LETTERS

Programming cells by multiplex genome engineering and accelerated evolution

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The breadth of genomic diversity found among organisms in nature allows populations to adapt to diverse environments^{1,2}. However, genomic diversity is difficult to generate in the laboratory and new phenotypes do not easily arise on practical timescales³. Although in vitro and directed evolution methods⁴⁻⁹ have created genetic variants with usefully altered phenotypes, these methods are limited to laborious and serial manipulation of single genes and are not used for parallel and continuous directed evolution of gene networks or genomes. Here, we describe multiplex automated genome engineering (MAGE) for large-scale programming and evolution of cells. MAGE simultaneously targets many locations on the chromosome for modification in a single cell or across a population of cells, thus producing combinatorial genomic diversity. Because the process is cyclical and scalable, we constructed prototype devices that automate the MAGE technology to facilitate rapid and continuous generation of a diverse set of genetic changes (mismatches, insertions, deletions). We applied MAGE to optimize the 1-deoxy-Dxylulose-5-phosphate (DXP) biosynthesis pathway in Escherichia coli to overproduce the industrially important isoprenoid lycopene. Twenty-four genetic components in the DXP pathway were modified simultaneously using a complex pool of synthetic DNA, creating over 4.3 billion combinatorial genomic variants per day. We isolated variants with more than fivefold increase in lycopene production within 3 days, a significant improvement over existing metabolic engineering techniques. Our multiplex approach embraces engineering in the context of evolution by expediting the design and evolution of organisms with new and improved properties.

With the advent of next-generation fluorescent DNA sequencing¹⁰, our ability to sequence genomes has greatly outpaced our ability to modify genomes. Existing cloning-based technologies are confined to serial and inefficient introduction of single DNA constructs into cells, requiring laborious and outdated genetic engineering techniques. Whereas *in vivo* methods such as recombination-based genetic engineering (recombineering) have enabled efficient modification of single genetic targets using single-stranded DNA (ssDNA)¹¹⁻¹⁴, no such attempts have been made to modify genomes on a large and parallel scale. MAGE provides a highly efficient, inexpensive and automated solution to simultaneously modify many genomic locations (for example, genes, regulatory regions) across different length scales, from the nucleotide to the genome level (Fig. 1).

Efficiency of the MAGE process was characterized using a modified *E. coli* strain (EcNR2). Mediated by the bacteriophage λ -Red ssDNA-binding protein β , allelic replacement is achieved in EcNR2 by directing ssDNA or oligonucleotides (oligos) to the lagging strand of the replication fork during DNA replication¹⁴. We optimized a number of parameters (see Supplementary Information, Supplementary Fig. 2 and Supplementary Table 1) to maximize efficiency of oligo-mediated allelic replacement. To generate sequence diversity in any region of the chromosome by allelic replacement, a pool of targeting oligos is repeatedly introduced into a cell. Under optimized conditions, we can successfully introduce new genetic modifications in >30% of the cell population (Supplementary Fig. 2d) every 2–2.5 h.

Oligo-mediated allelic replacement is capable of introducing a variety of genetic modifications at high efficiency. The efficiency of generating a mismatch or insertion modification is correlated to the amount of homologous sequence between the oligo and its chromosomal target (Fig. 2a, b); the efficiency of producing a deletion modification is correlated to the size of the deletion (Fig. 2c). Figure 2d shows that the predicted two-state hybridization free energy ΔG (ref. 15) between the oligo and target chromosomal sequence is a predictor of the allelic replacement efficiency. Thus, in a pool of oligos with degenerate sequences, oligos with more homology to the target will be incorporated in the chromosome at a higher frequency than those with less homology. This feature of MAGE enables tunable generation of divergent sequences along favourable evolutionary paths.



Figure 1 Multiplex automated genome engineering enables the rapid and continuous generation of sequence diversity at many targeted chromosomal locations across a large population of cells through the repeated introduction of synthetic DNA. Each cell contains a different set of mutations, producing a heterogeneous population of rich diversity (denoted by distinct chromosomes in different cells). Degenerate oligo pools that target specific genomic positions enable the generation of a diverse set of sequences at each chromosomal location.

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Figure 2 | Characterization of allelic replacement efficiency as a function of the type and scale of genetic modifications. a, Introducing mismatch mutations of up to 30 bp. b, Inserting exogenous sequences of up to 30 bp. c, Removing up to 45 kbp of chromosomal sequence using a single oligo. d, Correlation of replacement efficiency and two-state hybridization energy ΔG between the oligo and the targeted complement region in the genome. See Supplementary Fig. 1 for an illustration of oligo interaction with genomic targets and Supplementary Table 3 for a list of oligos and corresponding ΔG values. Dashed line is the linear regression correlation (y = -0.288x - 13.7, $R^2 = 0.799$). All oligos used were 90 bp with two phosphorothioate bonds at the 3' and 5' ends. All error bars indicate \pm s.d.; n = 3.

To determine the rate at which MAGE generates sequence diversity, we used three different 90-mer oligos to produce mismatch changes in a targeted region of the *lacZ* gene in three distinct cell populations. The *cN6* and *cN30* oligos contained 6 and 30 consecutive degenerate bases, respectively; the *iN6* oligos contained 6 degenerate bases interspersed across a 30-bp region (Fig. 3). For these cell populations, the targeted *lacZ* region was sequenced in 96 random clonal isolates after MAGE cycles 2, 5, 10 and 15 that provided a snapshot of the genotypic variation in each population. Through successive cycles of MAGE, the chromosomal sequence of the *lacZ* region increasingly diverged away from wild type (Fig. 3). For *cN6*, after five MAGE cycles, we detected an average change of 3.1 bp per cell across the population (Fig. 3 inset, blue line), which equates to the generation of more than 4.3×10^9 bp



Figure 3 | Sequence diversity generated across three separate cell populations as a function of the number of MAGE cycles. Three 90-mer oligo pools were investigated: cN30, iN6 and cN6. cN30 contains oligos with 30 bp of consecutive degeneracy; iN6 contains oligos with 6 bp of degeneracy spaced every 5 bp; cN6 contains oligos with 6 bp of consecutive degeneracy. Frequency of strains in each population that contains 0 to 7+ bp of differences from the wild-type *lacZ* sequence are colour-coded. The inset shows average number of base pairs changed from wild type across the whole cell population as a function of the number of MAGE cycles using the three oligo pools cN30 (orange line), cN6 (blue line) and iN6 (red line).

of variation per day (3.1 bp changes per five cycles in 7×10^8 cells at 10 cycles per day). Within 15 cycles, cell populations containing all possible N6 genotype combinations were generated using either *cN6* or *iN6* oligos. Because the replacement efficiency for a 30-bp mismatching oligo is lower (1.5% from Fig. 2a), only 21.8% of the *cN30* cell population had undergone allelic replacement after 15 cycles. We detected an average change of 5.6 bp per cell from the wild-type sequence across the whole *cN30* population (Fig. 3 inset, orange line).

The depth at which MAGE generates diversity is determined by a combination of three factors: (1) the degree of sequence variation desired at each locus; (2) the number of loci targeted; and (3) the number of MAGE cycles performed. When a single locus is targeted using a degenerate oligo pool, genetic diversity is generated across the population at that locus and is a function of the oligo pool complexity only. If more than one locus is targeted simultaneously, diversity is generated through the combinatorial arrangement of the different





Figure 4 | MAGE automation. a, Detailed schematic diagram of MAGE prototype including climate-regulated growth chambers with real-time cell density monitors (green), anti-fouling fluidics for transfer of cells between growth chambers and exchange of media and buffers (blue), and real-time generation of competent cells for transformation with synthetic DNA (yellow). Cultures are carried through different chambers at different temperature regimes (30 °C, 42 °C, 4 °C) depending on the necessary MAGE steps (that is, cell growth, heat-shock, cooling). Cells are made electrocompetent by concentration onto a filter membrane and resuspension with wash buffer. Oligos are delivered into cells by electroporation. b, Step-by-step diagram of MAGE cycling steps at a total run time of 2–2.5 h per cycle. Owing to high voltage (18 kV cm⁻¹) electroporation, ~95% of cells are killed at each cycle. Hence, the electroporation event serves to both introduce oligos into cells and to dilute the cell population, cells are then recovered and grown to mid-log phase $(7 \times 10^8 \text{ cells ml}^{-1})$ in liquid medium for the subsequent cycle.

modified loci. The frequency at which each locus is modified can be computationally predicted through a binomial distribution (Supplementary Fig. 3). Although the cell population at any cycle may reflect only a subset of all variants theoretically possible, we can cumulatively generate more variants than the actual size of the cell population ($\sim 7 \times 10^8$ cells) through successive MAGE cycles. Thus, we can generate all variants regardless of population size through computational predictions and continuous cycling by MAGE.

Given the cyclical and scalable nature of our approach, we constructed an integrated prototype device that automates the MAGE process to enable fast and reliable cellular programming. The device contains growth chambers to maintain healthy cell cultures and electroporation modules to repeatedly deliver DNA into the cells, thereby facilitating genome engineering and evolution (Fig. 4, Methods and Supplementary Information). Complex culturing conditions can be programmed into the device for growth of a diverse set of organisms and ecosystems.

To demonstrate an application of the MAGE process, we optimized metabolic flux through the DXP biosynthesis pathway to overproduce the isoprenoid, lycopene, in an *E. coli* strain (EcHW2) that contained the pAC-LYC plasmid that is necessary for the final steps of lycopene production. Twenty endogenous genes (*dxs, dxr, ispD, ispE, ispG, ispH, idi, ispA, appY, rpoS, crl, elbA, elbB, yjiD, purH, rnlA, yggT, ycgZ, ymgA, ariR*) documented to increase lycopene yield^{16,17} were targeted to tune translation (Fig. 5a). Specifically, for each of the 20 genes, 90-mer oligos containing degenerate ribosome binding site

(RBS) sequences (DDRRRRRDDDD; D = A, G, T; R = A, G) flanked by homologous regions on each side were used, with a total pool complexity of 4.7×10^5 ($3^6 \times 2^5 \times 20$). The replaced RBS regions were designed to be more similar to the canonical Shine–Dalgarno sequence (TAAGGAGGT)¹⁸, giving rise to enhanced translation efficiency. Additionally, four genes (*ytjC*, *fdhF*, *aceE*, *gdhA*) from secondary pathways¹⁹ were targeted for inactivation by oligos that introduced two nonsense mutations in the open reading frame, further improving flux through the DXP pathway. In contrast to prior strategies^{20–23} that were experimentally limited by the number of genetic components that could be manipulated at once, here we optimized 24 genes simultaneously to maximize lycopene production.

As many as 15 billion genetic variants $(4.3 \times 10^8 \text{ bp} \text{ variations per})$ cycle for 35 MAGE cycles) were generated. Screening of variants was done by isolating colonies that produced intense red pigmentation on Luria–Bertani agar plates. Variants were isolated from $\sim 10^5$ colonies screened after 5–35 cycles of MAGE (see Supplementary Information), some exhibiting as much as a fivefold increase in lycopene production relative to the EcHW2 ancestral strain (Fig. 5b). Under similar experimental conditions, our highest lycopene yield of ~9,000 p.p.m. (µg per g dry cell weight) is better than documented yields^{17,19}. Sequencing of six variants (EcHW2a–EcHW2f) revealed RBS convergence towards consensus-like Shine–Dalgarno sequences in genes localized at the beginning and end of the biosynthesis pathway (*dxs, dxr, idi, ispA*) as well as various gene knockouts from secondary pathways ($\Delta ytjC$, $\Delta gdhA$, $\Delta fdhF$) (Fig. 5c and Supplementary Table 2).





bars indicate \pm s.d.; n = 3. **c**, Modifications to the lycopene biosynthesis pathway of isolated variants EcHW2a–f with relevant genes highlighted by rectangular boxes. Blue labels represent knockout targets, red labels represent RBS tuning targets. AcCoA, acetyl-CoA; CDP-ME, 4-diphosphocytidyl-2-C-methyl-D-erythritol; CDP-MEP, 4-diphosphocytidyl-2C-methyl-D-erythritol-2-phosphate; DMAPP, dimethylallyl diphosphate; FPP, farnesyl diphosphate; G3P, glyceraldehyde 3-phosphate; HMBPP, (*E*)-4-hydroxy-3-methylbut-2-enyl-diphosphate; IPP, isopentenyl diphosphate; MEP, 2-C-methyl-D-erythritol-4-phosphate; MEC, 2C-methyl-D-erythritol-2,4-cyclodiphosphate.

Different tuning parameters of the lycopene pathway can be individually and combinatorially assessed in the isolated variants. For example, translation optimization of *idi* alone (EcHW2a) increased lycopene production by 40%. Whereas optimizing dxs and idi increased lycopene production by 390% (EcHW2e), additional optimization at rpoS and dxr, along with inactivation of ytjC, improved the growth rate of EcHW2f to that of the EcHW1 ancestor. Interestingly, rpoS is the alternative RNA polymerase subunit sigma factor σ^{s} , the master stress response regulator, and its upregulation can increase stress resistance to the accumulation of lycopene, a very hydrophobic molecule¹⁷. *vtiC* is an uncharacterized phosphoglycerate mutase enzyme and is thought to increase metabolic flux through the DXP pathway by increasing the accumulation of glyceraldehyde 3-phosphate intermediates¹⁹. Inactivation of glutamate dehydrogenase (gdhA) increases lycopene production, but causes a 32% decrease in growth rate of EcHW2b relative to wild type. Combinations of genetic modifications can also be assessed against each other. For example, an optimized dxs and idi strain (EcHW2e) produces 12% more lycopene than a strain with dxs and $\Delta f dhF$ (EcHW2d) and 27% more than a strain with dxs and ispA modifications (EcHW2c). The optimized DXP biosynthesis pathway presented here can be used to produce many other isoprenoid compounds of industrial and pharmaceutical relevance²⁴.

The diversity (that is, the degree of sequence change per target) generated by MAGE is adjustable and the specificity of targeting is always high. Oligos with defined sequences produce well-defined modification, whereas oligos with degenerate sequences produce high-diversity modifications tailored for exploring a vast sequence space. In this study, we used well-defined oligos to inactivate protein-coding sequences and high-diversity degenerate oligos to modify and sample different RBS sequences. We have also used the MAGE platform to perform whole-genome recoding of *E. coli* to enhance the incorporation of non-natural amino acids into proteins²⁵ and construct safer and multi-virus-resistant strains (F.J.I. *et al.*, manuscript in preparation). MAGE is thus a complementary technology to *de novo* genome synthesis²⁶, allowing the tuning of synthetic and natural genomes *in vivo* for various applications.

MAGE is also an accelerated evolution platform that permits the repeated introduction and maintenance of many neutral (or deleterious) mutations in the cell population. Although these mutations would normally disappear in the population via genetic drift or natural selection, MAGE accelerates the rate of their accumulation in any individual cell, thus increasing the likelihood of finding sets of mutations that may interact synergistically to produce a surprisingly beneficial phenotype. Using this technology, we could engineer or evolve cells with higher transfection efficiency (for example, harnessing natural competence systems²⁷), increased allelic replacement efficiency (for example, expressing higher levels or mutants²⁸ of the λ -Red β protein) and perform large-scale bacterial artificial chromosome engineering. The simple allelic replacement mechanism could make this method amenable for use in other organisms, given that other ssDNA-binding protein homologues are functional²⁹. Currently, 30 US dollars of commercially synthesized oligos can introduce up to 27 bp of modification at full degeneracy for a single genomic target. To target many loci, obtaining oligos from programmable DNA microchips³⁰ can significantly decrease the cost in comparison to traditional oligo synthesis. We envision that large-scale pipelines to program synthetic organisms and ecosystems¹⁰ will greatly benefit from integration of hardware, software and wetware to engineer and evolve microbial, plant and animal systems.

METHODS SUMMARY

Strains. The EcNR2 strain is MG1655 with λ -prophage::*bioA/bioB* and *cmR*::*mutS*. The EcHW2 strain is MG1655 with λ -prophage::*bioA/bioB*, *kanR*::*mutS* and pAC-LYC plasmid. EcHW2a–f strains were MAGE-cycled EcHW2 derivatives containing DXP pathway modifications. Details of strains used are described in Methods.

Allelic replacement. Liquid cell cultures were inoculated from colonies obtained from a plate and grown at 30 $^\circ$ C to an absorbance (600 nm) of ~0.7 $(\sim 7 \times 10^8 \text{ cells ml}^{-1})$ in a rotor drum at 200 r.p.m. To induce expression of the λ -Red recombination proteins (Exo, Beta and Gam), cell cultures were shifted to 42 °C for 15 min and then immediately chilled on ice. In a 4 °C environment, 1 ml of cells was centrifuged at 16,000g for 30 s. Supernatant medium was removed and cells were resuspended in dH2O (Gibco catalogue number 15230). This process was repeated twice with water. Supernatant water was removed and oligos suspended in water were added to the cell pellet. The oligos/cells mixture was transferred to a pre-chilled 1 mm gap electroporation cuvette (Bio-Rad) and electroporated with a Bio-Rad GenePulser electroporation system under the following parameters: 1.8 kV, 200Ω and 25μ F. LB-min medium (1-3 ml) was immediately added to the electroporated cells. The cells were recovered from electroporation and grown at 30 °C for 2–2.5 h. Once cells reached mid-logarithmic growth they were used in additional MAGE cycles, isolated or assayed for genotype and/or phenotype analysis.

Lycopene assay. EcHW2 and derivatives (EcHW2a–f) were quantitatively assessed for lycopene production after 24 h of growth in LB-min-chloramphenicol liquid medium at 30 $^{\circ}$ C. Lycopene was extracted and quantified at an absorbance of 470 nm (see Methods). Lycopene yield was calculated by normalizing the amount of lycopene extracted to the dry cell weight.

Full Methods and any associated references are available in the online version of the paper at www.nature.com/nature.

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METHODS

Strains and culture conditions. The λ prophage was obtained from strain DY330³¹, modified to include the *bla* gene and introduced into wild-type MG1655 *E. coli* by P1 transduction at the *bioA/bioB* gene locus and selected on ampicillin to yield the strain EcNR1 (λ -Red⁺). Replacement of *mutS* with the chloramphenicol resistance gene (*cmR* cassette) in EcNR1 produced EcNR2 (mutS⁻, λ -Red⁺). EcNR2 was grown in low salt LB-min medium (10 g tryptone, 5 g yeast extract, 5 g NaCl in 11 dH₂O) for optimal electroporation efficiency. A premature stop codon was introduced into the *cmR* gene of EcNR2 with oligo cat_fwd_stop (Supplementary Table 3) to produce EcFI5, thus inactivating the *cmR* gene. An oligo (cat_fwd_restore) containing the wild-type sequence was used to restore the CmR phenotype. The pAC-LYC plasmid³² containing genes *crtE*, *crtB* and *crtI* was electroporated into EcNR1 to generate EcHW1, which produces lycopene at basal levels. Replacement of *mutS* with a kanamycin resistance gene in EcHW1 produced EcHW2.

Oligonucleotides and DNA sequencing. All oligonucleotides were obtained from Integrated DNA Technologies with standard purification. Oligonucleotides used in the MAGE process contained the following modifications: (1) 30–110 bp in length for optimization experiments; (2) different numbers of phosphorothioated bases; and (3) degenerate nucleotides as described elsewhere in this paper. Additional primers were purchased to amplify relevant genetic regions of the lycopene pathway to sequence strains that expressed high levels of lycopene. DNA sequencing to confirm allelic replacements was performed by Agencourt Bioscience.

LacZ and chloramphenicol replacement efficiency assays. Replacement efficiency was characterized by performing the allelic replacement protocol on EcNR2 cells using 90-mer oligos (Supplementary Table 3) that produced a premature stop codon in the chromosomal *lacZ* gene. In general, 250–500 cells were plated on LB-min containing 5-bromo-4-chloro-3-indoyl- β -D-galactoside and isopropyl- β -D-thiogalactoside (USB Biochemicals) agar plates. Efficiency of allelic replacement was calculated by taking the ratio of the number of white colonies to the total number of colonies on plates. A similar strategy was used in the *cmR* gene recovery experiments with strain EcFI5 where 30–110mer oligos were used to determine optimal oligo length for allelic replacement (Supplementary Fig. 2a). These oligos contained two phosphorothioate bonds at both the 5' and 3' termini. Cells were plated on LB-min-chloramphenicol and LB-min agar plates and grown overnight. Efficiency of allelic replacement was calculated by taking the ratio of the number of colonies on LB-min chloramphenicol plates to the number of colonies on LB-min plates.

MAGE automation device. Cells were grown in sterilized 10-ml glass vials placed in thermally conductive blocks. The growth chambers were climate-regulated through temperature controllers that actuate Peltier heating/cooling elements. Cultures acclimatized quickly (<15 s) in chambers held at different temperatures due to small volumes (for example, 3 ml). Aeration of the culture was achieved through agitation of the chambers at 300 r.p.m. using an orbital shaker. Real-time monitoring of growth rates was achieved by detecting changes in light transmittance across the chamber from light-emitting diodes emitting at 600 nm. Cultures were transferred between chambers through solenoid isolation valves using syringe pumps. Cells were made electrocompetent by a filtration system, which uses a syringe pump to concentrate cells onto a filter membrane (0.22 μ m pore size) and resuspend them off the membrane with appropriate electroporation buffer (for example, dH₂O). Single-stranded oligos (or PCR products) were electroporated into cells in a conductive cuvette with an electric pulse (18 kV cm⁻¹). After each cycle, the entire system (chambers, syringes, filters) was washed with 70% ethanol followed by dH₂O three times to reduce contamination and biofilm formation. All instruments were digitally controlled through software written in the LabView programming environment (National Instruments).

Cultures were initially inoculated into 3 ml of medium in the growth chamber. The device then executed repeatedly and continuously through the following steps: (1) grow cells at 30 °C to a pre-set density (that is, $OD_{600 \text{ nm}}$ of 0.7); (2) induce cells for allelic replacement via 42 °C heat shock for 15 min; (3) chill cells at 4 °C to halt cellular metabolism; (4) wash cells through 15–20 iterations of filtration and resuspension with dH₂O; (5) mix cell suspension and synthetic DNA; (6) deliver DNA into cells by electroporation; and (7) resuspend electroporated cells with growth media.

Colorimetric screen and assay for lycopene production. Cells from the cycled EcHW2 population were plated on LB-min-chloramphenicol agar plates and grown for 1 day at 30 °C and 2 days at room temperature to produce red colonies. The 24 gene targets were divided into three oligo pools, one containing 10 targets, a second containing 14 targets and a third containing all 24 targets, which were cycled through the MAGE process in three separate cultures. In total, 10^5 colonies with increased colour intensity by visual inspection were screened after cycles 5, 10, 15, 20, 25, 30 and 35. From 127 isolates, six strains (EcHW2a-f, with representation from each pool) were selected for direct DNA sequencing across all gene targets and quantitatively measured for lycopene production. For lycopene quantification, these isolated colonies were grown in LB-min medium for 24 h. Lycopene was extracted from 1 ml of cells as follows: centrifuged at 16,000g for 30 s, removal of supernatant media and resuspended with water, and recentrifuged at 16,000g for 30 s. Once the supernatant was removed, the cells were resuspended in 200 µl acetone and incubated in the dark for 15 min at 55 °C with intermittent vortexing. The mixture was centrifuged at 16,000g for 30 s and the supernatant lycopene solution was transferred to a clean tube for quantification. Absorbance at 470 nm of the extracted lycopene solution was measured using a spectrophotometer and calibrated against a known lycopene standard (Sigma-Aldrich, catalogue number L9879) to determine the lycopene content. Dry cell weight was determined by baking a washed cell pellet at 105 °C for 24 h in the dark. Lycopene yield of different EcHW2 derivatives (EcHW2a-f) was calculated by normalizing the amount of lycopene extracted to the dry cell weight.

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