DNA-catalyzed reactivity of a phosphoramidate functional group and formation of an unusual pyrophosphoramidate linkage†

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During in vitro selection for DNA-catalyzed lysine reactivity, we identified a deoxyribozyme that instead catalyzes nucleophilic attack of a phosphoramidate functional group at a 5′-triphosphate-RNA, forming an unusual pyrophosphoramidate (N–PⅤ–O–PⅤ) linkage. This finding highlights the relatively poor nucleophilicity of nitrogen using nucleic acid catalysts, indicating a major challenge for future experimental investigation.

Introduction

Deoxyribozymes are specific DNA sequences that have catalytic activity.1–5 We have recently focused on expanding deoxyribozyme catalysis to include reactions of amino acid side chains,6–8 with the long-term goal of DNA-catalyzed modification of large proteins. Our initial effort in this direction demonstrated robust DNA catalysis (>70% yield in 2 h) of nucleopeptide linkage formation between the nucleophilic tyrosine (Tyr) phenolic OH side chain and an electrophilic 5′-triphosphate-RNA.6 In parallel, however, catalysis involving the serine (Ser) aliphatic hydroxyl side chain was extremely poor (only ~0.2% yield), and reactivity of the lysine (Lys) amine side chain was not observed. Lys side chain reactivity has never previously been observed with either DNA or RNA enzymes, suggesting that this reactivity is a particular challenge. One report found RNA-catalyzed reaction of a peptide N-terminal α-amino group even when the Lys ε-amino side chain was available as a competing nucleophile.9

Our initial study on DNA-catalyzed side chain reaction6 used a substrate that presented a single amino acid residue in a highly preorganized three-helix-junction (3HJ) architecture (Fig. 1, Z = single amino acid).10,11 Subsequently, we found that expanding the substrate to contain a tripeptide (Z = Ala-Ser-Ala) enabled robust Ser reactivity by newly identified deoxyribozymes.7 Separately, by discarding the 3HJ preorganization altogether, we identified deoxyribozymes that function with free peptide substrates.8 In the present report, we sought DNA-catalyzed Lys reactivity using the 3HJ preorganized architecture (Z = Ala-Lys-Ala). However, we instead identified a DNA-catalyzed reaction between a phosphoramidate functional group in the substrate and 5′-triphosphate-RNA, forming an unusual pyrophosphoramidate linkage. The results provide a clear calibration point on the relative (un)reactivity of amines using nucleic acid catalysts, which is important information for our ongoing efforts to expand DNA catalysts to include the Lys side chain.
Results and discussion

In vitro selection of new DNA catalysts that covalently modify the Lys-containing substrate

We began our efforts to seek DNA-catalyzed Lys reactivity by retaining the 3HJ architecture of Fig. 1 while expanding the substrate to include an Ala-Lys-Ala tripeptide rather than the single Lys amino acid used previously,6 noting that analogous expansion successfully enabled Ser reactivity.7 In vitro selection12,13 was used to identify individual DNA sequences with catalytic ability, following our established approach.7,14,15 Briefly, the 5′-triphosphate-RNA electrophile was covalently attached at its 3′-terminus to the DNA pool, which initially incorporated a total of 40 fully random nucleotide positions partitioned between two loops. Specific DNA sequences that can join the Lys-containing substrate to the 5′-triphosphate-RNA were separated by PAGE from the vast majority of nonfunctional DNA sequences and amplified by PCR. Attachment of the resulting DNA pool, now enriched in functional sequences, to a fresh sample of 5′-triphosphate-RNA enabled the next selection round to be performed. By iteration of this process for many rounds, catalytically active deoxyribozymes dominate the pool and can be cloned and characterized in detail.

In each selection round’s key step, during which the Lys-containing substrate can become attached to the 5′-triphosphate-RNA, we used incubation conditions of 50 mM HEPES, pH 7.5, 150 mM NaCl, 2 mM KCl, 20 mM MnCl2, and 40 mM MgCl2 at 37 °C for 2 h. Mn2+ was included because of its previous utility in supporting both Tyr and Ser reactivity.6,7 and Mg2+ was used because of its value as a cofactor for a wide range of deoxyribozymes.14 The selections were initiated for seven rounds using an Ala-Lys-Ala substrate that included a 3′-phosphate that was intended to block nucleophilic reactivity of the 3′-terminus. However, this tactic proved ineffective, leading the majority of the pool to catalyze unwanted nucleophilic reaction of the 3′-phosphate with the 5′-triphosphate-RNA (Fig. S1 in the ESI†). We therefore redirected the selection effort16 by replacing the 3′-phosphate group with a 2′,3′-dideoxytideidine nucleotide (3′-ddC), which lacks any strong nucleophile. The selection process was resumed from round 8, and by round 13, 12% ligation activity of the pool was observed. On the basis of PAGE migration rates, the uncloned products have a linkage to the RNA substrate that is either within or very near to the tripeptide region (Fig. S2A in the ESI†). However, an Ala-Ala-Ala substrate was comparable in reactivity to Ala-Lys-Ala, demonstrating that the linkage to the RNA was created not directly at the Lys side chain but instead at a nearby functional group. Upon cloning, a single deoxyribozyme sequence named 13LS3 was identified and found to form a product assignment was made by excluding all other reasonable candidates. Treatment with 80% acetic acid is known to hydrolyze phosphoramidate (P–N) bonds,18 and the tripeptide-containing substrate has a phosphoramidate linkage directly to the N-terminal side of the tripeptide moiety (Fig. 1). Therefore, treatment of the 13LS3 ligation product with 80% acetic acid should lead to a readily distinguishable PAGE pattern that depends sensitively on the site of nucleophilic reactivity within the tripeptide-containing substrate. Overall, the observations in Fig. 2 compel the conclusion that the product has the structure of pyrophosphoramide 1. The data exclude the alternative, isomeric diphasphoramidate 2, which would have been formed by nucleophilic attack of the phosphoramidate N atom at the 5′-triphosphate. Product 2 would be susceptible to cleavage by 80% acetic acid, considering that a simple phosphoramidate is itself highly reactive under such conditions,19 and 2 should be even more reactive.

The 13LS3 ligation product 1 was observed to regenerate a variable amount of the original tripeptide-containing substrate by cleavage of its pyrophosphate linkage. Note that this linkage is different from the P–N bond that is hydrolyzed by 80% AcOH within the phosphoramidate substrate (but not within 1). In particular, after separation by PAGE, the 13LS3 product 1 was partially cleaved during extraction from the polyacrylamide gel. Extraction in the conventional TEN buffer (10 mM Tris, pH 8.0, 300 mM NaCl, 1 mM EDTA) led to 40–50% substrate regeneration by pyrophosphate cleavage, whereas extraction in buffers of lower pH value (e.g., 10 mM HEPES, pH 7.0 or 10 mM NaOAc, pH 4.6) led to only 10–20% substrate regeneration. The extent of pyrophosphate cleavage during TEN extraction was time-dependent; e.g., 10–15% after 30 min extraction, but 40–50% after 2 h extraction. The regenerated substrate could itself be isolated by PAGE and was equally as competent for 13LS3-catalyzed reaction as fresh substrate (data not shown). Curiously, the substrate regeneration reaction does not occur when the
Reaction is directly analogous to formation of phosphoromorpholidate; of $3\textsuperscript{-32}$P-ATP and T4 PNK or either the Ala-Ala-Ala or Ala-Lys-Ala substrate, with connectivity. S for stability properties. which was found by Moffatt and Khorana to have similar extraction process.

The newly formed linkage is related to linkage 3 (Fig. 3), which was found by Moffatt and Khorana to have similar stability properties. These authors reported the formation of 3 by slow, uncatalyzed self-condensation of adenosine 5'-phosphoromorpholidate; i.e., attack of a phosphoramidate into an activated 5'-phosphorus (reaction A in Fig. 3). This uncatalyzed reaction is directly analogous to formation of 1 by DNA-catalyzed attack of the phosphoramidate linkage of the tripeptide-containing substrate into 5'-triphosphate-RNA (reaction B in Fig. 3), thereby providing a chemical precedent for the DNA-catalyzed reaction leading to 1. Moffatt and Khorana also reported that a simple phosphodiester is much less nucleophilic towards adenosine 5'-phosphoromorpholidate than is the phosphoromorpholidate itself. Consistent with this, we do not observe any DNA phosphodiester group within the tripeptide-containing substrate attacking the 5'-triphosphate-RNA.

Phosphoramidate linkages are found naturally as the lysine-adenylated intermediates of DNA ligase enzymes, among other instances. The 13LS3-catalyzed reaction of its substrate to form 1 rather than 2 indicates that the amphoteric phosphoramidate group within the substrate is more reactive at oxygen than at nitrogen, although 13LS3 necessarily offers only a single data point in this regard.

### Broader implications of the results

A key implication of the finding that 13LS3 forms pyrophosphoramide is that at least under the evaluated incubation conditions, which are typical for in vitro selection experiments with ribozymes and deoxyribozymes, the phosphoramidate functional group within the Ala-Lys-Ala substrate is much more nucleophilic (at oxygen) than is the originally intended Lys amine side chain (at nitrogen). This result highlights the relative unreactivity of nitrogen as a nucleophile with the assistance of nucleic acid catalysts. Our ongoing efforts towards DNA-catalyzed Lys reactivity include use of more reactive electrophiles than 5'-triphosphate, which is intended to counterbalance the relatively poor nucleophilicity of the Lys amine. More broadly, in combination with our observation that DNA-catalyzed peptide bond cleavage is disfavored relative to phosphodiester hydrolysis, the results indicate that any kind of DNA-catalyzed reactivity of nitrogen-containing substrates (where the N atom is either nucleophile or leaving group) is a major experimental challenge.
Experimental section

Materials were prepared and procedures were performed as described in our previous report. The reductive amination procedure is described in the ESI.† The 13LS3 sequence for the in trans (intermolecular) assays was 5′-CCGTCGCCATCTCCAAATAGGGAGGGACAAATGA-AAAAATGCAGCAGCAGTGGTTACCAGTTATCCGATTT-ATCC-3′, where the two underlined sequences are those of loops A and B derived from the N33+N7 random regions, and the three non-underlined sequences are those of the fixed P3, P1, and P4+P2 regions (in that order), all as shown in Fig. 1. Note that loop A of 13LS3 contains 34 rather than 33 nucleotides, presumably due to an insertion by Taq polymerase during one of the selection rounds. The two substrate sequences are shown in Fig. 1.

MALDI mass spectrometry was performed on 13LS3 reaction products that were each prepared as follows. A 560 µL sample containing 10 nmol of tripeptide-containing substrate, 10.5 nmol of 13LS3 deoxyribozyme, and 11 nmol of 13LS3-catalyzed reactions and had the expected mass: calcd. 5686.5, found 5685.4 (Δ = −0.02%).

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Notes and references