again. Water-insoluble CGP was isolated from cell debris by resuspension in 0.1 M HCl. The now dissolved CGP was separated from insoluble material by centrifugation (30 min, 5000×g), and it was subsequently precipitated from the supernatant by neutralization (pH 7) using 0.1 to 4 M NaOH. From the supernatant of the insoluble CGP again soluble CGP was isolated. The washing steps were repeated three times for insoluble CGP to further purify the polymer. The precipitated insoluble CGP was then again sedimented by centrifugation and washed twice with H$_2$O. Persistent protein impurities were removed after the first

**Table 2** Details about the used cyanophycin synthetases

| CphA       | Length | Origin | CGP solubility      | Reference          |
|------------|--------|--------|---------------------|--------------------|
| CphA$_{6308}$Δ1 | 873 AS | CphA from Synechocystis sp. strain PCC 6308 shortened by one AS at the C-terminus | Insoluble and soluble | Steinle et al. (2010) |
| CphA$_{6308}$Δ1_C595S | 873 AS | CphA from Synechocystis sp. strain PCC 6308 with point mutation and shortened by one AS at the C-terminus (see Fig S1) | Insoluble and soluble | Steinle et al. (2010) |
| CphADh     | 885 AS | CphA from Desulfotobacterium hafniense DSM 10664 | Only soluble | Ziegler et al. (2002) |
precipitation by adding 100 μg proteinase K per ml and by incubating the CGP for 3 h at 60 °C.

Analysis of isolated cyanophycin

The purity and the molecular masses of isolated CGP were determined by sodium dodecylsulphate polyacrylamide gel electrophoresis (SDS-PAGE) in 11.5% (wt/vol) polyacrylamide gels, according to Laemmli (1970). The apparent molecular weight was estimated using the PageRuler Prestained Protein Ladder (Thermo Scientific).

The amino acid compositions of CGP were determined by high-performance liquid chromatography (HPLC) using a Waters B801 column (300 × 4 mm) after the polymer was converted into derivatives of the constituents as described by Aboulmagd (2001b) and Steinle (2009). Pre-column ortho-phthalaldehyde (OPA) derivatization was performed using a Smartline autosampler 3900 according to the manual (Knauer GmbH, Berlin, Germany). A reference kit (Kollektion AS-10 from Serva Feinbiochemica, Heidelberg, Germany) was used for calibration. Ornithine and citrulline were purchased as monohydrochloride from Fluka or Merck, respectively. CGP was hydrolyzed in 6 M HCl (100 μl/mg) at 95 °C overnight, neutralized, and lyophilized before measurement.

Results

Cyanophycin synthesis in C. glutamicum ATCC 13032

The CGP synthesis in the wild type of C. glutamicum was studied with three different CphAs (CphA6308Δ1, CphA6308Δ1_C595S and CphADh). For CGP synthesis the strains were cultivated for 72 h in CGXII medium. Optimizations in cultivation conditions improved cell growth when compared to the previous study of Wiefel et al. (2019a). As a result, the cell density could be increased up to 16.6 g dry mass (DW)/l (see Table 3 and Additional file 1: Fig. S2). All three strains grew similarly well, but in terms of CGP content there were marked differences between the cells harboring different genes for CphAs. With CphADh a CGP content over 36% of the cells dry mass was achieved, which corresponds to a CGP concentration of about 6 g/l. In contrast, cells expressing CphA6308Δ1 and CphA6308Δ1_C595S synthesized

Table 3  Comparison of different C. glutamicum strains for CGP synthesis

| Strains                          | DW [g/l] | Cyanophycin content [%DW] | Cyanophycin solubility | Amino acid composition [mol%] * |
|----------------------------------|---------|---------------------------|------------------------|--------------------------------|
| ATCC 13032                        |         |                           |                        |                                |
| pVWEx1::cphA6308Δ1               | 15.9 ± 0.1 | Insoluble                 | 1.9 ± 0.5              | Asp 50.1, Arg 38.9, Lys 8.7, Glu 0.5, Orn 1.0, Cit 0.8 |
| pVWEx1::cphA6308Δ1_C595S         | 15.6 ± 0.2 | Insoluble                 | 1.4 ± 0.3              | Asp 54.7, Arg 30.0, Lys 13.6, Glu 0.3, Orn 0.7, Cit 0.6 |
| pVWEx1::cphADh                    | 16.6 ± 0.4 | Insoluble                 | 0.0 ± 0.0              | Asp 49.3, Arg 40.1, Lys 6.9, Glu 2.3, Orn 0.6, Cit 0.8 |
| DM1279                           |         |                           |                        |                                |
| pVWEx1::cphA6308Δ1               | 11.6 ± 0.3 | Insoluble                 | 12.2 ± 2.1             | Asp 52.9, Arg 30.3, Lys 16.2, Glu 0.2, Orn 0.0, Cit 0.4 |
| pVWEx1::cphA6308Δ1_C595S         | 11.7 ± 0.2 | Insoluble                 | 13.1 ± 2.3             | Asp 48.0, Arg 29.8, Lys 21.6, Glu 0.2, Orn 0.0, Cit 0.4 |
| pVWEx1::cphADh                    | 14.4 ± 0.6 | Insoluble                 | 0.0 ± 0.0              | Asp 42.7, Arg 19.8, Lys 33.5, Glu 1.0, Orn 2.6, Cit 0.4 |
| ORN2(P_gdh)                      |         |                           |                        |                                |
| pVWEx1::cphA6308Δ1               | 11.6 ± 0.2 | Insoluble                 | 1.9 ± 0.2              | Asp 47.6, Arg 42.7, Lys 0.0, Glu 0.3, Orn 9.1, Cit 0.3 |
| pVWEx1::cphA6308Δ1_C595S         | 11.7 ± 0.1 | Insoluble                 | 2.4 ± 0.2              | Asp 42.8, Arg 47.9, Lys 0.0, Glu 0.3, Orn 8.7, Cit 0.3 |
| pVWEx1::cphADh                    | 12.9 ± 0.3 | Insoluble                 | 0.0 ± 0.0              | Asp 41.7, Arg 44.2, Lys 0.0, Glu 0.7, Orn 12.6, Cit 0.8 |

The strains were cultivated for 72 h in 100 ml CGXII medium in a 500 ml Erlenmeyer flask with two baffles at 130 rpm and 30 °C. The cultures were inoculated with an OD600 of about 0.2 and after 4 h IPTG was added. For the ORN2(P_gdh) strains an additional 15 mM arginine was added to the medium. The cells dry mass was determined, and the contained cyanophycin was isolated and analysed. The values indicate the mean and standard deviation of at least three independent cultivations.

* The values have been rounded to one decimal place.
significantly less CGP, but the CGP accumulated contained a higher amount of alternative amino acids. The point mutation CphA\textsubscript{6308\Delta1_C595S} led to a 4.9 mol\% higher lysine content in the insoluble CGP. The lysine content in the CGP of CphA\textsubscript{6308\Delta1_C595S} was 13.6 mol\% and therefore almost twice as high as that of CphA\textsubscript{Dh}.

The non-biotin limiting growth conditions in this study had an impact on the composition of the synthesized CGP. The alternative amino acids ornithine and citrulline were found for the first time in the CGP accumulated in \textit{C. glutamicum}. These amino acids were mainly found in the soluble CGPs accumulated in cells expressing CphA\textsubscript{6308\Delta1} or CphA\textsubscript{6308\Delta1_C595S}, respectively. This CGP thus contains all the alternative amino acids known to date that are used for in vivo CGP synthesis (see Fig. 1).

\textbf{Cyanophycin synthesis in a \textit{C. glutamicum} mutant for lysine production}

To increase the lysine content in CGP, CGP synthesis was investigated in \textit{C. glutamicum} strain DM1729 (Georgi et al. 2005), which was developed for the production of lysine. Although the strains did not reach as high cell densities as the cells of \textit{C. glutamicum} ATCC 13032, up to three times more lysine was incorporated into the accumulated CGP (see Table 3 and Additional file 1: Fig. S2). It is remarkable that the strain grew better with the CphA\textsubscript{Dh} and reached a higher CGP content with 24.2\% of cells dry mass. The lysine content in the CGP was 15.4 mol\% and was therefore significantly higher than in the CGP from the cells of \textit{C. glutamicum} ATCC 13032 pVWEx1::cphA\textsubscript{Dh}. With the other two CphAs, lysine contents of up to 33.5 mol\% were measured in the soluble CGP, and even in the insoluble CGP higher lysine contents were found. The higher lysine concentrations in the cells of these strains resulted in a strong increase in insoluble CGP with up to 13.1\% of cells dry mass at a lysine content of up to 21.6 mol\%. It was previously thought that the higher lysine concentration would have a greater effect on soluble CGP, as high lysine is often a characteristic of soluble CGP (Frommeyer and Steinbüchel 2013). The insoluble CGP synthesised with CphA\textsubscript{6308\Delta1_C595S} showed a 5.4 mol\% higher lysine content in comparison to the CGP accumulated by the single mutant cyanophycin synthetase. The content of ornithine and citrulline was lower in all these strains than in the \textit{C. glutamicum} ATCC 13032 strains.

\textbf{Cyanophycin synthesis in a \textit{C. glutamicum} mutant for ornithine production}

\textit{Corynebacterium glutamicum} ORN2(P\textsubscript{gdh4}) is a strain developed for the increased production of ornithine (Jensen et al. 2015). In this strain, genes of the arginine biosynthesis pathway were deleted to prevent the further metabolism of ornithine. As a result, the strain is auxotrophic for arginine. In a preliminary experiment it was shown that a concentration of 15 mM arginine in the CGXII medium is a suitable concentration for CGP synthesis, as this concentration allows good growth.

The CGP content of these strains is nevertheless lower than that of the other \textit{C. glutamicum} strains investigated. With these strains ornithine contents of 10 mol\% on average were achieved (see Table 3 and Additional file 1: Fig. S2). These results represent a slight improvement when compared to the previous results with \textit{Saccharomyces cerevisiae}, where ornithine contents of only 8 mol\% were achieved (Steinle et al. 2009). There were only slight differences in the amino acid composition between the different isolates. The SDS-PAGE analysis showed that there are large differences in the molecular masses of the accumulated soluble and insoluble CGPs (see Fig. 2). Most of the soluble CGP exhibited molecular masses below 15 kDa (lane 1,

![HPLC chromatogram of hydrolysed soluble cyanophycin from C. glutamicum ATCC 13032 pVWEx1::cphA6308\Delta1_C595S with all known alternative amino acids](image_url)
4, 7 and 8). In the case of CphADh it is clearly visible that most CGP has a molecular mass of only 10 kDa. The soluble CGP isolated from the supernatant of the insoluble CGP is slightly larger with 15 to 25 kDa (lane 2 and 5). The insoluble CGP, on the other hand, exhibited a molecular weight of 35 to 55 kDa (lane 3 and 6).

Cyanophycin synthesis in a C. glutamicum mutant for cadaverine production

In a previous in vitro study, it was shown that agmatine (biogenic amine of arginine) has a negative effect on the incorporation of arginine into the polymer. Therefore, it was suspected that it could be incorporated into CGP instead of arginine (Aboulmagd 2001a). Cadaverine is also a biogenic amine and is produced by decarboxylation of lysine. To enable the synthesis of cadaverine in C. glutamicum, the lysine decarboxylase LdcC was additionally introduced into the DM1729 strains using the vector pEC-XT99A (Sgobba et al. 2018). This is the first study on the effect of cadaverine on CGP synthesis. The conversion of lysine to cadaverine results in a significant decrease in the lysine content of CGP (see Table 4 and Additional file 1: Fig. S2). The CGP content in the cells also decreases in comparison to the DM1729 strains not harboring pEC-XT99A::ldcC.

Noticeable was the slower growth of the strain expressing CphADh. Cadaverine could not be detected in any of the extracted CGP. Possibly cadaverine has an inhibiting effect on CGP synthesis.

Cultivation in stirred tank reactors and scale up

In the first fermentation of C. glutamicum DM1729 pVWEx1::cphA6308 Δ1_C595S in CGXII medium CGP biosynthesis was induced with 0.1 mM IPTG after a cultivation period of 4 h. The fermentation took 53 h in total. A maximum cells dry mass of 11 g/l was obtained. Only insoluble CGP could be isolated. This reached an average of 2.5% of the cells dry mass. The lysine content was 16.7 mol%.

In the second fermentation with addition of 5 g/l CASO broth, 0.1 mM IPTG was used directly at the start of the fermentation to induce cells for production of CGP. Based on the OD$_{600}$ it could be seen that the cells grew significantly faster and they reached a higher cells dry mass of 15 g/l than in the flask. Although soluble and insoluble CGP could be isolated, the total CGP content was again only 2.5% of the cells dry mass. The amino acid composition was comparable to that of the CGP formed in the flask experiment.
**Discussion**

Based on the study of Wiefel et al. (2019a), the cultivation conditions were further optimized to improve cell growth and to increase CGP yield. The omission of biotin limitation had the greatest impact, even if this means that less glutamic acid is incorporated into CGP. The growth of the wild type strains could thus be increased to a cells dry mass of 16 g/l, and a CGP content of 36.2% of the cells dry mass was achieved with CphADh. In comparison, the cultivation of C. glutamicum ATCC 13032 pVWEx1::cphA with biotin limitation in Wiefel et al. (2019a) achieved only a CGP content of 17.1% of cells dry mass. For the CphA variants of Synechocystis sp. PCC6308 it was possible for the first time to isolate water soluble CGP from cells cultivated in CGXII medium. The CGP contents of cells harboring these CphAs were significantly lower than that of cells harboring CphA_Dh. However, due to the soluble CGP, a slight increase compared to previous studies (Aboulmagd et al. 2001b; Wiefel et al. 2019a) was observed. Due to the changes in cultivation conditions, ornithine and citrulline could be detected in CGP from C. glutamicum for the first time. However, the amounts of citrulline were very low. With S. cerevisiae, for example, citrulline contents of 20 mol% have been already achieved (Steinle et al. 2009).

Experiments with the mutants of C. glutamicum showed that the proportion of alternative amino acids in CGP can be increased in these strains. The lysine contents were more than doubled by DM1729 in many samples and much more insoluble CGP was synthesized. This could possibly be related to the intracellular pH increasing due to the higher lysine concentration and thus being closer to the optimal pH of 8.2 of CphA6308 (Aboulmagd et al. 2001a). It is noticeable that due to the point mutation C595S in the insoluble CGP the lysine content is 5.4 mol% higher. Also in P. pastoris and E. coli the lysine contents were slightly higher if the point mutation C595S of CphA was used (Steinle et al. 2010). However, the increase was not as high as in this case. This effect has already been observed with C. glutamicum ATCC 13032. The point mutation seems to have a positive influence on the incorporation of lysine.

An increase of the alternative amino acid was also observed in the ornithine-forming strains. Here, a maximum ornithine content of 12.6 mol% was reached. The proportion is thus higher than in the study of Steinle et al. 2009, where with S. cerevisiae an ornithine proportion of 8 mol% was achieved. The differences in the composition of soluble and insoluble CGP were very small in these strains, which suggests that the size of the polymer also has an influence on the solubility behavior, as this was the main difference. This assumption was already made in Wiefel et al. (2019a).

The decarboxylation of lysine to cadaverine has resulted in a lower cellular cyanophycin content and thus in less CGP synthesised and in a drastic reduction in the lysine content of the polymer. However, no cadaverine could be detected in cyanophycin. The strains for cadaverine production have therefore proved unsuitable for cyanophycin synthesis.

Although the growth of the mutants is lower than that of C. glutamicum ATCC 13032, mutants that produce more lysine in particular have shown great potential for CGP synthesis. It would therefore be useful to investigate this further and to optimize the scale up, because until now growth has been improved on a larger scale, but the CGP content is lower than in flasks.

### Table 4 Comparison of CGP synthesis in C. glutamicum strains for cadaverine production

| Strains                | DW [g/l] | Cyanophycin | Amino acid composition [%DW] | Asp | Arg | Lys | Glu | Orn | Cit | Cad |
|------------------------|----------|-------------|-----------------------------|-----|-----|-----|-----|-----|-----|-----|
|                        |          | Solubility  | CGP content [%DW]           |     |     |     |     |     |     |     |
| DM1279 pEC-XT99A::ldcC |          | Insoluble   | 3.5 ± 0.9                   | 46.6| 46.2| 6.2 | 0.4 | 0.1 | 0.5 | 0.0 |
| DM1279 pEC-XT99A::ldcC |          | Soluble     | 3.5 ± 1.4                   | 44.1| 38.0| 13.6| 1.6 | 2.2 | 0.5 | 0.0 |
| pVWEx1::cphA6308Δ1     | 13.3 ± 0.1| Insoluble   | 3.2 ± 1.2                   | 47.4| 45.8| 5.9 | 0.4 | 0.1 | 0.4 | 0.0 |
| pVWEx1::cphA6308Δ1_C595S| 12.7 ± 0.3| Insoluble   | 1.5 ± 1.2                   | 44.2| 37.1| 16.0| 1.9 | 0.0 | 0.8 | 0.0 |
| pVWEx1::cphA6308Δ1_C595S| 9.2 ± 0.1 | Insoluble   | 0.0 ± 0.0                   | –   | –   | –   | –   | –   | –   | –   |
| pVWEx1::cphA6308Δ1_C595S| 20.4 ± 0.7| Soluble     | 20.4 ± 0.7                  | 44.1| 41.9| 11.0| 2.2 | 0.5 | 0.3 | 0.0 |

The strains were cultivated for 72 h in 100 ml CGXII medium in 500 ml Erlenmeyer flasks with two baffles at 130 rpm and 30 °C. The cultures were inoculated with an OD₆₀₀ of about 0.2 and after 4 h IPTG was added. The cells dry mass was determined and the contained cyanophycin was isolated and analyzed. The values indicate the mean and standard deviation of at least three independent cultivations.

* The values have been rounded to one decimal place.
Abbreviations

Ang: Arginine; AS: Amino acids; Asp: Aspartic acid; Cad: Cadaverine; CGP: Cyanophycin grana peptide; Cit: Citrulline; CphA: Cyanophycin synthetases; CphB: Intracellular cyanophycinase; CphE: Extracellular cyanophycinase; DW: Dry weight; Glu: Glutamic acid; LB: Lysogeny broth; Lys: Lysine; OD$_{600}$: Optical density at 600 nm; OPA: ortho-Phthalaldehyd; Orn: Ornithine; RBS: Ribosome binding site; WT: Wild type.

Supplementary Information

The online version contains supplementary material available at https://doi.org/10.1016/j.ymben.2020.05.001

Additional file 1. Fig. S1: Sequence alignment of cpha$_{AS}$ and cpha$_{AS}$d1, C. glutamicum. Fig. S2: Graphical representation of the results obtained with the different C. glutamicum strains for CPG synthesis.

Authors’ contributions

RW carried out this study and drafted the manuscript. LW and AS supervised the study and revised the manuscript. VFW provided strains and helped with molecular genetic methods. All authors read and approved the final manuscript.

Funding

No funding was received.

Availability of data and materials

All datasets on which the conclusions of the manuscript are presented in the main paper.

Declarations

Ethics approval and consent to participate

Not applicable.

Consent for publication

Not applicable.

Competing interests

On behalf of all authors, the corresponding author states that there is no conflict of interest.

Author details

1 Institut für Molekulare Mikrobiologie und Biotechnologie (IMMB), Westfälische Wilhelms-Universität Münster, Corrensstraße 3, 48149 Münster, Germany. 2 Faculty of Biology & CeBiTec, Bielefeld University, Universitätsstraße 25, 33615 Bielefeld, Germany. 3 Environmental Science Department, King Abdulaziz University, Jeddah, Saudi Arabia.

Received: 31 January 2021 Accepted: 7 April 2021 Published online: 15 April 2021

References

Aboulmagd E, Oppermann-Sanio FB, Steinbüchel A (2001a) Purification of Synecocystis sp. strain PCC6308 cyanophycin synthetase and its characterization with respect to substrate and primer specificity. Appl Environ Microbiol 67:2176–2182. https://doi.org/10.1128/AEM.67.5.2176-2182.2001

Aboulmagd E, Voss I, Oppermann-Sanio FB, Steinbüchel A (2001b) Heterologous expression of cyanophycin synthetase and cyanophycin synthetase in the industrial relevant bacteria Corynebacterium glutamicum and Ralstonia eutropha and in Pseudomonas putida. Biomacromol 2:1338–1342. https://doi.org/10.1021/bm010075a

Berg H, Ziegler K, Piotukh K, Baier K, Lockau W, Vollmer-Engert R (2000) Biosynthesis of the cyanobacterial reserve polymer multi-L-arginyl-poly-L-aspartic acid (cyanophycin). Eur J Biochem 267:5561–5570. https://doi.org/10.1046/j.1432-1327.2000.01622.x

Frommeyer M, Steinbüchel A (2013) Increased lysine content is the main characteristic of the soluble form of the polypeptide cyanophycin synthesized by recombinant E. coli. Appl Environ Microbiol 79:4474–4483. https://doi.org/10.1128/AEM.00986-13

Frommeyer M, Bergander K, Steinbüchel A (2014) Guanidination of soluble lysine-rich cyanophycin yields a homoarginine-containing polypeptide. Appl Environ Microbiol 80:2381–2389. https://doi.org/10.1128/AEM.04013-13

Georg T, Rittmann D, Wendisch VF (2005) Lysine and glutamate production by Corynebacterium glutamicum on glucose, fructose and sucrose. Roles of malic enzyme and fructose-1,6-bisphosphatase. Metab Eng 7:291–301. https://doi.org/10.1016/j.meb.2005.05.001

Jensen JX, Eberhardt D, Wendisch VF (2015) Modular pathway engineering of Corynebacterium glutamicum for production of the glutamate-derived compounds ornithine, proline, putrescine, citrulline, and arginine. J Biotechnol 214:85–94. https://doi.org/10.1016/j.jbiotec.2015.09.017

Keilhauer C, Eggeling L, Sahm H (1993) Isocitruline Synthesis in Corynebacterium glutamicum: molecular analysis of the ilvB-ilvN-ilvC operon. J Bacteriol 175:5595–5603. https://doi.org/10.1128/jb.175.17.5595-5603.1993

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

Persicke M, Plassmeier J, Neuweger H, Rückert C, Pühler A, Kalinowski J (2011) Size exclusion chromatography. An improved method to harvest Corynebacterium glutamicum cells for the analysis of cytosolic metabolites. J Biotechnol 154:171–178. https://doi.org/10.1016/j.jbiotec.2010.08.016

Peters-Wendisch PG, Schiel B, Wendisch VF, Katsoulidis E, Möckel B, Sahm H, Eikmanns BJ (2001) Pyruvate carboxylase is a major bottleneck for glutamate and lysine production by Corynebacterium glutamicum. J Mol Microbiol Biotechnol 3:295–300

Sallam A, Kast A, Przybilla S, Mieswinkel T, Steinbüchel A (2009) Biotechnological process for production of beta-di-dipeptides from cyanophycin on a technical scale and its optimization. Appl Environ Microbiol 75:29–38. https://doi.org/10.1128/AEM.00644-08

Sallam A, Steinbüchel A (2010) Dipeptides in nutrition and therapy. Cyanophycin-derived dipeptides as natural alternatives and their biotechnological production. Appl Microbiol Biotechnol 87:815–828. https://doi.org/10.1007/s00253-010-1261-0

Sambrook J, Fritsch E, Maniatis T (1989) Molecular cloning: a laboratory manual, 2nd edn. Cold Spring Harbor Laboratory Press, New York.

Sgobba E, Blöbaum L, Wendisch VF (2018) Production of food and feed additives from non-food-competing feedstocks. Valorizing N-acetylmuramic acid production. Metab Eng 58:17–34. https://doi.org/10.1016/j.ymben.2015.08.003

Siebert D, Wendisch VF (2015) Metabolic pathway engineering for production of 1,2-propanediol and 1-propanol by Corynebacterium glutamicum. Biotechnol Biofuels 8:91. https://doi.org/10.1186/s13068-015-0269-0

Steinle A, Bergander K, Steinbüchel A (2009) Metabolic engineering of S. cerevisiae for production of novel cyanophysins with an extended range of constituent amino acids. Appl Environ Microbiol 75:3437–3446. https://doi.org/10.1128/AEM.00383-09

Steinle A, Witthoff S, Krause JP, Steinbüchel A (2010) Establishment of cyanophycin biosynthesis in Pichia pastoris and optimization by use of engineered cyanophycin synthetases. Appl Environ Microbiol 76:1062–1070. https://doi.org/10.1128/AEM.01659-09

Tauch A, Kirchner O, Löffler B, Göttker S, Pühler A, Kalinowski J (2002) Efficient electrotransformation of Corynebacterium glutamicum on glucose, fructose and sucrose. Roles of malic enzyme and fructose-1,6-bisphosphatase. Metab Eng 7:291–301. https://doi.org/10.1016/j.meb.2005.05.001

van der Rest ME, Lange C, Molenaar D (1999) A heat shock following electroporation induces highly efficient transformation of Corynebacterium glutamicum with xenogeneic plasmid DNA. Appl Microbiol Biotechnol 52:541–545. https://doi.org/10.1007/s002530051557

Wendisch VF (2020) Metabolic Engineering advances and prospects for amino acid production. Metab Eng 58:17–34. https://doi.org/10.1016/j.ymben.2019.03.008
Wiefel L, Bröker A, Steinbüchel A (2011) Synthesis of a citrulline-rich cyanophycin by use of Pseudomonas putida ATCC 4359. Appl Microbiol Biotechnol 90:1755–1762. https://doi.org/10.1007/s00253-011-3224-4
Wiefel L, Steinbüchel A (2014) Solubility behavior of cyanophycin depending on lysine content. Appl Environ Microbiol 80:1091–1096. https://doi.org/10.1128/AEM.03159-13
Wiefel L, Steinbüchel A (2016) Enzymatic modification of soluble cyanophycin using the type II peptidyl arginine deiminase from Oryctolagus cuniculus. Macromol Biosci 16:1064–1071. https://doi.org/10.1002/mabi.201500433
Wiefel L, Wöhlers K, Steinbüchel A (2019a) Re-evaluation of cyanophycin synthesis in Corynebacterium glutamicum and incorporation of glutamic acid and lysine into the polymer. Appl Microbiol Biotechnol 103:4033–4043. https://doi.org/10.1007/s00253-019-09780-5
Wiefel L, Bachmann F, Terwort J, Steinbüchel A (2019b) In vitro modification of bacterial cyanophycin and cyanophycin dipeptides using chemical agents towards novel variants of the biopolymer. Earth Syst Environ 3:637–650. https://doi.org/10.1007/s41748-019-00107-y
Ziegler K, Deutzmann R, Lockau W (2002) Cyanophycin synthetase-like enzymes of non-cyanobacterial eubacteria: characterization of the polymer produced by a recombinant synthetase of Desulfitobacterium hafniense. Z Naturforsch 57:522–529. https://doi.org/10.1515/znc-2002-5-621

Publisher’s Note
Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.