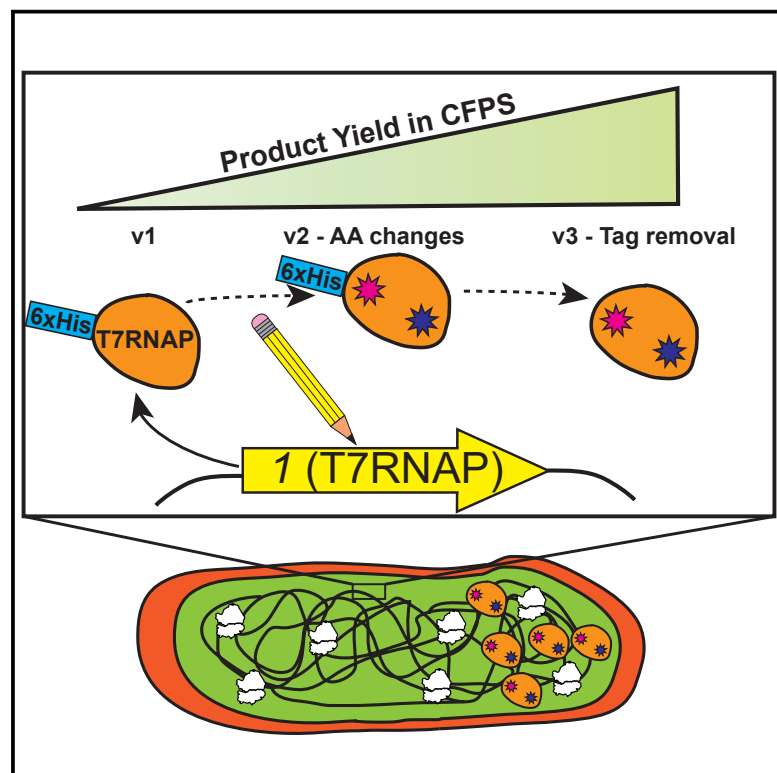


Cell Chemical Biology

A Highly Productive, One-Pot Cell-Free Protein Synthesis Platform Based on Genomically Recoded *Escherichia coli*

Graphical Abstract



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In Brief

Des Soye et al. create and optimize a strain of *Escherichia coli* that expresses T7 RNA polymerase so that lysates prepared from the strain are enriched with sufficient polymerase to catalyze high-yielding cell-free transcription and translation reactions. Using the resulting platform, the authors synthesize products containing up to 40 non-canonical amino acids.

Highlights

- A one-pot, high-yield CFPS system was developed from a recoded strain of *E. coli*
- Genomic edits improved the function of a T7 RNA polymerase expressed in the strain
- Lysates prepared from the strain function without exogenous biological components
- The platform is capable of incorporating up to 40 non-canonical amino acids per protein



A Highly Productive, One-Pot Cell-Free Protein Synthesis Platform Based on Genomically Recoded *Escherichia coli*

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SUMMARY

The site-specific incorporation of non-canonical amino acids (ncAAs) into proteins via amber suppression provides access to novel protein properties, structures, and functions. Historically, poor protein expression yields resulting from release factor 1 (RF1) competition has limited this technology. To address this limitation, we develop a high-yield, one-pot cell-free platform for synthesizing proteins bearing ncAAs based on genomically recoded *Escherichia coli* lacking RF1. A key feature of this platform is the independence on the addition of purified T7 DNA-directed RNA polymerase (T7RNAP) to catalyze transcription. Extracts derived from our final strain demonstrate high productivity, synthesizing 2.67 ± 0.06 g/L superfolder GFP in batch mode without supplementation of purified T7RNAP. Using an optimized one-pot platform, we demonstrate multi-site incorporation of the ncAA *p*-acetyl-L-phenylalanine into an elastin-like polypeptide with high accuracy of incorporation and yield. Our work has implications for chemical and synthetic biology.

INTRODUCTION

A burst of recent development has transformed cell-free protein synthesis (CFPS) from a niche tool for molecular biology into a new technology platform with promise for manufacturing proteins at scale and for accelerating biological design (Carlson et al., 2012; Dudley et al., 2015; Garamella et al., 2016; Hodgman and Jewett, 2012; Karim et al., 2015). This rapid advancement has been spurred by the desire to take advantage of the beneficial features unique to CFPS systems, which include easy sys-

tem access and manipulation, the elimination of competition with cellular growth and adaptation objectives, and a dilute reaction environment that can facilitate folding of complex eukaryotic protein products (Caschera and Noireaux, 2014; Jewett and Swartz, 2004). Batch CFPS reactions now persist for up to a day with yields exceeding 1.5 g/L (Caschera and Noireaux, 2014; Martin et al., 2018), and improvements in scalability culminated recently with the successful completion of a 100-L reaction (Zawada et al., 2011). These impressive advances can be largely attributed to extensive efforts to engineer CFPS systems via chassis organism development, usually by the targeted genetic deletion of genes whose products are known to destabilize key biological substrates (e.g., DNA, mRNA, amino acids, and energy) in cell-free reactions (Hong et al., 2015; Martin et al., 2018; Michel-Reydellet et al., 2004; Yang et al., 2009). As a result of these transformative efforts, CFPS platforms can now be used to complement protein overexpression *in vivo*, with particular utility in rapid prototyping (Chappell et al., 2013; Dudley et al., 2016; Karim and Jewett, 2016; McManus et al., 2019; Schinn et al., 2017; Takahashi et al., 2015), synthesis of toxic products (Martemyanov et al., 2001; Renesto and Raoult, 2003; Watanabe et al., 2010; Xu et al., 2005; Yim et al., 2018), the production of proteins that are difficult to solubly express *in vivo* (Heinzelman et al., 2015; Li et al., 2016; Sullivan et al., 2016; Zawada et al., 2011), manufacturing of glycoproteins (Jaroentomeechai et al., 2018; Kightlinger et al., 2018; Schoborg et al., 2018), detection of disease (Chen et al., 2018; Gootenberg et al., 2017; Pardee et al., 2016a; Slomovic et al., 2015; Takahashi et al., 2018), on-demand biomanufacturing (Adiga et al., 2018; Hunt et al., 2017; Karig et al., 2017; Pardee et al., 2016a, 2016b; Smith et al., 2014; Sullivan et al., 2016), and education (Huang et al., 2018; Stark et al., 2018, 2019; Wandra et al., 2019).

One particularly appealing application of CFPS is the production of proteins containing non-canonical amino acids (ncAAs) (Amiram et al., 2015; d'Aquino et al., 2018; Des Soye et al., 2015; Hong et al., 2014a, 2014b, 2015; Liu et al., 2018; Liu et al., 2017; Martin et al., 2018). To date, more than 150 different



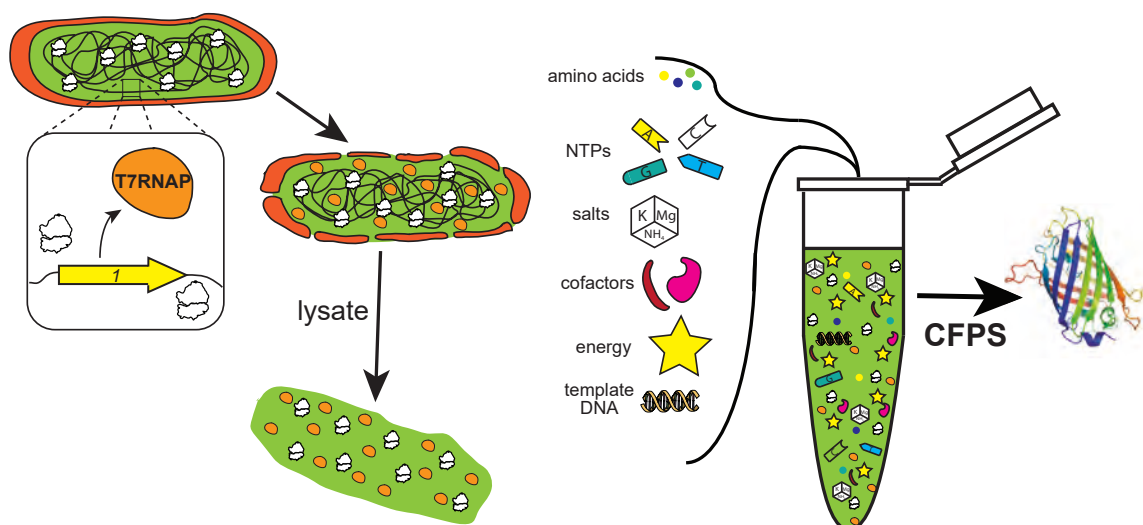


Figure 1. Simplified Schematic of the Production and Utilization of Crude Lysates from *E. coli* Cells to Catalyze Cell-Free Protein Synthesis

Reactions are supplemented with enzymatic cofactors, energy, and other substrates required for protein synthesis as well as plasmid DNA template directing the system toward the production of a product of interest. The strain illustrated is shown endogenously expressing T7RNAP to enable orthogonal transcription *in vitro* and generate a one-pot system independent of supplementation with purified protein components. CFPS, cell-free protein synthesis; NTPs, nucleoside triphosphates.

ncAAs have been incorporated into (poly)peptides (Dumas et al., 2014), enabling the synthesis of proteins featuring novel structures and functions that would otherwise be difficult or even impossible to obtain using only the 20 canonical amino acids. Typically, site-specific ncAA incorporation into proteins is enabled by amber suppression, whereby the amber stop codon (UAG) is recoded as a sense codon designating a ncAA of interest (Liu and Schultz, 2010). This process is mediated by orthogonal translation systems (OTSs), which generally consist of the ncAA, an orthogonal suppressor tRNA (o-tRNA) that has been modified to associate with UAG in the ribosomal A site, and an ncAA-specific aminoacyl-tRNA synthetase (ncAA-RS) that has been evolved to covalently load the ncAA onto the o-tRNA (Des Soye et al., 2015; Santoro et al., 2002), without recognizing natural amino acids. The cytotoxicity of many OTSs (Martin et al., 2018; Nehring et al., 2012), membrane impermeability of some ncAAs (Bundy and Swartz, 2010), and the ability to overcome the relatively poor incorporation efficiencies of OTSs via direct supplementation with OTS components (Des Soye et al., 2015; Hong et al., 2014a, 2015; Martin et al., 2018) makes CFPS an attractive method for the synthesis of peptides featuring ncAAs.

Unfortunately, efforts to apply amber suppression for the incorporation of ncAAs into proteins have long been limited by competition with release factor 1 (RF1), which is responsible for terminating translation in response to the ribosome encountering a UAG codon (Young and Schultz, 2010). In attempting amber suppression, functional RF1 can outcompete ncAA-bearing o-tRNAs at UAG codons, leading to the production of errant truncated products (Hong et al., 2014a, 2014b). Historically, this competition has led to poor protein expression yields, which limits applications in both basic and applied science. Recently, we addressed this limitation. Specifically, we developed a CFPS system derived from a genomically recoded strain of *Escherichia coli* in which all native instances of the amber codon were changed to

the synonymous ochre codon (UAA) followed by elimination of RF1 from the genome (C321.ΔA) (Lajoie et al., 2013; Martin et al., 2018). Extracts derived from the resulting strain (C321.ΔA.759) resulted in CFPS yields of ~1,700 mg/L and 99% suppression efficiency for superfolder green fluorescent protein (sfGFP) with 2 ncAAs, which outperform the best expression of proteins with single or multiple ncAAs *in vivo* (Martin et al., 2018). While this strain demonstrates high productivity for ncAA incorporation into proteins *in vitro*, it is limited by its dependence on the addition of purified viral T7 DNA-directed RNA polymerase (T7RNAP) to catalyze transcription. This adds another step to reaction assembly and increases the cost of the system by requiring the addition of purified polymerase to catalyze robust transcription. In principle, one could create a one-pot CFPS system if T7RNAP could be integrated into the genome, and overexpressed in the source strain prior to lysis. Indeed, extracts derived from T7RNAP-expressing strains (most notably BL21(DE3) (Studier and Moffatt, 1986) and its derivatives) are innately enriched in polymerase activity and generally do not require (or even benefit from) supplementation (Kwon and Jewett, 2015). These one-pot CFPS systems, containing all of the biological components necessary to support transcription and translation, are highly attractive due to their convenient plug-and-play nature.

In this study, we developed a high-yielding one-pot CFPS platform for ncAA incorporation into proteins derived from a genomically recoded RF1-deficient strain of *E. coli* that has been optimized for productivity in CFPS (C321.ΔA.759) (Martin et al., 2018) (Figure 1). Since C321.ΔA.759 does not express T7RNAP, we applied λ-Red-mediated homologous recombination (Datsenko and Wanner, 2000; Mosberg et al., 2010) (λHR) to genomically integrate a series of synthetic constructs featuring the T7RNAP-encoding *1* gene (Studier and Moffatt, 1986) and then assessed the ability of extracts derived from the resulting transformants to catalyze CFPS in the absence of exogenous

polymerase supplementation. While native bacterial RNA polymerases and associated sigma factors can be used to catalyze transcription in CFPS reactions (Shin and Noireaux, 2010, 2012), we chose to pursue T7RNAP because of its high productivity, orthogonality, and strong sequence preference (Shin and Noireaux, 2010; Studier and Moffatt, 1986). Two different genomic loci were targeted for integration, with *1* placed under the regulation of three promoters of different strengths. A high-performing strain, C321.ΔA.759.T7, was capable of synthesizing ~1.4 g/L of sfGFP without purified T7RNAP supplementation. We next exploited multiplex automated genome engineering (MAGE) (Wang et al., 2009) to install mutations in the *1* gene of C321.ΔA.759.T7 and make it resistant to proteolytic cleavage during lysate preparation. The resulting strain, C321.ΔA.759.T7.D, yielded ~1.6 g/L sfGFP without T7RNAP supplementation and ~2.2 g/L with supplementation, but remained transcriptionally limited as evidenced by the increase in yields when purified polymerase was supplemented to the system. To address this, we applied a combination of CRISPR/Cas9 and MAGE (CRMAGE) (Ronda et al., 2016) to remove an N-terminal His tag from the polymerase gene in an effort to improve the function of the T7RNAP expressed by the cells. Lysates from the resulting strain, 759.T7.Opt, demonstrate no improvement when supplemented with additional purified polymerase, yielding ~2.7 g/L sfGFP using only the T7RNAP synthesized in-cell. Using an optimized system, we were able to synthesize proteins (elastin-like polypeptides) bearing up to 20, 30, and 40 ncAAs with yields up to ~70 mg/L in the absence of supplemental T7RNAP. When compared with BL21(DE3) and its derivative strains, one-pot CFPS systems derived from 759.T7.Opt are highly productive and superior for applications involving ncAAs.

RESULTS

CFPS Activity of C321.ΔA.759 and BL21 Star (DE3) with and without Supplemental T7RNAP

We first set out to establish the extent to which C321.ΔA.759 lysates could perform T7-based transcription. To test this, we prepared batches of crude S12 lysates from C321.ΔA.759 as well as BL21 Star (DE3) which had T7RNAP expression induced with 1 mM isopropyl β-D-1-thiogalactopyranoside (IPTG). Batch CFPS reactions were performed using these lysates, directed to synthesize superfolder green fluorescent protein (sfGFP) both with and without direct supplementation of 16 μg/mL purified T7RNAP (Martin et al., 2018) (Figure 2A). As expected based on our previous work (Martin et al., 2018), the yield from C321.ΔA.759 lysates with T7RNAP added was ~30% higher than either BL21 Star (DE3) condition. There was no observable benefit to supplementing additional T7RNAP into reactions utilizing polymerase-enriched BL21 Star (DE3) lysates. Unsurprisingly, essentially no sfGFP was synthesized by the C321.ΔA.759 lysates when no T7RNAP was supplemented. Thus, we hypothesized that introducing the *1* gene into C321.ΔA.759 would imbue the strain with the ability to synthesize T7RNAP and eliminate its dependence on supplemental polymerase *in vitro*.

T7RNAP Insert Design and Integration

A large body of work has explored various ways of enabling bacteria to produce T7RNAP (Davanloo et al., 1984; McAllister et al.,

1981; Studier and Moffatt, 1986). Plasmid-based approaches are simple and effective, but expression levels are high enough to impair plasmid maintenance or otherwise place a significant metabolic burden on host cells, manifesting itself in the form of increased doubling time (Studier and Moffatt, 1986). As increases in doubling time are often indicative of reduced ribosome abundance, this phenomenon is undesirable for CFPS chassis strains (Bremer and Dennis, 1996; Zawada and Swartz, 2006). Another common scheme for *1* gene introduction is via lysogenization with synthetic DE3 bacteriophage (as in BL21(DE3) and its derivatives) (Studier and Moffatt, 1986), but as phage insertion occurs site-specifically at a fixed genomic locus and the viral *1* gene is under the control of a fixed set of *cis*-regulatory sequences, this approach suffers from a lack of tunability and control. An attractive alternative method for genomic integration is λHR, which site-specifically integrates linear DNA constructs into target genomes using flanking-sequence homology to direct insertion at the desired site (Datsenko and Wanner, 2000; Mosberg et al., 2010). As C321.ΔA.759 natively expresses the requisite λ-Red recombination machinery (Lajoie et al., 2013; Martin et al., 2018), we elected to proceed via λHR.

Several design criteria were considered for genome integration. First, a challenge in protein expression is tuning the expression level: enough protein must be synthesized to adequately perform the desired function, but aggressive overexpression can place too high a metabolic burden on the host organism and/or lead to production of inhibitory levels of the protein. Lacking a priori knowledge as to how to achieve an ideal level of T7RNAP production in C321.ΔA.759, we decided to test a variety of different expression levels. We designed a series of synthetic constructs that placed the *1* gene under the regulation of IPTG-inducible promoters of varying transcriptional strengths, with lacUV5 (Stefano and Gralla, 1979), Ptacl (de Boer et al., 1983), and Lpp5 (Inouye and Inouye, 1985) representing relatively low, medium, and high strength, respectively (Figure 2B). Promoter-specific synthetic ribosome binding sites (RBSs) designed for maximal translation using the Ribosome Binding Site Calculator v2.0 were employed for the regulation of translation initiation. In this way, any differences in T7RNAP expression between strains could be predominantly attributed to differences in transcription (Espah Borujeni et al., 2014; Salis et al., 2009). Second, in the interest of easy visualization via western blotting, we added a 6-His tag to the N terminus of the polymerase (a modification that had previously been suggested to have little to no effect on polymerase activity [Ellinger and Ehrlich, 1998]). Third, each construct also included the kanamycin kinase (*kanR*) gene from pKD4 (Datsenko and Wanner, 2000) (which confers resistance to the antibiotic kanamycin) for selection of successful integrants. Finally, to explore influences of genome position on expression, we designed each construct with 50 bp of flanking-sequence homology at each end to facilitate integration at one of two genomic loci: the *asl* locus, selected because it was previously identified as a highly expressing locus in the *E. coli* genome (Bryant et al., 2014), and the *int* locus, selected because it is analogous to the DE3 lysogenization site in BL21(DE3) (Studier and Moffatt, 1986).

A total of six T7RNAP-expressing constructs were assembled (*int.lacUV5*, *int.Ptacl*, *int.Lpp5*, *asl.lacUV5*, *asl.Ptacl*, and

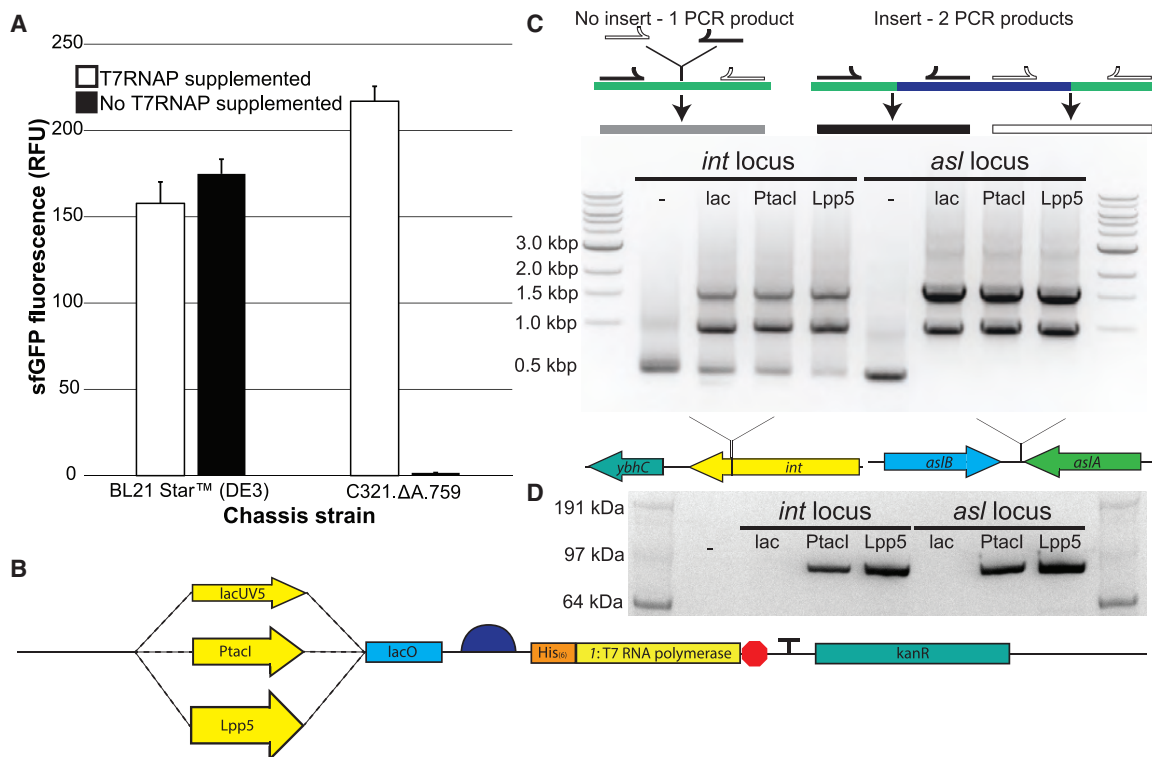


Figure 2. Engineering a Genomically Recoded *E. coli* Strain for T7RNAP Overexpression

(A) sfGFP fluorescence *in vitro* from cell extracts derived from induced BL21 Star (DE3) cells as well as C321.ΔA.759 cells, both with and without supplementation with purified T7RNAP.

(B) Schematic of the synthetic genomic insert used in this study to introduce the gene encoding the T7RNAP into the genome of C321.ΔA.759.

(C) Top: diagram illustrating the PCR-based detection scheme for successful genomic integration of the synthetic T7RNAP cassette into strain C321.ΔA.759. Middle: shown are the MASCC PCR products generated from C321.ΔA.759 and the six T7RNAP-expressing strains generated in this study, run on an agarose gel. The dash indicates the unaltered strain without any insert incorporated. Bottom: depiction of the two insertion loci used.

(D) α-His western blot analysis of protein samples derived from IPTG-induced populations of C321.ΔA.759 and the six T7RNAP-expressing strains generated in this study. The His-tagged T7RNAP version used has a molecular weight of ~100 kDa. The dash indicates the unaltered strain without any insert incorporated.

asl.Lpp5) and transformed individually into C321.ΔA.759 for site-specific genomic integration. Potential integrants were identified by the ability to survive in the presence of kanamycin and verified via screening by multiplex allele-specific colony (MASC) PCR (Figure 2C). Sanger sequencing of all insert loci confirmed that each construct was integrated at the correct locus, fully intact and free of any unwanted mutations. Finally, western blotting with antibodies against the polymerase's N-terminal 6-His tag verified that each insert was indeed promoting expression of T7RNAP (Figure 2D). Polymerase expression as determined by western blot band intensity tracked as expected with promoter strength.

Characterization of T7RNAP-Expressing Strains in CFPS

To assess the ability of these strains to independently catalyze T7RNAP-dependent transcription in CFPS, we prepared crude S12 lysates from all six strains for use in cell-free reactions. To promote robust T7RNAP overexpression, we induced all strains with 1 mM IPTG during exponential cell growth. Batch CFPS reactions using each lysate were directed to synthesize sfGFP over 20 h at 30°C both with and without addition of 16 μg/mL of purified T7RNAP to the reactions (Figure 3A). With polymerase supplemented, lysates from all six strains performed within 15% of

one another. The strains featuring Ptacl- and Lpp5-driven T7RNAP expression demonstrated the ability to perform transcription using only the polymerase expressed by the chassis strain. Not surprisingly, the amount of sfGFP fluorescence appears to be related to the amount of T7RNAP produced in the cells (Figure 3A). At both insertion loci, the amount of fluorescence increases with increasing promoter strength, and for each promoter more fluorescence was observed from the strains featuring inserts at the highly expressing *asl* locus. The strain capable of generating the most sfGFP fluorescence without T7RNAP supplementation, C321.ΔA.759.*asl*.Lpp5, achieved ~85% as much sfGFP production without supplementation as with. This strain, hereafter referred to as C321.ΔA.759.T7, was selected for further characterization and development.

Curiously, western blot analysis of samples derived from C321.ΔA.759.T7 revealed that the T7RNAP produced by the strain is cleaved during lysis near the N terminus to yield a ~21-kDa fragment (Figure 3B). This cleavage is well documented in the literature (Davanloo et al., 1984; Tabor and Richardson, 1985), and previous work has identified the membrane-bound periplasmic protease OmpT as the responsible agent in *E. coli* (Grodberg and Dunn, 1988). The cleaved polymerase itself has been heavily characterized, and prior work

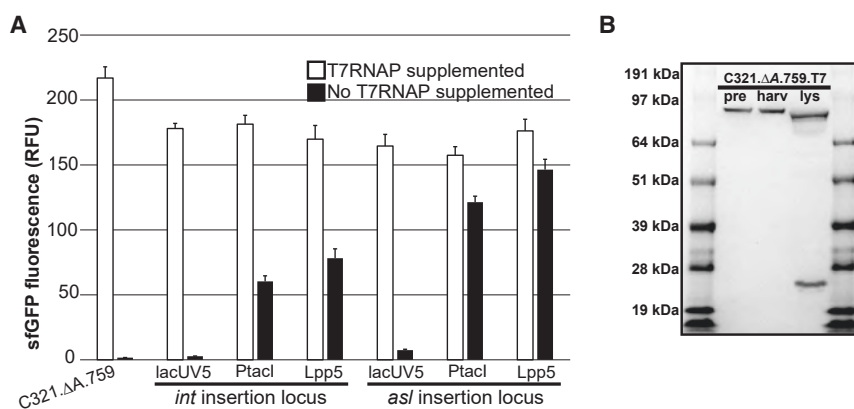


Figure 3. Characterization of C321.ΔA.759 T7RNAP-Expressing Variants

(A) Characterization of the six C321.ΔA.759 T7RNAP-expressing variants generated in this study. Extracts derived from C321.ΔA.759 and its T7RNAP-expressing derivatives were directed to synthesize sfGFP in CFPS both with and without supplementation with purified T7RNAP, and fluorescence was measured after incubation for 20 h at 30°C. Three independent CFPS reactions were performed for each condition, and one standard deviation is shown.

(B) α-His western blot characterization of C321.ΔA.759.T7. pre: samples derived from cells immediately prior to induction. harv: samples derived from mid-exponential phase cells immediately prior to harvest. lys: samples derived from final clarified lysate.

has concluded that the nicked enzyme is impaired by a loss in polymerase activity and efficiency (Ikeda and Richardson, 1987a, 1987b; Muller et al., 1988; Tabor and Richardson, 1985). Thus, we reasoned that OmpT-mediated proteolysis of the T7RNAP expressed by C321.ΔA.759.T7 during cell lysis (when the periplasm and cytoplasm mix) contributed to the reduced capacity of the resulting lysates to support transcription independent of supplemental T7RNAP. To assess this hypothesis, we next sought to inactivate this OmpT activity to protect T7RNAP from proteolysis.

ompT Inactivation to Protect T7RNAP during C321.ΔA.759.T7 Crude Lysate Preparation

BL21(DE3) and its derivative strains all feature a deletion at the *ompT* locus, which presumably prevents the proteolytic degradation of the T7RNAP produced by those strains (Gottesman, 1996). Based on this, we hypothesized that a deletion at the *ompT* locus of C321.ΔA.759.T7 would similarly protect strain-synthesized T7RNAP and thus eliminate the strain's partial dependence on supplemental polymerase in CFPS. To test this, we first "looped" *kanR* out of the C321.ΔA.759.T7 genome using MAGE (Wang et al., 2009). Next, we applied λHR to replace a ~12-kbp region of the C321.ΔA.759.T7 genome analogous to the spontaneous *ompT* deletion in BL21(DE3) with a *kanR* cassette to select for successful integrants. MASC PCR verified the knockout, yielding strain C321.ΔA.759.T7.Δ*ompT*.

To assess the CFPS capabilities of the *ompT*-deficient strain, we prepared crude S12 extracts from culture induced with 1 mM IPTG for analysis via both western blot and batch CFPS reactions. As expected, a western blot revealed that in the absence of OmpT the T7RNAP is no longer cleaved (i.e., we did not observe the expected 21-kDa band) (Figure 4A, inset). Unfortunately, batch sfGFP CFPS reactions demonstrated that the strain's ability to perform CFPS suffered significantly overall in response to the *ompT* knockout (Figure 4A). Compared with C321.ΔA.759.T7 lysate, C321.ΔA.759.T7.Δ*ompT* lysates show a 2- to 3-fold reduction in CFPS yields both with and without T7RNAP supplementation. This is consistent with earlier work demonstrating that functional OmpT is critical for robust protein synthesis in lysates derived from C321.ΔA and its descendants (Martin et al., 2018). Given our interest in designing a one-pot, high-yield CFPS system, we concluded that this was not a viable

strategy for improving C321.ΔA.759.T7 and discontinued our pursuit of this scheme for preventing T7RNAP cleavage during cell lysis.

Engineering a Protease-Resistant T7RNAP

We next considered a chemical biology approach to protecting the T7RNAP produced by C321.ΔA.759.T7 from proteolysis during lysate preparation. We reasoned that since the source of the degradation could not be removed without deleterious effects on the strain's productivity *in vitro*, perhaps the T7RNAP could be mutated such that it would no longer be an efficient substrate for OmpT. OmpT binds its substrates at pairs of adjacent basic residues and catalyzes hydrolysis of the amide bond linking them (Hwang et al., 2007). The requirement of basic residues for OmpT activity at the cleavage site is fairly rigid—in particular, the 1' residue residing immediately upstream of the polypeptide cut site *must* be basic in order for OmpT to facilitate hydrolysis (Hwang et al., 2007). In T7RNAP, two such sites have been identified proximal to the enzyme's N terminus at K172/R173 (Ikeda and Richardson, 1987a) (K183/R184 in His-tagged mutant polymerase) and K179/K180 (Muller et al., 1988) (K190/K191 in His-tagged mutant polymerase). These sites are relatively close together such that OmpT proteolysis at either would liberate a ~21-kDa N-terminal fragment, consistent with what was observed on our C321.ΔA.759.T7 western blot.

Because K183/R184 was previously identified as the primary site of OmpT activity in T7RNAP (Ikeda and Richardson, 1987a), we hypothesized that mutating K183 to a non-basic residue would abolish the target site and thus protect the polymerase from proteolysis despite the presence of fully functional OmpT in the lysate. To test this, we used MAGE to edit the sequence of the *1* gene on the genome of C321.ΔA.759.T7 to mutate K183 to either glycine or leucine, as these mutants had previously been shown to retain robust polymerase activity (Tunitskaya and Kochetkov, 2002). Mutations were detected using allele-specific primers in MASC PCR and confirmed by Sanger sequencing. When extracts prepared from each mutant strain were directed to synthesize sfGFP in batch CFPS reactions both with and without supplemental T7RNAP, neither performed better than C321.ΔA.759.T7 (Figure S1A). Analysis of the extracts revealed that despite the installed mutations, the polymerase was still being cleaved during cell lysis (Figure S1B).

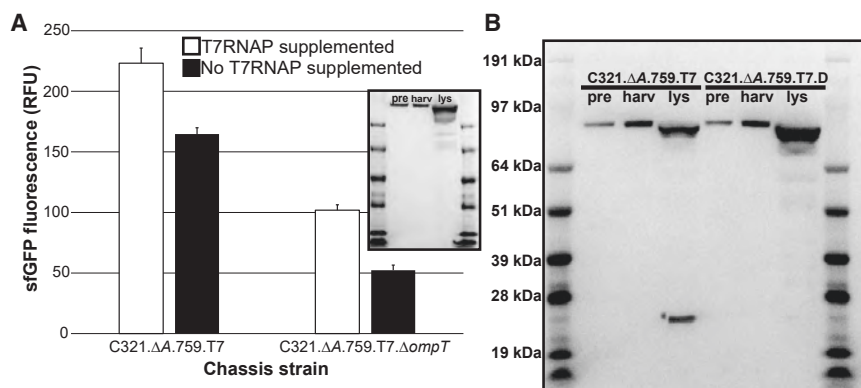


Figure 4. Engineering an OmpT-Resistant T7 Polymerase

(A) Characterization of C321.ΔA.759.T7 and C321.ΔA.759.T7.ΔompT. Extracts from each strain were directed to synthesize sfGFP in CFPS both with and without supplementation with purified T7RNAP, and fluorescence was measured after incubation for 20 h at 30°C. Three independent CFPS reactions were performed for each condition, and one standard deviation is shown. Inset: α-His western blot characterization of C321.ΔA.759.T7.ΔompT.

(B) α-His western blot comparison of C321.ΔA.759.T7 and C321.ΔA.759.T7.D. pre: samples derived from cells immediately prior to induction. harv: samples derived from mid-exponential phase cells immediately prior to harvest. lys: samples derived from final clarified lysate.

See also Figure S1.

Next, we reasoned that while the K183/R184 site may be the preferential site for proteolysis when both sites are present, when this site is unavailable OmpT may simply cleave at K190/K191 instead. We hypothesized that the simultaneous elimination of both sites may be necessary to fully prevent the ability of OmpT to bind and cleave the polymerase. To test this, we again exploited MAGE to edit the sequence of *1* on the C321.ΔA.759.T7 genome and install the mutations K183G and K190L. Mutations were detected using allele-specific primers in MASC PCR and confirmed via Sanger sequencing. We prepared crude cell lysates from the resulting strain, C321.ΔA.759.T7.D, for western blot and CFPS analysis. The western blot revealed that the double mutant T7RNAP expressed by C321.ΔA.759.T7.D is not cleaved despite the presence of active OmpT in the cellular lysate (Figure 4B). In batch mode CFPS reactions, C321.ΔA.759.T7.D lysates exhibit a ~15% increase in productivity over C321.ΔA.759.T7, producing ~2.2 g/L and ~1.6 g/L of sfGFP with and without T7RNAP supplementation, respectively (Figure 5).

Optimization of Endogenous T7RNAP Productivity

Noting that CFPS reactions using C321.ΔA.759.T7.D lysates were still partially dependent on supplementation with purified T7RNAP to achieve maximal productivity, we next sought to identify and address the fundamental feature limiting the productivity of the T7RNAP expressed endogenously by the strain. Because the presence of an associated His tag has been shown to impair the function of other recombinantly expressed proteins (Ledent et al., 1997; Sabaty et al., 2013), we hypothesized that the N-terminal tag used throughout this effort as a way of detecting T7RNAP on blots could be reducing the productivity of the T7RNAP expressed by C321.ΔA.759.T7.D, thus causing lysates derived from the strain to be transcription limited. To test this, we applied a combination of CRISPR/Cas9 and MAGE (Ronda et al., 2016) to further edit the sequence of the *1* gene on the genome of C321.ΔA.759.T7.D, removing the His tag from the N terminus of the polymerase to yield strain C321.ΔA.759.T7.D.ΔHis (hereafter referred to simply as 759.T7.Opt). Batch CFPS reactions using 759.T7.Opt demonstrated that, consistent with our hypothesis, lysates derived from this strain no longer benefit from the supplementation of

purified polymerase; indeed, in the absence of additional polymerase the system yields ~2.7 g/L of sfGFP, dropping to ~2.1 g/L when the enzyme is supplemented (Figure 5), an observation we do not fully understand. 759.T7.Opt lysates also significantly outperform those derived from BL21 Star (DE3), C321.ΔA.759, and C321.ΔA.759.T7.D regardless of T7RNAP supplementation, establishing 759.T7.Opt as a robust one-pot CFPS system and one of the most productive batch CFPS platforms developed to date.

Demonstration of Capacity for Multiple ncAA Incorporations Using T7RNAP-Expressing Strains

With 759.T7.Opt lysates in hand, we next assessed the capacity of these lysates to produce proteins featuring ncAAs. Because the parent strain had RF1 removed (Lajoie et al., 2013), we expected that ncAA incorporation via amber suppression would be highly efficient—indeed, our previous effort using a recoded strain showed up to 40 ncAA incorporations into a single polypeptide (Martin et al., 2018). To assess ncAA incorporation into proteins, we transformed both 759.T7.Opt and BL21 Star (DE3) with a pEVOL plasmid encoding the OTS components for the ncAA *p*-acetyl-L-phenylalanine (pAcF) (Young et al., 2010); namely, the pAcF-specific aminoacyl-tRNA synthetase (pAcFRS) and an orthogonal suppressor tRNA engineered to decode the amber codon (o-tRNA) (Wang et al., 2003). We then quantitatively assessed the incorporation of pAcF into sfGFP variants with up to five in-frame amber codons. CFPS reactions were supplemented with additional OTS components based on our previous work (Martin et al., 2018).

As an initial demonstration of ncAA incorporation, we directed the lysates derived from these pEVOL-bearing strains to synthesize amber mutant variants of sfGFP. We first established the optimal concentrations of pAcF OTS components to be supplied to these CFPS reactions via a series of CFPS reactions directed to synthesize an sfGFP variant featuring two amber codons (sfGFP-2UAG) (Figure S2). Using these conditions, the 759.T7.Opt and BL21 Star (DE3) pEVOL-pAcF lysates were used in CFPS to synthesize wild-type sfGFP (sfGFPwt), sfGFP with a single amber codon (sfGFP-T216X), sfGFP with two amber codons (sfGFP-2UAG), or sfGFP with five amber codons (sfGFP-5UAG). Reactions were performed without supplementation with

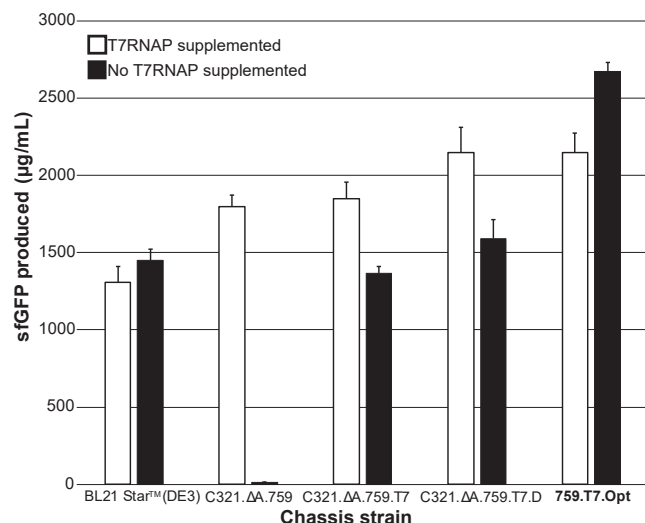


Figure 5. 759.T7.Opt Is a Highly Productive, One-Pot CFPS System

A side-by-side comparison of sfGFP produced in CFPS using crude lysates derived from IPTG-induced BL21 Star (DE3), C321.ΔA.759, C321.ΔA.759.T7, C321.ΔA.759.T7.D, and 759.T7.Opt cells. Shown are results from CFPS reactions performed both with and without supplementation with purified T7RNAP. At least three independent reactions were performed per condition, and one standard deviation is shown.

purified T7RNAP (Figure 6A). As expected, the RF1-deficient 759.T7.Opt lysates exhibit a significantly higher capacity for pAcF incorporation than the BL21 Star (DE3) lysates, with the difference becoming more pronounced as the number of pAcF incorporations increases. Indeed, 759.T7.Opt yields 781 ± 33 µg/mL of sfGFP-5UAG, a more than 5-fold improvement over BL21 Star (DE3). 759.T7.Opt lysates remain highly productive for sfGFP variants bearing up to two pAcFs, yielding >2.1 g/L for sfGFPwt, sfGFP-T216X, and sfGFP-2UAG. To assess the degree of pAcF incorporation in these samples, we performed high-resolution top-down mass spectrometry (i.e., mass spectrometric analysis of whole proteins) (Figure 6B). The results clearly indicate the mass shifts associated with the incorporation of one, two, and five pAcF residues. Analysis of these data shows that site-specific incorporation of pAcF was $\geq 90\%$ in all samples, with ≤ 2 ppm difference between experimental and theoretical masses for sfGFPwt, sfGFP-T216X, and sfGFP-2UAG, and ~ 16 ppm difference for sfGFP-5UAG. These results confirm that the dominant species produced in each reaction features a pAcF at every amber codon.

We next explored the synthesis of large polypeptides containing multiple identical ncAAs utilizing our one-pot CFPS platform derived from 759.T7.Opt. For our model protein, we sought to produce elastin-like polymers (ELPs) (Martin et al., 2018) containing ncAAs using 759.T7.Opt lysates. ELPs are biocompatible and stimuli-responsive biopolymers that can be applied for drug delivery and tissue engineering (Despanie et al., 2016; Raucher and Ryu, 2015). Previously, we have introduced multiple, identical ncAAs into ELPs by substituting natural amino acids with ncAAs at a guest position in the repeating pentapeptide unit (VPGVG) that can be modified while maintaining ELP structure and function (Amiram et al., 2015; Martin et al., 2018).

Our ELP construct consisted of three pentapeptide repeats per monomer unit with a single valine codon per monomer changed to UAG in amber mutants (Martin et al., 2018) (Figure 7A). Lysates derived from 759.T7.Opt bearing pEVOL-pAcF were directed in the absence of supplemental purified T7RNAP to synthesize wild-type (ELP-WT) and amber mutant (ELP-UAG) ELPs with 20, 30, and 40 monomer units in the presence of pAcF. Products were visualized using an autoradiogram, demonstrating that a high percentage of the protein produced in the presence of the ncAA is full length. Full-length ELP-UAG protein was no longer observed when reactions were performed without the addition of pAcF (Figure 7C). Absolute yields for each of the various ELPs were quantified via [^{14}C]-glycine radioactive scintillation counting (Figure 7B). The lysates generated ~ 55 – 70 mg/L of each ELP-WT. For each of the ELP-UAG constructs, the 759.T7.Opt lysates yielded >65 mg/L of product operating in one-pot mode. Of note, we observe some read-through of the UAG codon in the absence of pAcF. This is likely a result of near-cognate suppression via incorporation of natural amino acids, as we and others have observed before (Aerni et al., 2015; Lajoie et al., 2013; Oza et al., 2015). To verify pAcF incorporation in the ELP products, we performed top-down mass spectrometry on intact ELPs to characterize the efficiency of multi-site pAcF incorporation. This analysis confirmed that for 20-, 30-, and 40-mer ELP-UAG constructs, the main products feature pAcF incorporations at all amber codons (Figures 7D–7F, S3, and S4). For example, we observe that site-specific incorporation of pAcF was $\geq 90\%$ in the 20-mer ELP-UAG construct. Taken together, these results demonstrate that 759.T7.Opt lysates are capable of catalyzing the production of proteins bearing multiple ncAAs independent of supplementation with T7RNAP.

DISCUSSION

One-pot systems, such as those derived from the state-of-the-art protein overexpression strain BL21 Star (DE3), are highly desirable for CFPS due to their enrichment with critical enzymes such as T7RNAP. In the specific case of T7RNAP, such systems reduce the cost of CFPS and make the system easier to put together. While robust and versatile, existing one-pot platforms based on BL21 Star (DE3) struggle with the production of proteins containing multiple ncAAs due to the competitive action of RF1 in the reaction environment. In this study, we describe the generation and utilization of a highly productive one-pot CFPS platform beginning with C321.ΔA.759, a genomically re-coded RF1-deficient strain that was previously optimized for CFPS. We integrated a series of DNA constructs into the genome of C321.ΔA.759, each of which featured the T7RNAP-encoding gene 1 under the control of one of three different promoter sequences of varying potency. The construct featuring 1 regulated by strong promoter Lpp5 integrated at a previously identified high-expression genomic locus as/ (Bryant et al., 2014) yielded C321.ΔA.759.T7, which was capable of supporting *in vitro* transcription independent of supplementation with purified T7RNAP. When used in CFPS, C321.ΔA.759.T7 lysates yielded $\sim 85\%$ as much sfGFP without T7RNAP supplementation as with supplementation. In an effort to address the continuing partial dependence of the system on polymerase supplementation, we

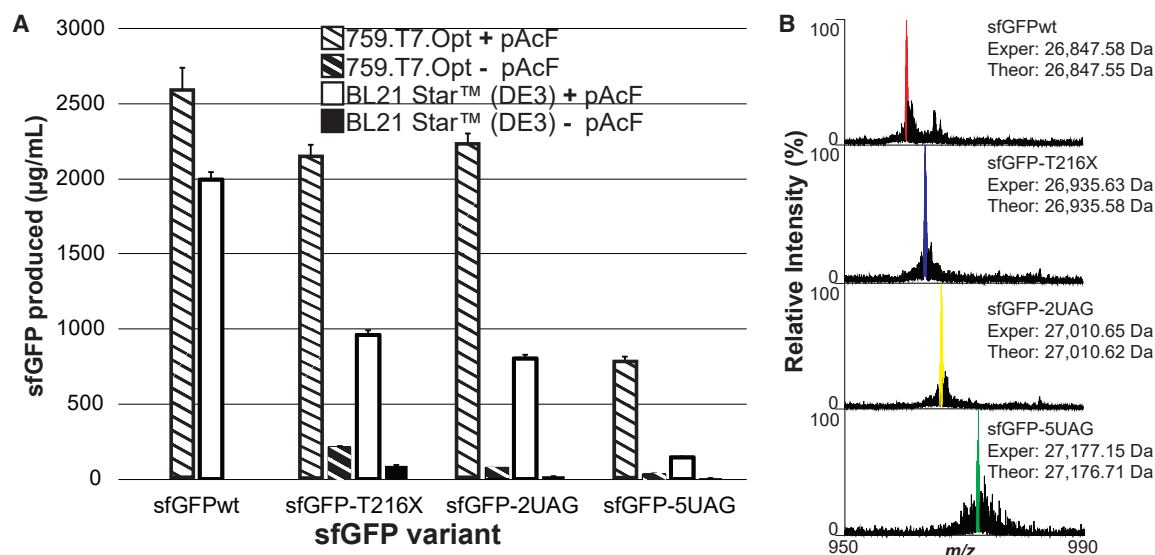


Figure 6. One-Pot ncAA Incorporation into sfGFP Using 759.T7.Opt Lysates

(A) sfGFP produced *in vitro* from cell extracts derived from induced BL21 Star (DE3) cells as well as 759.T7.Opt cells, without supplementation with purified T7RNAP. The indicated sfGFP amber mutant variants were synthesized in the presence of the complete pAcF OTS. At least three independent reactions were performed per condition, and one standard deviation is shown.

(B) Mass spectra of the +28 charge states of the indicated sfGFP variants obtained by top-down mass spectrometry illustrating site-specific incorporation of one to several pAcF residues. Deconvoluted experimental (Exper) and theoretical (Theor) monoisotopic mass values for each variant are listed. Major peaks are highlighted in each spectrum with color.

See also Figure S2.

explored different strategies to protect the T7RNAP expressed in C321.ΔA.759.T7 from OmpT-mediated proteolysis during lysate preparation. By mutating two lysine residues proximal to the N terminus of 1, we were able to abolish the putative OmpT target sites and establish an OmpT-resistant mutant version of T7RNAP in strain C321.ΔA.759.T7.D. We finally removed the N-terminal His tag from 1 in C321.ΔA.759.T7.D to yield strain 759.T7.Opt. 759.T7.Opt lysates compose a one-pot system, yielding ~2.7 g/L sfGFP in batch mode reactions without supplementation with purified biological components. We also demonstrated the merits of RF1-deficient systems for ncAA incorporation, highlighting the significantly increased capacity for amber suppression in 759.T7.Opt lysates as compared with BL21 Star (DE3). Furthermore, we were able to confirm the synthesis of full-length polypeptides containing up to 40 ncAAs without the addition of purified T7RNAP using 759.T7.Opt lysates. This joins an emerging number of reports pushing the bounds of multiple, identical ncAA incorporations using genomically recoded organisms (Mukai et al., 2015).

Looking forward, one intriguing avenue is the continued development of 759.T7.Opt for improved productivity and enhanced functionality in CFPS. This might be achieved by correcting some of the potentially harmful off-target mutations incurred during the initial recoding of the strain. Additionally, upregulation of other positive effectors of CFPS (e.g., chaperones, elongation factors, energy regeneration enzymes) could be achieved via genomic integration using a strategy similar to that employed in this study. In particular, the development of orthogonal synthetases with improved kinetics and substrate specificity is a critical hurdle that must be overcome before these enzymes

can be overexpressed in CFPS chassis strains without deleterious effects on cellular health and lysate performance (Nehring et al., 2012).

Developing efficient CFPS systems specialized for ncAA incorporation is important for synthetic biology for various emerging applications. Powerful one-pot production platforms will support the large-scale synthesis of protein products featuring novel structures and functions, in turn promoting the mass production of potent therapeutics and materials. We anticipate that CFPS platforms such as that derived from 759.T7.Opt are promising for these and other synthetic biology applications.

SIGNIFICANCE

The addition of non-canonical amino acids (ncAAs) to the genetic code has been transformative for protein design and engineering, unlocking protein properties, structures, and functions that would be otherwise be difficult or even impossible to achieve. To better leverage the power of these ncAAs, a cell-free protein synthesis (CFPS) platform comprising crude lysates derived from a genomically recoded strain of *Escherichia coli* that is capable of high-level expression of proteins bearing ncAAs was recently developed. While an extremely potent synthesis tool, this platform is dependent upon supplementation with purified polymerase enzyme to catalyze transcription, limiting its use in both academic laboratories and industrial-scale syntheses. To address this limitation, our work sought to develop this existing platform into a “one-pot” system containing all of the biological components required for transcription and

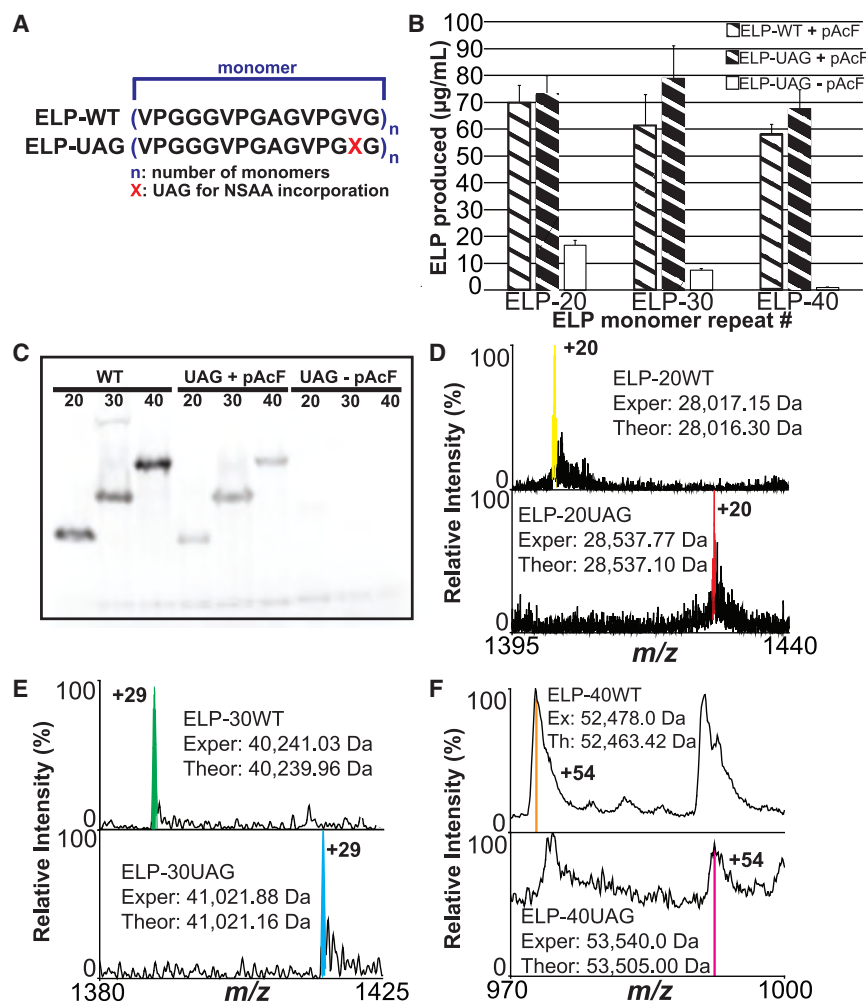


Figure 7. Multi-Site One-Pot ncAA Incorporation Using 759.T7.Opt Lysates

(A) Illustration of both the wild-type (WT) and amber mutant (UAG) ELP monomers used in this study. NSAA, non-standard amino acid.

(B) [¹⁴C]Glycine radioactive count quantification of ELPmers produced by 759.T7.Opt lysates under the indicated conditions. At least three independent reactions were performed per condition, and one standard deviation is shown.

(C) Autoradiogram of ELPmers produced by 759.T7.Opt lysates under the indicated reaction conditions.

(D–F) Mass spectra of the indicated charge states for (D) ELP-20, (E) ELP-30, and (F) ELP-40 constructs. Each set of spectra indicates that 759.T7.Opt lysates are catalyzing the site-specific incorporation of the indicated number of pAcF residues. Deconvoluted experimental (Exper, Ex) and theoretical (Theor, Th) average mass values for each variant are listed. Major peaks are highlighted in each spectrum with color. Reactions for all figure panels were performed without supplementation with purified T7RNAP. (D) displays high-resolution data, (E) displays mid-resolution data, and (F) displays low-resolution data. Resolution was constrained by instrumentation limits based on the increasing size of the targets.

See also [Figures S3 and S4](#).

translation, with a specific focus on the viral T7 RNA polymerase (T7RNAP) that is commonly used to perform transcription in CFPS systems. By genomically incorporating the gene 1 (encoding T7RNAP) into the genome of the source strain and editing its coding sequence, we developed a novel source strain, which expresses a mutant T7RNAP variant that is highly active and resistant to proteolysis. Crude lysates derived from this strain are enriched with this polymerase and are capable of high levels of protein synthesis *in vitro* independent of supplementation with purified enzymes. This platform facilitates the use of CFPS systems and presents a strategy for the introduction of new capabilities to *E. coli* strains for use both *in vivo* and *in vitro*.

STAR★METHODS

Detailed methods are provided in the online version of this paper and include the following:

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- [EXPERIMENTAL MODEL AND SUBJECT DETAILS](#)
 - Microbe Strains

• METHOD DETAILS

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- DNA Gel Electrophoresis
- Plasmid Assembly Using Gibson Assembly
- T7RNAP Linear Insert Construction
- Strain Transformation and Insert Verification
- Cell Extract Preparation
- CFPS Reaction
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- Detection of His-tagged T7RNAP by Western Blot
- Multiplex Advanced Genome Engineering (MAGE) Cycling
- Replica Plating of Bacterial Colonies
- Knockout of *ompT* Locus
- Generation and Verification of OmpT-resistant T7RNAP-expressing Strains
- pMAZ Plasmid Assembly
- CRMAGE Removal of N-terminal His-tag from C321.ΔA.759.T7.D
- Expression and Purification of His-tagged Orthogonal pAcF tRNA Synthetase

- ELP Radioactive Quantitation
- Autoradiogram Analysis
- Purification and Top Down Mass Spectrometry of sfGFP and ELP Constructs
- DNA Sequencing
- **QUANTIFICATION AND STATISTICAL ANALYSIS**
- **DATA AND CODE AVAILABILITY**

SUPPLEMENTAL INFORMATION

Supplemental Information can be found online at <https://doi.org/10.1016/j.chembiol.2019.10.008>.

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AUTHOR CONTRIBUTIONS

B.J.D. designed research, performed all genomic engineering and CFPS reactions, performed top-down mass spectrometry of ELP-40 constructs, analyzed all data, and wrote the paper. V.R.G. performed top-down mass spectrometry of all sfGFP constructs as well as ELP-20 and ELP-30 constructs. P.M.T. and N.L.K. supervised and advised on all top-down mass spectrometry. M.C.J. directed research, analyzed data, and wrote the paper.

DECLARATION OF INTERESTS

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STAR★METHODS

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		
Monoclonal Anti-polyhistidine antibody produced in mouse	Sigma-Aldrich	Cat#H1029; RRID: AB_260015
Goat Anti-Mouse IgG (H+L)-HRP Conjugate	Bio-Rad	Cat#1721011; RRID: AB_11125936
Bacterial and Virus Strains		
<i>E. coli</i> BL21 Star™ (DE3)	ThermoFisher	Cat#C601003
<i>E. coli</i> BL21 (DE3)	New England Biolabs	Cat#C25271
<i>E. coli</i> DH5 α	ThermoFisher	Cat#18265017
<i>E. coli</i> C321. Δ A.759	Martin et al., 2018	N/A
<i>E. coli</i> 759.T7.int.lacUV5	This paper	N/A
<i>E. coli</i> 759.T7.int.PtacI	This paper	N/A
<i>E. coli</i> 759.T7.int.Lpp5	This paper	N/A
<i>E. coli</i> 759.T7.asl.lacUV5	This paper	N/A
<i>E. coli</i> 759.T7.asl.PtacI	This paper	N/A
<i>E. coli</i> 759.T7.asl.Lpp5	This paper	N/A
<i>E. coli</i> 759.T7. Δ kanR	This paper	N/A
<i>E. coli</i> 759.T7. Δ ompT	This paper	N/A
<i>E. coli</i> 759.T7.K172L	This paper	N/A
<i>E. coli</i> 759.T7.K172G	This paper	N/A
<i>E. coli</i> 759.T7.D	This paper	N/A
<i>E. coli</i> 759.T7.D. Δ AbR	This paper	N/A
<i>E. coli</i> 759.T7.Opt	This paper	N/A
Chemicals, Peptides, and Recombinant Proteins		
T7 RNA polymerase	Prepared in-house	N/A
pAcF aminoacyl-tRNA synthetase	Prepared in-house	N/A
Potassium glutamate	Sigma-Aldrich	Cat#G1501
Magnesium glutamate	Sigma-Aldrich	Cat#49605
Ammonium glutamate	MP Biomedicals	Cat#02180595.1
HEPES Buffer	Sigma-Aldrich	Cat#H3375
PEG-8000	Sigma-Aldrich	Cat#202452-500G
Dithiothreitol	Sigma-Aldrich	Cat#43816
Putrescine	Sigma-Aldrich	Cat#P5780
Spermidine	Sigma-Aldrich	Cat#S0266
Glycerol	Sigma-Aldrich	Cat#G5516
Folinic acid	Sigma-Aldrich	Cat#47612
tRNA from <i>E. coli</i> MRE600	Roche	Cat#10109541001
Phosphoenol pyruvate	Roche	Cat#10108294
NAD	Sigma-Aldrich	Cat#N8535
CoA	Sigma-Aldrich	Cat#C3144
Oxalic acid	Sigma-Aldrich	Cat#P0963
Nuclease-free water	Ambion	Cat#AM9937
ATP	Sigma-Aldrich	Cat#A2383
GTP	Sigma-Aldrich	Cat#G8877
UTP	Sigma-Aldrich	Cat#U6625

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REAGENT or RESOURCE	SOURCE	IDENTIFIER
CTP	Sigma-Aldrich	Cat#C1506
L-Valine	Sigma-Aldrich	Cat#V0500
L-Tryptophan	Sigma-Aldrich	Cat#T0254
L-Phenylalanine	Sigma-Aldrich	Cat#P2126
L-Isoleucine	Sigma-Aldrich	Cat#I2752
L-Leucine	Sigma-Aldrich	Cat#L8000
L-Cysteine	Sigma-Aldrich	Cat#C7352
L-Methionine	Sigma-Aldrich	Cat#M9625
DL-Alanine	Sigma-Aldrich	Cat#A7627
L-Arginine	Sigma-Aldrich	Cat#A8094
L-Asparagine	Sigma-Aldrich	Cat#A0884
L-Aspartic Acid	Sigma-Aldrich	Cat#A9256
L-Glutamic acid	Sigma-Aldrich	Cat#G1501
L-Glycine	Sigma-Aldrich	Cat#G7126
L-Glutamine	Sigma-Aldrich	Cat#G3126
L-Histidine	Sigma-Aldrich	Cat#H8000
L-Lysine	Sigma-Aldrich	Cat#L5501
L-Proline	Sigma-Aldrich	Cat#P0380
L-Serine	Sigma-Aldrich	Cat#S4500
L-Threonine	Sigma-Aldrich	Cat#T8625
L-Tyrosine	Sigma-Aldrich	Cat#T3754
Phusion™ polymerase	New England Biolabs	Cat#M0530L
SYBR® Safe	ThermoFisher	Cat#S33102
Quick-Load® 1 kb DNA ladder	New England Biolabs	Cat#N0468S
Quick-Load® 100 bp DNA ladder	New England Biolabs	Cat#N0467S
T5 Exonuclease	New England Biolabs	Cat#M0363S
Taq Ligase	New England Biolabs	Cat#M0208L
dNTP Solution Mix	New England Biolabs	Cat#N0447L
Phusion™ HF Buffer	New England Biolabs	Cat#B0518S
Kanamycin	Sigma-Aldrich	Cat#K4000-25G
Carbenicillin	IBI Scientific	Cat#IB02025
Chloramphenicol	Sigma-Aldrich	Cat#C0378-25G
IPTG	Sigma-Aldrich	Cat#I6758-10G
Tris-acetate	Sigma-Aldrich	Cat#T1258-250G
Tris	Sigma-Aldrich	Cat#T1503-500G
¹⁴ C Glycine	Perkin Elmer	Cat#NEC276E050UC
Scintillation fluid	MP Biomedicals	Cat#0188245304
Bacto™ Tryptone	ThermoFisher	Cat#211705
Bacto™ Yeast Extract	ThermoFisher	Cat#211929
Sodium Chloride	Sigma-Aldrich	Cat#S9888
Bacto™ Agar	VWR	Cat#90000-762
Potassium phosphate monobasic	Sigma-Aldrich	Cat#P0662-500G
Potassium phosphate dibasic	Sigma-Aldrich	Cat#P3786-500G
Magnesium acetate	Sigma-Aldrich	Cat#M5661-250G
Potassium acetate	Sigma-Aldrich	Cat#P1190-1KG
L- <i>p</i> -acetylphenylalanine	Chem-Impex International Inc.	Cat#24756
4x NuPAGE LDS Sample Buffer	ThermoFisher	Cat#NP0007
SeeBlue™ Plus2 Pre-stained Protein Standard	ThermoFisher	Cat#LC5925

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REAGENT or RESOURCE	SOURCE	IDENTIFIER
Methanol	Sigma-Aldrich	Cat#34860-1L-R
2x Colony PCR Master mix	ThermoFisher	Cat#K0172
Cutsmart® buffer	New England Biolabs	Cat#B7204S
USER™ Enzyme	New England Biolabs	Cat#M5505S
anhydrotetracycline	Sigma-Aldrich	Cat#37919-100MG-R
rhamnose	Sigma-Aldrich	Cat#83650-10G
Trichloroacetic acid	Sigma-Aldrich	Cat#T6399-500G
Sodium Hydroxide	Sigma-Aldrich	Cat#221465-500G
200-proof ethanol	Sigma-Aldrich	Cat#E7023-6X500ML
Gel drying solution	Bio-Rad	Cat#1610752
LC-MS Grade methanol	Fisher Scientific	Cat#A456-4
LC-MS Grade water	Fisher Scientific	Cat#W6-4
LC-MS Grade acetonitrile	Fisher Scientific	Cat#A955-4
LC-MS Grade formic acid	Fisher Scientific	Cat#A1171-AMP
NuPAGE MOPS SDS running buffer	ThermoFisher	Cat#NP0001
Critical Commercial Assays		
HiSpeed® Plasmid Maxi Kit	Qiagen	Cat#12643
Ni-NTA Agarose	Qiagen	Cat#30210
MagStrep "type3" XT beads 5% suspension	IBA Life Sciences	Cat#2-4090-002; Cat#2-1003-100; Cat#2-1042-025
E.Z.N.A.® Plasmid Mini Kit	Omega Biotek	Cat#D6943-01
E.Z.N.A.® Gel Extraction Kit	Omega Biotek	Cat#D2500-01
Immun-Blot® Opti-4CN™ Colorimetric Kit	Bio-Rad	Cat#1708235
Oligonucleotides		
See Table S1		
Recombinant DNA		
pKD4	Datsenko and Wanner (2000).	N/A
pAR1219	Davanloo et al., 1984.	N/A
pDPtacIacRSTT1	de Boer et al., 1983.	N/A
pDTT1-Lpp5-EF-Tu	Inouye and Inouye (1985). Gan et al., 2017.	N/A
pY71-sfGFP	Bundy and Swartz (2010).	N/A
pY71-sfGFP-T216amb	Bundy and Swartz (2010).	N/A
pY71-sfGFP-2amb	Hong et al., 2014b.	N/A
pY71-sfGFP-5amb	Hong et al., 2014b.	N/A
pY71-pAcFRS	Hong et al., 2014b.	N/A
pEVOL-pAcF	Young et al., 2010.	N/A
pY71-T7-tz-o-tRNA	Hong et al., 2014b.	N/A
pUC-T7-int.lacUV5	This paper	N/A
pUC-T7-asl.lacUV5	This paper	N/A
pUC-T7-asl.PtacI	This paper	N/A
pUC-T7-asl.Lpp5	This paper	N/A
p15a-T7-asl.Lpp5	This paper	N/A
p15a-T7-asl.PtacI	This paper	N/A
p15a-T7-int.Lpp5	This paper	N/A
pMA7CR_2.0	Ronda et al., 2016	N/A
pMAZ-SK	Ronda et al., 2016	N/A
pMAZ-His	This paper	N/A
pMAZ-Cure	This paper	N/A

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REAGENT or RESOURCE	SOURCE	IDENTIFIER
pY71-FI-ELP20	Martin et al., 2018	N/A
pY71-FI-ELP30	Martin et al., 2018	N/A
pY71-FI-ELP40	Martin et al., 2018	N/A
pY71-FI-ELP20X	Martin et al., 2018	N/A
pY71-FI-ELP30X	Martin et al., 2018	N/A
pY71-FI-ELP40X	Martin et al., 2018	N/A

LEAD CONTACT AND MATERIALS AVAILABILITY

Plasmids and strains generated in this study are available to interested parties pending a Materials Transfer Agreement (MTA). Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Michael C. Jewett (m-jewett@northwestern.edu).

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Microbe Strains

Escherichia coli Strains

BL21 Star™(DE3): grow in Luria-Bertoni media at 37°C

DH5α: grow in Luria-Bertoni media at 37°C

C321.ΔA.759: *E. coli* K-strain derivative in which all endogenous amber stop codons (TAG) have been recoded to TAA and release factor 1 has been deleted (ΔA). Strain also features functional deactivations of the following genes: *endA*, *gor*, *rne*, *mazF*. Grow in Luria-Bertoni media at 34°C, select with 100 μg/mL ampicillin or carbenicillin.

759.T7.int.lacUV5: C321.ΔA.759 with a synthetic construct encoding T7 RNA polymerase under control of the lacUV5 promoter inserted into the genome at the *int* locus. Grow in Luria-Bertoni media at 34°C, select with 50 μg/mL kanamycin.

759.T7.int.Ptacl: C321.ΔA.759 with a synthetic construct encoding T7 RNA polymerase under control of the Ptacl promoter inserted into the genome at the *int* locus. Grow in Luria-Bertoni media at 34°C, select with 50 μg/mL kanamycin.

759.T7.int.Lpp5: C321.ΔA.759 with a synthetic construct encoding T7 RNA polymerase under control of the Lpp5 promoter inserted into the genome at the *int* locus. Grow in Luria-Bertoni media at 34°C, select with 50 μg/mL kanamycin.

759.T7.asl.lacUV5: C321.ΔA.759 with a synthetic construct encoding T7 RNA polymerase under control of the lacUV5 promoter inserted into the genome at the *asl* locus. Grow in Luria-Bertoni media at 34°C, select with 50 μg/mL kanamycin.

759.T7.asl.Ptacl: C321.ΔA.759 with a synthetic construct encoding T7 RNA polymerase under control of the Ptacl promoter inserted into the genome at the *asl* locus. Grow in Luria-Bertoni media at 34°C, select with 50 μg/mL kanamycin.

759.T7.asl.Lpp5: C321.ΔA.759 with a synthetic construct encoding T7 RNA polymerase under control of the Lpp5 promoter inserted into the genome at the *asl* locus. Grow in Luria-Bertoni media at 34°C, select with 50 μg/mL kanamycin.

759.T7.ΔkanR: 759.T7.asl.Lpp5 with kanamycin resistance gene removed. Grow in Luria-Bertoni media at 34°C, select with 100 μg/mL ampicillin or carbenicillin.

759.T7.ΔompT: 759.T7.asl.Lpp5 with deletion at genomic *ompT* locus. Grow in Luria-Bertoni media at 34°C, select with 50 μg/mL kanamycin.

759.T7.K172L: 759.T7.asl.Lpp5 with a K172L mutation installed in the strain's genomic 1 gene (encoding T7 RNA polymerase). Grow in Luria-Bertoni media at 34°C, select with 50 μg/mL kanamycin.

759.T7.K172G: 759.T7.asl.Lpp5 with a K172G mutation installed in the strain's genomic 1 gene (encoding T7 RNA polymerase). Grow in Luria-Bertoni media at 34°C, select with 50 μg/mL kanamycin.

759.T7.D: 759.T7.asl.Lpp5 with mutations K172G and K179A installed in the strain's genomic 1 gene (encoding T7 RNA polymerase). Grow in Luria-Bertoni media at 34°C, select with 50 μg/mL kanamycin.

759.T7.D.ΔAbR: 759.T7.D with resistances to ampicillin/carbenicillin and kanamycin removed. Grow in Luria-Bertoni media at 34°C.

759.T7.Opt: 759.T7.D.ΔAbR with the N-terminal 6xHistag removed from the strain's genomic 1 gene (encoding T7 RNA polymerase). Grow in Luria-Bertoni media at 34°C. Strain is not resistant to ampicillin/carbenicillin or kanamycin.

Cell lines produced in this study have not been authenticated.

METHOD DETAILS

Strains and Plasmids

The bacterial strains and plasmids used in this study are listed in [Table S2](#). Carbenicillin (50 μg/mL) was used for culturing C321.ΔA.759 and to maintain plasmid pMA7CR_2.0, kanamycin (50 μg/mL) was used for culturing C321.ΔA.759 T7RNAP linear

insert transformants and maintaining pY71/pJL1/pMAZ-based plasmids, and chloramphenicol (34 $\mu\text{g/mL}$) was used to maintain the pEVOL-pAcF plasmid.

PCR Reactions

Reactions using purified DNA species as template were performed using Phusion™ polymerase along with dNTP solution mix and Phusion™ HF Buffer (all from New England Biolabs, Ipswich, MA). Reactions from cellular genomic DNA (colony PCR, multiplex allele-specific colony (MASC) PCR) were performed using 2x colony PCR master mix (Thermo Fisher Scientific, Inc., Waltham, MA).

Plasmid DNA Purification

For applications requiring small amounts of plasmid DNA (PCR, cloning, moving between strains, etc.) plasmid DNA was purified from cells using E.Z.N.A.® Plasmid Mini Kit (Omega Bio-tek, Norcross, GA). For applications requiring large amounts of plasmid DNA (templating CFPS reactions) plasmid DNA was purified from cells using Hi-Speed® Plasmid Maxi Kit (Qiagen, Venlo, The Netherlands).

DNA Gel Electrophoresis

Unless otherwise stated, all DNA electrophoresis was done in 1% agarose gels stained with SYBR® Safe (Thermo Fisher Scientific, Inc., Waltham, MA). Samples were run at 100V for 30–60 minutes on a Mini Gel II Complete Electrophoresis System (VWR, Radnor, PA). 100 bp and 1 kb Quick-load® DNA Ladders (New England Biolabs, Ipswich, MA) were used for fragment size reference.

Plasmid Assembly Using Gibson Assembly

To assemble plasmids from linear DNA pieces, throughout this work we use the method of Gibson *et al.* (Gibson *et al.*, 2009). Individual pieces were synthesized by PCR such that adjacent fragments had ~20 bp of flanking homology with each neighboring fragment. Fragments were assembled by incubating 50 ng of each fragment with assembly mix (6.7 mM PEG-8000, 107 mM Tris-HCl pH 7.5, 10.7 mM magnesium chloride, 213 μM dATP, 213 μM dGTP, 213 μM dCTP, 213 μM dTTP, 10.7 mM dithiothreitol, 1 mM nicotinamide adenine dinucleotide, 0.0043 U/ μL T5 exonuclease (New England Biolabs, Ipswich, MA), 4.3 U/ μL Taq ligase (New England Biolabs, Ipswich, MA), and 0.023 U/ μL Phusion polymerase (New England Biolabs, Ipswich, MA)) for 1 hr at 50°C. Following this incubation, 2 μL of reaction volume were transformed into electrocompetent DH5 α cells using a Micropulse electroporator (Bio-Rad, Hercules, CA). Transformed cells were recovered for 1 hr in LB media at 37°C and 250 rpm. 100 μL of recovered culture was spread onto selective plates and put at 37°C overnight. Resulting colonies contained the assembled plasmid of interest.

T7RNAP Linear Insert Construction

The six T7RNAP-encoding inserts used in this study were assembled from PCR products obtained using primers listed in Table S1. The insert loci were at coordinates 3,986,255 (*asI*) and 805,473 (*int*) in the genome of C321. Δ A.759. In brief, each insert was assembled from four segments of linear DNA – a promoter segment featuring 50 base pairs of sequence homology to the genome of C321. Δ A.759 upstream of the targeted insert site, a T7RNAP segment containing the *1* gene, a terminator segment encoding the synthetic terminator sequence L3S2P21 (Chen *et al.*, 2013), and a kanR segment featuring the kanamycin resistance cassette from pKD4 (Datsenko and Wanner, 2000) as well as 50 bp of sequence homology to the genome of C321. Δ A.759 downstream of the targeted insert site. All constructs were designed such that the coding strand would be integrated into the leading strand during genome replication. Adjacent segments featured at least 20 bp of sequence homology to one another to facilitate their assembly into a single unit of DNA, and novel sequence elements (e.g. 6His-tag and synthetic RBS sequences) were built into the 5' tails of primers. Inserts featuring the promoter Lpp5 were assembled via overlap assembly PCR (SOEing) (Horton, 1995) and amplified with end primers. End primer PCR reactions for these species generated a large number of off-target sequences, so full-length insert DNA was separated from other products via electrophoresis and extracted from 1% agarose gel using a DNA gel extraction kit (Product No. D2500; Omega Bio-tek, Norcross, GA). Inserts featuring promoters Ptacl and lacUV5 were assembled together with plasmid origins of replication (p15a and pUC, respectively) via Gibson assembly (described above) to yield plasmid DNA, and these plasmids were used as template with end primers to yield the linear insert DNA via PCR. We found that the PCR products generated using this approach had a significant reduction in the prevalence of off products observed for the other inserts, accelerating our workflow.

Strain Transformation and Insert Verification

The T7RNAP cassettes were inserted into the genome of C321. Δ A.759 via λ HR following the protocol of Datsenko and Wanner (Datsenko and Wanner, 2000). In brief, for each cassette a 5 mL culture of C321. Δ A.759 was grown in LB media (10 g/L tryptone, 5 g/L yeast extract and 10 g/L NaCl) (Wang and Church, 2011) to an OD₆₀₀ of 0.6, after which it was incubated at 42°C for 15 min. 1.5 mL of this culture was washed twice in ice cold, sterile nuclease-free water and resuspended in 30 μL of insert DNA at a concentration of 70 ng/ μL . The cell suspension was transferred to a 2 mL electroporation cuvette and DNA was introduced into cells using a Micropulse electroporator (Bio-Rad, Hercules, CA). Immediately following electroporation, cells were resuspended in 1 mL sterile LB media and recovered for 3 hrs at 34°C at 250 rpm. The recovered cell culture was plated on kanamycin selective plates and permitted to grow overnight at 34°C. The following day, colonies to be screened were picked and inoculated into 100 μL of kanamycin media on a 96-well plate (Costar 3370; Corning, Corning, NY) and cultured for 3 hrs at 34°C at 250 rpm. 1 μL of each miniature culture was used as template to detect successful genomic integration of each insert by colony PCR using Colony PCR Master Mix (Thermo Fisher

Scientific, Inc., Waltham, MA) with primers listed in [Table S1](#). Two primer pairs were used for detection such that if no insert was present, the outermost pair of primers would anneal to the flanking genomic sequence and generate a single ~500 bp product; however, if the insert was present intact at the locus both pairs of detection primers would be able to anneal and generate two products of ~1250 and ~1750 bp ([Figure 2C](#)).

Cell Extract Preparation

For rapid prototyping of engineered strains, cells were grown in 1 L of 2xYTPG media (16 g/L tryptone, 10 g/L yeast extract, 5 g/L NaCl, 7 g/L K_2HPO_4 , 3 g/L KH_2PO_4 , pH 7.2) in a 2.5 L Tunair® shake flask and incubated at 34°C at 220 rpm. Unless otherwise stated, cultures were inoculated with 1 mM IPTG at an OD_{600} of 0.6 and permitted to continue to grow to an OD_{600} of 3.0. Cells were pelleted by centrifuging for 15 min at 5000 × g at 4°C, washed three times with cold S30 buffer (10 mM tris-acetate pH 8.2, 14 mM magnesium acetate, 60 mM potassium acetate, 2 mM dithiothreitol (DTT)) ([Swartz et al., 2004](#)), and stored at -80°C. To make cell extract, cell pellets were thawed and suspended in 0.8 mL of S30 buffer per gram of wet cell mass and 1.4 mL of cell slurry was transferred into 1.5 mL microtubes. The cells were lysed using a Q125 Sonicator (Qsonica, Newtown, CT) with 3.175 mm diameter probe at a 20 kHz frequency and 50 % amplitude for three cycles of 45s ON/59s OFF. To minimize heat damage during sonication, samples were placed in an ice-water bath. For each 1.4 mL sample, the input energy was ~844 Joules and was monitored during sonication. Immediately following sonication, DTT was added to each tube to a final concentration of 3 mM. Extract was then centrifuged at 12,000 × g at 4°C for 10 min. For strain derivatives of C321.ΔA.759, a run-off reaction (37°C at 250 rpm for 1 h) and second centrifugation (10,000 × g at 4°C for 10 min) were performed ([Kwon and Jewett, 2015](#)). The supernatant was flash-frozen using liquid nitrogen and stored at -80°C until use.

CFPS Reaction

A modified PANOX-SP system was utilized for CFPS reactions ([Jewett and Swartz, 2004](#)) ([Jewett et al., 2008](#)). Briefly, each 15 μL CFPS reaction was assembled in a 2.0 mL microtube by mixing the following components: 1.2 mM ATP; 0.85 mM each of GTP, UTP, and CTP; 34 μg/mL folinic acid; 170 μg/mL of *E. coli* tRNA mixture; 13.3 μg/mL plasmid; 16 μg/mL T7 RNA polymerase; 2 mM for each of the 20 standard amino acids; 0.33 mM nicotinamide adenine dinucleotide (NAD); 0.27 mM coenzyme-A (CoA); 1.5 mM spermidine; 1 mM putrescine; 4 mM sodium oxalate; 130 mM potassium glutamate; 10 mM ammonium glutamate; 12 mM magnesium glutamate; 57 mM HEPES, pH 7.2; 33 mM phosphoenolpyruvate (PEP), and 27% v/v of cell extract. For ncAA incorporation, 2 mM pAcF, 0.5 mg/mL pAcFRS, and 30 μg/mL of o-tz-tRNA linear DNA were supplemented to cell-free reactions. o-tRNA linear DNA was amplified from pY71-T7-tz-o-tRNA plasmid via PCR and transcribed during the cell-free reaction. Furthermore, the o-tRNA was expressed in the source strain prior to extract preparation. Each CFPS reaction was incubated for 20 h at 30°C unless noted otherwise. *E. coli* total tRNA mixture (from strain MRE600) and phosphoenolpyruvate was purchased from Roche Applied Science (Indianapolis, IN). ATP, GTP, CTP, UTP, 20 amino acids and other materials were purchased from Sigma (St. Louis, MO) without further purification. T7RNAP was purified in house as described below.

Quantification of Active sfGFP

CFPS reactions were diluted 1:25 in nanopure water and active full-length sfGFP protein yields were quantified by measuring fluorescence using a Synergy 2 plate reader (BioTek, Winooski, VT) with excitation at 485 nm, emission at 528 nm, and cut-off at 510 nm in 96-well half area black plates (Costar 3694; Corning, Corning, NY). sfGFP fluorescence units were converted to concentration using a standard curve established with ^{14}C -Leucine quantification.

Detection of His-tagged T7RNAP by Western Blot

To visualize T7 RNA polymerase overexpression *in vivo*, cell samples were collected during harvest. Pre-induction cell samples were derived from 1 mL of culture at OD_{600} of 0.6, harvest samples were derived from 200 μL of culture at OD_{600} of 3.0. To prepare samples for gel electrophoresis, cells were pelleted and resuspended in 200 μL of nuclease-free water. 100 μL of this suspension was mixed with 34 μL of 4x NuPAGE® LDS Sample Buffer (Thermo Fisher Scientific, Inc., Waltham, MA) and boiled for 10 minutes. Following the boil, samples were spun at >13,500 × g. Samples derived from lysates were prepared by diluting 1 μL of extract in 8 μL of nuclease-free water and boiling for 10 minutes with 3 μL of 4x NuPAGE® LDS Sample Buffer. 12 μL of each sample was loaded into 12% Bis-Tris NuPAGE® gel (Thermo Fisher Scientific, Inc., Waltham, MA) and run at 130 V for 90 min using 1X MOPS running buffer (diluted from 20X MOPS SDS Running Buffer, Thermo Fisher Scientific, Inc., Waltham, MA). For reference, SeeBlue® Plus2 Pre-Stained Protein Standard (Thermo Fisher Scientific, Inc., Waltham, MA) was loaded into wells flanking the samples. Following electrophoresis, gels were washed in nanopure water. Proteins were transferred to Immobilon-P® PVDF membrane (Bio-rad, Hercules, CA) using a semi-dry protocol in 20% methanol/80% 1x MOPS. Transfer proceeded at 80 mA per gel for 55 min using a Trans-Blot® SD Semi-Dry Transfer Cell (Bio-rad, Hercules, CA). Blots were blocked overnight in 5% (m/v) fat-free dry milk at 4°C. Primary antibody (Sigma, Cat. #H1029, St. Louis, MO) was diluted 10,000x in PBS and applied to blots for 2 hrs. Secondary antibody conjugated to horseradish peroxidase (Bio-rad, Cat. #1701011, Hercules, CA) was diluted 3,000x in PBS-T and applied to blots for 1 hr. Finally, His-tagged proteins were visualized using the Immobilon-P® Opti-4CN™ Colorimetric kit (Bio-rad, Hercules, CA).

Multiplex Advanced Genome Engineering (MAGE) Cycling

To perform a single cycle of MAGE, a 5 mL liquid culture of the strain of interest was grown to an OD₆₀₀ of 0.6–0.8 in LB media at 32°C and 250 rpm, after which the culture was transferred to a 42°C water bath for 15 minutes. Next, the culture was immediately put on ice for at least 5 minutes. 1.5 mL of culture was transferred to a microcentrifuge tube and washed 3x in ice-cold nuclease-free water by pelleting cells for 1 min at 13,500 x *g* followed by resuspension in water. After the third wash, cells were resuspended directly in a solution of MAGE oligo (5–20 μM). Oligo was introduced into cells using a Micropulse electroporator (Bio-Rad, Hercules, CA). Immediately following electroporation, cells were resuspended in 5 mL sterile LB media – the resulting 5 mL culture was used to begin the next cycle.

Replica Plating of Bacterial Colonies

To identify colonies that had regained sensitivity to a given antibiotic, plates were replicated. To begin, cells from a mixed population (some of which would be sensitive to the antibiotic, and some of which would not be) were spread onto LB agar plates *lacking* the antibiotic and grown overnight at 34°C to produce a series of source plates. Then, for each source plate a sterile velvet was fastened to a replica plating tool (VWR, Radnor, PA). A source plate was gently pressed down onto the surface of the velvet such that the velvet picked up some cells from each colony present. Next, a new plate *lacking* the antibiotic was pressed onto the velvet, after which a new plate *containing* the antibiotic was pressed onto the velvet. All plates were returned to 34°C overnight. Ultimately, colonies that were present on the source plate as well as the replica plate lacking antibiotic but absent from the replica plate containing antibiotic were identified as having regained sensitivity to the antibiotic.

Knockout of *ompT* Locus

In order to use kanamycin resistance to select for successful knockout of the *ompT* locus in C321.ΔA.759.T7, the kanR cassette first employed to select for integration of the T7RNAP insert needed to be removed from the genome. This DNA was physically looped out of the genome using the oligo listed in [Table S1](#) for MAGE. Cultures were grown in LB media at 32°C and 250 rpm throughout 8 MAGE cycling steps as described above. To identify colonies that regained sensitivity to kanamycin, cells were replica plated as described above. Colony PCR using the protocol described above confirmed that the kanamycin resistance cassette DNA was no longer present in the genome. The kanamycin resistant cassette from pKD4 was then amplified with primers containing up- and downstream homology to the genomic region targeted for deletion in their 5' tails. The knockout construct was given flanking homology such that coordinates 580,650–592,260 in the genome of C321.ΔA.759.T7 would be replaced by the resistance cassette. ΔHR followed by colony PCR detection of the knockout were performed as described above to yield C321.ΔA.759.T7.Δ*ompT*.

Generation and Verification of OmpT-resistant T7RNAP-expressing Strains

Nucleotide changes designed to introduce mutations of K183 to glycine/leucine and K190 to alanine were installed into C321.ΔA.759.T7's genomic copy of the N-terminally 6His-tagged *1* gene via MAGE (described above) using the oligos listed in [Table S1](#). Cultures were grown in selective LB media at 32°C and 250 rpm throughout 6 MAGE cycling steps. Putative mutant colonies were picked and cultured as described above prior to screening for the desired mutation. Multiplex allele-specific colony (MASC) PCR was performed with Colony PCR Master Mix (Thermo Fisher Scientific, Inc., Waltham, MA) to verify mutations ([Wang and Church, 2011](#)) using wild-type forward or mutant forward primers and reverse primers ([Table S1](#)). Wild-type and mutant forward primers were identical except at the 3'-ends of the oligonucleotides which featured allele-specific sequence such that stable annealing of the end of the primer should only be possible when paired with the corresponding genomic allele. In this way the mutant allele could be amplified using the mutant forward and reverse primer set but not amplified by the wild-type forward and reverse primer set, and vice versa. The reverse primers were used for detection of both wild-type and mutant alleles.

pMAZ Plasmid Assembly

All primers and oligos described in this section are presented in [Table S1](#). pMAZ-ΔHis was assembled from two parts: linear plasmid backbone derived from PCR amplification of pMAZ-SK (primers pMAZbb_F/pMAZbb_R), and annealed oligos encoding the target gRNA sequence (T7delHis_oligo1 and T7delHis_oligo2). To anneal, 10 μL of a 100 μM stock for each oligo were combined with 10 μL CutSmart® buffer (New England Biolabs, Ipswich, MA) and 70 μL nuclease-free water, and the mix was heated to 95°C for 5 mins before being returned to room temperature. Insertion of the annealed oligos into the plasmid backbone was performed with USER® Enzyme (New England Biolabs, Ipswich, MA). To assemble pMAZ-Cure, pMAZ backbone DNA was amplified using primers pMAZ-Curebb_F/pMAZCurebb_R. DNA insert encoding gRNAs against the origin of replication and ampicillin resistance gene of pMA7CR_2.0 was purchased from Life Technologies (Carlsbad, CA). This pMAZ backbone and gRNA cassette insert were assembled via Gibson assembly to yield pMAZ-Cure. Sanger sequencing confirmed correct sequences of both pMAZ plasmids.

CRMAGE Removal of N-terminal His-tag from C321.ΔA.759.T7.D

All primers and oligos described in this section are listed in [Table S1](#). In preparation for CRMAGE, genomically-encoded resistances to ampicillin and kanamycin were removed from C321.ΔA.759.T7.D to allow for the CRMAGE plasmids to be selected for. This was achieved via MAGE to loop the resistance genes out of the genome followed by replica plating to identify sensitized colonies, yielding strain C321.ΔA.759.T7.D.Δ*AbR*. To begin CRMAGE, pMA7CR_2.0 was transformed into C321.ΔA.759.T7.D.Δ*AbR* where it was stably maintained throughout the process. For each cycle of CRMAGE, the strain was grown to an OD₆₀₀ of ~0.6–0.8, incubated for

15 minutes at 42°C, and made electrocompetent via washing with cold nuclease-free water. Next, pMAZ-ΔHis and MAGE oligo T7delHis_MAGE were transformed into the strain simultaneously. After electroporation, cells were allowed to recover for 1 hour in 5 mL SOC media at 34°C at 250 RPM. After 1 hr, 50 μg/mL kanamycin was added to the recovery media to begin selecting for the pMAZ plasmid and the culture was left to continue shaking for 2 hours. Next, a total of 3 hours post-transformation, 1 mL of recovery culture was transferred to 4 mL of fresh SOC media with 50 μg/mL carbenicillin, 50 μg/mL kanamycin, and 400 ng/mL anhydrotetracycline (aTc), and put back at 34°C, 250 RPM for another 3 hours. Finally, the culture was diluted 1:100 in nanopure water and plated on LB plates with 50 μg/mL carbenicillin, 50 μg/mL kanamycin, and 400 ng/mL anhydrotetracycline. Plates recovered at 34°C overnight. Individual colonies were sequenced to verify incorporation of T7delHis_MAGE and, by extension, removal of the N-terminal His-tag from the genomic *1* gene in the strain. Once removal of the His-tag was detected, cells were grown to an OD₆₀₀ of ~0.6–0.8 and induced with 400 ng/mL aTc and 2% (m/v) rhamnose to induce destruction of pMAZ-ΔHis. Replica plating identified cells that had lost pMAZ-ΔHis via their regained sensitivity to kanamycin. To remove all CRMAGE-associated plasmids from the strain, plasmid pMAZ-Cure was introduced into the cells. pMAZ-Cure encodes a gRNA cassette targeting the selectable marker and origin of replication of pMA7CR_2.0. such that induction of the system with both aTc and rhamnose triggers elimination of both plasmids. pMAZ-Cure-bearing cells were grown to an OD₆₀₀ of ~0.6–0.8 and induced with 400 ng/mL aTc and 2% (m/v) rhamnose to trigger the simultaneous destruction of both pMAZ-Cure and pMA7CR_2.0 in the cells. Replica plating identified colonies that had lost both plasmids to yield strain 759.T7.Opt.

Expression and Purification of His-tagged Orthogonal pAcF tRNA Synthetase

BL21 (DE3) harboring a pY71 plasmid encoding pAcFRS were grown in 1L of 2xYTPG to an OD₆₀₀ of 0.6 at 250 rpm and 37°C. At that point, synthetase expression was induced by adding 1mM IPTG and the culture was moved to 30°C and allowed to continue to grow for 4 hrs. Cells were pelleted by centrifuging for 15 min at 5000 × g at 4°C, washed three times with cold S30 buffer, and stored at -80°C overnight. To purify pAcFRS, cell pellets were thawed on ice and resuspended in 5 mL of 50 mM Tris/300 mM NaCl per gram of wet cell pellet. Cells were lysed using an EmulsiFlex-B15 homogenizer (Avestin, Ottawa, Canada) with three passes at a pressure of 12,000 psig and centrifuged at 10,000 × g for 10 mins. To purify pAcF synthetase from the supernatant, Ni-NTA agarose (Qiagen, Venlo, The Netherlands) was used following manufacturer protocol. Following elution, purified synthetase was dialyzed overnight at 4°C in a Slide-a-Lyzer™ cassette (10 kDa MWCO; Thermo Fisher Scientific, Inc., Waltham, MA) against S30 buffer with 25% glycerol. After dialysis, synthetase was concentrated with Amicon® Ultracel centrifugal spin filters (3 kDa MWCO; MilliporeSigma, Burlington, MA). Final synthetase concentration was determined with a NanoDrop™ spectrophotometer (Thermo Fisher Scientific, Inc., Waltham, MA) using an extinction coefficient of 20985, determined using ExPASy ProtParam tool (Gasteiger et al., 2005).

ELP Radioactive Quantitation

Radioactive ¹⁴C-Glycine was added into 15 μL CFPS reactions. After incubation, yields were quantified by determining radioactive ¹⁴C-Gly incorporation into trichloroacetic acid (TCA) -precipitated protein (Swartz et al., 2004). Briefly, following CFPS reactions were quenched with 100 μL 0.1N sodium hydroxide and put at 37°C for 30 minutes. Two small tabs of Whatman 3MM chromatography paper (GE Healthcare Life Sciences, Little Chalfont, United Kingdom) were prepared for each individual radioactive CFPS reaction and suspended on pins above a block of Styrofoam wrapped with aluminum foil. Following the sodium hydroxide incubation, reactions were mixed by pipetting. Next, 50 μL of material from each reaction vessel was deposited onto both paper tabs prepared for that vessel. The population of loaded paper tabs was split in half, into a “washed” subpopulation and an “unwashed” subpopulation, such that a single tab derived from each CFPS reaction was present in each subpopulation. Tabs were allowed to dry under a heat lamp for ~2 hr. After drying, all “washed” subpopulation tabs were combined in a beaker and washed 3x with ice-cold 5% TCA at 4°C. Following the third wash, these tabs were washed with 100% ethanol for 10 mins at room temperature. Washed tabs were remounted on Styrofoam and dried under a heat lamp for ~2 hr. Finally, all tabs (“washed” and “unwashed”) were individually deposited into 1.5 mL microcentrifuge tubes and submerged into 1 mL of scintillation fluid. Radioactivity of TCA-precipitated samples in terms of counts per minute (cpm) was measured using liquid scintillation counting (MicroBeta2, PerkinElmer, Waltham, MA). With radioactive counts in hand, protein yield for each sample could be calculated according to the following equation:

$$\text{Yield} \left(\frac{\mu\text{g}}{\text{mL}} \right) = \frac{\frac{\text{Washed cpm} - \text{Background cpm}}{\text{Unwashed cpm}} * 2010 \mu\text{M Glycine} * \text{Protein Molecular Weight} \left(\frac{\mu\text{g}}{\mu\text{mol}} \right)}{\# \text{ Glycine residues per protein} * \frac{1000 \text{ mL}}{1\text{L}}}$$

Autoradiogram Analysis

For autoradiogram analysis of ELP constructs, CFPS reactions were performed supplemented with 10 μM of radioactive ¹⁴C-glycine. Samples were prepared for polyacrylamide gel electrophoresis as described above from 4 μL of each reaction and run on a 12% Bis-Tris NuPAGE® gel (Thermo Fisher Scientific, Inc., Waltham, MA). After electrophoresis, the gel was soaked in Gel Drying solution (Bio-Rad, Hercules, CA) for 30 min, fixed with cellophane films, dried overnight in GelAir Dryer (Bio-Rad, Hercules, CA), and exposed for 3 days on Storage Phosphor Screen (GE Healthcare Biosciences, Pittsburgh, PA). Autoradiograms were scanned using Typhoon FLA 7000 Imager (GE Healthcare Biosciences, Pittsburgh, PA).

Purification and Top Down Mass Spectrometry of sfGFP and ELP Constructs

To prepare sfGFP and ELP-20mer/30mer samples for top-down mass spectrometry, products were purified out of CFPS reactions using Strep-Tactin®XT magnetic beads (IBA Lifesciences, Göttingen, Germany) precipitated in methanol/chloroform and water (Toby et al., 2019), dried, and resuspended in Buffer A (95% water, 5% acetonitrile, 0.2% formic acid). ELP-40mers were purified using a modified inverse transition cycling (ITC) method as previously described (Martin et al., 2018), which was sufficient to purify ELP-40WT. ELP-40UAG products required further enrichment, and for this construct ITC was followed up with affinity purification using Strep-Tactin®XT magnetic beads following manufacturer's instructions to generate a highly pure sample of the protein. Purified proteins were injected onto a trap column (150 μ m ID x 3 cm) coupled with a nanobore analytical column (75 μ m ID x 15 cm). Trap and column were packed with polymeric reverse phase (PLRP-S, Phenomenex, Torrance, CA) media (5 μ m, 1,000 Å pore size). Samples were separated using a linear gradient of Buffer A and Buffer B (5% water, 95% acetonitrile, 0.2% formic acid). Samples were loaded for 10 min onto the trap column and subsequently separated using a linear gradient from 5% to 95% of solvent B (80 min for sfGFP samples and ELP-20mer/30mer, 60 min for ELP-40mer). Mass spectrometric data for all sfGFP and ELP-20mer/30mer samples were obtained on a Orbitrap Fusion™ Lumos™ (Thermo Fisher Scientific, Inc., Waltham, MA) instrument fitted with a custom nanospray ionization source. The acquisition method was essentially a full scan FTMS experiment, with data obtained from 500–2000 m/z at a resolving power of 120,000 at m/z 400. sfGFP data were deconvoluted using Xtract (Thermo Fisher) and monoisotopic masses were reported. ELP-20mer/30mer data were deconvoluted using Esiprot (Winkler, 2010) and average masses were reported. Mass spectrometric data for ELP-40mer samples were obtained on a Velos Pro ion trap (Thermo Fisher Scientific, Inc., Waltham, MA) fitted with a custom nanospray ionization source. The acquisition method was a full scan using the ion trap. ELP-40mer data were deconvoluted using UniDec (Marty et al., 2015).

DNA Sequencing

To sequence the genomic T7RNAP inserts, the entire region was PCR amplified using end primers listed in Table S1. Amplified linear insert DNA was submitted to the NUSeq Core facility along with forward primers spaced ~700 bp apart, and the sequence for each region was determined using traditional Sanger sequencing. pMAZ plasmids were sequenced by submitting purified plasmid samples as well as primer pMAZ_seq to the NUSeq Core facility.

QUANTIFICATION AND STATISTICAL ANALYSIS

Quantified data in this work consist exclusively of protein yields from various CFPS platforms. As outlined in figure legends, these data present average yields from at least three independent CFPS reactions along with one standard deviation. All calculations were performed using Microsoft Excel.

DATA AND CODE AVAILABILITY

Data for this study will be provided upon reasonable request and agreement. Such requests will be fulfilled by the Lead Contact, Michael C. Jewett (m-jewett@northwestern.edu).