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Letter

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Birth of a photosynthetic chassis: a MoClo toolkit enabling synthetic biology in the microalga Chlamydomonas reinhardtii

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1	Birth of a photosynthetic chassis: a MoClo toolkit enabling synthetic biology in the
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27 Abstract

28 Microalgae are regarded as promising organisms to develop innovative concepts based on 29 their photosynthetic capacity that offers more sustainable production than heterotrophic hosts. However, to realize their potential as green cell factories, a major challenge is to make 30 31 microalgae easier to engineer. A promising approach for rapid and predictable genetic 32 manipulation is to use standardized synthetic biology tools and workflows. To this end we have developed a Modular Cloning toolkit for the green microalga Chlamydomonas 33 reinhardtii. It is based on Golden Gate cloning with standard syntax, and comprises 119 34 35 openly distributed genetic parts, most of which have been functionally validated in several 36 strains. It contains promoters, UTRs, terminators, tags, reporters, antibiotic resistance genes, 37 and introns cloned in various positions to allow maximum modularity. The toolkit enables 38 rapid building of engineered cells for both fundamental research and algal biotechnology. 39 This work will make *Chlamydomonas* the next chassis for sustainable synthetic biology.

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41 Keywords: Algal biotechnology, *Chlamydomonas reinhardtii*, modular cloning, synthetic
42 biology.

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45	There is an urgent need to decarbonize the world economy due to depletion of fossil fuel
46	reserves coupled with accumulation of greenhouse gases produced by their combustion.
47	One alternative to the use of fossil fuels is to use photosynthetic microorganisms, such as
48	microalgae, as green cell factories to produce fuels and chemicals from atmospheric CO_2 in a
49	sustainable process driven by sunlight ^{1, 2} . The fixed carbon can be redirected towards
50	compounds that can be used in the fuel, food, cosmetic and pharmaceutical industries, such
51	as proteins, alcohols, alkanes, lipids, sugars, pigments or terpenes ³⁻⁵ . By contrast with land
52	plant-based photoproduction, microalgae do not compete with agriculture and can be grown
53	at high yields even at large scale ^{4, 6} , including on waste streams, thus minimizing inputs ³ . The
54	green microalga Chlamydomonas reinhardtii (referred to hereafter as "Chlamydomonas")
55	has been extensively engineered for basic research and industrial biotechnology ^{4, 6-8} . Its
56	nuclear and organellar genomes are sequenced and annotated, molecular biology
57	techniques and culture conditions are highly developed, and its physiology and metabolism
58	are well understood ⁹⁻¹³ . Moreover, the metabolic plasticity and cellular compartments of
59	Chlamydomonas offer great potential for advanced metabolic engineering strategies ^{14, 15} .
60	Chlamydomonas has already been engineered for production of the biodiesel precursor
61	bisabolene ⁸ , the terpene patchoulol ⁷ , and recombinant proteins as well as enzymes such as
62	an HIV antigen ¹⁶ and xylanase ¹⁷ . Despite these proofs of concept however, engineering of
63	Chlamydomonas is still slow due to a lack of standardized resources and tools ¹¹ .
64	Development of the field of algal synthetic biology offers the means to enable design and
65	construction of microalgal cells with defined and predictable properties ¹⁸ . Besides
66	biotechnological applications, the transition from empirical to synthetic approaches also
67	provides the opportunity to answer fundamental biological questions using new concepts
68	and approaches based on understanding by construction rather than deconstruction.

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Synthetic biology approaches, predicated on the Design-Build-Test-Learn cycle¹⁹, make organisms easier to engineer through the use of standardized parts and their assembly to simplify the building of designed DNA molecules¹⁹. Among available standards²⁰, the Golden Gate Modular Cloning (MoClo) technology, based on Type IIS restriction enzymes, offers extensive standardization and allows the assembly of complex multigenic DNA from basic gene parts (*e.g.* promoters, CDS, terminators) in just two steps^{21, 22}. The method accelerates and multiplies the possibilities to permute multiple genetic elements, and makes facile the building of multigene constructs for full metabolic pathways²³. MoClo is efficient and versatile, but relies on intensive upfront generation of a standardized library of basic building blocks, the gene parts, that have been domesticated to remove Type IIS sites, and codon optimized for the host as appropriate. MoClo toolkits have already been developed for a few model organisms $^{24-29}$ although not yet for microalgae.

Here, we report the generation of a MoClo toolkit composed of more than 100 gene parts codon-optimized for the Chlamydomonas nuclear genome. These genetic parts were designed to provide maximum modularity to end-users, and to facilitate the development of engineered strains for fundamental and green biotechnological applications, through iterative design and testing. We provide functional validation and characterization of many gene parts in several Chlamydomonas strains. This kit is available to the community, to allow Chlamydomonas to become the next chassis for sustainable synthetic biology approaches.

RESULTS

89 Standard and content of the Chlamydomonas MoClo kit

90 Standardization is the key to efficient building. The Chlamydomonas MoClo kit adopts the 91 syntax proposed by the plant synthetic biology community including the OpenPlant

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92	Consortium ³⁰ (Fig. 1). This syntax is defined for level 0 plasmids containing standard gene
93	parts (promoters, coding sequences, untranslated regions, etc.) and assigns strict fusion sites
94	for 10 cloning positions. In a single step, standardized parts can be assembled into modules
95	(Transcriptional Unit, TU, level 1) and modules into devices (multigenic construct, level M or
96	2) according to the original MoClo syntax ²² (Supplementary Fig. 1). Our Chlamydomonas
97	MoClo toolkit is composed of a set of 119 parts representing 67 unique genetic elements
98	available at different positions within the standard, thereby providing maximum modularity
99	to designers (Fig. 1, Fig. 2). The kit recapitulates most of the standard genetic elements
100	previously developed for Chlamydomonas which we "domesticated" by removing Bpil and
101	<i>Bsa</i> l restriction sites (the two enzymes used by the MoClo strategy ²² , Supplemental Figure 1)
102	from their sequences by DNA synthesis or PCR-based mutagenesis. The available gene parts
103	encompass 7 promoters coupled or not to their original 5'UTR, the corresponding 5'UTR and
104	the CrTHI4 riboswitch, 8 immunological or purification tags in positions leading to N- or C-
105	terminal translational fusions, 9 signal and targeting peptides, 12 reporters, 5 antibiotic
106	resistance genes, the foot and mouth virus (FMDV) 2A peptide which allows expression of
107	two or more proteins from a single transcriptional unit ^{17, 31, 32} , 2 micro RNA (miRNA)
108	backbones and associated controls, and six 3'UTR-terminators (Fig. 1b, Fig. 2 and
109	Supplementary Table 1). All sequences and plasmids are available through the public
110	Addgene repository (http://www.addgene.org/).

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Constitutive promoters and reporter genes

Five antibiotic resistance genes are used as selectable markers for Chlamydomonas but also can function as reporter genes^{33, 34}. We assembled three modules that allow control of the expression of the *aadA* gene, conferring spectinomycin resistance, by three constitutive

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promoters: P_{PSAD} and P_{BTUB2} with or without the first intron of $\beta TUB2$ (pCM1-1 to 3, Supplementary Table 3). The transformation efficiency of the three modules in UVM4³⁵ cells was estimated by counting spectinomycin resistant colonies and showed resistance frequencies within the same range (Fig. 3a). The presence of the first $\beta TUB2$ intron significantly increased the transformation efficiency as previously observed with the presence of RBCS2 introns in the ble marker^{33, 36, 37}. Alternative reporters are bioluminescent proteins, which allow more sensitive and quantitative analysis of gene expression. The kit contains Gaussia princeps luciferase, the brightest luciferase established in Chlamydomonas³⁸, as well as the redesigned Nanoluciferase (NanoLuc) which provides a stable and strong luminescence signal³⁹. Chlamydomonas NanoLuc was specifically developed for our MoClo kit through recoding to match the codon bias of Chlamydomonas, and cloned at 6 different positions within the standard. This new part was first tested with the most widely used promoter/terminator combination (P_{AR} promoter / T_{RBCS2} terminator) for strong constitutive expression in Chlamydomonas. The corresponding module (pCM1-04) was assembled with another module conferring paromomycin resistance (Supplementary Fig. 2) into a device (pCMM-1) that was introduced into the genome of the D66 strain (CC-4425, Fig. 3b). Among paromomycin resistant colonies, 34.8% ± 8.3 (N=48, mean ± SEM) were luminescent. The signal was variable between clones due to genomic position effects^{40,} ⁴¹ but was linear from 50 to 5 x 10⁵ cells (Fig. 3b and Supplementary Fig. 2). By contrast, non-expressing transformants (resistant to paromomycin only) or the D66 recipient strain displayed only a faint signal, 3 orders of magnitude lower, and saturating swiftly (Fig. 3b, inset). The modularity of the MoClo strategy allows rapid assessment of combinations of multiple parts. For example, we assembled 4 modules where NanoLuc expression is controlled by all possible combinations of the two most common constitutive promoters (P_{AB}

and P_{PSAD} and terminators (T_{RBCS2} and T_{PSAD}) (Fig. 3b, pCM1-4 to 7, Supplementary Table 3). Each module was assembled with the paromomycin resistance module (pCMM-1 to 4, Supplementary Table 4) and introduced into the Chlamydomonas genome. Bioluminescence levels were averaged over hundreds of transformants to account for the genome position effect^{40, 41}. The strengths of the two promoters were found to be comparable, whilst T_{PSAD} appeared to confer robust expression from both promoters, 10-fold higher than T_{RBCS2} (Fig. 3c). In a distinct context (strain, reporter sequence, culture conditions, etc.), the same genetic element may perform differently^{31, 35}. Such context sensitivity can be overcome by taking advantage of the modularity of the Chlamydomonas MoClo kit, which allows for the rapid characterization of all possible parts combinations. These results also confirmed the performance of the Chlamydomonas NanoLuc reporter and its employability for detailed understanding and characterization of genetic circuits especially if coupled with automated cell-sorting microfluidic devices⁴².

152 Control of gene expression

To build genetic circuits, the fine-tuning of gene expression is a prerequisite. Multiple parts enabling controlled gene expression have therefore been implemented. The activity of the P_{NIT1} promoter can be controlled by switching the nitrogen source since it is strongly repressed by ammonium and highly induced on nitrate^{34, 43, 44}. A module where P_{NIT1} controls expression of the ble-GFP gene (pCM1-8) conferred strong zeocin resistance in the CC-1690 strain but only when ammonium was replaced by nitrate as nitrogen source. By contrast, the P_{PSAD} promoter (pCM1-9) conferred strong antibiotic resistance on both nitrogen sources (Fig. 4a and Supplementary Fig. 3a-c). The vitamin B_{12} -repressible promoter P_{METE}^{45} allowed conditional functional complementation of the photosynthetic mutant nac2-26 (CC-4421),

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which lacks photosystem II due to the absence of the TPR-like protein NAC2 required for stability of the *psbD* mRNA encoding the D2 reaction center protein⁴⁶. *nac2-26* mutant cells engineered with a module harboring the *NAC2* coding sequence under the control of the P_{METE} promoter (pCM1-10) could grow photoautotrophically in the absence of vitamin B₁₂, but growth was compromised by increasing its concentration by amounts as low as 5 ng/L (Fig. 4b).

Regulation of gene expression can also be controlled by vitamin B_1 (thiamine) at the level of the transcript through riboswitches^{47, 48}. Binding of thiamine pyrophosphate to the THI4 riboswitch (RS) results in alternative splicing and retention of an 81 bp upstream open reading frame, ultimately interfering with translation^{47, 48}. The RS also responds when cells are grown in the presence of the thiamine biosynthetic intermediate 4-methyl-5-(2-hydroxyethyl) thiazole (HET), but not with 4-amino-5-hydroxymethyl-2-methylpyrimidine $(HMP)^{47}$. A module combining P_{AB} and THI4 (RS) to drive expression of the ble-GFP gene (pCM1-11) conferred conditional zeocin sensitivity in the UVM4 strain³⁵. Resistance was compromised by thiamine or HET but not HMP (Fig. 4c and Supplementary Fig. 3d), thereby demonstrating the efficient repression of the transgene through the *THI4* riboswitch.

Finally, to allow targeted repression of gene expression, a microRNA precursor sequence derived from the pre-miR1157 and used for the generation of artificial miRNAs (amiRNA)⁴⁹ was re-designed for compatibility with the Golden Gate cloning method. To demonstrate its effectiveness in driving gene repression, a specific amiRNA sequence directed against the MAA7 gene, whose repression provides resistance to 5'-fluoroindole (5'-Fl)⁵⁰, was inserted into the microRNA precursor. A control random sequence ("scrambled") amiRNA was inserted into the same backbone. These parts were placed under the control of P_{PSAD} and T_{PSAD} (pCM1-12 and 13) and assembled with a paromomycin resistance module (pCM1-27).

The same amiRNA sequences were introduced into the previously established pChlamiRNA3 vector⁴⁹ as controls. After transformation of the CC-1690 strain, 36% of paromomycin-resistant cells displayed resistance to 5'-FI with the device targeting MAA7 (pCMM-5) but not with the scrambled amiRNA (pCMM-6) (Fig. 4d and Supplementary Fig. 3f). A modified 5' rapid amplification of cDNA ends (5'-RACE) assay revealed that the MAA7 transcript was most frequently cleaved at a site corresponding to positions 10 and 11 of the amiRNA, as expected for a specific action of the miRNA (Fig. 4d). The properties of controllable parts can also be combined as shown for P_{NIT1} control of amiRNA-dependent gene repression³⁴. An amiRNA strategy recently proved useful for concerted metabolic engineering of a biodiesel precursor in Chlamydomonas⁸. The versatility of the MoClo kit opens new possibilities for sophisticated metabolic engineering strategies, *e.q.* the specific downregulation of up to six target genes with one level M assembly. Multiple fusion tags for detection and purification of gene products. Protein fusion tags are indispensable tools used to improve protein expression yields, enable

protein purification, and accelerate the characterization of protein structure and function⁵¹. Our MoClo kit includes multiple epitope and affinity tags known to be functional in Chlamydomonas. The modularity of the MoClo assembly allows rapid assessment of the best tagging strategy through a rapid design/build/test/learn cycle. We took advantage of the well characterized *rap2* mutant (Δ *FKBP12*), which is insensitive to rapamycin⁵², to test the functionality of five tags (Fig. 5a,b). We designed and built 5 devices allowing strong constitutive expression of N- or C-terminal tagged FKBP12 coupled to a paromomycin module (pCMM-7 to 11, Fig. 5c-h and Supplementary Table 4). The engineered strains were selected on paromomycin and the functionality of the fusion protein was tested by assessing

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sensitivity to rapamycin. Protein extracts were probed by immunoblotting using FKBP12-specific and tag-specific antibodies (Fig. 5d-h). All tags allowed detection (Fig. 5d-h) or purification (Fig. 5i) of FKBP12 even though some were not functional for restoring rapamycin sensitivity. The test revealed that pCMM-9 outperforms other devices since it provides a WT-like phenotype and expression level coupled to a strong and specific Myc signal with no significant processing of the protein. These results demonstrate the importance of the modularity provided by the Chlamydomonas MoClo toolkit for designing optimal fusion proteins.

Visualization and targeting of proteins in living cells

Fluorescent protein tags allow the temporal and spatial monitoring of dynamic expression patterns at cellular and subcellular scales⁵³. Natural and synthetic metabolic pathways can be optimized through spatial organization since cell compartments offer many advantages, such as isolation of metabolic reactions and generation of concentration gradients¹⁴. In a eukaryotic chassis like Chlamydomonas, organelles such as microbodies, mitochondria and chloroplasts can be engineered to implement or improve metabolic pathways¹⁵. The Chlamydomonas MoClo kit includes 11 targeting and signal peptides that allow the targeting of fusion proteins to mitochondria, chloroplast, nucleus, secretory pathway, ER and peroxisome-like microbodies. The functionality of the targeting and signal peptides and of the five fluorescent proteins (mVenus - yellow, mCherry - red, mRuby2 - red, Clover - green, mCerrulean3 - cyan) included in the toolkit was tested. Eight modules (pCM1-19 to 26, Supplementary Table 3) combining diverse fluorescent proteins and targeting sequences were assembled into devices with an antibiotic resistance module (pCMM-12 to 19). All devices were found to behave as expected and provided the expected fluorescent signal in

the targeted compartment (Fig. 6). The fluorescent and targeting parts of the Chlamydomonas MoClo toolkit, most of which have been validated here, enable engineering in the third dimension¹⁴ *i.e.* isolation and organization in multiple cellular compartments, and offer new tools for biological design/build/test cycles.

236 Discussion

The Chlamydomonas MoClo toolkit presented here provides more than 100 domesticated gene parts to allow advanced synthetic biology in microalgae. Numerous parts of multiple types have been characterized and validated in different genetic backgrounds¹⁰ and culture conditions, and can be readily used for biological design without further development. With the efficiency and modularity of the MoClo strategy, molecular cloning is no longer a limiting step for engineering Chlamydomonas cells. Indeed, from design to building, a complex device of up to six different genes/modules can be obtained within a week using the standardized parts provided in our kit. The modularity will also enable combinatorial assembly by shuffling part libraries⁵⁴ and determine *a posteriori* which combination is the most relevant. The development of gene-editing technologies in Chlamydomonas, including Zinc-finger nucleases^{55, 56} and several CRISPR-Cas9 approaches^{55, 57-59}, together with the development of high-throughput microfluidics⁴² are beginning to gather pace. Coupling these resources to our standardized MoClo toolkit will facilitate the use of Chlamydomonas as the photosynthetic chassis for innovative synthetic biology approaches aimed at fundamental and biotechnological applications. We expect that the creativity of designers, released from the time constraints associated with classical cloning strategies, will allow rapid expansion of the standard gene parts, modules and devices through open distribution, notably using the Addgene repository. We invite the community to openly share their parts

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through Addgene and/or our consortium (contact M. Schroda). The development of the 255 256 Chlamydomonas MoClo toolkit constitutes a complete step-change in the fields of microalgal biology and biotechnology. The parts developed for the MoClo toolkit may also be employed 257 in other microalgal species since the orthogonality of several Chlamydomonas 258 transcriptional units has been demonstrated in multiple hosts, including the industrially 259 relevant species Chlorella ellipsoidea, Nannochloropsis sp. and Dunaliella salina⁶⁰. Synthetic 260 approaches will allow engineering of microalgae in a predictable and efficient manner and 261 thereby offer great potential to couple environmental protection, energy transition and 262 bioeconomic growth⁴. 263

264

265 Methods

All chemicals were obtained from Sigma-Aldrich, unless otherwise specified.

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268 *Escherichia coli* and *Chlamydomonas reinhardtii* strains, transformation and growth 269 conditions.

Bacterial growth was performed at 37°C in LB broth supplemented with agar (20% m/V),
spectinomycin (50 μg/mL), ampicillin or carbenicillin (50 or 100 μg/mL, respectively) and Xgal (40 μg/mL) when required. Chemically competent *E. coli* DH10β (New England Biolabs)
were used for transformation (by heat shock following the manufacturer's instructions) and
maintenance of plasmids. All plasmids of the kit were maintained and amplified in TOP10 *E. coli* strain prior to submission to Addgene.

C. reinhardtii strains^{35, 46, 52, 61, 62}, culture and transformation conditions are recapitulated in
 Supplementary Table 5. They were grown in Tris-Acetate-Phosphate (TAP) medium⁶³
 supplemented with agar (1.6 % m/V), spectinomycin (100 μg/mL), paromomycin (15 μg/mL),

zeocin (ThermoFisher Scientific, 10 to 15 μ g/mL), 5-fluoroindole (20 μ M) or rapamycin (LC Laboratories, 1 µM) when required. For NIT1 promoter characterization (Figure 4a), a modified TAP medium lacking nitrogen source (TAP-N) was used instead, and was supplemented with 4 mM KNO₃ (nitrate) or 7.5 mM NH_4Cl (ammonium). For NAC2 autotrophy test (Figure 4b), cells were grown in minimal media (HSM) for selection of complemented strains. The responsiveness to B12 was assessed on plate and then in liquid. Cells were grown for 15 days in HSM until 1-5 x10⁷ cells/mL concentration prior to inoculation in a 96-well plate at a concentration of 10^5 cells/mL in 200 μ L of HSM. For response assays (Figure 4c) thiamine (Melford Laboratories Ltd.), 4-methyl-5-(2-hydroxyethyl) thiazole (HET) and 4-amino-5-hydroxymethyl-2-methylpyrimidine (HMP, Fluorochem UK) were added to TAP media at a final concentration of 10 μ M. For transformation by electroporation (see Supplementary Table 5), a TAP culture of $1-5 \times 10^{-5}$ 10⁶ cells/mL was concentrated 100 times in TAP complemented with 60 mM sucrose or the MAX Efficiency Transformation reagent for Algae (ThermoFisher scientific) and 25-250 µL were incubated with 80-300 ng of DNA for 10-30 min on ice in a 0.4 cm gapped cuvette (BioRad) prior to electroporation (BioRad Gene Pulser Xcell). The cells were then left to recover in TAP complemented with 40-60 mM sucrose for 16 h under appropriate light and shaking conditions (typically 50 μ mol photon m⁻² s⁻¹ at 100 rpm) prior to plating on TAP-agar plates with adapted antibiotics. Transformation by glass-beads method followed previously published protocols^{7, 64}. Briefly, after growth in TAP until 5 x 10^6 cells/mL, cells were concentrated 30 times and 5 x 10^7 cells were mixed with DNA using glass beads. After 2-fold

spectinomycin. Plates were incubated for 16 h in the dark prior to light exposition (30 µmol photon.m⁻².s⁻¹). When colony counting was performed (Figure 3a), it was 8 days after the

 dilution with TAP, 2.5 x 10^7 cells were spread onto TAP agar plates containing 100 µg ml⁻¹

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beginning of light. In both cases, the transformation protocol leads to insertion of a linearDNA in a random location within the nuclear genome.

Design. All *in silico* sequence designs and analysis were performed with Serial Cloner, 307 Benchling, SnapGene, ApE or Genome Compiler. For exogenous parts, reverse translation 308 was performed with Serial Cloner using *C. reinhardtii* nuclear genome codon frequency 309 (http://www.kazusa.or.jp/codon/cgi-bin/showcodon.cgi?species=3055).

amiRNAs can be generated using DNA parts pCM0-068 and pCM0-069. Both are derived from the endogenous pre-miR1157⁶⁵, but differ in the way in which the amiRNA specific sequence is introduced. pCM0-069/pCMM-20 is analogous to pChlamyRNA3⁶⁵, and a dsDNA fragment containing the amiRNA/loop/amiRNA* sequence is introduced into a Spel site inside the miRNA precursor sequence. pCM0-068 presents two divergently oriented Bpil sites, allowing the cloning of the dsDNA fragment by Golden Gate. In this last case, the dsDNA fragment is formed by the annealing of two oligos with the following sequence: 1) sense oligo (5' AGTA-(MIRNA*SEQ)-oligo (5'CAGT-A-(rev MIRNA SEQ)com TAGCGCTGATCACCACCACCCCCATGGTGCCGATCAGCGAGA-(rev com MIRNA*SEQ) 3'). There online tools that help with the design of the amiRNA sequence are (http://wmd3.weigelworld.org/cgi-bin/webapp.cgi).

Parts repository. All sequences listed in Supplementary Table 2 were deposited in Addgene.
Physical distribution of the DNA is performed through Addgene. We invite the community to

share their future parts through Addgene and/or with our consortium (contact M. Schroda)which will make them available to the community.

Parts cloning. All PCR reactions were performed using the Phusion DNA polymerase, KOD Xtreme Hot Start DNA polymerase (Merck) or Q5 DNA polymerase purchased from New England Biolabs (NEB) following the manufacturer's instructions adapted to GC-rich DNA, typically duration of hybridization and polymerization was doubled and/or GC enhancer solution was used. Molecular biology kits were purchased from Macherey-Nagel, peqLab, NEB or QIAgen (gel extraction and miniprep kits). Primers were produced by Eurofins Genomics or Sigma-Aldrich while synthesized parts were obtained from Genecust, DC Biosciences, IDTDNA or Sigma-Aldrich.

MoClo Assembly Conditions. All Restriction/ligation reactions were performed using *BbsI* or BbsI-HF (Bpil is an isoschizomer) or Bsal-HF (NEB or ThermoFisher) together with T4 ligase (NEB) in a medium containing the NEB CutSmart buffer and 1 mM ATP (with stock of 10 mM solubilized in 0.1 M Tris-HCl, pH 7.9). Typical ratio between destination plasmid (100 fmol) and entry plasmid/parts was 1:2. To facilitate handling of the kit for end-users, we provide detailed protocols and reaction mix calculators for each type of assembly: level 0 for parts (Supplementary Table 6), level 1 for modules (Supplementary Table 7) and level M for devices (Supplementary Table 8).

347 Quality Control of generated DNAs. All plasmids were controlled by differential restriction.
348 In addition, all level 0 plasmids were sequenced with specific primers. Sequencing was

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performed by Eurofins Genomics, Source BioSciences UK, Seqlab, Macrogen, Microsynth,
 GATC Biotech or Core Facility (CeBiTec, Bielefeld University).

NanoLuc activity determination. Reagents were purchased from Promega (ref. N1110) and activity was determined as previously described³⁹. For screening, *C. reinhardtii* colonies were transferred into a 96-well plate containing 100 µL of TAP in each well. After gentle resuspension, 50 µL was transferred into a solid white 96-well plate to which 50 µL of Nano-Glo substrate diluted in the provided buffer (2% V/V) was added and gently mixed by pipetting. Luminescence was measured at 460 nm with a CLARIOstar plate reader (BMG Labtech). For promoter/terminator combination assessment experiment (Figure 3c), all C. reinhardtii colonies from a transformation event were pooled and resuspended in TE buffer (50 mM Tris-HCl pH 7.9, 1 mM EDTA) complemented with anti-protease (1 tablet per 50 mL, Sigma-Aldrich: S8830). The cells were lysed by vortexing 10 seconds twice in the presence of glass beads (about 1:5 ratio beads/cells V/V) prior to two centrifugations (20000 g for 10 min at 4°C) to clarify the supernatant. The protein concentration was then determined using Bradford reagent with a Bovine Serum Albumin standard curve and the concentration was standardized to 0.5 g/L. The activity was determined in a 96-well plate in a final volume of 50 μL (1:1 with nano-Glo resuspended in provided lysis buffer) per well. NanoLuc activity was determined on 6 different increasing protein quantities (0.1 to 2.5 μ g) for each assay, allowing to assess linearity of the signal.

370 Absorbance measurement of cultures growing in microtiter plates

Growth in microtiter plates was determined by measuring the optical density of each well at
730 nm. Microtiter plates containing 180-200 μL culture were incubated under constant light

(125 μmol photon.m⁻².s⁻¹) at 25°C and 40 rpm orbital shaking. For density determination,
cultures were resuspended by pipetting and 100 μL of cell suspension was transferred to a
new microtiter plate containing 50 μL TAP 0.03% Tween-20. Optical density of each well was
determined at 730 nm in a CLARIOstar plate reader (BMG Labtech). Plates were shaken for
6-10 sec at 600 rpm before measurement.

379 RNA extraction and miRNA-mediated cleavage mapping

RNA isolation was carried out as previously described⁴⁹ (a detailed protocol can be found at http://www.plantsci.cam.ac.uk/research/davidbaulcombe/methods/downloads/smallrna.pd f/view), with the following modifications: Cells were centrifuged and resuspended in 0.25 mL of water and mixed with 0.25 mL Lysis buffer (50 mM Tris-HCl pH 8.0, 200 mM NaCl, 50 mM EDTA, 2% SDS, 1 mg/mL proteinase K). Lysis buffer was incubated at 50°C for 5 min prior mixing with cells. Cell suspension was then incubated at 25°C for 20 min. Finally, 2 mL of PureZol (Biorad) was added and samples were snap-frozen. RNA quality was assessed in gel and quantified in Nanodrop (ThermoFisher scientific).

miRNA cleavage site determination was performed as previously described⁶⁶. Briefly, 10 μ g of total RNA was ligated with an RNA oligo (5' CGACUGGAGCACGAGGACACUGACAUGGAC UGAAGGAGUAGAAA 3') using T4 RNA ligase for 1 h at 37°C. RNA was extracted with phenol:chloform and precipitated with ethanol and sodium acetate. The precipitated RNA was retrotranscribed into cDNA by SuperScript IV reverse transcriptase (ThermoFisher scientific), using random hexamers and following manufacturer's recommendations. Two µL of the cDNA was used as template of a PCR using primers FJN456 (5'-CGACTGGAGCACGAGGACACTGA) and FJN495 (5'- TGGGGTAGGGGTGGGGGCCAG). Two µL of this PCR was used as template of a second PCR with primers FJN457 (5'-

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397 GGACACTGACATGGACTGAAGGAGTA) and FJN496 (5'- TGACCCAGTCGCGGATGGCCT). PCR 398 was resolved in a 2% agarose gel and the specific band was isolated from the gel and cloned 399 into pGEM-T easy (Promega) for sequencing.

Immuno-blotting. Chlamydomonas cells expressing FKB12 fusion proteins from liquid
cultures were collected by centrifugation 4000 g for 5 min at room temperature (RT),
washed in 50 mM Tris-HCl pH 7.5, and resuspended in a minimal volume of the same
solution. Cells were lysed by two cycles of slow freezing to -80°C followed by thawing at RT.
The soluble cell extract was separated from the insoluble fraction by centrifugation (15000 g
for 20 min at 4°C). Total protein extracts (15 µg) were then subjected to 15% SDS-PAGE.

407 mCherry-expressing cells were harvested at 3500 rpm for 2 min (4 °C) and resuspended in 408 60 μ L of DTT-carbonate buffer (0.1 M DTT, 0.1 M Na₂CO₃). After freezing at -20 °C and 409 thawing, 55 μ L of SDS-Sucrose buffer were added (5 % SDS, 30 % sucrose). Samples were 410 then boiled for 45 s at 95 °C, followed by 2 min incubation on ice and 13000 g centrifugation 411 for 2 min at RT. Protein extracts corresponding to 2 μ g of Chlorophyll were then separated 412 using 12% SDS-PAGE.

For immunoblot analyses, proteins were then transferred to nitrocellulose membranes (Bio-Rad, 162-0115 or Amersham Protran). After blocking with 3 to 5% low-fat Milk in PBS for 1 h at RT, membranes were incubated with primary antibody in 5% low-fat Milk in PBS for 16 h at 4°C. After 4 washes in PBS - 0.1% tween-20 (TPBS), the membranes were incubated with secondary antibody in 5% low-fat Milk in PBS for 1 h at RT, and subsequently washed 4 time in TPBS prior to chemi-luminescence revelation using ECL. Primary antibodies used were anti-FKBP12 ⁵² (1/5000 dilution; secondary was anti-rabbit 1/10000), anti-FLAG (Sigma-Aldrich F1804, 1/5000 dilution; secondary was anti-mouse 1/5000), anti-STREP (IBA, Catalog

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> N. 2-1509-001, 1/5000 dilution; conjugated to HRP), anti-cMYC (Sigma-Aldrich M4439, 421 422 1/2500 dilution; secondary was anti-mouse 1/5000), anti-HA (Sigma-Aldrich H9658, 1/5000 dilution; secondary was anti-mouse 1/2500) and anti-PRPL1⁶⁷. For mCherry serum, rabbits 423 were immunized against purified full-length mCherry protein containing an N-terminal His₆-424 425 tag. 426 Microscopy. For mCherry experiments (Figure 6b-e), images were taken at 100x 427 magnification with a BX53F microscope (Olympus). Fluorescence images for the detection of 428 429 mCherry were taken using a TRITC filter. For other fluorescent proteins (Figure 6f-h), microscopy was performed as previously described ^{7, 68}. 430 431 Accession numbers. All parts accession numbers and the corresponding references are listed 432 433 in Supplementary Table 2. 434 435 ASSOCIATED CONTENT 436 Supporting Information. Supplementary Figure 1 - MoClo assembly workflow reflecting the abstraction hierarchy 437 438 Supplementary Figure 2 - Variability of Nanoluc expression in pCMM-1 transformants. 439 Supplementary Figure 3 - Control of gene expression, complementary data. Supplementary Table 1 - list of all unique parts of the Chlamy MoClo kit 440 441 Supplementary Table 2 - list of all parts of the Chlamy MoClo kit: level 0 plasmids 442 Supplementary Table 3 - list of all modules used for the Chlamy MoClo kit validation: level 1 plasmids 443 444 Supplementary Table 4 - list of all modules used for the Chlamy MoClo kit validation: level M plasmids 445

446 Supplementary Table 5 - list of Chlamydomonas reinhardtii strains and associated 447 transformations

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2 3	448	Supplementary Table 6 - level 0 ligation file: protocol and reaction mix calculator to clone
4	449	parts.
5	450	Supplementary Table 7 - level 1 ligation file: protocol and reaction mix calculator to
6 7	450 451	assemble modules.
8	451	assemble modules.
9	452	Supplementary Table 8 - level M ligation file: protocol and reaction mix calculator to assemble
10 11	453	devices.
12	454	
13	434	
14	455	Abbreviation.
15 16		
17	456	MoClo: Modular Cloning, TU: Transcriptional Unit, RBCS2: Ribulose Bisphosphate
18		
19	457	Carboxylase oxygenase Small subunit 2,HSP70: Heat Shock Protein 70, AR: HSP70A/RBCS2,
20		
21 22	458	TUB2: Tubulin 2, PSAD: Photosystem I reaction center subunit II, HET: 4-methyl-5-(2-
23		
24	459	hydroxyethyl) thiazole, HMP: 4-amino-5-hydroxymethyl-2-methylpyrimidine, amiRNA:
25	460	artificial micro RNA, TAP: Tris Acetate Phosphate
26 27	460	artificial micro RNA, TAP. This Acetate Phosphate
28	461	
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30	462	Author Information.
31 32		
33	463	K. Vavitsas current address is: Australian Institute for Bioengineering and Nanotechnology
34	464	(AIBN), The University of Queensland, Australia
35 36		
30 37	465	
38	100	Author Contribution
39	466	Author Contribution.
40 41	467	SDL, AGS, MS, PEJ, OK, JLC and GP created the consortium that led this study.
42	107	
43	468	PC, FJN, FW, PM, DCB, GP, JLC, OK, PEJ, MS, AGS and SDL designed the study and wrote the
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45 46	469	manuscript.
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48	470	PC, FJN, FW, PM, KB, KJL, MEPP, PA, AGR, SSG, JN, BS, JT, RT, LW, KV, TB, KS, MC, FdC, AD,
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50 51	471	MdM, JH, WH, CHM designed parts, modules and devices, performed the experiments,
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53	472	and/or analyzed data.
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474 Competing Financial Interests statement.

- The authors declare no competing financial interest.
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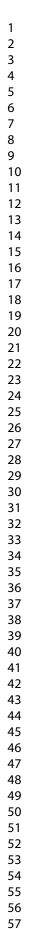
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702 Graphical Abstract

Design — ➤ Build > Test 119 PARTS (I) MODULES MoClo ---Harb, CALSHOLD BAD



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visual⁶⁹ representation described in **b**. Each of the 11 fusion sites defining a part position is represented with a

(b) Table summarizing unique and total gene parts available. The SBOL2.0 symbols are indicated for each type.

color and a number. Positions presented are representative of the whole set of each part type. Parts in

italicized letters are non-transcribed, parts in regular letters are transcribed and parts in bold letters are

When the SBOL2.0 standard was not existing for a part type, the symbol proposed before²⁸ was used, or

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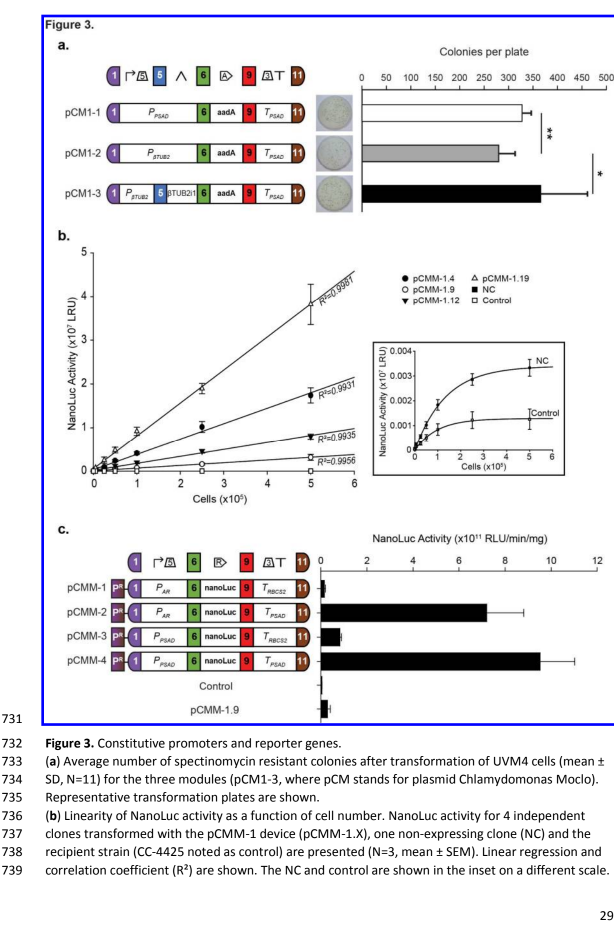
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а		Promot	ter	5'UTR	5'UTR/CD	S	CDS			3'UTR +	Ter
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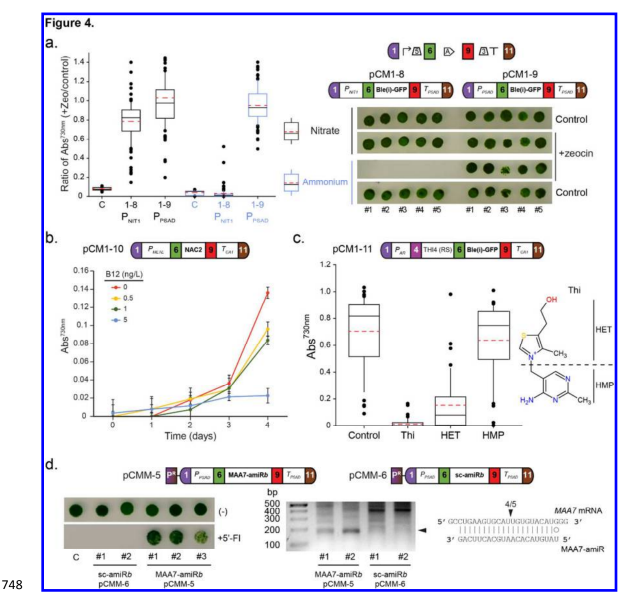
717 Figure 2. List of parts in function of their type and assembly position.

718 (a) Plant MoClo syntax³⁰ indicating the color code for fusion sites used in this figure.

(b) All parts in the Chlamydomonas MoClo kit are classified primarily by their function, indicated by 719 SBOL2.0 visual code⁶⁹ as in Fig. 1 (from left to right: promoters, promoter+5'UTR, 5'UTR, introns, 720 721 antibiotic resistance genes, reporter genes, artificial microRNA, immunological and purification tags, 722 2A peptide, and 3'UTR+terminators). Colored stripes on the left and right sides of each box represent the fusion sites in 5' and 3' of the position, respectively, and follow the color code on top. AR and A'R 723 stand for HSP70A/RBCS2 and HSP70A467/RBCS2, respectively³⁶. A star (*) indicates that the part 724 contains extra restriction sites as in pOpt vectors⁶⁸ while the same part unmarked does not. An (i) 725 726 indicates the presence of an intron within the part (cf. Supplementary Table 2). For amiRNA (amiR) 727 backbones, b and s mean that Bpil and Spel site are within the backbone for amiR cloning, 728 respectively, while M and sc mean that the target amiR sequence for MAA7 and the control 729 scrambled sequence were introduced into the miR1157 backbone, respectively (cf. Fig. 4). mSTOP 730 stands for multi-STOP.



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3	740	(c) Average NanoLuc activity of D66 (CC-4425) cells transformed with 4 devices (pCMM-1 to 4)
4	741	harboring promoter/terminator combinations to drive NanoLuc expression coupled to a
5 6	742	paromomycin resistance module (represented as P ^R , left, Supplementary Fig. 2). Luminescence levels
7	743	are represented as mean ± SEM (average of a total of more than 400 clones from 3 biological
8	744	replicates). The negative and positive controls are the recipient strain and the pCMM-1.9 strain
9	745	(shown in b), respectively.
10	746	a,c *p<0.05; **p< 0.01 assessed by Student's t-test, SBOL2.0 ⁶⁹ visual of module designs are shown
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749 Figure 4. Control of gene expression.

(a) Control of gene expression by the nitrogen source. Zeocin resistant colonies (conferred by Ble(i)-GFP) selected after transformation of CC-1690 cells with each of the two represented modules ("1-8" for pCM1-8 and "1-9" for pCM1-9) were grown in TAP-nitrogen ± zeocin (15 µg/mL) supplemented with either 7.5 mM (NH₄)Cl (ammonium, blue) or 4 mM KNO₃ (nitrate, black) and their growth was followed (Absorbance at 730 nm). The plot shows the ratio between the growth in the presence and absence of zeocin (C is the non-transformed CC-1690 strain). The right panel shows cells grown in similar conditions but on solid media. Results presented (N=16 for control CC-1690 and N=86 for each other conditions) correspond to one out of three independent transformations (for the other two, see Supplementary Fig. 3). (b) Control of gene expression by vitamin B_{12} . Conditional complementation of *nac2-26* cells with the pCM1-10

759 module expressing NAC2 under P_{METE} control. Complemented strains were selected for photoautotrophic 760 growth on solid minimal medium and the cells were grown in liquid minimal medium supplemented with the 761 indicated amount of vitamin B₁₂. Data are mean ± SD (N=3).

(c) Control of gene expression by vitamin B₁. Average growth (absorbance at 730 nm after 7 days of growth, N=40) of UVM4 cells transformed with the pCM1-11 module designed to express constitutively *Ble(i)-GFP* transcripts containing the *THI4* riboswitch in the 5'UTR. After culture in TAP, the cells were transferred to

- 56 765 TAP+zeocin (10 μ g/mL) supplemented with 10 μ M thiamine (Thi), 10 μ M 4-methyl-5-(2-hydroxyethyl) thiazole

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3	766 767	(HET) or 10 μ M 4-amino-5-hydroxymethyl-2-methylpyrimidine (HMP) or not (control). The chemical structure
4	767	of Thi is represented on the right and the HET and HMP moieties are indicated (See also Supplementary Fig. 3). (d) Targeted gene knockdown with artificial miRNA. Paromomycin resistant cells selected after transformation
5	768	of CC-1690 cells by each of the two represented devices (pCMM-5 and pCMM-6), carrying an amiRNA cloned
6 7	770	with <i>Bpi</i> l and directed against <i>MAA7</i> (MAA7-amiRb) or a random sequence ('scrambled': sc-amiRb), were
8	771	grown in the absence (denoted (-)) or presence of 5'-fluoroindole (+5'-Fl) (left panel). C indicates non-
9	772	transformed cells. Clones resistant to 5'FI were analyzed by a modified 5'-RACE assay. A specific 173 bp PCR
10	773	band (black arrowhead) was amplified only from the 5'-FI resistant transformants and not from ones expressing
11	774	the amiRNA with scrambled sequence (middle panel and Supplementary Fig. 3). Sequencing revealed that the
12	775	most frequent cleavage occurred at positions opposed to positions 10 and 11 of the amiRNA (right panel, black
13	776	arrowhead). P ^R represents the paromomycin resistance module (pCM1-27, Supplementary Fig. 2a).
14	777	a, c The box and whisker plots show the 10 th (lower whisker), 25 th (base of box), 75 th (top of box) and 90 th (top
15	778	whisker) percentiles. The line within the box is the median, the dashed red line is the mean. Outliers are
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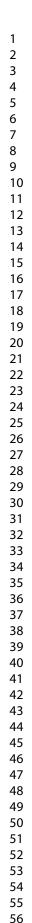
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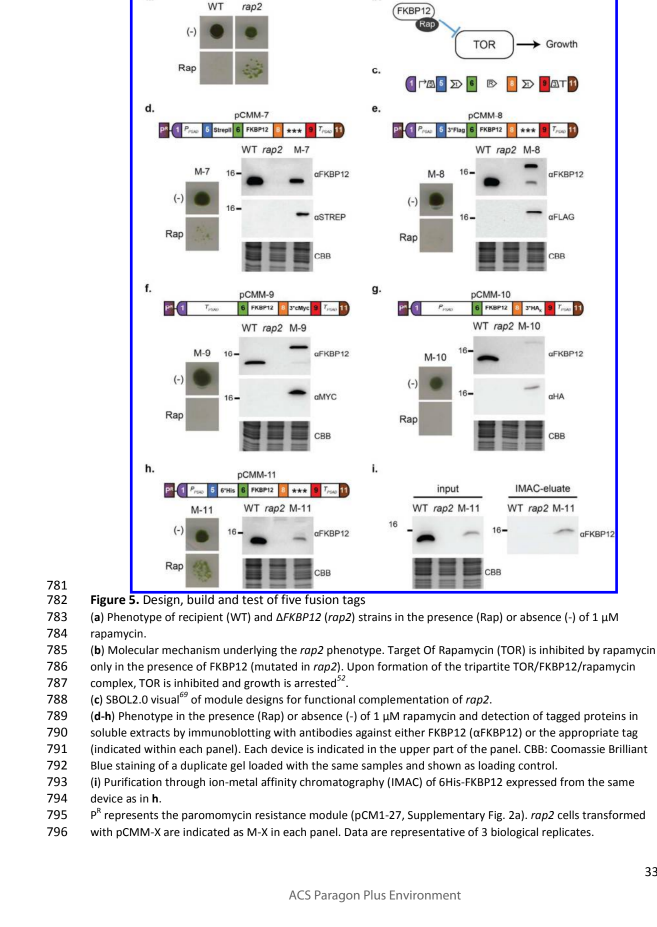
Figure 5.

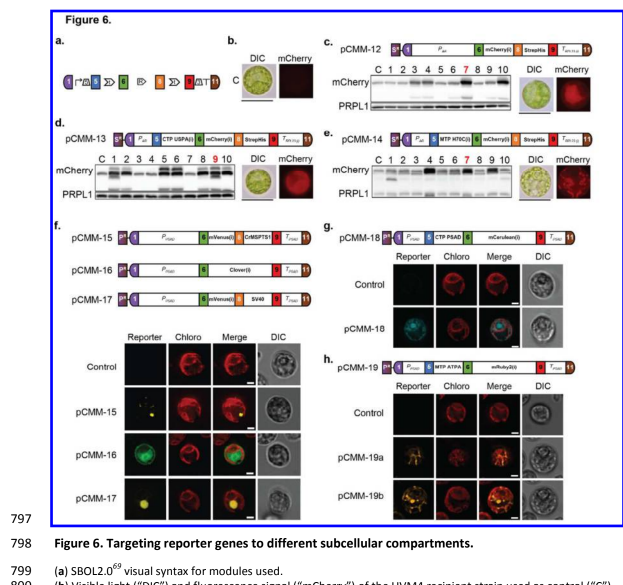
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800 (b) Visible light ("DIC") and fluorescence signal ("mCherry") of the UVM4 recipient strain used as control ("C")
801 for panels c-e.

(c-e) mCherry targeting to the (c) cytosol with no transit peptide, (d) chloroplast with CTP USPA (Chloroplast Transit Peptide of Universal Stress Protein A) or (e) mitochondria with MTP H70C (Mitochondrial Transit Peptide of HSP70C) in UVM4 cells transformed with the indicated devices (pCMM-12 to 14). In each panel, an anti-mCherry immunoblot analysis of transformants is shown. Note that the anti-mCherry antibody cross-reacts with a protein of similar size present in control cells (C). An anti-PRPL1 immunoblot is shown as loading control. The transformant strain number indicated in red corresponds to the images (bars are $10 \,\mu m$) presented on the right. (f-h) Fluorescent marking of (f) microbodies with mVenus-CrMSPTS1 (Malate Synthase PTS1-like sequence), cytosol with Clover or the nucleus with mVenus-SV40 (Simian Virus 40 nuclear localization signal), (g) the chloroplast with CTP PSAD-mCerulean (Chloroplast Transit Peptide of PSAD), (h) mitochondria with MTP ATPA-mRuby2 (Mitochondrial Transit Peptide of ATPA) after transformation of UVM4 cells with the indicated devices (pCMM-15 to 19). Images of representative transformants are grouped with the corresponding control image

- 51 (recipient strain) according to the filter used. pCMM-19a and pCMM-19b show two images taken on different 52 915 The State Provide The
- 52 815 z-axis on the same cell. "Chloro" refers to chlorophyll autofluorescence. The Scale bars represent 2 μ m. 53 816 S^R and P^R represent respectively modules conferring resistance to spectinomycin (S^R=pCM1-1, Fig. 3a and
- 817 Supplementary Fig. 2a) and paromomycin (P^R=pCM1-27, Supplementary Fig. 2a).