A semisynthetic organism engineered for the stable expansion of the genetic alphabet

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All natural organisms store genetic information in a four-letter, two-base-pair genetic alphabet. The expansion of the genetic alphabet with two synthetic unnatural nucleotides that selectively pair to form an unnatural base pair (UBP) would increase the information storage potential of DNA, and semisynthetic organisms (SSOs) that stably harbor this expanded alphabet would thereby have the potential to store and retrieve increased information. Toward this goal, we previously reported that *Escherichia coli* grown in the presence of the unnatural nucleoside triphosphates dNaMTP and d5SICS TP, and provided with the means to import them via expression of a plasmid-borne nucleoside triphosphate transporter, replicates DNA containing a single dNaM-d5SICS UBP. Although this represented an important proof-of-concept, the nascent SSO grew poorly and, more problematically, required growth under controlled conditions and even then was unable to indefinitely store the unnatural information, which is clearly a prerequisite for true semisynthetic life. Here, to fortify and vivify the nascent SSO, we engineered the transporter, used a more chemically optimized UBP, and harnessed the power of the bacterial immune response by using Cas9 to eliminate DNA that had lost the UBP. The optimized SSO grows robustly, constitutively imports the unnatural triphosphates, and is able to indefinitely retain multiple UBPs in virtually any sequence context. This SSO is thus a form of life that can stably store genetic information using a six-letter, three-base-pair alphabet.

unnatural base pair | CRISPR | Cas9 | DNA replication | nucleotide transporter

The natural genetic alphabet is composed of four letters whose selective pairing to form two base pairs underlies the storage and retrieval of virtually all biological information. This alphabet is essentially conserved throughout nature, and has been since the last common ancestor of all life on Earth. Significant effort has been directed toward the development of an unnatural base pair (UBP), formed between two synthetic nucleotides, that functions alongside its natural counterparts (1–3), which would represent a remarkable integration of a man-made, synthetic component into one of life’s most central processes. Moreover, semisynthetic organisms (SSOs) that stably harbor such a UBP in their DNA could store and potentially retrieve the increased information, and thereby lay the foundation for achieving the central goal of synthetic biology: the creation of new life forms and functions (4).

For over 15 years, we have sought to develop such a UBP (1), and these efforts eventually yielded a family of predominantly hydrophobic UBPs, with that formed between dNaM and d5SICS (dNaM-d5SICS; Fig. 1A) being a particularly promising example (5–7). Despite lacking complementary hydrogen bonding, we demonstrated that the dNaM-d5SICS UBP is well replicated by a variety of DNA polymerases in vitro (7–10), and that this efficient replication is mediated by a unique mechanism that draws upon interbase hydrophobic and packing interactions (11, 12). These efforts then culminated in the first progress toward the creation of an SSO in 2014, when we reported that *Escherichia coli* grown in the presence of the corresponding unnatural nucleoside triphosphates (dNaMTP and d5SICSTP), and provided with a plasmid-encoded nucleoside triphosphate transporter (NTT2) from *Phaeodactylum tricornutum* (which we denote as *PtNTT2*) (13), is able to import the unnatural triphosphates and replicate a single dNaM-d5SICS UBP on a second plasmid (14).

Although this first SSO represented an important proof-of-concept, the generality of the expanded genetic alphabet remained unclear, as retention of the UBP was explored at only a single locus and in only a single sequence context. True expansion of the genetic alphabet requires the unrestricted retention of multiple UBPs at any loci and in any sequence context. Moreover, several limitations were already apparent with the nascent SSO (14). First, although expression of the nucleoside triphosphate transporter enabled *E. coli* to import dNaMTP and d5SICSTP, its expression caused the SSO to grow poorly, with doubling times twice that of the parental strain. Second, the UBP was not well retained during high-density liquid growth or during growth on solid media, presumably due to the secretion of phosphatases that degrade the unnatural triphosphates. Finally, even under optimal conditions, the nascent SSO was unable to retain the UBP with extended growth. Clearly, the ability to robustly grow under the standard repertoire of culture conditions and indefinitely retain the UBP is a prerequisite for true semisynthetic life. Here, we used genetic and chemical approaches to optimize different components of the SSO, ultimately resulting in a simplified and optimized SSO that grows robustly and is capable of the virtually unrestricted storage of increased information.

Significance

The genetic alphabet encodes all biological information, but it is limited to four letters that form two base pairs. To expand the alphabet, we developed synthetic nucleotides that pair to form an unnatural base pair (UBP), and used it as the basis of a semisynthetic organism (SSO) that stores increased information. However, the SSO grew poorly and lost the UBP under a variety of standard growth conditions. Here, using chemical and genetic approaches, we report the optimization of the SSO so that it is healthy, more autonomous, and able to store the increased information indefinitely. This SSO constitutes a stable form of semisynthetic life and lays the foundation for efforts to impart life with new forms and functions.


Conflict of interest statement: Y.Z. and B.M.L. contributed equally to this work.

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Pt600 across a 1-h window of uptake, with the uptake in DM1 induced with the total number of radioactive counts per minute normalized to the average otherwise indicated; r.d.u. is relative decay units, which corresponds to the transporter (see Dataset S1). Although increasing expression of E. coli ΩM IPTG set to 1 (see Materials and Methods). Expression of PtNTT2(66-575) (gray); pSC and constitutive PtNTT2(66-575) (blue), and chromosomally integrated and constitutive PtNTT2(66-575) (green). The promoters from which PtNTT2 is expressed are indicated by the labels next to their corresponding markers. Open triangles denote corresponding control strains without PtNTT2. The pCDF plasmids are in E. coli C41(DE3); pSC plasmids and integrants are in E. coli BL21(DE3). All PtNTT2 strains are non-codon-optimized for plasmid-based expression and codon-optimized for chromosomal expression unless otherwise indicated; r.d.u. is relative decay units, which corresponds to the total number of radioactive counts per minute normalized to the average OD600 across a 1-h window of uptake, with the uptake in DM1 induced with 1,000 μM IPTG set to 1 (see Materials and Methods for additional details). Error bars represent SD of the mean, n = 3 cultures grown and assayed in parallel; the error bars on some data points are smaller than their marker.

Results and Discussion

In our previously reported SSO (hereafter referred to as DM1), the transporter was expressed from a T7 promoter on a multi-copy plasmid (pCDF-1b) in E. coli C41(DE3), and its toxicity mandated carefully controlled induction (14). In its native algal cell, PtNTT2’s N-terminal signal sequences direct its subcellular localization and are ultimately removed by proteolysis. However, in E. coli, they are likely retained, and could potentially contribute to the observed toxicity. Using the cellular uptake of [α-32P]-dATP as a measure of functional transporter expression and as a proxy for the uptake of the unnatural triphosphates, we found that removal of amino acids 1 to 65 and expression of the resulting N-terminally truncated variant PtNTT2(66-575) in E. coli C41(DE3) resulted in significantly lower toxicity, but also significantly reduced uptake (Fig. S1 A and B), possibly due to reduced expression (15). Expression of PtNTT2(66-575) in E. coli BL21(DE3) resulted in significant levels of [α-32P]-dATP uptake with little increase in toxicity relative to an empty vector control, but the higher level of T7 RNAP in this strain (16) was itself toxic (Fig. S1 A and C).

We next explored constitutive expression of PtNTT2(66-575) from a low-copy plasmid or a chromosomal locus, which we anticipated would not only eliminate the need to use T7 RNAP but would also impart the SSO with greater autonomy (eliminating the need to induce transporter production), and, importantly, would result in more homogeneous transporter expression and triphosphate uptake across a population of cells, which we reasoned might improve UBP retention. We explored expression of PtNTT2 (66-575) in E. coli BL21(DE3) with the promoters PlacI, Pbhr, and Pbr from a pSC plasmid, and with Pbhr, Pacc, Puvv, P1207-P4, Pacc, and P25 from the chromosomal lacZYA locus (Dataset S1). We also explored the use of a codon-optimized variant of the truncated transporter (see Dataset S1). Although increasing expression of PtNTT2(66-575) (as measured by uptake of [α-32P]-dATP) was correlated with increasing doubling time, indicating that expression of PtNTT2(66-575) still exhibited some toxicity (uptake of [α-32P]-dATP is itself not toxic), each strain exhibited an improved ratio of uptake to fitness compared with DM1 (Fig. 1B). Strain YZ3, which expresses the codon-optimized, chromosomally integrated PtNTT2(66-575) from the PlacUV5 promoter, exhibited an optimal compromise of robust growth (<20% increased doubling time relative to the isogenic strain without the transporter), and [α-32P]-dATP uptake, and was thus selected for further characterization.

To determine whether the optimized transporter system of YZ3 facilitates high UBP retention, we constructed three plasmids that position the UBP within the 75-nt TK1 sequence (14) [with a local sequence context of d(A-NaM-M)]. These include two high-copy pUCX1-derived plasmids, pUCX1 [referred to in previous work as pINF (14)] and pUCX2, as well as one low-copy pBR322-derived plasmid, pBRX2 (Fig. S2). In addition to allowing us to examine the effect of copy number on UBP retention, these plasmids position the UBP at proximal (pUCX1) and distal (pUCX2 and pBRX2) positions relative to the origin of replication, which we previously speculated might be important (14). Strains YZ3 and DM1 were transformed with pUCX1, pUCX2, or pBRX2 and directly cultured in liquid growth media supplemented with dαM, TPT3 and dαM and isopropyl β-D-1-thiogalactopyranoside (IPTG) for DM1 to induce the transporter, and growth and UBP retention were characterized at an OD600 of ~1 (Fig. 2A and Fig. S3A; see also Materials and Methods). Although DM1 showed variable levels of retention and reduced growth, especially with the high-copy plasmids, YZ3 showed uniformly high levels of UBP retention and robust growth with all three UBP-containing plasmids (Fig. 2B and Fig. S3A). Given that no plasmid locus or copy number biases on UBP retention were observed in YZ3, we chose pUCX2 as a representative UBP-containing plasmid to explore the effect of local sequence context on UBP retention, and we constructed 16 pUCX2 variants in which the UBP was flanked by each possible combination of natural base pairs within a fragment of the GFP gene (see Dataset S1). Under the same growth conditions as above, we observed a wide range of UBP retentions, with some sequence contexts showing complete loss of the UBP (Fig. 2C). However, since the development of DM1 with the dαM-SSICS UBP, we have determined that ring contraction and sulfur derivatization of dSSICS, yielding the dαM-dTPT3 UBP (Fig. 1A), results in more efficient replication in vitro (17). To explore the in vivo use of dαM-dTPT3, we repeated the experiments with YZ3 and each of the 16 pUCX2 plasmids, but with growth in media supplemented with dαMTP and dTTP3. UBP retentions were clearly higher with dnαM-dTPT3 than with dαM-SSICS (Fig. 2C).

Although dαM-dTPT3 is clearly a more optimal UBP for the SSO than dαM-SSICS, its retention is still moderate to poor in some sequence contexts (Fig. 2C). Moreover, several sequences that show good retention in YZ3 cultured in liquid media show poor retention when growth includes culturing on solid media (Fig. S3B). To further increase UBP retention with even these challenging sequences and/or growth conditions, we sought to selectively eliminate plasmids that lose the UBP. In prokaryotes, the clustered regularly interspaced short palindromic repeats (CRISPR)-CRISPR-associated (Cas) system provides adaptive immunity against viruses and foreign plasmids (18–20). In type II CRISPR-Cas systems, such as that from Streptococcus pyogenes (21, 22), the endonuclease Cas9 uses encoded RNAs [or their artificial mimics known as single-guide RNAs (sgRNAs) (23)] to introduce double-strand breaks in cDNA upstream of a 5′-NGG-3′ protospacer adjacent motif (PAM) (24), which then results in DNA degradation by exonucleases (25) (Fig. 3A). In vitro, we found that the presence of a UBP in the target DNA generally reduces Cas9-mediated cleavage relative to sequences that are fully complementary to the provided sgRNA (Fig. S4). We thus
hypothesized that, within a cell, Cas9 programmed with sgRNA(s) complementary to natural sequences that arise from UBPs would enforce retention in a population of plasmids (by eliminating those that lose the UBP), which we refer to as immunity to UBP loss. To test this, we used a p15A plasmid to construct pCas9, which expresses Cas9 via an IPTG-inducible lacO promoter, as well as an 18-nt sgRNA that is complementary to the TK1 sequence (negative control). The bars correspond, respectively, to growth and regrowth with an sgRNA that targets the dT mutation and a single nucleotide deletion mutation (Fig. S5A). With a pCas9 plasmid that expresses two sgRNAs, one targeting the most common substitution mutation and one targeting the single nucleotide deletion mutation (Fig. S5D), and the same growth and regrowth assay, loss of the UBP was undetectable (Fig. 3B).

To more broadly explore Cas9-mediated immunity to UBP loss, we examined retention using 16 pUCX2 variants with sequences that flank the UBP with each possible combination of natural base pairs but also vary its position relative to the PAM, and vary which unnatural nucleotide is present in the strand recognized by the sgRNAs (Table S1). We also constructed a corresponding set of 16 pCas9 plasmids that express two sgRNAs, one targeting a substitution mutation and one targeting the single nucleotide deletion mutation, for each pUCX2 variant. Strain YZ2 carrying a pCas9 plasmid was transformed with its corresponding pUCX2 variant and grown in the presence of the unnatural triphosphates and IPTG (to induce Cas9), and UBP retention was assessed after cells reached an OD600 of ~1. As a control, the 16 pUCX2 plasmids were also propagated in YZ2 carrying a pCas9 plasmid with a nontarget sgRNA. For 4 of the 16 sequences explored, UBP loss was already minimal without immunity (nontarget sgRNA), but was undetectable with expression of the correct sgRNA (Fig. 4D). The remaining sequences showed moderate to no retention without immunity, and significantly higher retention with it, including at positions up to 15 nt from the PAM.

To further simplify and streamline the SSO, we next constructed strain YZ4 by integrating an IPTG-inducible Cas9 gene at the arsR locus of the YZ3 chromosome, which allows for the use of a single plasmid that both carries a UBP and expresses the sgRNAs that enforce its retention. Sixteen such “all-in-one” plasmids (pAIO) were constructed by replacing the Cas9 gene in each of the pCas9 variants with a UBP sequence from the corresponding pUCX2 variant (Fig. S2 and Table S1). YZ4 and YZ3 (included as a no-Cas9 control due to leaky expression of Cas9 in YZ4) were transformed with a single pAIO plasmid and cultured on solid growth media supplemented with the unnatural triphosphates and with or without IPTG to induce Cas9. Single colonies were used to inoculate liquid media of the same composition, and UBP retention was assessed after cells reached an

Image descriptions:
- **Fig. 2.** UBP retention assay and the effects of transporter and UBP optimization. (A) Schematic representation of the biotin shift assay used to determine UBP retention. The plasmid DNA to be analyzed is first amplified in a PCR supplemented with the unnatural triphosphates, and the resulting products are then incubated with streptavidin and subjected to PAGE analysis. X = dNaM, or, in the PCR, its biotinylated analog dTMMPNaM. Y = dSSICs in the PCR, whereas Y = dTPT3 or d5SSICs in the plasmid DNA, depending on the experimental conditions. Lane 1 is the product from the oligonucleotide analogous to that used to introduce the UBP during plasmid assembly, but with the UBP replaced by a natural base pair (negative control). This band serves as a marker for DNA that has lost the UBP. Lane 2 is the product from the synthetic oligonucleotide containing the UBP that was used for plasmid assembly. The shift of this band serves as a marker for the shift of DNA containing the UBP. Lane 3 is the product from the in vitro-assembled plasmid before SSO transformation (positive control). The unshifted band results from DNA that has lost the UBP during in vitro plasmid assembly. Lane 4 is the product from an in vivo replication experiment. (B) UBP retention of plasmids pUCX1, pUCX2, and pBRX2 in strains DM1 and YZ3. Error bars represent SD of the mean, n = 4 transformations for pUCX1 and pUCX2, n = 3 for DM1 pBRX2, and n = 5 for YZ3 pBRX2. (C) UBP retention of pUCX2 variants, wherein the UBP is flanked by all possible combinations of natural nucleotides (NNX, where N = dG, C, A, or T) and X = dNaM, in strain YZ3 grown in media supplemented with either dNaMTP and d5SSICsCTP (gray bars) or dNaMTP and dTPT3CTP (black bars).
OD_{600} of ~1 to 2 (Fig. 4B). Despite variable levels of retention in the absence of Cas9 (YZ3), with induction of Cas9 expression in YZ4, loss was minimal to undetectable in 13 of the 16 sequences. Although retention with the three problematic sequences—d(C-NaM-Ć), d(C-NaM-A), and d(C-NaM-G)—might be optimized, for example, through alterations in Cas9 or sgRNA expression, the undetectable loss of the UBP with the majority of the sequences after a regimen that included growth both on solid and in liquid media, which was not possible with our previous SSO DM1, attests to the vitality of YZ4.

Finally, we constructed a pAIO plasmid, pAIO2X, containing two UBPs: dNaM paired opposite dTPT3 at position 453 of the sense strand of the GFP gene and dTPT3 paired opposite dNaM at position 36 of the sense strand of the SerT tRNA gene, as well as encoding the sgRNAs targeting the most common substitution mutation expected in each sequence (Fig. S2). YZ4 and YZ3 (again used as a control) were transformed with pAIO2X and subjected to the challenging growth regime depicted in Fig. 5, which included extensive high-density growth on solid and in liquid growth media. Plasmids were recovered and analyzed for UBP retention (Fig. S6) when the OD_{600} reached ~1 to 2 during each liquid outgrowth. In YZ3, which does not express Cas9, or in the absence of Cas9 induction (no IPTG) in YZ4, UBP retention steadily declined with extended growth (Fig. 5). With induction of immunity (20 or 40 μM IPTG), we observed only a marginal reduction in growth rate (less than 17% increase in doubling time; Fig. S7), and, remarkably, virtually 100% UBP retention (no detectable loss) in both genes.

Conclusion
Since the last common ancestor of all life on Earth, biological information has been stored with the same four-letter, two-base-pair genetic alphabet. By combining chemical optimization with genetic and immunological engineering, we have created an SSO that is more autonomous (it is naturally competent to import the unnatural triphosphates) and which stores increased information with a fidelity approaching that of natural information. However, unlike any natural organism, the SSO includes an innimate, man-made component: a UBP that allows it to store information with a virtually unrestricted ability to maintain increased information, the optimized SSO now provides a suitable platform for efforts to retrieve the increased information and create organisms with wholly unnatural attributes and traits not found elsewhere in nature.

Materials and Methods
Strains, Plasmids, and Oligonucleotides. A complete list of strains and plasmids and the sequences of oligonucleotides used in this work can be found in Dataset S2; for information regarding strain construction and plasmid cloning, as well as additional experimental details, see SI Materials and Methods. Unless otherwise stated, liquid bacterial cultures were grown in 2×YT (casein peptone 16 g/L, yeast extract 10 g/L, NaCl 5 g/L) supplemented with potassium phosphate (50 mM, pH 7), referred to hereafter as “media,” and incubated at 37 °C in a 48-well flat-bottomed plate (CELLSTAR; Greiner Bio-One) with shaking at 200 rpm. Solid growth media was prepared with
overnight to a storage phosphor screen (BAS-IP MS; GE Healthcare Life Sciences), sample were normalized to the length of time and average OD$_{600}$ during dATP metric analysis using Image Studio Lite (LI-COR). Raw image intensities of each

with IPTG (0 mM to 1 mM, pCDF strains only) or grown (all other strains) for icol for pSC plasmids and integrants) in 500 μL

strepotmycin for pCDF plasmids and chloramphenicol,

synthesized as described previously (5, 7, 10, 17) or kindly provided by Synthorx. Sequencing was performed by GeneArt Gene Synthesis (Thermo Fisher) and desalting. Gene synthesis of the codon-optimized

α32P-dATP [final concentration ~37 °C 20:00

400 ng), PCR insert(s) (3:1 insert:plasmid molar ratio), T4 DNA ligase (200 U), BsaI-HF

was described previously (5, 7, 10, 17) or kindly provided by Synthorx. Sequencing was per-

multiplex kit (QiAprep; Qiagen, or ZR Plasmid Miniprep Classic; Zymo Research).

The [α-32P]-dATP (3,000 Ci/mmol, 10 μCi/mL) was purchased from PerkinElmer. Triphosphates of nDNA, dSCIS, dPTP3, and dMMOD2α were synthesized as described previously (5, 7, 10, 17) or kindly provided by Synthorx. The dATP-containing TK1 oligonucleotide was described previously (14). All other unnatural oligonucleotides containing dNαM were synthesized by Biosearch Technologies with purification by reverse phase cartridge and kindly provided by Synthorx.

daATP Uptake Assay. Radioactive uptake assays were conducted as described (26), with the following modifications: C41(DE3) and BL21(DE3) strains carrying phasing-transporter vectors and their appropriate empty plasmids containing corresponding transporter proteins were electroporated with a 96-well 0.65-

following modifications: All media was additionally supplemented with streptomycin and IPTG (1 mM) to maintain expression of the transporter.

In Vivo Plasmid Replication Experiments with Cas9 (Liquid Culture Only). Electrocortom Y22 cells were transformed with various pCas9 guide plasmids, and single clones were used to inoculate overnight cultures. Cells were then grown, prepared, and electroporated as described above for Y22, with the following modifications: Media for growing cells after electroporation only contained chloramphenicol (i.e., no zeocin), zeocin (to select for pCas9) and 0.2% glucose, electrocompetent cells were stored in 10% (vol/vol ddH2O) DMSO at ~80 °C until use, and recovery and growth media were supplemented with dNαM (250 μM) and dSSICSTP (75 μM). Varying concentrations of IPTG (0 μM to 100 μM) were added to the growth media (but not the recovery media) to induce Cas9 expression. The sgRNA corresponding to the d(AXT) sequence were used for EMR experiments. In Vivo Plasmid Replication Experiments with Cas9 (Plating and Liquid Culture). Electrocortom Y22 cells were grown, prepared, and electroporated as described in In Vivo Plasmid Replication Experiments with Cas9 (Liquid Culture Only) for Y22, with the following modifications: Media for growing cells before electroporation only contained chloramphenicol (i.e., no zeocin), zeocin was used to select for pAIO (i.e., no carbenicillin), and recovery and growth media were supplemented with dNαM (250 μM) and dSSICSTP (75 μM). After transformation with pAIO, dilutions of the recovery mixture were spread onto solid media containing chloramphenicol, zeocin, dNαM (150 μM), dSSICSTP (37.5 μM), 0.2% glucose, and various concentrations of IPTG (0–50 μM). Following overnight growth (37 °C, ~14 h), individual colonies were used to inoculate liquid media of the same composition as the solid media. Experiments performed with pAIOZX were conducted as described above for Y24 without using frozen electroporant cells or glucose. The second plating depicted in Fig. 5 was performed by streaking cells from liquid culture onto solid media of the same composition as the liquid medium, and growth at 37 °C (7–14 h). Six random colonies were selected to continue propagation in liquid culture.

Cell Doubling Calculation. Cell doublings for liquid culture growth–duration–regrowth experiments were calculated by log$_2$ of the dilution factor (30,000 or 300,000) between growths, except for growths inoculated from a plated colony, the cell doublings for which were calculated by averaging, for each individual clone, the time from inoculation to target OD$_{600}$ (0.4 ± 0.11 absorbance [1 L (1 SD) for the first plating inoculation, 10.2 ± 3 h for the second plating inoculation]) and dividing these averages by an estimated doubling time of 40 min. Growth times vary for each clone because colonies were isolated when they were barely visible to the naked eye, and thus we did not attempt to control for variability in the number of cells inoculated into the liquid culture. Note that the reported number of cell dou-

volumes, ~10 mL of media per transformation), and growth to OD$_{600}$ of ~0.3 to 0.4. Cells were then rapidly chilled in an ice water bath with shaking, pelletted (2,500 x g, 10 min), and washed twice with one culture of ice-cold ddH$_2$O. Electrocompetent cells were then resuspended in ice-cold ddH$_2$O (50 μL per transformation), mixed with a Golden Gate assembled plasmid (~1 μL, ~1 ng) containing the UBP, and transferred to a prechilled 0.2-cm-gap electroporation cuvette. Cells were electroporated (Gene Pulser II; Bio-Rad) according to the manufacturer’s recommendations (voltage 25 kV, capacitor 2.5 μF, resistor 200 Ω), then immediately diluted with 950 μL of prevarmed media supplemented with chloramphenicol. An aliquot (10 μL to 40 μL) of this dilution was then immediately diluted fivefold with the same pre-

warmed media, but additionally supplemented with dNαM (250 μM) and dSSICSTP (250 μM). The samples were incubated (37 °C, 1 h), and then ~15% (vol/vol) of the sample was used to inoculate media (final volume 250 μL to 300 μL) supplemented with chloramphenicol, carbenicillin, dNαM (250 μM), and dSSICSTP (250 μM). Cells were then monitored for growth, collected at the density (OD$_{600}$) indicated in Results and Discussion, and subjected to plasmid isolation.

Dilutions of the recovery mixture were also spread onto solid media with chlor-

amphenicol and carbenicillin to ascertain transformation efficiencies. Experiments with dNαM (150 μM) and dSSICSTP (37.5 μM) were performed analogously. Experiments with DM1 were performed analogously using media supple-

mented with streptomycin, with the additional step of inducing trans-

porter expression with IPTG (1 mM, 1 h) before pelleting the cells. All media following electrocompetent cell preparation was also supplemented with streptomycin and IPTG (1 mM) to maintain expression of the transporter.

inoculated into the liquid cultures. Note that the reported number of cell dou-

blings is only an estimate of doublings in liquid culture, which underreports the
doubling rate, and thus we did not attempt to control for variability in the number of cells

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growth at 37 °C (14 h). Six random colonies were selected to continue propagation in liquid culture.

Fig. 5 was performed by streaking cells from liquid culture onto solid media of the same composition as the liquid medium, and growth at 37 °C (~14 h). Six random colonies were selected to continue propagation in liquid culture.

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Biotin Shift Assay. The retention of the UBP(s) in isolated plasmids was determined as previously described and validated (14), with the following modifications. Long-gap or Golden Gate assembled plasmids (0.5 μL, 0.5 ng/μL to 5 ng/μL), or dmNA containing oligonucleotides (0.5 fmol), were PCR-amplified with dNTPs (400 μM), 1× SYBR Green, MgSO4 (2.2 mM), primers (500 nM each), d5SSICSTP (65 μM), dIMMO2AMTP (65 μM), OneTaq DNA polymerase (0.018 U/μL), and DeepVent DNA polymerase (0.007 U/μL) in 1× OneTag standard reaction buffer (final volume 15 μL) under the following thermocycling conditions: [20 × (95 °C : 015 × x′ : 015 / 68 °C : 4:00); see Dataset 2S for a list of primers and their corresponding annealing temperatures (x′ °C) used in this assay. After amplification, 1 μL of each reaction was mixed with streptavidin (2.5 μL, 2 μg/μL Promega) and briefly incubated at 37 °C. After incubation, samples were mixed with loading buffer and run on a 6% (wt/vol) polyacrylamide (29:1 acrylamide: bis-acrylamide) Tris/borate/ EDTA (TBE) gel, at 120 V for ~30 min. Gels were then stained with 1× SYBR Gold dye (Thermo Fisher) and imaged using a Molecular Imager Gel Doc XR+ (Bio-Rad) equipped with a 520Df3 filter (Bio-Rad).

Calculation of UBP Retention. UBP retention was assessed by densitometric analysis of the gels (image) or Image Studio Lite (LICOR) from the biotin shift assay and calculation of a percent raw shift, which equals the intensity of the streptavidin-shifted band divided by the sum of the intensities of the shifted and unshifted bands for the carrion bands. 2A for representative gels. Reported UBP retentions are normalized values.

Unless otherwise indicated, for experiments not involving plating on solid media, UBP retention was normalized by dividing the percent raw shift of each propagated plasmid sample by the percent raw shift of the Golden Gate assembled input plasmid. We assume that the starting UBP content of the media, UBP retention was normalized by dividing the percent raw shift of the dNMA-containing oligonucleotide template used in the assembly of the input plasmid. Plating enables clonal isolation of UBP-containing plasmids from fully natural plasmids that arose during plasmid construction [some of which may contain sequences that are not recognized by the sgRNA(s) used]. Because there is no PCR-mediated loss of the UBP in the oligonucleotide template, normalization to the oligonucleotide template is a better indicator of absolute UBP retention than normalization to the input plasmid. Under the conditions used in the biotin shift assay, changes in oligonucleotide templates and sequence contexts give >90% raw shift, with <2% shift for a cognate fully natural template (i.e., UBP misincorporation during the biotin shift assay is negligible).

Plating allows for the differentiation between UBP loss that occurs in vivo from loss that occurs in vitro, with the exception of clonally derived samples that give <2% shift, for which we are unable to differentiate between whether the UBP was completely lost in vivo or if the sample comes from a transformant that originally received a fully natural plasmid. Such samples are excluded from reported average values when other samples from the same transformation give higher shifts.

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