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1 Generation of synthetic shuttle vectors enabling modular genetic 2 engineering of cyanobacteria

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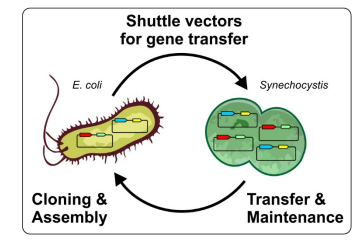
16 Abstract

17 Cyanobacteria have raised great interest in biotechnology due to their potential for a sustainable, 18 photosynthesis-driven production of fuels and value-added chemicals. This has led to a 19 concomitant development of molecular tools to engineer the metabolism of those organisms. In 20 this regard, however, even cyanobacterial model strains lag behind compared to their 21 heterotrophic counterparts. For instance, replicative shuttle vectors that allow gene transfer 22 independent of recombination into host DNA are still scarce. Here, we introduce the pSOMA 23 shuttle vector series comprising ten synthetic plasmids for comprehensive genetic engineering of 24 Synechocystis sp. PCC 6803. The series is based on the small endogenous plasmids pCA2.4 and 25 pCB2.4 each combined with a replicon from E. coli, different selection markers as well as features 26 facilitating molecular cloning and the insulated introduction of gene expression cassettes. We 27 made use of genes encoding green fluorescent protein (GFP) and a Baeyer-Villiger 28 monooxygenase (BVMO) to demonstrate functional gene expression from the pSOMA plasmids 29 in vivo. Moreover, we demonstrate the expression of distinct heterologous genes from individual 30 plasmids maintained in the same strain and thereby confirmed compatibility between the two 31 pSOMA sub-series as well as with derivatives of the broad-host-range plasmid RSF1010. We also 32 show that gene transfer into the filamentous model strain Anabaena sp. PCC 7120 is generally 33 possible, which is encouraging to further explore the range of cyanobacterial host species that 34 could be engineered via pSOMA plasmids. Altogether, the pSOMA shuttle vector series displays 35 an attractive alternative to existing plasmid series and thus meets the current demand for the 36 introduction of complex genetic setups and to perform extensive metabolic engineering of 37 cyanobacteria.

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Keywords: synthetic biology, shuttle vectors, cyanobacteria, (photo)-biotechnology, genetic
 engineering, molecular tools

42 Graphical Table of Content



44 Introduction

Cyanobacteria are a monophyletic but extraordinary diverse group of phototrophic bacteria.¹ They 45 46 are the only prokaryotes able to perform oxygenic photosynthesis, a process that makes use of 47 solar energy to oxidize water, which thereby produces dioxygen and protons. The obtained 48 electrons are used to drive an autotrophic metabolism, i.e. to fix atmospheric CO₂ into organic 49 carbon molecules and biomass. Accordingly, cyanobacteria have a huge environmental impact as 50 key players in global biogeochemical cycles.² Due to their photosynthetic lifestyle, cyanobacteria 51 have also drawn huge interest as promising biotechnological hosts, as they could be exploited for 52 a sustainable and CO₂-neutral production of fine chemicals or fuels.^{3–8}

53 However, the biotechnological utilization of cyanobacteria requires the concomitant development 54 of molecular tools for metabolic engineering. Several cyanobacteria have been used as microbial 55 chassis, whereby the most prominent example is the unicellular model strain Synechocystis sp. PCC 6803 (hereafter Synechocystis). Meanwhile, an increasing number of biological parts and 56 building blocks, i.e. BioBricks,⁹ and other standardized genetic elements have become available. 57 These include, for instance, a modular cloning system¹⁰ to assemble and introduce genes as well 58 59 as promoters, optimized ribosome binding sites, various transcription terminators or regulatory RNAs to control their expression.^{11–14} Nevertheless, the current molecular toolset and the 60 61 metabolic engineering capacity of Synechocystis as well as other cyanobacteria are still somewhat 62 limited.

63 Some cyanobacteria, including Synechocystis, are naturally competent, enabling the uptake of 64 exogenous DNA and to integrate it into their genome via homologous recombination.¹⁵ Although 65 neutral chromosomal sites for the integration of heterologous genes devoid of pleiotropic effects are available,¹⁶ the method is impaired by chromosome polyploidy,¹⁷ which requires time-66 67 consuming genetic segregation. Moreover, the number of neutral sites is inevitably limited, which 68 requires additional alternatives to introduce genes. In this regard, replicative plasmids could be 69 used. These extrachromosomal, circular DNA elements are autonomously maintained, either 70 based on the endogenous replication machinery or in a host-independent manner, i.e. using 71 factors encoded on the plasmids itself. However, plasmids that are commonly used for cloning in 72 Escherichia coli (hereafter E. coli) are usually not suitable for cyanobacteria. Their origins of 73 replication are not supported by the intrinsic replication machinery and hence, cannot be maintained as extrachromosomal element.¹⁶ In contrast, the broad-host-range shuttle vector 74 RSF1010^{18,19} belongs to the IncQ plasmids and encodes its own replication factors for host-75 76 independent maintenance in a variety of gram-negative bacteria,²⁰ including several 77 cyanobacterial strains.^{21,22} In 1990, the earliest utilization of a RSF1010 derivative in *Synechocystis* was reported.²³ Since then, replicative plasmids based on RSF1010 have been extensively used in cyanobacteria. Thereby, several optimization and size reduction steps resulted in different plasmid series, including e.g., pVZ,²⁴ pPMQAK1,²⁵ pSHDY,²⁶ and the pSEVAx5n series.^{27,28} Additionally, an alternative broad-host-range vector that is instead based on RK2²⁹ has recently been shown to replicate in *Synechocystis* as well.¹⁰ However, the limited availability of different, i.e. compatible replicative plasmids impedes comprehensive genetic engineering of cyanobacteria and thus, to fully exploit their biotechnological potential.

85 To overcome this problem, a number of chimeric shuttle vectors have been generated that exhibit two fused replicons to enable maintenance and cloning in *E. coli* as well as extrachromosomal 86 87 replication in certain cyanobacteria. This comprises for instance pDC1 from Nostoc sp. MAC PCC 8009,³⁰ pDU1 from *Nostoc* sp. PCC 7524,³¹ pFDA from *Fremyella diplosiphon*,³² and pANS (also 88 called pUH24) from Synechococcus elongatus PCC 7942.33 However, none of these have been 89 90 described to be maintained in Synechocystis cells. Synechocystis harbors seven extrachromosomal elements³⁴ subdivided into four large plasmids, ranging from about 44 to 120 91 kilobase pairs (kb),³⁵ and three small ones: pCA2.4 (2.4 kb),³⁶ pCB2.4 (2.4 kb),³⁷ and pCC5.2 92 93 (5.2 kb).³⁸ These small plasmids were predicted to replicate via rolling-circle amplification^{36–38} and have already been investigated concerning copy numbers^{39,40} as well as potential open reading 94 frames.⁴¹ Furthermore, initial attempts using synthetic derivatives of pCA2.4,⁴² pCB2.4,⁴² and 95 pCC5.2^{41,43} indicated stable maintenance and exhibited reporter gene expression in 96 97 Synechocystis. However, to implement complex genetic setups, a prerequisite for extensive 98 metabolic engineering, entire series of compatible plasmids would be advantageous.

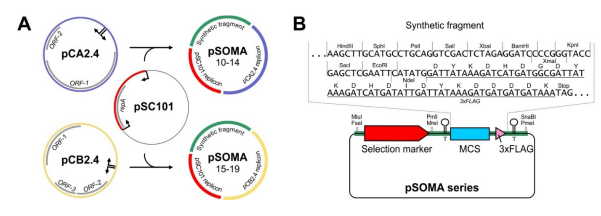
99 In this study, we introduce the pSOMA shuttle vector series to expand the available molecular 100 toolset to genetically engineer cyanobacteria. This library of ten individual plasmids is based on 101 chimeric fusions of the two smallest endogenous plasmids from Synechocystis, either pCA2.4 or 102 pCB2.4, as well as the pSC101 replicon originating from *E. coli.*⁴⁴ It further features multiple 103 selection markers and allows customized cloning in E. coli as well as the transfer of insulated gene 104 expression cassettes into Synechocystis. Moreover, we demonstrate compatibility of the two 105 subseries with each other and further shuttle vectors, i.e. RSF1010-based replicative plasmids in 106 Synechocystis. As demonstrated here, the pSOMA series enables the introduction and 107 maintenance of distinct heterologous genes by individual plasmids and hence, provides further 108 options for extensive and flexible metabolic engineering of Synechocystis.

110 Results

111 Generation of the pSOMA shuttle vector series

Among the seven endogenous plasmids of Synechocystis, especially the three small ones 112 pCA2.4,³⁶ pCB2.4,³⁷ and pCC5.2³⁸ display attractive targets for genetic modifications due to their 113 higher copy numbers relative to the chromosome or larger plasmids like pSYSM.^{39,40} Moreover, 114 115 their small size of 2.4-5.2 kb in principle allows molecular cloning and thus, synthetic combination 116 with other elements in vitro. Therefore, we made use of pCA2.4 and pCB2.4 to generate a 117 synthetic shuttle vector series that can be utilized in both E. coli as well as Synechocystis. Each 118 replicon, i.e. the sequence harboring a vegetative origin of replication as well as two (pCA2.4) or three (pCB2.4) open reading frames (ORFs), whose products (potentially) enable replication, was 119 fused to the pSC101 backbone⁴⁵ that is feasible for the maintenance in *E. coli*. The overall design 120 121 of the pSOMA shuttle vector series is shown in **Figure 1**.

122



124 Figure 1. Generation and structure of the pSOMA shuttle vector series. (A) Different replicons were amplified by 125 PCR and fused to each other in the illustrated order (arrows indicate primer binding sites). The replicon of pSC101 for 126 maintenance in E. coli was fused to the entire sequences from either pCA2.4 or pCB2.4 to enable replication in 127 Synechocystis. Moreover, a synthetic fragment harboring one of five antibiotic resistance genes (selection marker) and 128 further features was added. (B) For easy-to-use cloning, all plasmids feature a multiple cloning site (MCS, sequence 129 and restriction sites are highlighted). Genes can be cloned in frame with a sequence encoding a 3xFLAG-tag, which in 130 turn allows detection of recombinant fusion proteins via a specific antibody (sequence underlined). The MCS is flanked 131 by transcriptional terminators (T) to avoid read-through from inserted units.

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In a previous report by Liu and Pakrasi (2018),⁴² the usability of pCA2.4 and pCB2.4 as basis for synthetic plasmids has already been demonstrated by combining them individually with pUC118⁴⁶ or pSC101 replicons. The two generated shuttle vectors pCA-UC118 and pCB-SC101 harbor a chloramphenicol and streptomycin resistance cartridge, respectively. In addition to that, the pSOMA library is especially aiming for modularity and compatibility and hence, consists of two subseries, each exhibiting five different selection markers. The pSOMA vectors were further 139 equipped with a multiple cloning site (hereafter MCS) accommodating distinct restriction 140 endonuclease sites as well as a sequence encoding a 3xFLAG-tag, which potentially allows fusion 141 with coding sequences and finally the synthesis of easily detectable, i.e. tagged proteins (Figure 142 **1B**). To avoid undesired read through from the gene of interest that is potentially inserted later on, the MCS module was flanked by *oop* transcription terminators,⁴⁷ which has been applied in 143 144 Synechocystis before.^{48,49} Moreover, the surrounding unique restriction sites enable modular 145 exchange of selection markers and the MCS module between all derivatives of the pSOMA series, 146 which substantially increases cloning flexibility from a user perspective. Detailed features of the 147 individual plasmids are given in Table 1.

148

149 Table 1. Plasmids generated or used in this study. Feature description: Amp^{R} = ampicillin resistance, Cm^{R} = 150 chloramphenicol resistance, Gen^R = gentamicin resistance, Km^R/Neo^R = kanamycin/neomycin resistance, Strep^R/Spec^R 151 = streptomycin/spectinomycin resistance, Tet^{R} = tetracycline resistance, MCS = multiple cloning site, P_{J23101} = BioBrick 152 BBa_J23101 synthetic promoter,⁹ P_{nrsB} = nrsB promoter from Synechocystis,⁵⁰ atpE 5'UTR = untranslated region upstream of *E. coli atpE* gene,⁵¹ RBS* = synthetic ribosome binding site,⁵² *bvmo* = Baever-Villiger monooxvgenase 153 154 gene from Acidovorax sp. CHX100.⁵³ sfqfp = superfolder green fluorescent protein gene,⁵⁴ 3xFLAG-tag = sequence 155 coding for triple FLAG protein tag, Step-tag = sequence coding for streptavidin protein tag, BBa_B0015 = BioBrick BBa B0015 double transcription terminator.⁹ $T_{oop} = oop$ transcription terminator⁴⁷ 156

Plasmid	Features	Replicons	References
pSOMA10	Km ^R /Neo ^R , MCS, 3xFLAG-tag	pSC101, pCA2.4	This study
pSOMA11	Cm ^R , MCS, 3xFLAG-tag, Toop	pSC101, pCA2.4	This study
pSOMA12	Gen ^R , MCS, 3xFLAG-tag, Toop	pSC101, pCA2.4	This study
pSOMA13	Strep ^R /Spec ^R , 3xFLAG-tag, <i>T_{oop}</i>	pSC101, pCA2.4	This study
pSOMA14	Tet ^R , MCS, 3xFLAG-tag, Toop	pSC101, pCA2.4	This study
pSOMA15	Km ^R /Neo ^R , MCS, 3xFLAG-tag, Toop	pSC101, pCB2.4	This study
pSOMA16	Cm ^R , MCS, 3xFLAG-tag, Toop	pSC101, pCB2.4	This study
pSOMA17	Gen ^R , MCS, 3xFLAG-tag, Toop	pSC101, pCB2.4	This study
pSOMA18	Strep ^R /Spec ^R , MCS, 3xFLAG-tag, <i>T</i> _{oop}	pSC101, pCB2.4	This study
pSOMA19	Tet ^R , MCS, 3xFLAG-tag, Toop	pSC101, pCB2.4	This study
pPMQAK1- <i>bvmo</i>	Amp ^R , Km ^R , <i>P_{nrsB}</i> , RBS*, <i>bvmo</i> , Strep- tag, BBa_B0015	RSF1010	Ref. [55]
pSOMA10-bvmo	<i>P_{nrsB}</i> , RBS*, <i>bvmo</i> , Strep-tag, BBa_B0015	pSOMA10	This study
pSEVA351	Cm ^R , MCS	RSF1010	Ref. [28]
pSEVA351-sfgfp	<i>P</i> _{J23101} , <i>atpE</i> 5'UTR, <i>sfgfp</i> , BBa_B0015	pSEVA351	This study
pSOMA10-sfgfp	<i>P</i> _{J23101} , <i>atpE</i> 5'UTR, <i>sfgfp</i> , BBa_B0015	pSOMA10	This study
pSOMA11-sfgfp	<i>P</i> _{J23101} , <i>atpE</i> 5'UTR, <i>sfgfp</i> , BBa_B0015	pSOMA11	This study
pSOMA15-sfgfp	<i>P</i> _{J23101} , <i>atpE</i> 5'UTR, <i>sfgfp</i> , BBa_B0015	pSOMA15	This study
pSOMA16-sfgfp	<i>P</i> _{J23101} , <i>atpE</i> 5'UTR, <i>sfgfp</i> , BBa_B0015	pSOMA16	This study

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The pSOMA derivatives can be used as shuttle vectors due to the maintenance in *E. coli* and *Synechocystis*

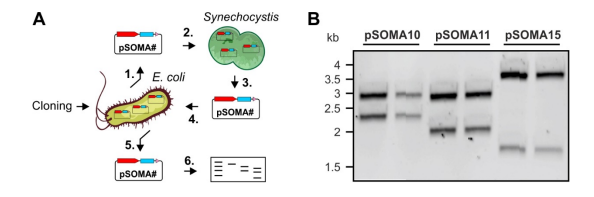
160 The successful cloning already proved the maintenance of the pSOMA plasmids in E. coli.

161 However, to verify their usability as shuttle vectors, i.e. their replication and maintenance in both

162 E. coli and Synechocystis, we performed a transformation-isolation-analysis circuit with selected 163 plasmids (Figure 2A). This included isolation of the plasmids from *E. coli* (step 1), which were 164 then used to transform Synechocystis via electroporation (step 2). In general, colony forming units 165 (CFUs) were obtained on selective agar plates after transformation of Synechocystis with 166 pSOMA10-13 and pSOMA15-17. In case of pSOMA14 and pSOMA19, harboring a tetracycline 167 resistance cartridge, a green cell lawn instead of single colonies was obtained, likely due to the 168 light-sensitivity of this antibiotic. These plasmids were therefore excluded from the following 169 experiments. Several Synechocystis colonies were randomly checked by colony PCR, which 170 confirmed the presence of the recombinant plasmids (Supplemental Figure S1). However, it 171 should be noted that for each pSOMA plasmid only low transformation efficiencies, i.e. 4 - 600 172 CFU µg_{DNA}⁻¹ were achieved. In comparison, RSF1010 derivatives, such as pSEVA351, yielded by 173 average around ten times more colonies from 180 up to 2,500 CFU µg_{DNA}-1.

174 After initial selection, plasmids (including also the endogenous ones) were isolated from monoclonal Synechocystis cultures harboring pSOMA10, pSOMA11, or pSOMA15 (step 3) and 175 176 E. coli cells were transformed using these mixtures (step 4). Again, colonies formed by cells 177 resistant against the corresponding antibiotics were selected and plasmids re-isolated (step 5), 178 followed by control digestion (step 6). Finally, a comparative digestion of the initially obtained 179 pSOMA plasmid and the one which has been transferred into Synechocystis and subsequently 180 back into *E. coli* was performed (Figure 2B). This result, in combination with Sanger sequencing 181 (not shown) ultimately proved that both molecules were identical. Thus, the pSOMA plasmids can 182 indeed be maintained as extrachromosomal elements and facilitate gene transfer into 183 Synechocystis independent from recombination into the host DNA. Moreover, they allow 184 consecutive transfer of genetic information between E. coli and Synechocystis without any 185 observed sequence deviations or decisive structural changes, making them valuable molecular 186 tools.

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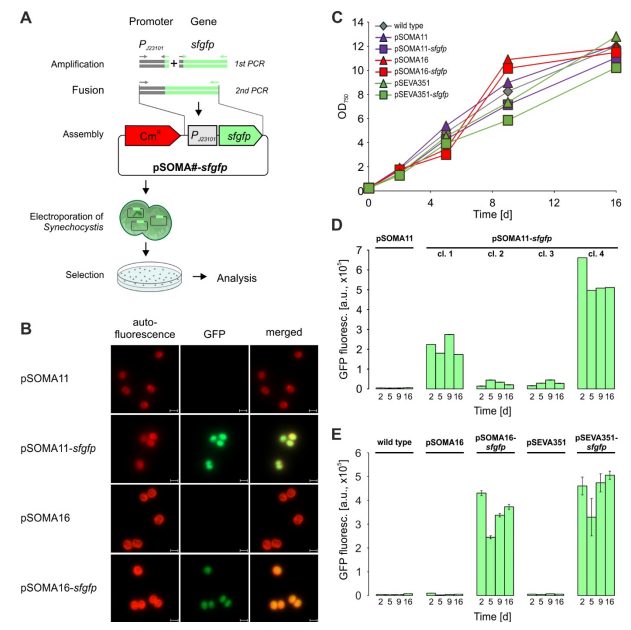
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191 Figure 2. The pSOMA plasmids can be utilized as shuttle vectors. (A) Verification circuit of consecutive 192 transformation of and plasmid isolation from E. coli and Synechocystis to confirm the pSOMA series as true shuttle 193 vector system for genetic engineering of two different species. In vitro assembly of DNA vector parts was followed by 194 (1.) selection and plasmid isolation from E. coli, (2.) transformation of Synechocystis, (3.) selection and plasmid 195 preparation from Synechocystis, (4.) transformation of E. coli with plasmid mixture, (5.) selection and plasmid isolation 196 form E. coli, and finally (6.) control digestion and Sanger sequencing. (B) Representative agarose gel showing restriction 197 analyses of pSOMA10, pSOMA11, and pSOMA15 of both the initially obtained version (step 1) and after passing through 198 the entire circuit (step 5). In each case, the left lane represents the plasmid after step 1, the right lane the same plasmid 199 but after step 5, i.e. after it has already been maintained in Synechocystis. Enzymes used: PstNI and KpnI for pSOMA10 200 and pSOMA11, Bcul and Kpnl for pSOMA15.

201

202 Utilizing the pSOMA shuttle vectors as molecular tool for genetic engineering of 203 *Synechocystis*

204 To further demonstrate the applicability of the pSOMA shuttle vectors to engineer cyanobacteria. 205 we introduced heterologous genes and analyzed their products in Synechocystis. First, we 206 introduced the *sfgfp* gene encoding the superfolder green fluorescent protein⁵⁴ (hereafter GFP) 207 into pSOMA11 and pSOMA16 that are based on two different replicons, pCA2.4 and pCB2.4, 208 respectively. The pSOMA series does not contain promoters to drive the expression of inserted 209 genes by default, preserving flexibility for future applications. However, favored promoters can 210 easily be interconnected to the gene of interest beforehand, e.g. by a fusion PCR approach, to 211 subsequently introduce the synthetic fragment into the pSOMA plasmids via classical assembly 212 procedures (Figure 3A). Accordingly, the sfafp coding sequence was fused to the artificial 213 BioBrick promoter BBa_J23101⁹ (hereafter P_{J23101}) that was found to promote high and constitutive expression in Synechocystis⁵⁶ and Synechococcus elongatus PCC 7942⁵⁷ (Figure 3A). Indeed, a 214 215 specific GFP signal could be observed in cells of two recombinant Synechocystis strains harboring 216 the plasmids pSOMA11-sfgfp or pSOMA16-sfgfp. Via fluorescence microscopy these strains 217 could clearly be distinguished from cells accommodating the respective empty vectors (Figure 218 3B).



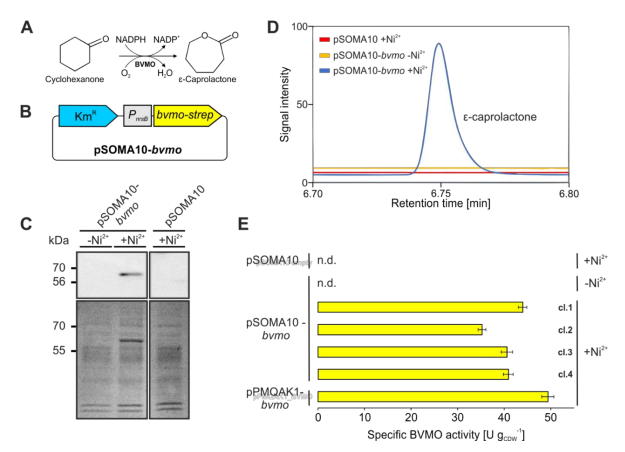
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220 Figure 3. Expression of a reporter gene from pSOMA11 and pSOMA16. (A) Cloning strategy for the genetic 221 constructs. A gene encoding the superfolder green fluorescent protein (sfgfp) was fused to a synthetic minimal promoter 222 (BioBrick BBa_J23101, = P_{J23101}) that was found to mediate constitutive and high expression in Synechocystis.⁵⁶ The 223 construct was inserted into respective pSOMA variants conferring chloramphenicol resistance (Cm^R), particularly 224 pSOMA11 (pCA2.4 backbone) and pSOMA16 (pCB2.4 backbone). As control, the same sfgfp expression cassette was 225 also inserted into the RSF1010 derivative pSEVA351. (B) Fluorescence microscopy pictures of Synechocystis cells 226 expressing sfgfp from pSOMA11 or pSOMA16. Autofluorescence and GFP were visualized separately or as merged 227 pictures (scale bar 2 µm). (C) Cell growth as a measure of OD750. The wild type parental strain lacks a synthetic plasmid 228 and was cultivated without chloramphenicol. (D, E) Spectrofluorimetrical quantification of GFP fluorescence in vivo. 229 Shown are GFP signals normalized to OD₇₅₀ (arbitrary units, a.u.) obtained from Synechocystis transformed with either 230 pSOMA11-sfgfp (D), pSOMA16-sfgfp (E), pSEVA351-sfgfp (E), or the respective empty vectors without sfgfp, as well 231 as the parental strain (wild type). Panel D shows individual data obtained from independent clones (cl.1-4), each 232 measured in three technical replicates. In panel E, data are the mean ±SD of at least three biological replicates, i.e. 233 clones.

235 To evaluate the stability of the genetic setup, the GFP fluorescence signal was quantitatively monitored over a period of 16 days. For a comparison with widely used RSF1010 plasmids, the 236 237 same *P*_{J23101}::sfgfp cassette was also inserted into pSEVA351,²⁸ giving pSEVA351-sfgfp, which 238 was also introduced into Synechocystis. In general, maintenance of the pSOMA plasmids did not 239 impair cellular fitness as displayed by similar growth behavior of the recombinant strains under 240 selection pressure compared to the wild type cultivated without antibiotics (Figure 3C). A GFP 241 signal considerably exceeding fluorescence of control strains could continuously be detected in 242 the same time range, consistent with a constitutive expression of the sfqfp gene (Figure 3D, E). 243 However, it should be noted that individual transformants harboring pSOMA11-sfafp showed high 244 biological variation, whereas strains harboring pSOMA16-sfgfp and pSEVA351-sfgfp behaved 245 more homogenous (Figure 3D, E). Furthermore, the maximum relative fluorescence intensities 246 for strains carrying pSOMA11-sfgfp or pSOMA16-sfgfp were comparable to those harboring 247 pSEVA351-sfgfp.

248 In addition to GFP fluorescence, we illustrate the utilization of pSOMA shuttle vectors to implement 249 heterologous metabolic pathways and hence, to design cyanobacterial photobiocatalysts. Here 250 we made use of previously demonstrated concepts of biotransformations using electrons derived 251 from photosynthesis.^{55,58} In particular, we introduced the gene encoding Baeyer-Villiger 252 monooxygenase (hereafter BVMO) from Acidovorax sp. CHX100 into Synechocystis. This 253 cyclohexanone monooxygenase is NADP⁺/NADPH-dependent and uses molecular oxygen (O₂) 254 as co-factor to oxidize cyclic ketones to lactones or esters while forming water as by-product.⁵⁹ 255 BVMO is part of a multistep process for the reaction cascade transforming cyclohexane to the 256 value-added chemical ε-caprolactone, in which BVMO catalyzes the conversion of cyclohexanone 257 to ε -caprolactone (**Figure 4A**). The product ε -caprolactone is a chemical commodity and can be 258 used as precursor for the production of adipic acid and its derivatives, e.g. nylon polymers.⁶⁰ To 259 achieve *bvmo* expression from pSOMA plasmids in *Synechocystis*, a previously assembled construct⁵⁵ was amplified and transferred into pSOMA10. Accordingly, the resulting plasmid 260 261 pSOMA10-bvmo contained the bvmo gene in fusion with a Strep-tag and the nickel ion (Ni²⁺)-262 dependent *nrsB* promoter (Figure 4B). The latter is native to *Synechocystis*⁶¹ and shows tight repression as well as a high dynamic range in this strain.⁵⁰ After transformation, the recombinant 263 264 Synechocystis strains carrying either pSOMA10 or pSOMA10-bvmo were analyzed for sufficient 265 bymo expression and in vivo activity of the corresponding enzyme. First, soluble proteins were 266 separated by SDS-PAGE, whereby a Ni²⁺-inducible *bvmo* expression was already indicated by a 267 discrete protein band at ~60 kDa that did not appear in the lane representing the control strain 268 with empty vector. Furthermore, the Strep-tagged fusion protein was immunologically detected via

269 Western Blot and the obtained signal corresponds to the band that became visible in the SDS gels (Figure 4C). Consistently, specific BVMO activity, i.e. *e*-caprolactone formation from added 270 cyclohexane, could only be detected in presence of Ni²⁺, whereas background signals without 271 inducer and a control strain lacking the *bvmo* gene were negligible (Figure 4D). As a positive 272 control, we made use of the previously generated plasmid pPMQAK1-bvmo⁵⁵ that harbors the 273 274 same gene cassette but is instead based on the broad-host-range vector RSF1010. Thereby, a 275 maximum of ~44 U g_{CDW}^{-1} was observed for cells carrying pSOMA10-*bvmo*, exhibiting ~90% productivity relative to the positive control (~49 U g_{CDW}⁻¹; Figure 4E). 276





279 Figure 4. Implementation of a heterologous enzyme reaction in Synechocystis using pSOMA10. (A) Reaction 280 catalyzed by Baeyer-Villiger monooxygenase (BVMO) from Acidovorax sp. CHX100, which performs the oxygen- and 281 NADPH-dependent biotransformation of cyclohexanone to ε-caprolactone.⁶⁰ (B) Schematic illustration of the genetic 282 constructs. The *bvmo* gene encoding a C-terminally Strep-tagged fusion protein was inserted into pSOMA10. Ni²⁺ 283 serves as inducer for the nrsB promoter (PnrsB).⁵⁰ (C) Western Blot for the detection of a BVMO-Strep fusion protein 284 (~60.2 kDa). Soluble proteins separated by SDS-PAGE are shown as loading control (lower panel). (D) Representative 285 gas chromatograms for Synechocystis extracts revealing ε-caprolactone product formation from cyclohexanone via 286 BVMO-driven biotransformation. (E) Specific BVMO whole-cell activity of four Synechocystis clones accommodating 287 pSOMA10-bvmo (cl.1-4) or indicated controls (n.d. = not detectable). Cyclohexanone conversion units (in µmol min⁻¹) 288 were normalized to cell dry weight (CDW). Data are the mean ± SD of four technical replicates.

The two pSOMA subseries allow combination with each other and further compatible plasmids

292 Next, we introduced two compatible plasmids with different origins of replication and selection 293 markers into one cell lineage of Synechocystis to investigate, whether the pSOMA shuttle vectors 294 can be used in combination with other plasmids, such as the widely used RSF1010 derivatives 295 (Figure 5A). Therefore, we subsequently transformed our obtained Synechocystis strain carrying pSOMA10-bvmo either with pSEVA351²⁸ or pSEVA351-sfgfp, which harbored the same 296 297 P_{J23101} ::sfgfp cassette as used before (**Figure 3**). This strategy permitted us to analyze the 298 presence of the two distinct plasmids by measuring the GFP signal and BVMO activity. As 299 expected, specific GFP fluorescence as well as ε -caprolactone production were detected again, 300 yet, both in a single strain (Figure 5B, C). Analogously, we introduced representatives of the 301 pCB2.4-based pSOMA subseries, namely pSOMA16 or pSOMA16-sfgfp into the same 302 Synechocystis host strain that already carried pSOMA10-bvmo. Consistently, this strain also 303 showed both activities, illustrating the possibility of combining both pSOMA subseries based on 304 pCA2.4 or pCB2.4 (Figure 5D, E).

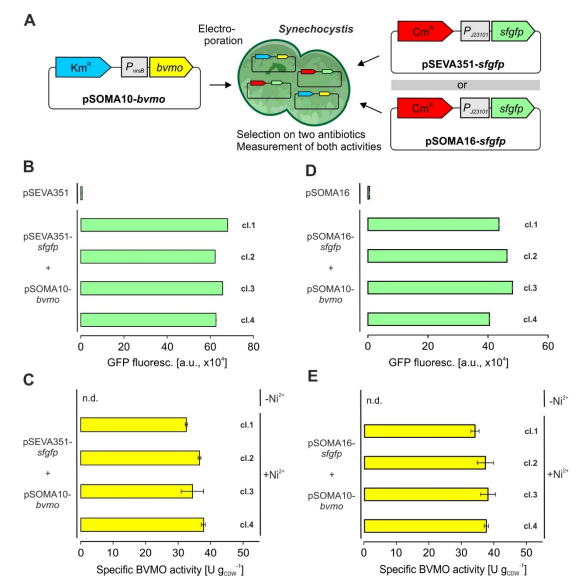


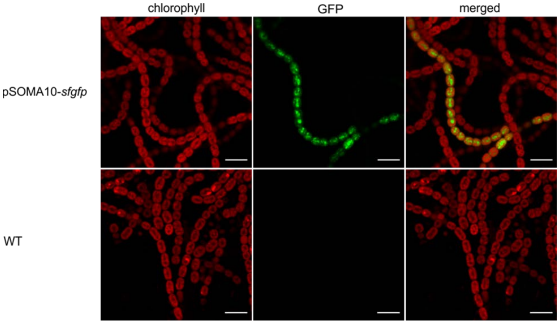
Figure 5. Expression of distinct heterologous genes from individual shuttle vectors. (A) Generated Synechocystis
 strain with two replicative plasmids. Wild type cells were transformed with the pCA2.4 derivative pSOMA10-bvmo
 mediating kanamycin resistance (Km^R) as well as either the RSF1010-based pSEVA351-sfgfp or the pCB2.4-derived
 pSOMA16-sfgfp (plus respective empty vector without sfgfp) giving chloramphenicol resistance (Cm^R). (B,
 D) Spectrofluorimetrical quantification of GFP fluorescence of cells carrying the given plasmid combination. Data are
 the mean ±SD of three technical replicates. (C, E) Specific BVMO whole-cell activity of the same strains. Data are the

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305

314 The pSOMA shuttle vectors show a limited host range

As the application in different species might be of interest, we tested if the pSOMA plasmids could also be transferred into other cyanobacteria. These comprised the filamentous strains *Anabaena* sp. PCC 7120 and *Trichormus* (previously known as *Anabaena*) *variabilis* ATCC 29413, as well as two fast-growing strains that have attracted attention as potential production systems: *Synechococcus elongatus* UTEX 2973 and the recently discovered marine strain *Synechococcus* 320 sp. PCC 11901. In particular, pSOMA10 and pSOMA15, as well as respective derivatives harboring the sfgfp reporter gene, pSOMA10-sfgfp and pSOMA15-sfgfp, were utilized. For 321 322 T. variabilis, Synechococcus elongatus UTEX 2973, and Synechococcus sp. PCC 11901 no CFUs 323 could be obtained after transformation via electroporation. However, several colonies were gained 324 on selective plates in case of the filamentous model strain Anabaena sp. PCC 7120 following 325 transformation with pSOMA10-sfgfp. Remarkably, reporter gene expression could also be 326 illustrated by fluorescence microscopy, whereby several independent filaments that clearly show 327 GFP fluorescence were observed (Figure 6). In contrast, a similar pattern was not detected in any 328 of the wild type filaments. Moreover, GFP fluorescence was distributed over various individual 329 cells of the same filament. This indicates that the respective pSOMA plasmid was maintained in 330 all these cells, due to the fact that the reporter protein cannot diffuse through the septal junctions.⁶² 331 However, it should be noted that the fluorescence intensity between different filaments varied and 332 even a variation between cells within the same filament could be observed, which necessitates 333 further characterization and optimization. Nonetheless, with this proof of principle we suggest that 334 the host range of pSOMA shuttle vectors based on the pCA2.4 replicon can be extended to at 335 least one more cyanobacterium in addition to Synechocystis.



WT

337 Figure 6. Representative fluorescence microscopy pictures of Anabaena sp. PCC 7120 wild type (WT) and cells 338 harboring pSOMA10-sfgfp. It should be noted that none of the WT filaments showed a similar GFP fluorescence 339 pattern as observed for various individual filaments harboring pSOMA10-sfgfp. Samples were excited at 488 nm and 340 fluorescence emission detected separately for chlorophyll (red) and GFP (green). Both channels together are shown in 341 the merged images (scale bar 10 µm).

342 **Discussion**

343 Due to the restricted number of replicative plasmids, integration of (heterologous) genes into 344 endogenous DNA is still the predominant strategy for metabolic engineering of Synechocystis. 345 Accordingly, this has been accompanied by the development of recombination and selection strategies as well as the search for suitable integration sites.^{16,40,63} However, genetic segregation 346 347 in a polyploid organism, such as Synechocystis, is a laborious procedure and all the more time-348 consuming when carried out sequentially, even if neutral sites are targeted. Moreover, targeting endogenous plasmids via homologous recombination directly is accompanied by the same 349 350 problems and appears to be even more difficult compared to chromosomal sites.^{40,64,65} For 351 instance, when targeting pCC5.2, even after several attempts and multiple rounds of selection 352 wild type alleles were still detectable, i.e. only partially segregated mutant strains were 353 obtained.^{40,64} In addition, once a recombinant strain was generated by homologous recombination, 354 it cannot be easily altered, e.g. a selection marker cannot be changed without implementing 355 genetic modifications within the host DNA. Therefore, this strategy has a rather low flexibility, in 356 particular for extensive engineering. The latter might be required to implement complex genetic 357 scenarios such as for sophisticated metabolic pathways and/or heterologous enzyme complexes 358 which also include various maturation factors.

359 To overcome these problems, we designed a chimeric shuttle vector series to expand the 360 molecular toolset of cyanobacteria, in particular Synechocystis. As the limited number of 361 replicative plasmids currently presents a bottleneck for the introduction of genes independent from 362 homologous recombination, similar studies were conducted previously.^{41–43} An overview of so far 363 reported synthetic shuttle vectors derived from small endogenous plasmids of Synechocystis, 364 including this study, is given in Table 2. For example, Jin et al. (2018) generated the pCC5.2derived shuttle vector pSCBe that was fused to the replicon of pMB1⁶⁶ from *E. coli*. Their study 365 366 confirmed heterologous expression of a reporter gene from pSCBe in Synechocystis as well as 367 stable maintenance of the plasmid for at least 50 generations even without antibiotic selection. 368 Moreover, similar to our study, stable co-existence of pSCBe with an RSF1010-based plasmid 369 was reported.⁴¹ The principle usability of pCA2.4 and pCB2.4 to generate synthetic shuttle vectors 370 for Synechocystis has also been demonstrated previously.⁴² Albeit the two obtained plasmids 371 pCA-UC118 and pCB-SC101 could generally be used as expression platforms for heterologous 372 genes, several aspects might require optimization to finally enable user-friendly cloning and a 373 flexible combination of customized plasmids with genetic setups that might already exist in a 374 particular lab. Therefore, our study expands these initial attempts by introducing a whole library of 375 replicative plasmids based on pCA2.4- and pCB2.4. It comprises a broad spectrum of selection markers as well as suitable cloning features, such as a MCS and a 3XFLAG-tag for the subsequent
 synthesis of proteins that can easily be detected. Moreover, it is known that efficient promoters
 can interfere with plasmid replication and diminish expression of plasmid-specified genes.⁶⁷
 Therefore, the MCS module was additionally flanked by transcription terminators that prevent
 transcriptional read-through and hence, shield the potentially inserted gene expression cassettes
 to ensure plasmid stability and support high gene expression.

382

Table 2: **Overview of reported chimeric shuttle vectors for** *Synechocystis.* All plasmids were derived from small endogenous plasmids pCA2.4, pCB2.4, and pCC5.2 (or ORFb minimal replicon), which were individually fused to a replicon from *E. coli.* Feature description: ori = origin of replication, Cm^R = chloramphenicol resistance, Gen^R = gentamicin resistance, Km^R = kanamycin resistance, Strep^R/Spec^R = streptomycin/spectinomycin resistance, oriT = origin of transfer for conjugation, MCS = multiple cloning site

Plasmid	Synechocystis	Further relevant	Plasmid	Application	Reference
	replicon	features	introduction		
pCA-UC118	pCA2.4	pUC118 ori, Cm ^R ,	transformation	shuttle vector,	Ref. [42]
		fluorescent reporter	based on	reporter verification,	
		expression cassette	natural	compatibility analysis	
			competence		
pSOMA10-13	pCA2.4	pSC101 ori,	electroporation	shuttle vector	This study
		Km ^R /Cm ^R /Gen ^R /			
		Strep ^R /Spec ^R , MCS,			
		3xFLAG-tag, flanking			
		transcription			
		terminators			
pCB-SC101	pCB2.4	pSC101 ori,	transformation	shuttle vector,	Ref. [42]
		Strep ^R /Spec ^R ,	based on	reporter verification,	
		fluorescent reporter	natural	compatibility analysis	
		expression cassette	competence		
pSOMA15-18	pCB2.4	pSC101 ori,	electroporation	shuttle vector,	This study
		Km ^R /Cm ^R /Gen ^R /		compatibility analysis	
		Strep ^R /Spec ^R , MCS,			
		3xFLAG-tag, flanking			
		transcription			
		terminators			
pSCB	pCC5.2 (ORFb)	pMB1 ori,	conjugation	shuttle vector	Ref. [41]
		Strep ^R /Spec ^R , oriT,			
		MCS			
pSCB-YFP	pCC5.2 (ORFb)	pMB1 ori,	conjugation	shuttle vector,	Ref. [41]
		Strep ^R /Spec ^R , oriT,		reporter verification,	
		MCS, fluorescent		stability assay,	
				compatibility analysis	

		reporter expression cassette			
pSCBe	pCC5.2 (ORFb)	pMB1 ori, Strep ^R /Spec ^R , oriT, MCS, expression cassette (<i>P</i> _{trc10} , FLAG-tag, His-tag)	conjugation	shuttle/expression vector	Ref. [41]
pCCM1-FbFP	pCC5.2	pMB1 ori, Strep ^R /Spec ^R , fluorescent reporter expression cassette, encoded single guide RNA targeting native pCC5.2	transformation based on natural competence	shuttle vector, reporter verification, simultaneous curing of native pCC5.2 via CRISPR/Cas9	Ref. [43]

388

389 Albeit the mechanisms of transformation based on natural competence are not experimentally 390 verified for Synechocystis, the anticipated process likely results in the linearization of supplied plasmids,⁶⁸ which are then taken up as single-stranded DNA.^{15,69} Consequently, the linearized, 391 392 single-stranded DNA needs to be repaired to maintain a replicative plasmid within the cell. This, 393 however, might interfere with the overall transformation procedure and/or raise the possibility of 394 recombination events, e.g. into the endogenous plasmids pCA2.4, pCB2.4, or pCC5.2. Therefore, 395 the pSOMA series should be introduced into cells via one-step transformation by electroporation. 396 This approach has been proven to work well in case of replicative plasmids for Synechocystis 397 while maintaining their structural integrity²⁷ and it also spares the need for time-consuming 398 microbial segregation to obtain axenic cultures following conjugation (triparental mating).⁷⁰ 399 Previous studies in fact reported the successful transformation with synthetic shuttle vectors via 400 natural competence (see **Table 2**). However, re-isolation of these plasmids followed by molecular 401 investigation had not been performed yet. Here, we demonstrate that the pSOMA vectors can be 402 re-isolated from recombinant cells without showing any variation from the plasmid that was initially 403 supplied for transformation. Thus, they serve as true shuttle vectors and could be used as 404 standard tools for DNA exchange between the molecular "work horse" E. coli and Synechocystis.

The introduction of multiple plasmids into one cell increases flexibility of genetic combinations and hence, supports customized engineering. The maintenance of several shuttle vectors in one cell lineage and compatibility with the broad-host-range plasmid RSF1010 have already been indicated for *Synechocystis*.^{41,42} However, the simultaneous expression of distinct heterologous genes and the determination of their associated activities from two different plasmids have not been demonstrated. To the best of our knowledge, this is shown here for the first time in a cyanobacterium by confirming BVMO activity (gene maintained on pSOMA10) and GFP fluorescence (gene provided either by pSOMA16 or an RSF1010 derivative, see **Figure 5**). Likewise, other combinations, e.g. also with pCC5.2 derivatives, or even more than two compatible plasmids might be possible. Such coexistence facilitates the insertion of multiple genes or pathway clusters of large size, which may be difficult to assemble and to be maintained on just one plasmid or to be even recombined into the host DNA.

417 However, genetic engineering via pSOMA plasmids still requires further optimization, as we 418 achieved only low and fluctuating transformation efficiencies for Synechocystis. This may be 419 caused by the electroporation procedure itself. The competition with the native endogenous 420 plasmids and their overall regulation, e.g. abundance control or the differential expression of the 421 encoded genes with unknown function, also present possible hurdles. For example, the copy 422 number of pCA2.4 and pCB2.4 was shown to vary depending on the growth phase and nutritional 423 conditions in Synechocystis.³⁹ To avoid this problem, a Synechocystis host strain cured from the 424 native counterparts could be used in the future. This should in principle be possible, as both 425 endogenous plasmids were reported to be dispensable under standard phototrophic growth 426 conditions.^{36–38,43} Simultaneous curing of the native plasmids by a CRISPR/Cas approach could 427 be another option, as already described for a pCC5.2-based shuttle vector.⁴³ The eliminated 428 competition could also enhance gene expression levels while reducing the observed variation in 429 GFP fluorescence between individual clones, which might be due to different pSOMA plasmid 430 copy numbers. Controversially, individual clones harboring a similar pSOMA plasmid based on 431 pCA2.4 but instead the gene encoding BVMO showed rather low variation. This contrast points 432 towards the influence of the respective reporter gene and its expression driven either by a 433 constitutive or inducible promoter, rather than a general pCA2.4-related effect. Nevertheless, 434 variation between different transformants and gene expression levels obtained from various 435 derivatives of the pSOMA series are worthwhile to become evaluated in prospective studies. 436 Another factor to be considered in the future is plasmid stability. Therefore, maintenance should 437 be investigated without the unpleasant supplementation of antibiotics after initial selection, as already shown for RSF1010^{27,40} and pCC5.2 derivatives.⁴¹ 438

Cyanobacterial plasmids typically show a rather narrow host range and only replicate in closely
related strains, e.g. plasmid pMA4 isolated from the thermophilic strain *Synechococcus* sp. MA4
could be introduced into *Synechococcus* sp. MA19.⁷¹ However, shuttle vectors originating from
the endogenous plasmid pANS of *Synechococcus elongatus* PCC 7942 were successfully used
to transform phylogenetically distant *Anabaena* sp. PCC 7120, whereas it was not possible to

444 introduce them into Leptolyngbya BL0902, Synechocystis WHSYN, and Synechocystis.33 445 Moreover, derivatives of pDU1 from Nostoc sp. PCC 7524 could be maintained in a number of 446 rather distantly related strains, including Anabaena sp. PCC 7120 and Anabaena sp. M-131³¹; 447 Fischerella muscicola sp. PCC 7414 and Chlorogloeopsis fritschii sp. PCC 6912⁷²; Fischerella thermalis (also known as Mastigocladus laminosus) SAG 4.84⁷³, Chroococcidiopsis spp. CCMEE 448 029, 057, and 12374; as well as Oscillatoria MKU 277.75 Similar to the mentioned studies, we 449 450 investigated the cyanobacterial host range of pSOMA plasmids. Therefore, four different 451 cyanobacterial species other than Synechocystis were tested, whereby transformants were 452 obtained for Anabaena sp. PCC 7120 carrying the pCA2.4 derivative pSOMA10-sfgfp (Figure 6). 453 Therefore, it is likely that at least pCA2.4-based vectors could in principle be used for the 454 transformation of filamentous strains related to Anabaena. Interestingly, Anabaena species seem 455 to have relaxed requirements for supported heterologous plasmids, even from phylogenetically distant strains, as they are able to maintain plasmids based on pANS³³ and pDU1,³¹ besides the 456 457 acceptance of pCA2.4 derivatives reported in this study. However, it should be noted that the 458 transformation efficiency was again rather low, similar to the observations made for 459 Synechocystis. Therefore, in case of the other tested strains we cannot finally conclude if pSOMA 460 plasmid transfer and maintenance is indeed not possible or the transformation efficiency was 461 simply too low. Nevertheless, the results are promising for the development of novel 462 cvanobacterial shuttle vectors that have a broader host range.

463

464 Material and Methods

465 Strains and culture conditions

466 *E. coli* strain DH5α was grown at 30-37°C in liquid LB medium shaking at 180-200 rpm or LB agar 467 plates. The medium was supplemented with 35 μ g ml⁻¹ chloramphenicol, 10 μ g ml⁻¹ gentamicin, 468 50 μ g ml⁻¹ kanamycin, 25 μ g ml⁻¹ streptomycin, or 10 μ g ml⁻¹ tetracycline when necessary. The 469 RepA protein of pSC101 is temperature-sensitive,⁷⁶ which is why the incubation of *E. coli* 470 exhibiting pSOMA was performed at 30°C in order to maintain the plasmid.

471 Cyanobacterial freshwater strains *Synechocystis* (sp. PCC 6803) and *Synechococcus elongatus*472 UTEX 2973 were cultivated in BG11 liquid medium, containing 16 µM Na₂EDTA (yBG11),⁷⁷ or on
473 BG11⁷⁸ agar plates, both buffered with 10-50 mM HEPES to pH 7.2. The marine strain
474 *Synechococcus* sp. PCC 11901 was cultivated with a modified AD7 liquid or solid medium (MAD)
475 containing 96 mM NaNO₃, 240 µM FeCl₃, 1.2 mM KH₂PO₄, 18 g l⁻¹ NaCl, 0.6 g l⁻¹ KCl, and 3 pM
476 cobalamin (vitamin B₁₂).⁷⁹ Filamentous cyanobacteria, *Anabaena* sp. PCC 7120 and *Trichormus*477 *variabilis* ATCC 29413 were grown in BG11 liquid media or on solid BG11 agar plates⁷⁸ buffered

with 10 mM TES to pH 8.2. The media were supplemented with 2-10 μ g ml⁻¹ chloramphenicol, 2-10 μ g ml⁻¹ gentamicin, 10-50 μ g ml⁻¹ kanamycin, or 4-20 μ g ml⁻¹ streptomycin as needed. Conditions of growth were set to 28-30°C, ambient CO₂, 25-50 μ mol photons m⁻² s⁻¹, 120-200 rpm for shake flasks permitting gas exchange, and 75% humidity.

482 **Construction of plasmids**

483 The pSOMA plasmids were assembled using PCR-generated DNA fragments. PCR fragments 484 were obtained using Phusion[™] High-Fidelity DNA Polymerase (Thermo Scientific) and 485 interconnected via overlap extension PCR utilizing the same enzyme or Gibson Assembly⁸⁰ 486 according to the manufacturer's instructions. 5' extensions were added to the primers to create 487 homologous overhang sequences of the antibiotic resistance cartridges and the two replicons. All 488 primers for the amplification of PCR products and their respective templates are given in 489 Supplemental Table S1. The MCS module containing two flanking *oop* transcription 490 terminators,^{47,48} 11 unique restriction sites (HindIII, Sphl, Pstl, Sall, Xbal, BamHI, Xmal, Kpnl, 491 SacI, EcoRI, and NdeI), as well as a 3xFLAG-tag was synthesized by Eurofins Genomics. The 492 generated plasmids are given in **Table 1**. Control digestion was performed using FastDigest[™] 493 restriction endonucleases (Thermo Scientific). Purification of PCR products, plasmids, and 494 genomic DNA was performed using the respective kits NucleoSpin[™] Gel & PCR Clean-up, 495 NucleoSpin Plasmid QuickPure™ (MARCHEREY-NAGEL), and peqGOLD Bacterial DNA 496 (PEQLAB) according to the manufacturer's instructions. Verification of plasmids was done by control digestion by different FastDigest[™] restriction enzymes or colony PCR using DreamTaq[™] 497 498 DNA Polymerase (Thermo Scientific) or GoTag® DNA Polymerase (Promega). In silico work was 499 performed using the software Geneious (Biomatters).

500 Expression cassettes to be inserted into individual pSOMA plasmids were generated beforehand 501 including promoters, ribosome binding sites and the gene of interest. The plasmid pSEVA351-502 sfgfp was assembled by first amplifying two linear fragments with homologous overhangs suitable for insertion into PstI cut pIGA⁸¹: BioBrick BBa_B0015^{9,25} and the reporter gene sfafp^{54,82} with 503 504 additionally added 5' primer extension containing the BioBrick BBa_J23101 $(P_{J23101})^{9,56}$ promoter 505 (with $C \rightarrow G$ substitution at 3' end) and a KpnI restriction site, yielding the intermediate plasmid 506 pIGA_Ribo. The *E. coli atpE* 5' untranslated region⁵¹ was amplified with homologous overhangs 507 allowing subsequent introduction between promoter and gene through restriction at the 508 interconnecting Kpnl site, creating the intermediate plasmid pIGA_Ribo atpE. Promoter, atpE 5' 509 untranslated region, and gene are thereby fused by KpnI scars (5'-TAC-3' and 5'-GTACC-3', 510 respectively) and downstream of *sfgfp* is a BioBrick scar.⁹ The whole *P*_{J23101}::*sfgfp* cassette was 511 afterwards amplified to comprise homologous ends for insertion into KpnI-linearized pSEVA351.28

The same P_{J23101} ::sfgfp reporter construct was equipped with homologous sequences for insertion into pSOMA11 and pSOMA16 cut by KpnI. The P_{nrsB} ::bvmo gene expression unit; consisting of P_{nrsb} ,⁵⁰ RBS*,⁵² bvmo,⁵³ linker 5'- AGCGCT-3', Strep-tag II, and BioBrick BBa_B0015; was obtained by PCR amplification from the template pEERM3_BVMO⁵⁵ with suitable homologous overhangs for the subsequent insertion into KpnI-linearized pSOMA10. The complete DNA sequences of the pSOMA plasmids are provided as genbank files in the supplementary material.

518 Generation of recombinant cyanobacterial strains

519 Synechocystis sp. PCC 6803, Synechococcus elongatus UTEX 2973, and Synechococcus sp. 520 PCC 11901 cells were made electro-competent, transformed via electroporation, and selected as described elsewhere.⁸³ Briefly, cultivation was performed as described above until logarithmic 521 522 growth phase of an optical density at λ =750 nm (OD₇₅₀) from ~0.5-1. Afterwards, cells were 523 harvested and the resulting pellet washed three times with ice-cold HEPES (1 mM, pH 7.5) and 524 prepared as 60 µl aliguots. For transformation via electroporation, 100-500 ng of plasmid DNA 525 were added to an aliquot of electro-competent cells. An electroporation pulse was performed in 526 respective cuvettes by applying 2.5 kV for 5 ms (12.5 kV cm⁻¹) in an Eporator® 4309 (Eppendorf). 527 Afterwards, cells were resuspended in 1 ml respective media, i.e. yBG11 for freshwater strains or 528 MAD for the marine strain Synechococcus sp. PCC 11901, and added to fresh liquid medium for 529 24 h shaking standard incubation without antibiotics and subsequently collected to be spread on 530 selective agar plates containing appropriate antibiotics with concentration as given above. 531 Colonies appeared within ~10 days after further cultivation at 25-50 µmol photons m⁻² s⁻¹ and 30°C. 532 Transformants were checked for plasmid presence via colony PCR using suitable primers listed 533 in Supplemental Table S1. For Anabaena sp. PCC 7120 and T. variabilis electroporation was performed as described by Thiel and Poo (1989)⁸⁴ with minor modifications. pSOMA plasmids 534 535 were first transferred to E. coli strain HB101[pRL623], which encodes three methylases protecting the plasmid from the native restriction enzymes Aval, II and III in Anabaena sp. PCC 7120.85 The 536 537 plasmids were isolated from E. coli using the Qiagen Plasmid Midi kit and adjusted to a DNA 538 concentration of ~ 1 µg ml⁻¹. Cyanobacteria were grown as described above until they reached 539 OD_{750} of 0.5-1. Cells were than collected by centrifugation at 5,000 x g for 5 min and washed once 540 in TE buffer (2 mM Tricine, 2 mM EDTA pH 8.0) and twice in distilled water. Electroporation was 541 done in 40 µl aliguots containing 20 µg chlorophyll using a MicroPulser (Biorad). After addition of 542 10 µg plasmid DNA, the mixture was kept for 2 min on ice before applying a pulse of 6 kV cm⁻¹ for 543 5 ms. Cells were resuspended in 1 ml yBG11 and incubated for 48 h at 10-20 µmol photons m⁻² 544 s⁻¹ and 28°C before spreading on an Immobilon membrane (HATF, Millipore) on a selective BG11 545 agar plate supplemented with 15 μ g ml⁻¹ neomycin. Once colonies appeared, they were transferred 546 to fresh BG11 agar plates with 15 µg ml⁻¹ neomycin.

547 **GFP fluorescence determination**

The superfolder green fluorescent protein⁵⁴ (referred to as GFP) quantitative measurement was 548 performed with adapted settings as described previously.⁸⁶ Synechocystis pre-cultures, which 549 550 harbored plasmids that encode GFP or their particular empty vectors, were grown in yBG11 551 medium supplemented with 10 µg ml⁻¹ chloramphenicol as given above and diluted to an initial 552 OD₇₅₀ of ~0.25 for main cultures. Cells were then cultivated for 16 days and throughout this period, 553 samples were taken and diluted to an OD₇₅₀ of ~0.25 with yBG11 medium in a final volume of 554 1,200 µl. Samples were then transferred into an opaque black flat microtiter 96-well-plate (Nunc) 555 as technical triplicates (each 200 µl), followed by fluorescence measurements at 556 excitation/emission wavelengths of 485 nm/535 nm, respectively, using an Infinite 200 PRO 557 microplate reader (Tecan, gain: 124, integration time: 2000, excitation bandwidth: 9 nm, emission 558 bandwidth: 20 nm, z-position: 2000 µm, 25 flashes). Furthermore, an identical technical triplicate 559 of each sample was taken to measure the absorption at λ =750 nm in a transparent flat microtiter 560 96-well-plate (Nunc), also using the Infinite 200 PRO microplate reader (bandwidth: 9 nm, 25 561 flashes). Technical triplicates were combined as means and the blank of the yBG11 medium 562 background was subtracted. The fluorescence intensities were normalized by division through 563 respective OD₇₅₀. For confocal fluorescence microscopy analysis of Synechocystis, cells were 564 taken from an exponentially growing culture, cultivated as described above, and visualized by 565 using the Zeiss AxioObserver.Z1/7 microscope (Zeiss) together with the Plan-Apochromat 566 100x/1.40 Oil Ph 3 M27 objective lens (Zeiss), filters 474-528 as well as 650-4095, and Colibri.2 567 Illumination System (Zeiss). Fluorescence was detected using filter set 38, BP 470/40, FT 495, 568 BP 525/50, for GFP at excitation/emission wavelengths 488/509 nm and autofluorescence at 569 599/625 nm. Anabaena sp. PCC 7120 cells were visualized with a confocal laser scanning 570 microscope (CLSM) SP8 (Leica) using a 63x/ 1.4 HCPL APO CS2 objective. Samples were 571 excited at 488 nm and emission detected from 500-540 nm for GFP and 670-720 nm for 572 chlorophyll. Images were analyzed using Fiji.⁸⁷

573 Expression analyses

574 Synechocystis strains containing pSOMA10 or pSOMA10-bvmo were cultivated as described for whole-cell biotransformation assays⁵⁵ and samples of 750 µl with an OD₇₅₀ ~20 taken ~24 h after 575 576 10 µM NiSO₄ was added (except control -Ni²⁺). Cell disruption was performed buffered in TBS 577 (100 mM Tris, 150 mM NaCl, 1 mM PMSF protease inhibitor, pH 7.5) using a Precellys® Evolution 578 homogenizer (Bertin) equipped with a Cryolys® cooling system (Bertin). Therefore, cell 579 suspensions were transferred to 2 ml Precellys® tubes (Bertin) together with a mixture (0.09-0.15 580 mm, 0.17-0.18 mm, and 0.5 mm diameter) of glass beads (Sartorius[™]) and disruption performed 581 for 4x30 s at 10.000 rpm with 30 s breaks in between cooled with liquid N₂ to ~0-4°C. Supernatants 582 of soluble extract were separated by centrifugation and collected for protein concentration 583 determination using Bradford Dye Reagent (Thermo Scientific) according to manufacturer's 584 instructions. Samples of ~10 µg soluble protein were separated by SDS-PAGE (10% acrylamide 585 separation gel) and transferred to 0.45 µm pore size nitrocellulose membranes (GVS). 586 Membranes were subsequently treated and hybridized with Strep-tactin® horse radish peroxidase 587 conjugate (iba) and exposed to WesternSure® PREMIUM Chemiluminescent Substrate (LI-COR) 588 to detect chemiluminescence using a FluorChem FC3 System (ProteinSimple), according to 589 manufacturer's instructions.

590 Whole-cell biotransformation assays

591 Specific BVMO (cyclohexanone monooxygenase, EC 1.14.13.22) activity determination using the recombinant enzyme from Acidovorax sp. CHX100⁵³ and gas chromatography (GC) analysis of 592 product formation were performed as described elsewhere.⁵⁵ Briefly, different Synechocystis 593 594 strains were cultivated at standard conditions until reaching an OD₇₅₀ of ~1. Expression of *bvmo* 595 was induced using 10 µM NiSO₄ 24 h prior to the biotransformation. Therefore, 1 ml cell 596 suspension was adjusted to a cell dry weight (CDW) of 1 g_{CDW} l⁻¹ using a correlation factor of 597 0.225 g_{CDW} l⁻¹ for OD₇₅₀=1 as determined previously.⁸⁸ These samples were transferred into 10 ml 598 Pyrex® tubes (Pyrex®) and equilibrated at 30°C, 150 µmol photons m⁻² s⁻¹, and 200 rpm 10 min 599 before the assay. The assay was started by the supplementation of 3 mM cyclohexanone 600 substrate and stopped after 30 min by adding an equal volume of diethyl ether with 0.2 mM n-601 decane as internal standard. Samples for GC were taken in technical duplicates. One unit (U) is 602 defined as the production of 1 μ mol ϵ -caprolactone min⁻¹ normalized to q_{CDW} l⁻¹.

603

604 Supporting Information

Verification of different pSOMA plasmids in recombinant *Synechocystis* strains, table of used
oligonucleotides (SI File 1); GenBank files of the individual pSOMA plasmids (SI File 2)

607

608 Author contributions

609 S.K. designed the study. F.O., N.A.S., and S.K. generated the plasmids and performed 610 experiments. A.T., J.T., and B.B. contributed to BVMO activity determination by supplying non-611 published DNA constructs and experimental expertise for biotransformation assays. D.J.N. and 612 J.G.H. provided the experimental data for *Anabaena* sp. PCC 7120. F.O. & S.K. wrote the 613 manuscript with contributions from all co-authors. 614

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- 622
- 623 Conflicts of Interest: None.
- 624

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