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Chromosomal integration of the pSOL1 megaplasmid of *Clostridium acetobutylicum* for continuous and stable advanced biofuels production

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Muhammad Ehsaan^{1,3}, Minyeong Yoo^{1,3}, Wouter Kuit \mathbb{O}^1 , Céline Foulquier², Philippe Soucaille $\mathbb{O}^{12} \boxtimes \&$ Nigel P. Minton \mathbb{O}^1

Biofuel production by *Clostridium acetobutylicum* is compromised by strain degeneration due to loss of its pSOL1 megaplasmid. Here we used engineering biology to stably integrate pSOL1 into the chromosome together with a synthetic isopropanol pathway. In a membrane bioreactor continuously fed with glucose mineral medium, the final strain produced advanced biofuels, *n*-butanol and isopropanol, at high yield (0.31 g g⁻¹), titre (15.4 g l⁻¹) and productivity (15.5 g l⁻¹h⁻¹) without degeneration.

The Weizmann process for acetone and *n*-butanol production by Clostridium acetobutylicum was the second largest fermentation process (after ethanol) and of considerable industrial, social and historical importance¹⁻³. Beyond its use during the First World War to produce acetone for smokeless gunpowder (cordite) manufacture, it was used worldwide to produce these two industrial solvents from a variety of renewable substrate³⁻⁵. Its demise in the early 1960s was a consequence of superior petrochemical-process economics. The fermentation process suffered from low yield, titre and productivity⁶ and, unlike rival petrochemical processes, was not suited to continuous-process technologies due to the loss of C. acetobutylicum's capacity to produce solvents, known as 'degeneration'. This was associated with loss of the pSOL1 megaplasmid carrying genes essential to solvent production^{7,8}. Today, there is a resurgence of interest in C. acetobutylicum for the production of advanced fuels (1) after chemical transformation of the solvents mixture^{9,10} or (2) directly as isopropanol–n-butanol–ethanol (IBE) mixtures^{11,12}.

Here we successfully addressed the issue of strain degeneration in the Weizmann process by using an innovative synthetic biology approach. Our method involved integrating pSOL1 into the chromosome of *C. acetobutylicum* at the *pyrE* locus of an allele-coupled exchange (ACE)¹³ compatible strain with a deletion in the 5' region of *pyrE* (Fig. 1a).

This strategy involves four crossing-over events and the use of an ACE replicative plasmid¹³ introduced by electroporation (Supplementary Note 1) and containing the following: (1) homologous regions flanking the origin of replication of pSOL1 (repA), (2) a truncated pyrE-hydA locus in reverse orientation with a full-length but promoterless ermB gene that becomes functional only after the initial crossing-over of SHA (CA PO176) with pSOL1 and (3) a functional codA gene. The first double crossing-over occurs between the two homologous regions surrounding the origin of replication of pSOL1 also present on the ACE plasmid. Each homology arm size was specifically designed to control the integration order. As a result, the engineered strain contains a pSOL1 plasmid lacking an origin of replication but having a functional ermB now expressed under the natural CA PO176 promoter on pSOL1. However, this pSOL1 intermediate cannot replicate and is therefore unstable. Consequently, a second double crossing-over event at the pyrE locus is necessary to ensure its maintenance through integration (Fig. 1a), as the mutant having pSOL1 integration is selected on a uracil-lacking Clostridium basal medium (CBM) requiring functional pyrE restoration. Thus, mutants having a clean integration of pSOL1 after four crossing-over events (Fig. 1a) were directly obtained by spreading 10° of ACE plasmid containing cells on a CBM agar lacking uracil and supplemented with erythromycin. Colonies obtained were then replica plated on CBM agar containing erythromycin and 5-fluorocytosine (5FC) (Supplementary Note 2). Resistance to 5FC confirms that all the clones were cured of the original ACE plasmid, as well as the hybrid ACE plasmid with two origins of replication that was generated after the first double crossing-over event. The accurate

¹BBSRC/EPSRC Synthetic Biology Research Centre, School of Life Sciences, Biodiscovery Institute, University of Nottingham, Nottingham, UK. ²TBI, Université de Toulouse, CNRS, INRAE, INSA, Toulouse, France. ³These authors contributed equally: Muhammad Ehsaan, Minyeong Yoo. © e-mail: soucaille@insa-toulouse.fr

12

6

З

40

45

35

Biomass (g l⁻¹)

Biomass (g l⁻¹)

Butanol Butyrate Biomass

Degen



Fig. 1 | Construction of pSOL1 megaplasmid integrated C. acetobutylicum producing solvent stably in continuous culture. a, Schematic illustration of pSOL1 integrated strain by using original synthetic approach based on counter selection markers and antibiotic markers selection. Genes in blue boxes are originally located on the chromosome, whereas genes in green boxes are originally located on pSOL1. The black triangle in the gene box presents

genotype and phenotype of the mutants were validated through PCR and growth in batch cultures. The resultant CAB2018 strain showed the correct PCR and product profiles in batch culture, as well as an equivalent sporulation phenotype to the wild-type strain (Supplementary Note 4). Its specific growth rate, however, was around 30% lower than that of the wild-type strain (Supplementary Note 4). Although we have no explanation for this phenotype, it could explain why pSOL1 did not integrate evolutionarily into the chromosome of C. acetobutylicum. Whole-genome sequencing of the strain confirmed that pSOL1 had integrated as expected and that four single-nucleotide polymorphisms in non-relevant chromosomal genes had been acquired (Supplementary Table 2). To the best of our knowledge, we have not found any study reporting the stable integration of such a big piece of DNA into the chromosome of any bacteria. In Extended Data Fig. 1, we proposed a similar strategy for the integration of any large pieces of DNA into the chromosome of the CAB2019 strain, a CAB2018-derived strain with pyrE truncated and ermB removed (Supplementary Note 3) using the following: (1) an engineered yeast artificial chromosome (YAC) vector with the origin of replication of pSOL1, (2) a yeast strain to assemble the large synthetic DNA insert and (3) protoplast fusion to introduce the large vector into the CAB2019 strain. This extension of the method will be of particular value when the introduction of complex metabolic pathways comprising numerous genes is required, for example, the Wood-Ljungdahl pathway for autotrophic growth¹⁴. Finally, from a fundamental point of view, one of the benefits of having pSOL1 integration into the chromosome is that it now allows the rapid functional analysis of the pSOL1-encoded genes using CRISPR-Cas9 for gene editing¹⁵.

5' truncation, while the red triangle presents 3' truncation. P below the black arrow indicates the position of the natural promoter of CA P0176. b, Product concentration and degeneration rate of wild-type and pSOL1 integrated strains in continuous cultures under the same conditions. The butanol and butyrate concentrations and degeneration rates are plotted on the left y axis, with biomass plotted on the right y axis.

30

35 40 45

Monitoring of continuous cultures of the wild-type and CAB2018 strains showed that whereas the former degenerated within 30 days, losing the ability to produce solvents concomitant with loss of pSOL1. the latter strain remained stable over the 42 day fermentation period (Fig. 1b). Having a stable strain that produces *n*-butanol in continuous cultures is a prerequisite, but alone is not sufficient, for a commercial biofuel production process as both the yields, titres and productivities are too low due to production of acetone, a natural final product that is not suitable as a biofuel.

Accordingly, to improve the strain for continuous biofuels production, we further engineered the strain to continuously produce an advanced IBE fuel mixture at high practical yield and productivity. This was achieved in three steps involving two intermediate strains (CAB2019 and CAB2020) and the final CAB2021strain with integration of sadh and hydG genes (Fig. 2a and Supplementary Note 3) of Clostridium beijerinckii NRRL B593¹⁶ under the control of the natural thlA promoter. These encode a reduced nicotinamide adenine dinucleotide phosphate (NADPH)-dependent acetone reductase (sadh) and a potential ferredoxin-NADP⁺ reductase (hydG). In batch culture, the CAB2021 strain almost completely converted acetone to isopropanol (Supplementary Note 5). Furthermore, when grown in chemostat cultures at different dilution rates on a glucose mineral media (GMM), it was possible to produce biofuels at high yield (0.3 g s^{-1}) and productivity $(1.3 \text{ g l}^{-1} \text{ h}^{-1})$ for the highest dilution rate evaluated, although under this condition high residual glucose concentrations were observed (Fig. 2b and Extended Data Fig. 2). The best compromise between residual glucose, titre and productivity



Fig. 2 | Synthetic biofuel pathway integration into pSOL1 integrated strain and stable solvent production in GMM in continuous cultures of CAB2021 strain. a, sadh (encoding an NADPH)-dependent acetone reductase) and hydG (encoding putative ferredoxin-NADP+ reductase) of C. beijerinckii NRRL B593 for conversion of acetone to isopropanol were integrated downstream of thlA to be under the strong thlA promoter in CAB2021. b, Isopropanol, butanol, productivity and yield of four different dilution rates in continuous chemostat

culture of CAB2021. Biofuel is defined as (isopropanol + butanol + ethanol). D, dilution rate (h^{-1}) . c, Continuous production of advanced biofuel by CAB2021 in membrane cell-recycle bioreactor. Same symbols are used as in **b**. d, Membrane cell-recycle bioreactor used in this study. Data are shown as mean and s.d. of steady state period (day 4 to day 10, n = 12). DCW, dry cell weight (g); D_{i} , incoming dilution rate (h⁻¹); D_{o} , outcoming dilution rate (h⁻¹).

was observed at a dilution rate of 0.075 h⁻¹ with a biofuel titre of 13.4 g l^{-1} , a productivity of 1 g l^{-1} h⁻¹ and a yield of 0.31 g g^{-1} . These values were much higher than those obtained with C. beijerinckii NRRL B593, the natural producer, or a metabolically engineered mutant of *C. acetobutylicum* expressing *sadh* inserted at the *pyrE* locus¹⁷, when grown in chemostat cultures (Supplementary Table 4). Furthermore, contrary to the natural producer that loses its ability to produce solvent after 10 to 20 days in continuous cultures¹⁸, no instability was observed with CAB2021 for at least 30 days (Fig. 2b). Finally, to evaluate the potential of strain CAB2021 for industrial production of advanced biofuels, we used it in a membrane cell-recycle bioreactor system (Fig. 2c,d) to increase the cell density and then the productivities¹⁹. In GMM, the performances obtained in terms of yield (0.31 g s^{-1}) , titre (15.4 g l⁻¹) and productivities (15.5 g l⁻¹ h⁻¹) were the highest ever reported for the continuous production of an advanced IBE fuel mixture^{17,19-25} (Fig. 2d and Supplementary Table 4). We recently used an advanced extractive fermentation process to improve the performances of a metabolically engineered C. acetobutylicum optimised for *n*-butanol production at high yield²⁶. This process, using vacuum distillation and high cell density culture, could be used with the CAB2021 strain developed here to further increase the titre of the advanced biofuel mixture produced.

In summary, engineering biology was used to develop a C. acetobutylicum strain for the stable and continuous production of advanced biofuels from a low-cost GMM at higher yield and productivities than previously reported. To further develop an economical industrial process, performances still need to be improved. This might be achievable by the following: (1) further metabolic engineering to remove by-product pathway such as butyrate^{11,27} to increase the biofuels yield and/or (2) optimisation of the bioreactor by using extractive fermentation²⁶.

Methods

Reagents

All chemicals used in this study were purchased from Sigma-Aldrich unless otherwise noted. Q5 High-Fidelity DNA Polymerase (NEB) or KOD DNA Polymerase (Merck) was used for PCR. All PCR products and plasmids were purified and extracted with the PCR purification kit and plasmid extraction kit (Qiagen), respectively. All oligonucleotides were purchased from Sigma-Aldrich or Integrated DNA Technologies. DNA sequencing was performed at Eurofins. Genomic DNA of CAB2018 strain was isolated by phenol:chloroform extraction based on the method of Marmur²⁸. After measuring (1) the amount of genomic DNA with a NanoDrop lite spectrophotometer (Thermo Scientific) and (2) its quality via agarose gel electrophoresis, whole-genome sequencing was performed using an Illumina MiSeq benchtop sequencer (Deepseq, University of Nottingham). Sequence reads were mapped to the reference sequences NC_003030 (chromosome) and NC_001988 (pSOL1) in the NCBI database using the program CLC Genomics Workbench version 22.0.1 (Qiagen).

Bacterial strains and medium composition

Bacterial strains and plasmids used in this study are detailed in Supplementary Table 1. Escherichia coli TOP10 (Invitrogen) was cultured aerobically (37 °C; shaking at 200 r.p.m.) in Luria-Bertani medium supplemented with chloramphenicol (25 µg ml⁻¹) and tetracycline $(10 \ \mu g \ ml^{-1})$ where appropriate for plasmid cloning and propagation, and TOP10 containing pAN2 plasmid was used for in vivo methylation of plasmid DNA before transformation of C. acetobutylicum ATCC 824 and any other recombinant strains (Supplementary note 1). C. acetobutylicum strains were routinely grown anaerobically at 37 °C using pre-reduced overnight CBM or Clostridium Growth Medium (CGM) agar supplemented with thiamphenicol (15 μ g ml⁻¹) or erythromycin $(20 \,\mu g \,m l^{-1})$ where appropriate under an atmosphere of N₂:H₂:CO₂ (80:10:10, vol:vol) in an anaerobic workstation (Don Whitley) or in anoxic broth in a serum bottle. Recovery after transformation was carried out in anoxic 2×YTG (pH 5.2) broth in the anaerobic workstation. Uracil was supplemented at 20 µg ml⁻¹ when needed. 5FC was supplemented at a final concentration of 100 µg ml⁻¹ where *codA*-based selection was performed. C. acetobutylicum strain degeneration was evaluated in chemostat culture in Clostridium Rich Media (CRM). Performances of the engineered C. acetobutylicum strains both in chemostat and in a membrane cell-recycle bioreactor were evaluated in GMM.

The CBM contains (per litre) 50 g glucose, 0.5 g K_2HPO_4 , 0.5 g KH_2PO_4 , 0.2 g MgSO₄·7H₂O, 7.58 mg MnSO₄·H₂O, 0.01 g FeSO₄·7H₂O, 1 mg para-aminobenzoic acid (PABA), 0.002 mg biotin, 1 mg thiamine HCl, 4 g casein hydrolysate and 5 g CaCO₃ if needed as buffering agent. The CGM contains (per litre) 0.75 g KH_2PO_4 , 0.75 g K_2HPO_4 , 0.4 g MgSO₄·7H₂O, 0.01 g MnSO₄·H₂O, 0.01 g FeSO₄·7H₂O, 1 g NaCl, 2 g asparagine, 5 g yeast extract, 2 g (NH4)₂SO₄ and 60 g glucose. The 2×YTG medium contains (per litre) 16 g tryptone, 10 g yeast extract, 5 g NaCl and 10 g glucose. The CRM contains (per litre) 50 g glucose, 4 g yeast extract, 0.5 g K_2HPO_4 , 0.5 g KH_2PO_4 , 0.2 g MgSO₄·7H₂O, 10 mg MnSO₄·H₂O, 80 µg biotin and 8 mg PABA. The GMM medium contains (per litre) 50 g glucose, 0.5 g K_2HPO_4 , 0.5 g KH_2PO_4 , 1.5 g NH_4 Cl, 0.2 g $MgSO_4$ ·7H₂O, 10 mg FeSO₄·7H₂O, 80 µg biotin and 8 mg PABA.

Plasmids construction

Plasmid pMTL-pSOL1-int (Fig. 1, Supplementary Table 1 and Supplementary Fig. 1) was constructed using plasmid pMTL-SC7515 carrying codA, repL and catP²⁹ as a backbone. This vector was designed to contain four different sizes of homology arms: (1) 1,200 bp of CA P0173-4 homology arm (LHA, long homology arm for first crossing-over with the pSOL1), (2) 305 bp of CA_PO176 homology arm (SHA, short homology arm for second crossing-over with pSOL1 to excise replication origin repL encoded by CA_P0175 of pSOL1 and to introduce 3' truncated pyrE and 3' truncated hydA into pSOL1), (3) 900 bp of hydA homology arm (LHA for third crossing-over with chromosome) and (4) 635 bp of pyrE homology arm (40 bp from start codon truncated SHA for fourth crossing-over with 3' truncated pyrE carrying chromosome to restore full-length functional pyrE at the last step). As previously reported¹³, the use of asymmetrical homology arm sizes can control the order of recombination events, allowing the initial isolation of single crossing-over integrants involving the LHAs followed by the selection of recombinants arising from subsequent, double crossing-over, excision events involving the SHAs. The cassette that consists of four homology arms and promoterless *ermB* gene was synthesized (Biomatik) in the order of a 1,200 bp LHA *CA_P0173-4* for pSOL1 integration, terminator, reverse-oriented 900 bp *hydA* LHA for chromosome integration, reverse-oriented 635 bp of *pyrE* SHA for chromosome integration, terminator, *ermB* with its ribosome binding site but without promoter to be active after integration and 305 bp of *CA_P0176* SHA. This cassette was cloned at the Pmel site of pMTL-SC7515 to construct pMTL-pSOL1-int. Plasmid pMTL-pSOL1-int was used to construct the CAB2018 strain.

Plasmid pMTL-pSOL1-*pyrE*-Negative was constructed to remove (1) the erythromycin resistance gene *ermB* and (2) 335 bp of the *pyrE* gene of CAB2018 by double crossing-over.

Previously constructed plasmid pMTL-JH12¹³, which was made for *pyrE 5'* truncation, was used as a backbone. The LHA (*hydA* fragment in pMTL-JH12) was replaced by the PCR fragment for *CA_P0176-77* homology by using primers Pr1 and Pr2, whereas the remaining part, including the SHA (partial *pyrE* for truncation), *lacZ* (MCS) and the antibiotic resistance marker *catP* in the original plasmid was retained. DNA fragment replacement was carried by enzyme digestions with Nhel/AscI (NEB) and ligation with T4 DNA ligase (NEB). Plasmid pMTL-pSOL1-*pyrE*-Negative was used to construct the CAB2019 strain.

Plasmid pMTL-pSOL1-*pyrE*-Positive was constructed to restore the deleted 335 bp *pyrE* fragment in CAB2019 for a full-length functional *pyrE*.

To restore functional *pyrE*, a synthetic DNA fragment for full-length *pyrE*–a Rho-independent terminator–*CA_P0176-77* ended with Sbfl/Ascl (Biomatak) was cloned into Sbfl/Ascl (NEB) digested pMTL-pSOL1-*pyrE*-Negative plasmid to replace partial *pyrE-lacZ* fragment. Plasmid pMTL-pSOL1-*pyrE*-Positive was used to construct the CAB2020 strain.

Plasmid pMTL-JH16-*sadh-hydG* B593 was constructed to integrate a synthetic isopropanol production pathway into the CAB2020 strain for stable and continuous production of advanced biofuels. It carries part of *thlA* (thiolase encoding gene, left SHA for recombination to maintain strong expression of inserted genes under *thlA* promoter throughout growth), *ermB* (erythromycin, antibiotic resistance marker for double crossing-over selection) and heterologous *sadh* (encoding a primary/secondary alcohol dehydrogenase) and *hydG* (encoding a putative electron transfer protein) genes from *C. beijerinckii* NRRL B593. The *sadh-hydG* DNA fragment, amplified by using primers Pr3 and Pr4, was cloned into Notl/Nhel (NEB, UK) digested plasmid pMTL-JH16¹³. Plasmid pMTL-JH16-*sadh-hydG* B593 was used to construct the CAB2021 strain.

Cultivation conditions

All liquid cultures of *C. acetobutylicum* strains were performed in 30 or 60 ml serum bottles under strict anaerobic conditions in CGM or GMM in which glucose concentration was 60 g l^{-1} and NH₄Cl was replaced by ammonium acetate at 2.2 g l^{-1} . All serum bottle batch culture data are shown as mean ± s.d. from biological replicates (n = 3). Spores were germinated by immersing the serum bottles in a water bath at 80 °C for 15 min.

Bioreactor cultivation

All chemostat cultures were carried out under strict anaerobic conditions in a 500 ml jacketed bioreactor (BBI-Biotech) with a working volume of 300 ml, an agitation speed set at 200 r.p.m. and pH controlled by automatic addition of NH_4OH (5 N). Chemostat cultures, in CRM, at 35 °C, a dilution rate of 0.05 h⁻¹ and a pH of 4.8, were used to evaluate the strains' degeneration as it was previously shown that *C. acetobutylicum* continuous cultures were unstable under these conditions³⁰. Performances of the CAB2021 strain both in chemostat and in a membrane cell-recycle bioreactor were evaluated in GMM at a

pH of 4.4 and a temperature of 35 °C as it was previously shown to give the highest yield of solvent formation³¹, CAB2021 chemostat culture was carried out in multi-stage mode to test different dilution rates. Stage 1 was maintained at a dilution rate of $0.1 h^{-1}$ for 2 days, and after confirming stable metabolite patterns for 6 time points, the dilution rate was increased to 0.2 h⁻¹ to enter stage 2. After confirming high concentration of residual glucose for 6 time points, the dilution rate was decreased to $0.05 h^{-1}$ to enter stage 3. After confirming stable metabolite patterns and low concentration of residual glucose for 5 time points, the dilution rate was increased to 0.75 h^{-1} to enter stage 4. This stage was run for 13 time points. Continuous cultures in the cell recycle membrane bioreactor were carried out in a 500 ml bespoke glass bioreactor connected to an ultra-filtration flat module (INSIDE KeRAM, TAMI Industries, molecular weight cut-off 50 kDa, surface area 0.25 m²). The fermentation broth was recirculated in the membrane by a peristaltic pump (Masterflex 77965-00, Fisher Scientific). The total working volume was 450 ml, and the pH and the temperature were respectively regulated at 4.4 (with 5 N NH₄OH) and 35 °C. After sterilisation, N₂ was sparged in the reactor. After a 10% inoculation, the cell recycle membrane bioreactor was operated in batch mode for 9 h before switching to continuous mode with total cell recycle and a step-by-step increase in dilution rate of permeate from 0.04 to 0.96 h⁻¹. When an optical density at 600 nm (OD_{600nm}) of 200 was reached, a cell bleeding rate was applied at a dilution rate of $0.05 h^{-1}$.

Microscope image

Culture broth (1 ml) was centrifuged at 13,000 g for 1 min, and the supernatant was discarded. The cells were washed with 1 ml deionized water and centrifuged at 13,000 g for 1 min, and 950 μ l of supernatant was discarded. Pellets and the remaining supernatant were mixed by pipetting, and 10 μ l of the mixture was spotted on a glass microscope slide and covered with a 0.17 mm microscope cover. Immersion oil (10 μ l) for microscopy (Merck) was spotted onto the microscope cover glass. Microscopic images were obtained using a Nikon OPTIPHOT-2 microscope at ×1,000 magnification.

Analytical procedures

Biomass concentration was monitored both by OD_{600nm} using a spectrophotometer and by the dry cell weight method after centrifugation of 1.5 ml of culture broth in an Eppendorf tube (16,000 g, 5 min, room temperature), two washes with Milli-O water and drving under vacuum at 80 °C. The concentrations of glucose, glycerol, acetate, butyrate, lactate, pyruvate, acetoin, acetone, ethanol, isopropanol and *n*-butanol were determined based on high-performance liquid chromatography (HPLC) using H₂SO₄ at 0.5 mM as mobile phase. Metabolite concentrations were determined by HPLC analysis (Dionex UltiMate 3000 HPLC system, Thermo Fisher Scientific). Biorad Aminex HPX-87H column (300 mm × 7.8 mm) was used for separation, and a refractive index detector (to analyse glucose, ethanol, butanol and isopropanol) and UV detectors (absorbance at 210 nm for acids, 280 nm for acetone) were used for detection, respectively. Samples were diluted two times and run at a flow rate of 0.5 ml min⁻¹ at 35 °C (if applicable, 20 °C was used to better resolve peaks of acetone and isopropanol) in 5 mM H₂SO₄ mobile phase for an hour.

Reporting summary

Further information on research design is available in the Nature Portfolio Reporting Summary linked to this article.

Data availability

Data supporting the work are available in the paper and Supplementary Information. The raw whole-genome sequencing data of CAB2018 are available in the European Nucleotide Archive with accession number ERR12915945, and the assembly file in GenBank format is provided as supplementary data. Further information on materials of this study are available from the corresponding author. Source data are provided with this paper.

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Author contributions

P.S. and N.P.M. conceived, designed and coordinated the project. M.Y., M.E., W.K. and C.F. performed experiments and analysed data. M.E., M.Y., P.S. and N.P.M. wrote the manuscript, and all authors approved the final manuscript.

Competing interests

The authors declare no competing interests.

Additional information

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Correspondence and requests for materials should be addressed to Philippe Soucaille.

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CBM lacking uracil +erythromycin



Extended Data Fig. 1 | Strategy for integrating large synthetic DNA into the pSOL1 integrated strain. To integrate large, synthetic DNA regions carrying novel metabolic or biosynthetic capabilities for further engineering, the following strategy is proposed. This process involves five steps, including the use of Yeast Artificial Chromosome (YAC) to assemble the large synthetic DNA, protoplast fusion to bring about its transfer into the target cell and final integration using a similar approach as demonstrated in this study. More details are provided below. Step 1 Preparation of C. acetobutylicum and yeast strains and vectors. CAB2019 strain carrying truncated pyrE with deletions of CA_C1502/ CA_C3535 genes encoding restriction endonucleases³², a yeast strain that is auxotrophic for several nutrients including uracil (URA3-deficient), that is W303-1A (genotype: MATa leu2-3,112 his3-11,15 trpl-1 canl-100 ade2-7 ura3-1)33,34 as well as the 2 following plasmids: i) pYAC Cace, YAC derivative containing 1) a yeast origin of replication (ARSH4), a centromere (CEN6), an uracil-selectable marker (URA3), 2) the origin of replication of pSOL1 and a catP gene and 3) synthetic SLHA and SSHA regions for the crossing over events. ii) pMTL-YAC-int, a derivative of pMTLpSOL1 int, carrying tetA (replacing catP), CA PO176 (replacing hydA to be LHA) and the SSHA' (shorter than SHA) and SLHA that are homologous regions of pYAC Cace vector. Step 2 Synthetic DNA assembly using pYAC Cace as a backbone. Up to 25 fragments can be synthesized and transformed into yeast^{35,36}. A large DNA vector including pYAC Cace backbone is assembled in W303-1A veast strain to generate lengths of up to 250 kb. Commercial Yeast Nitrogen Base (YNB) with all essential except uracil is used for selection after complementing with URA3

on the pYAC Cace vector. Additionally, the resultant plasmid pYAC Cace 250 kb carries the *catP* antibiotic resistance marker for step 3 selection. *Step 3* Protoplast fusion between the yeast carrying pYAC Cace 250 kb and CAB2019 by treating PEG (Polyethylene glycol) to induce fusion. PEG treated C. acetobutylicum³³ and yeast for protoplast induction are mixed. After protoplast fusion, CBM supplemented with uracil (to select C. acetobutylicum by deficiency of amino acids such as histidine due to auxotrophy of W303-1A) and thiamphenicol (to maintain pYAC Cace 250 kb) is used for selection. Step 4 Introduction of pMTL-YAC-int into CAB2019 harbouring pYAC Cace 250 kb by electroporation. Transformant is selected on CGM supplemented with tetracycline (tetA in pMTL-YAC-int) and thiamphenicol. Step 5 Four crossing over events to integrate 250 kb large DNA into pSOL1 integrated C. acetobutylicum. First and second crossing over occurs between SLHAs (Synthetic Large Homology Arms)/SSHA (Synthetic Short Homology Arm) and SSHA' of pYAC Cace 250 kb and pMTL-YACint. The promoter-less ermB gene from pMTL-YAC-int becomes functional after integration as it is now located under the promoter of SSHA in pYAC Cace 250 kb while thiamphenicol sensitivity is expected due to loss of pYAC backbone by use of 5FC. Third and fourth crossing over occurs between CA P0176 and pyrE of pSOL1 integrated chromosome and the resultant plasmid after pYAC Cace 250 kb and pMTL-YAC-int integration. As the truncated pyrE is restored to full length after the 4th crossing over, the final transformant can grow on CBM without uracil supplementation.



■ Biofuel Titer (g/l)

□ Productivity (g/l.h) □ Yield (g/g)

Extended Data Fig. 2 | **Biofuel titre, productivity, and yield of CAB2021** strain grown in chemostat culture in GMM at different dilution rates. To determine the value of CAB2021 for a commercial biofuel production process, an economical mineral medium GMM was continuously fed and different dilution rates were tested. In terms of biofuel titre, the dilution rates of 0.05 and 0.075 h⁻¹ showed the highest titre. In terms of productivity, a dilution rate of 0.2 h⁻¹ showed the highest productivity while high residual glucose was seen (Fig. 2b). In terms of yield, no significant difference was seen among all the tested dilution rates. Data are shown as mean \pm SD from biological replicates (a replicate number for each dilution rate is as following, dilution rate $0.05 \,h^{-1}$, n = 5; $0.075 \,h^{-1}$, n = 13; $0.1 \,h^{-1}$, n = 6; $0.2 \,h^{-1}$, n = 6).

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Corresponding author(s): Philippe Soucaille

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Software and code

Policy information about <u>availability of computer code</u>					
Data collection	Not used				
Data analysis	For genome mutation analysis we used CLC Genomics Workbench version 22.0.1				

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Data on the genome sequence of the CAB2018 strain is now provided as source data

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Life sciences study design

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Sample size	In our study we have constructed very well defined mutants. During the fermentation, the concentration can reach 10 EE9 bacteria per ml. However as they originate from a single cell perfectly characterize and homogeneous from a genetic point of vue we do not have to deal sample size.
Data exclusions	No data exclusions
Replication	Only batch cultures in anaerobic flasks were replicated (at least three time). Previous experiments in our laboratory have shown that it is sufficient for a phenotypic characterization of the mutants. Culture in chemostat were only performed once but different dilution rates (Growth rates) were evaluated and the values given were the average of at least four time points with at least one residence time between each of them. For all these experiments, the standard deviation was below 10% for all the reported values (titer, yield and productivities).
Randomization	Not applicable, same reasons as sample size
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