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Translation of Diverse Aramid- and 1,3-Dicarbonyl-peptides by Wild Type Ribosomes in Vitro

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Supporting Information

ABSTRACT: Here, we report that wild type Escherichia coli ribosomes accept and elongate precharged initiator tRNAs acylated with multiple benzoic acids, including aramid precursors, as well as malonyl (1,3-dicarbonyl) substrates to generate a diverse set of aramid-peptide and polyketide-peptide hybrid molecules. This work expands the scope of ribozyme- and ribosome-catalyzed chemical transformations, provides a starting point for in vivo translation engineering efforts, and offers an alternative strategy for the biosynthesis of polyketide-peptide natural products.

INTRODUCTION

As far as we know, ribosomes have evolved for billions of years to perform a single reaction—formation of an amide bond between two α-amino acid substrates brought into proximity by tRNAs within the ribosome active site, the peptidyl transferase center (PTC). In cells and extracts, the chemistry possible within a wild type ribosome PTC has expanded to include reactions of more than 200 different nonproteinogenic α-amino and hydroxy acids;1−5 ribosomes containing remodeled PTCs support amide bond formation to and from a small number of β-amino acids6−7 and dipeptides8,9 with limited efficiency. The combination of cell-free in vitro translation systems and ribozyme-catalyzed tRNA acylation reactions offers the opportunity for even greater reaction diversity, including the introduction of multiple N-alkyl10, p-α-,11,12 α-hydroxy,13 and β-amino acids,14,15 the precursors of β-peptide foldamers.16−19

A second family of foldamer-like molecules are aramids, oligomers of substituted aminobenzoic acids.20 Aramids possess remarkably varied properties. Kevlar, a polymer of 1,4-phenylenediamine and terephthaloyl chloride, is a strong and heat-resistant fiber,21 whereas cystobactamids are DNA gyrase inhibitors active against Gram-negative bacteria.22 Many other aramid foldamers with diverse and significant properties have been reported.23−25 Recently, wild type E. coli ribosomes were shown to accept and elongate initiator tRNAs precharged with aromatic foldamer-dipeptide appendages.26 Notably, in this case the foldamer monomers did not themselves react within the PTC, being displaced from the reaction center by a Gly–Phe dipeptide spacer.26,27 Here, we report that wild type E. coli ribosomes accept and elongate precharged initiator tRNAs acylated directly with multiple benzoic acids, including aramid precursors, as well as malonyl (1,3-dicarbonyl) substrates. The result is a diverse set of aramid-peptide and polyketide-peptide hybrid molecules. This work provides new knowledge about the generality of flexizyme-promoted tRNA acylation reactions, expands the scope of ribosome-catalyzed chemical transformations, provides a starting point for in vivo translation engineering efforts, and offers an alternative strategy for biosynthesis of polyketide-peptide natural products.

RESULTS AND DISCUSSION

As the first step toward the ribosomal synthesis of aramid-like peptides, we made use of an established microhelix (MH) gel-shift assay26 and high-resolution mass spectrometry (Figure 1A) to evaluate whether the cyanomethyl esters of unsubstituted aminobenzoic acids were substrates for the flexizyme ribozyme eFx.20 Incubation of cyanomethyl esters 1−3 (5 mM) with 25 μM microhelix MH and 25 μM eFx (Table S1) in bicine buffer at pH 9 for 48 h showed little or no evidence of MH acylation when the reaction products were evaluated on an acid-urea PAGE gel (Figure 1B). A low level of MH acylation by the o- and m-analogues 1 and 2 (and a trace with 3) could be observed using a highly sensitive RNase A/ LC-HRMS assay30 that detects the acylated adenine nucleoside (Figure 1C). We also investigated the extent of tRNA acylation using the alternative ribozyme dFx31 and the 1,3-dinitrobenzyl esters of p- and o-aminobenzoic acid (4 and 5, respectively, as shown in Scheme S1). These substrates also failed to yield the expected MH products when incubated with dFx under standard conditions14 and analyzed using acid-urea gels or
RNase A/LC-HRMS (Figure S1), perhaps due to insolvability. Even the more soluble cyanomethyl ester of ortho-aminonicotinic acid analogue 6 reacted poorly in the presence of eFx (Figure S1).

The inability to efficiently acylate MH or tRNA with simple aminobenzoic acids in high yields using eFx or dFx led us to consider chemical acylation methods for the preparation of these materials. Isotopic anhydride can acylate the terminal 2'- or 3'-OH group of an unprotected tRNA, and the resulting anthranoyl-tRNA (o-AN-tRNA) retains the ability to associate productively with EF-Tu-GTP. Inspired by this reactivity, we incubated E. coli tRNAVal (ValT) or initiator tRNA (tMetT) with 8–80 mM isatoic anhydride in 90% CH3CN containing 2–5 mM NaOH for 3 h at 37 °C, digested the products with RNase A, and used LC-HRMS to detect the formation of nucleoside 7 (m/z = 387.1411, Scheme S1); this product will be observed only if reaction occurs at the tRNA 3'-end (Figure S2A). A peak corresponding to this mass was observed only in reactions containing tRNA, isotopic anhydride, and base; in the absence of base, the acylation efficiency dropped by 1–2 orders of magnitude (Figure S2B). Mindful of the fact that isatoic anhydride reagents can also modify RNA on the 2'-OH group of internal ribose residues in SHAPE reactions,53 we also evaluated the reaction using ultra-performance liquid chromatography (UPLC), which (as expected) showed evidence of multiple reaction products, whereas eFx-promoted reactions did not (Figure S2C).

We next made use of a commercial in vitro translation kit (PURExpress Δ; aa, tRNA) to evaluate if an initiator tRNA (tMetT) acylated with o- (prepared using isotopic anhydride) or m-aminobenzoic acid (prepared using eFx) would be accommodated by the P-site of wild type E. coli ribosomes and initiate translation. We supplemented the kit with the requisite amino acids and tRNAs, precharged initiator tRNA (o- or m-AN-tRNA) (50–100 μM), and a duplex DNA template (0.5–1 μg) encoding the FLAG-containing polypeptide MVFDYKDDDDK (MVF-FLAG). After a 6 h incubation, the reaction mixture was treated with Ni-NTA resin to remove all PURExpress Δ components (which are His6-tagged), and the remaining material was analyzed by LC-HRMS (Figure 2A). If the o- or m-AN-tRNA initiates translation in place of an
FLAG polypeptide was detected in the presence of DNA template and m-AN-tRNA.

Aminobenzoate esters hydrolyze exceptionally slowly, suggesting that the electron-rich aromatic ring contributes to the low reactivity of 1–3. In addition, the structure of the ethyl ester of L-phenylalanine bound to Fx (as an Fx-tRNA fusion) shows pi-stacking between Fx base guanine 24 and the L-phenylalanine aromatic ring; this stacking would be less favorable with an electron-rich arene. To investigate whether reactivity in eFx-promoted reactions was correlated with arene electron density, we prepared a diverse set of substituted benzoic acid cyanomethyl esters (Figure 3A) and evaluated the extent to which eFx reactivity correlated with the sign and magnitude of the relevant sigma factor, which measures the inductive effect of the aromatic substituent. As expected, benzoic acid cyanomethyl esters possessing strong electron-withdrawing substituents, such as penta-fluoro 8, p-nitro 9 (σ = +0.78), or p-Cl 10 (σ = +0.23), were excellent eFx substrates in model MH reactions, with 78-99% yields (Figure 3B,C). However, other factors are clearly important: a benzoic acid cyanomethyl ester possessing a weak electron-withdrawing substituent, such as p-azido 11 (σ = +0.08), was also an excellent substrate (yield of acylated MH = 74%), as were analogues possessing both strong and weak electron-donating substituents, such as p-methoxy 12 (σ = −0.27; yield of acylated MH = 62%) and p-methyl 13 (σ = −0.17; yield of acylated MH = 54%). Notably, the poorest yields were observed in eFx-promoted reactions of substrates 6 (yield of acylated MH = 25%) and 15 (yield of acylated MH = 23%), all of which contain one or more acidic protons/hydrogen bond donors, just like amino benzoic acids 1, 2, and 3. These results imply that the presence of hydrogen bond donors in certain positions contributes to the poor reactivity of amino benzoic acids 1–3. Consistent with this notion, p-hydroxybenzoic acid 16 [pKa = 8.3 (p-hydroxybenzoic acid methyl ester)] was a poor substrate, whereas alcohol 17 [pKa = 15 (benzyl alcohol)] and aldehyde 18 reacted well (Scheme S1 and Figure S4). It is possible that certain hydrogen bond donors alter the position of the aromatic ring in the eFx active site or coordinate and inactivate functional groups involved in catalysis. Determining the exact nature of these interactions is beyond the scope of this discussion but will be essential to effectively engineer new ribozymes that accept diverse foldamer building blocks in vitro and in vivo.

With a new set of aramid substrates in hand, we used the PURExpress Δ(aa, tRNA) in vitro translation kit to evaluate if initiator tRNAs acylated with diverse benzoic acids could be accommodated in the ribosomal P-site and initiate translation of an AR-VF-FLAG polypeptide carrying an aramid monomer (AR) at the N-terminus (Figure 4). Every benzoic acid cyanomethyl ester that acylated the microhelix MH with a yield >50% in an eFx-promoted reaction (Figure 3) was used to acylate fMetT, and translation reactions were performed and analyzed as described above. With one exception, every single AR-fMetT initiated translation of an AR-VF-FLAG peptide whose mass corresponded to incorporation of the prescribed substituted benzoic acid. The singular exception was p-azidobenzoic acid 11; in this case the mass of the isolated polypeptide was consistent with in situ reduction of the azide to an amine. These results demonstrate that diverse aramid-like monomers can be accommodated directly within the ribosomal P-site and act as acceptors for a natural α-amino acid in the A-site. They show further that use of p-azidobenzoic acid 11 effectively circumvents the poor reactivity of p-aminobenzoic acid 3 to generate a polypeptide with a p-aminobenzoic acid monomer at the N-terminus. The observation that wild type E. coli ribosomes can initiate translation using tRNAs acylated with diverse aramid-like monomers significantly expands the scope of in vitro translation reactions beyond that of Kawakami and lays the initial groundwork for the biosynthesis of genetically encoded, sequence-defined polyaramid oligomers.

We next sought to evaluate the relative efficiency of PURExpress reactions initiated with differentially acylated fMetT derivatives. To begin, we monitored the yield of fMet-VF-FLAG (approximated as the extracted ion abundance) as a function of time in PURExpress Δ reactions supplemented with either 50 μM precharged fMetT-fMet (charged using the dFx substrate fMet-DBE) or 50 μM L-methionine. The bulk of both reactions was complete within 100 min, but the yield of fMet-VF-FLAG in reactions supplemented with precharged fMetT-fMet was 1.5% of that obtained in reactions supplemented with L-methionine (Figure S5A). Next, we
compared the extracted ion abundance (after 30−90 min) of the AR-VF-FLAG peptide initiated with fMetT precharged with benzoic acid ester. The yield of this AR-VF-FLAG polypeptide (C6F5-VF-FLAG) was 25−30% of the yield of fMet-VF-FLAG (generated in reactions supplemented with precharged fMetT-fMet) (Figure S5B) and within the range observed when translation was initiated with fMetT precharged with natural amino acids. The relative yields of AR-VF-FLAG peptides initiated with other precharged fMetT derivatives were also comparable (Figure S5D), suggesting similar initiation efficiencies. We note that when ValT was precharged with β-Phe, the yield of fMet-β-Phe-F-FLAG was 5-fold higher than the yield of fMet-VF-FLAG generated with precharged fMetT (Figure S5C). As initiation complex assembly is the rate limiting step during translation, the higher yield of fMet-β-Phe-F-FLAG relative to fMet-VF-FLAG is likely due to the difficulty assembling the translation initiation complex using non-natural fMetT derivatives. Benzoic acid monomers that were poor endFx substrates (yields <50%) in model MH reactions, such as 6 and 15 (Figure 3), failed to detectably initiate peptide synthesis from WT ribosomes in vitro. This observation suggests that the ribosome is largely agnostic of aramid structure, and that the concentration of non-natural fMetT derivative, rather than monomer structure, determines the reaction yield in PURExpress reactions.

Like aramid natural products,27 polyketide-peptide hybrid molecules are biosynthesized by mega-assemblies of complex protein enzymes,41−43 the combination of peptide and polyketide-based functionality can translate into highly unique biological functions.46−48 To evaluate whether wild type E. coli ribosomes are capable of biosynthesizing a polyketide-peptide hybrid, we prepared malonate derivatives 19−23 (Figure 5A).
reactions led to the isolation of polypeptides carrying malonates 22 and 23 (22-VF-FLAG and 23-VF-FLAG, respectively), whose masses were confirmed by LC-HRMS (Figure 5D). The yield of 23-VF-FLAG, estimated as described above, was approximately 20% of the yield of fMet-VF-FLAG produced in reactions supplemented with precharged fMetT (Figure S3A). We conclude that extant E. coli ribosomes have the capacity to biosynthesize simple polyketide-peptide hybrid molecules.

In summary, here we report that wild type E. coli ribosomes accept precharged initiator tRNAs acylated with multiple substituted benzoic acids, including the monomeric unit of Kevlar, as well as malonyl (1,3-dicarbonyl) substrates. The ribosome then elongates these substrates to generate a diverse set of aramid-peptide and polyketide-peptide hybrid molecules. This work expands the scope of reactions catalyzed by both flexzyme and wild type ribosomes, provides a starting point for in vivo translation engineering efforts, and offers an alternative strategy for biosynthesis of polyketide-peptide natural products.

Safety Statement. No unexpected or unusually high safety hazards were encountered during the execution of these experiments.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available on the ACS Publications website at DOI: 10.1021/acscentsci.9b00460.

Synthesis and characterization of flexzyme substrates; and procedures for formation, characterization, purification, and analysis of tRNA acylation and IVT reactions (PDF)

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Notes

The authors declare no competing financial interest.

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