Sequence-function relationships provide new insight into the cleavage site selectivity of the 8–17 RNA-cleaving deoxyribozyme

Kenny Schlosser, Jimmy Gu, Lauren Sule and Yingfu Li*

Department of Biochemistry and Biomedical Sciences and Department of Chemistry, McMaster University, Hamilton, Ontario, Canada L8N 3Z5

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ABSTRACT

Many sequence variations of the 8–17 RNA-cleaving deoxyribozyme have been isolated through in vitro selection. In an effort to understand how these sequence variations affect cleavage site selectivity, we systematically mutated the catalytic core of 8–17 and measured the cleavage activity of each mutant deoxyribozyme against all 16 possible chimeric (RNA/DNA) dinucleotide junctions. We observed sequence-function relationships that suggest how the following non-conserved positions in the catalytic core influence selectivity at the dinucleotide junction: (i) positions 2.1 and 12 represent a primary determinant of the selectivity at the 3’ position (N1,i) of the cleavage site; (ii) positions 15 and 15.0 represent a primary determinant of the selectivity at the 5’ position (rN18) of the cleavage site and (iii) the sequence of the 3-bp intramolecular stem has relatively little influence on cleavage site selectivity. Furthermore, we report for the first time that 8–17 variants have the collective ability to cleave all dinucleotide junctions with rate enhancements of at least 1000-fold over background. Three optimal 8–17 variants, identified from ~75 different sequences that were examined, can collectively cleave 10 of 16 junctions with useful rates of ≥0.1 min⁻¹, and exhibit an overall hierarchy of reactivity towards groups of related junctions according to the order NG > NA > NC > NT.

INTRODUCTION

A small deoxyribozyme known as 8–17 catalyses the site-specific cleavage of RNA. This DNA enzyme has been the subject of much basic and applied research interest, due to several favourable characteristics including a simple secondary structure, relatively fast reaction rate and generalizable substrate recognition. The utility of the 8–17 deoxyribozyme has been demonstrated through a wide variety of applications including (i) the in vitro manipulation and analysis of RNA (1–3), (ii) fluorescent and colorimetric biosensors (4–6), (iii) DNA computational elements (7–9) and (iv) potential antiviral/gene control agents (10–12). Because of its simplicity, 8–17 has also been frequently employed as a model system in the demonstration of new concepts (13–21). Further investigation into 8–17’s structure and function is expected to facilitate the development of new and/or improved applications.

Thus far, several studies have provided various insights into the kinetic and mechanistic details (22,23), metal-ion dependency (24,25), folding (26–29) and sequence requirements of the 8–17 deoxyribozyme (25,30). Aside from the potential practical benefits, basic research on 8–17 has also been motivated by its widespread recurrence. Since 1997, this catalytic motif has been isolated from five independent in vitro selection experiments (24,30–34), making it the most common solution to deoxyribozyme-mediated RNA-cleavage in sequence space. Its recurrence can be largely attributed to the fact that only four of the ~15 nt that comprise its catalytic core are absolutely conserved, suggesting there may be thousands of active sequence variants. Understanding the functional significance of these many sequence variations represents a challenging, but important objective that has only recently begun to be addressed.

Peracchi and colleagues previously analysed the sequence requirements of the catalytic core to show how sequence variations influence the reaction rate (25). This study used a substrate that contains a 5’-AG-3’ dinucleotide junction as the target cleavage site, since 8–17 was originally selected to cleave this junction. However, Cruz et al. have recently demonstrated that sequence variations can also influence the selectivity of the cleavage site (34). In that study, 16 parallel in vitro selections were conducted, each using a substrate containing one of the

*To whom correspondence should be addressed. Tel: +1 90 55259149 ext. 22462; Fax: +1 90 55229033; Email: liying@mcmaster.ca

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16 possible dinucleotide junctions as the target cleavage site. Ultimately, more than a 100 different 8–17 variants were isolated at various frequencies in nearly all of the selections. The cleavage versatility of five representative 8–17 variants was tested, and together these variants can cleave 14 out of 16 junctions with rate enhancements ranging from $\sim 10^3$–$10^7$ fold over background. By comparison, the well-known 10–23 deoxyribozyme can reportedly cleave only four of the 16 junctions (33). These results prompted the hypothesis that 8–17 might actually have a latent capacity to cleave all 16 junctions proficiently, masked only by the right combination of nucleotides.

Herein, we wanted to investigate the functional limitations of 8–17 mediated catalysis. In particular, we sought to understand how the sequence of its catalytic core affects selectivity of the dinucleotide cleavage site. Towards this end, we conducted a systematic mutational analysis, and measured the effects against all 16 dinucleotide junctions. In addition to providing new insight into the structure and function of 8–17, we expect this study will also broaden its general utility.

**MATERIALS AND METHODS**

**Oligonucleotides and reagents**

Deoxyribozyme and substrate oligonucleotides were prepared by automated DNA synthesis using cyanethylphosphoramidite chemistry (Integrated DNA Technologies; Mobix Central Facility, McMaster University). Oligonucleotides were purified by 10% preparative denaturing (8 M urea) polyacrylamide gel electrophoresis (PAGE) and their concentrations were determined by spectroscopic methods. [$\gamma^{32}$P]ATP was purchased from Amersham Pharmacia. T4 polynucleotide kinase (PNK) was purchased from MBI Fermentas. All chemical reagents were purchased from Sigma.

**Kinetic methods**

To assess the cleavage ability of more than 75 different mutants against 16 different substrates, simple assays based on single-timepoint measurements were preferred to complete kinetic characterizations. A similar strategy has been used previously to determine the sequence requirements in the catalytic core of the 10–23 deoxyribozyme (35). Each 8–17 mutant was separately incubated with each 5',32P-labelled substrate in trans under single-turnover conditions (deoxyribozyme:substrate 800:1). Substrate and deoxyribozyme were heated together at 90°C for 30 s, and allowed to cool at room temperature for $\sim$10 min. A 2 x reaction buffer was added to initiate the reaction giving a final concentration of 1.66 $\mu$M deoxyribozyme to 0.002 $\mu$M substrate. The reaction was terminated after a designated period of time by the addition of quenching buffer containing 60 mM EDTA, 7 M urea and loading dye solution.

All reactions were conducted at room temperature ($\sim$23°C) for a designated period of time, using a reaction buffer (1 x concentration = 400 mM KCl, 100 mM NaCl, 7.5 mM MgCl$_2$, 7.5 mM MnCl$_2$ and 50 mM HEPES pH 7.0 at 23°C) previously demonstrated to support diverse 8–17 cleavage activity (34). Under these reaction conditions, the background rate of substrate cleavage was determined to be $\sim 10^{-7}$ min$^{-1}$, based on both theoretical (36) and experimental determination (data not shown). We therefore chose reaction times ranging from 15 s to 72 h depending on the dinucleotide junction. These reaction times provided detectable but unsaturated cleavage yields by the wild type (WT) 8–17, such that mutants with lower or higher activity could be subsequently identified. The reaction rate was estimated using the equation $k_{obs} = -\ln(1− fraction cleaved)/time$. We assumed the endpoint of reactions to be $\geq 80\%$ (although experimental observations often showed endpoints up to 95%). Therefore, any reactions that exceeded 75% cleavage product at an initial timepoint were subsequently assayed with a shorter timepoint(s) to ensure accuracy. When necessary, slower reactions were followed for at least 72 h (and up to 6 days when complete timecourses were conducted). If a distinct cleavage signal was not observed in this period of time, the rate was assumed to be the background rate of $\sim 10^{-7}$ min$^{-1}$.

All reaction rates for 8–17NG, 8–17NA and 8–17NC were also independently determined with complete timecourses (data fit to a single-exponential equation), and the variation between the two methods was generally within 30%, suggesting that our simplified assay method provided a very good approximation of the rate. In general, rate constants calculated from either the simple assay or full timecourse were determined from at least two independent measurements that typically differed by <30%. Negative control reactions were conducted with each experiment, in which each substrate was incubated with reaction buffer in the absence of the deoxyribozyme for the same period of time. Any cleavage products detected in the negative controls were subtracted from the results of the deoxyribozyme reactions prior to analysis. This background cleavage was typically <1%.

**RESULTS AND DISCUSSION**

8–17 sequence and secondary structure

Figure 1 illustrates the sequence and secondary structure of an 8–17 motif, which is typically characterized by a
3-bp stem-triloop and a single-stranded turn region of 4–5 nt. Two binding arms engage the substrate through Watson–Crick base pairs on either side of the cleavage site. Deoxyribozyme 8–17 does not have any reported sequence requirements in the substrate, other than the identity of the dinucleotide cleavage junction. The catalytic core is composed of 14–15 nt, of which only 4 nt (i.e. A_6, G_7, C_13 and G_14) have been shown to be absolutely conserved and essential for activity (25,34). The sequence requirements may be even more flexible, as Cruz et al. have previously isolated 8–17 variants that contained one or two mismatches in the stem, insertions, deletions and combinations thereof (34). However, the activities of these sequence variants have not been confirmed. For simplicity, we use the term 8–17 generically to describe both the canonical motif first described in 1997 (33), and all related sequence variants.

In an effort to understand how these sequence variations influence 8–17’s cleavage site selectivity, we comprehensively mutated the catalytic core and analysed the effects against each of the 16 possible dinucleotide junctions. In addition to systematically replacing each position with the other three nucleotides, we also assayed the effects of various deletions, insertions and combination mutations. The 8–17 variant depicted in Figure 1 was chosen as the WT sequence, against which the activity of each mutant was subsequently compared. This particular 8–17 sequence was the fastest variant reported in a previous study by our laboratory for cleavage of 5’-GG junctions (37). Sixteen chimeric substrates (i.e. composed entirely of DNA except for a single-embedded scissile ribonucleotide) were designed with the sequence shown in Figure 1, differing only in the identity of the two nucleotides at the target cleavage site (denoted as positions rN_18N_1.1 in Figure 1). Therefore, any respective difference in cleavage reactivity could be attributed exclusively to the identity of this dinucleotide junction. The nucleotide numbering system used in Figure 1 is analogous to the system originally proposed for the hammerhead ribozyme (38), and subsequently adopted for the 8–17 deoxyribozyme (25). For future reference, all dinucleotide junctions are written in the 5’–3’ direction, and N = G, A, C or T/U (deoxy)ribonucleotides.

One 8–17 sequence can cleave all 16 dinucleotide junctions

The canonical 8–17 deoxyribozyme was first reported to cleave AG junctions (33), and a sequence variant was subsequently shown to have the ability to cleave all four NG junctions (23). Most recently, our laboratory extended the list of known cleavage sites to include as many as 14 different dinucleotide junctions, based on a set of five different 8–17 sequence variants (34). Only the CT and UT junctions could not be cleaved by these 8–17 variants. However, for practical reasons these conclusions were based on a minimum observed reaction rate of ~0.0001 min⁻¹, which corresponds to a ~1000-fold enhancement over the background rate of ~10⁻⁷ min⁻¹. In the present study, we reasoned it would be instructive to determine if any detectable level of catalysis were possible. Even a 10-fold rate enhancement could provide better insight into understanding the functional limitations and potential of 8–17.

The cleavage site versatility of the WT 8–17 sequence is illustrated in Figure 2. Remarkably, all 16 dinucleotide junctions were in fact susceptible to cleavage. Rate enhancements varied by several orders of magnitude, from as low as ~40-fold (UT junction) to as high as ~5 × 10⁷-fold (GG junction) over background. As a group, the NG dinucleotide junctions were the most susceptible to cleavage, with rates ranging from ~0.1 to 5 min⁻¹. The remaining NA, NC and NT junctions were at least two orders of magnitude less reactive, with rates that did not exceed ~0.001 min⁻¹. Within each group of dinucleotide junctions defined by the same 3’-nt (i.e. the NG group, NA group, NC group and NT group), the reactivity consistently followed the order N = G > A > C > U.

Positions 2.1 and 12 influence selectivity at the 3’ nucleotide (N₁.₁) of the cleavage site

The relative activity of WT 8–17 variants mutated at positions 2.1 and 12 are summarized in Figure 3, along with all other positions excluding the 3-bp intramolecular stem. For clarity and comparison, the data has been organized into eight different groups of related dinucleotide junctions defined by either a common 3’-nt (i.e. the NG group, NA group, NC group and NT group) or common 5’-nt (i.e. the GN group, AN group, CN group and UN group). Mutations at position 2.1 had the same effect against junctions with the same 3’-nt, as reflected by the near superimposition of the relevant graphs (Figure 3, left panel of graphs). However, these mutations clearly exhibited different effects against junctions with a different
3'-nt (Figure 3, right panel of graphs), both in terms of the direction of the effect (i.e. increase or decrease in activity) and the magnitude. These observations suggest that position 2.1 influences the selectivity through the 3' position N_{1,1} of the dinucleotide cleavage site. Nucleotide T_{2,1} has the greatest functional significance for the NG group of dinucleotide junctions. Substitution with one of the three other nucleotides impairs catalysis by \(10^{-3}\) fold. Furthermore, deletion of T_{2,1} reduces the reaction rate by \(10^{-4}\) fold, to a level approximately at par with the WT activity exhibited against the other junctions. In contrast, substitution or deletion of T_{2,1} has only a modest effect (i.e. decrease or increase of <10-fold) on the cleavage of NA, NC and NT junctions. These results indicate that mutation of position 2.1 is required, but not sufficient to confer a significant change in the cleavage site selectivity.

Unlike position 2.1, substitutions at position 12 showed a similar pattern of effects against all dinucleotide junctions, initially suggesting that this position does not contribute to cleavage site selectivity. The magnitude of these effects was <10-fold. However, the deletion of T_{12} had a significant impact on the cleavage of NG junctions, reducing the reaction rate by \(10^{3}\)–\(10^{4}\) fold. Deletion of
T12 caused <10-fold change in activity against all other junctions. Therefore, position 12 likely plays some role in cleavage site selectivity. Interestingly, position 12 has also previously been implicated as functionally important for the metal-ion selectivity of 8–17 (25).

**Positions 15 and 15.0 influence selectivity at the 5′ nucleotide (rN18) of the cleavage site**

Mutations at positions 15 and 15.0 exhibited the same effects against junctions with the same 5′-nt, as reflected by near superimposition of the graphs at these positions in the right panel of Figure 3. However, different effects were observed against junctions with a different 5′-nt, suggesting that position 15 and 15.0 influence selectivity through the 5′ position of the dinucleotide cleavage junction. Substitutions at position 15 and 15.0 generally exhibited moderate effects (<100-fold). Deletion of A15.0 caused a modest decrease in activity (<10-fold). Deletion of both A15 and A15.0 simultaneously, also caused a relatively modest decrease in activity for most junctions (<50-fold), but up to ~1000-fold decrease for some of the CN junctions. Double mutations, in which position 15 was substituted with another nucleotide while position 15.0 was deleted, showed deleterious effects as high as ~1000-fold for some junctions.

Though not officially part of the catalytic core of 8–17, we decided to evaluate the functional significance of position 16.1 given its proximity to the cleavage site. A previous study conducted on the 10–23 RNA-cleaving deoxyribozyme, which engages its substrate in an analogous manner, suggested that this position may benefit from the conformational freedom afforded by a mismatch (39). However, mutations to this position in 8–17 exhibited either neutral or deleterious effects (up to ~100-fold) on the reaction rate.

**Positions 6–8, 13 and 14 show no influence on cleavage site selectivity**

Mutations at positions 6–8, 13 and 14 exhibited similar effects against all 16 dinucleotide junctions (Figure 3), which argues against their involvement in conferring selectivity. This result is logical, since these positions are very highly conserved (with the exception of position 8), and therefore probably form the basis of the catalytic strategies employed by 8–17.

Mutations at position 6 impaired catalysis by 10–10^4 fold, with the largest reduction observed with an A6T substitution. G7 was the most crucial nucleotide in the loop region. Substitution of G7 generally reduced the reaction rate to background levels. Position C8 was the most tolerable to mutation, with substitutions causing changes <50-fold. In agreement with prior observations (25), a C8T substitution was the most tolerable, and in some cases even led to a very modest increase in the reaction rate. As expected, C13 and G14 were crucial for activity. Substitution at these positions generally reduced the activity to background or near background levels for all junctions.

**Mutations to the core stem show little influence on selectivity**

The functional consequences of mutations to the 3-bp stem region were also addressed. Nucleotide positions 3, 4 and 5 representing half of the intramolecular stem of WT 8–17, were individually substituted with each of the other three nucleotides. Nucleotide positions 11, 10 and 9 were co-mutated with positions 3, 4 and 5 respectively, to maintain base pairing. \( k_{rel} \) is defined as \( k_{obs, mutant}/k_{obs, WT} \). The \( k_{obs, WT} \) values are provided in Figure 2B. Combination mutants are denoted by the sequence of positions 3, 4 and 5 in that order. For clarity, dinucleotide junctions are organized into groups of related junctions.
modest improvements in the rate from ~2-fold for the NG, NA and NT junctions, to as high as ~10-fold for the NC junctions. The largest deleterious effect was seen with the C5G substitution, which led to ~100-fold decrease in activity against GC, AC, CC, GT and AT junctions. These differential effects tentatively suggest that the sequence of the stem may play some small role in the cleavage site selectivity of 8–17. It should also be noted that the stability of the stem was not directly correlated with the catalytic activity, as there were several examples of stem mutants exhibiting higher activity with fewer GC pairs.

Single-nucleotide insertions/bulges show little influence on selectivity

Single-nucleotide insertions or bulges have been observed at various positions among in vitro selected 8–17 variants (34). An insertion of thymidine between positions 8 and 9 has been most frequently observed, followed by bulges (typically thymidine) between positions 9 and 12. The functional consequences of such mutations were investigated by systematically inserting thymidine residues between positions 3 and 13, with the results presented in Figure 5. Single-nucleotide T insertions generally exhibited the same pattern of effects among all dinucleotide junctions, suggesting little or no influence on the cleavage site selectivity. The reaction rate progressively decreased between positions 3–6 (by up to ~10^6-fold), then progressively increased up to the WT activity levels by position 9 or 10, before once again declining by ~10-fold at position 13.

Cleavage site selectivity versus mutational distance

Although the WT sequence used in this study had the ability to cleave all 16 dinucleotide junctions, it showed a marked selectivity for the group of NG junctions. Cleavage of these four junctions proceeded ~10^2–10^6-fold faster than the 12 remaining junctions. During the systematic mutation of the WT sequence, we did not observe any significant changes to this selectivity, despite testing more than 60 mutants within 1–2 mutations of the WT sequence. Nevertheless, previous data indicates that large changes in selectivity are accessible within a relatively short mutational distance. For instance, Cruz et al. described an 8–17 variant called E5112 (sequence 5’-3’: T21 GTC AGC 8 GAC TCGAA) that differed by only two mutations from our G1T6C5 stem mutant (sequence 5’-3’: T21 GTC AGC8 GAC TCGAA), but showed at least 1000-fold selectivity for the group of NA junctions versus NG. This large change in selectivity is somewhat unexpected since one of the mutations (i.e. an insertion of T at position 8), is not expected to confer selectivity based on the results from our systematic analysis. The insertion of a T residue was shown to have similar effects against all dinucleotide junctions (Figure 5). This discrepancy illustrates an important point. The observed functional significance of a given mutation may change depending on the context of the surrounding sequence. Thus, the right combination of mutations could act cooperatively to elicit a large change in cleavage site selectivity.

One inherent limitation of this study is the potential bias associated with our choice of starting WT sequence. We therefore wondered if the conclusions from our mutational analysis would still be relevant, even as the mutational distance from the WT sequence increased. To provide some additional insight into this issue, we decided to look for faster NA, NC and NT-cleaving 8–17 variants. Since the WT sequence appeared to be already optimized for the cleavage of NG junctions (as reflected by the fact that nearly all point mutations were deleterious), we reasoned that more proficient cleavage of the other 12 junctions might require a larger mutational distance to escape the specialized fold of the WT sequence. This hypothesis is supported by the results of a study conducted by Bartel and colleagues, in which they demonstrated that the probability of deriving new function from an existing ribozyme increases with the mutational distance (40). In addition, Cruz et al. previously isolated many 8–17 sequences that contained
various combinations of deletions, insertions and stem mismatches (34). The fact that these unusual sequence variants survived selection, suggests they may be functionally important. Fourteen such candidate sequences were chosen from pools of sequences that had been selected to cleave one of the less reactive NA, NC and/or NT dinucleotide junctions (34). Several of these sequences were very slow, or interestingly, exhibited higher activity against junctions for which they were not originally selected to cleave (data not shown). However, we did identify two sequences that provided ~4-40-fold higher activities against NA and NC junctions than the best sequences reported by Cruz et al. under the same buffer conditions. These two sequences, denoted as 8–17NA and 8–17NC, respectively, differed from the WT sequence by nine mutations. Figure 6A shows the sequence of 8–17NA, 8–17NC and the sequence of one of the fastest NG-cleaving 8–17 variants (denoted as 8–17NG) that were identified during our systematic mutational analysis. 8–17NG also showed the highest activity against the group of NT junctions. However, it should be noted that the double mutant A15TΔA15.0, actually provided the largest rate enhancement for the UT junction (~1000-fold over background). The rate constants for 8–17NG, 8–17NA and 8–17NC are presented in Figure 6B against each dinucleotide junction.

The selectivity of 8–17NA followed the approximate scheme NA > NC > NT ~NG. The decrease in activity against NG junctions (compared to 8–17NG), is expected with the deletion of T2.1. Within each of these groups, the selectivity at the 5'-nt followed the order G > A > U > C, rather than the G > A > C > U order exhibited by 8–17NG. This change is consistent with the observed A15.0G substitution, since position 15.0 is expected to influence the selectivity at the 5'-nt of the cleavage site. Similarly, 8–17NC also showed reduced activity against NG junctions (selectivity = NC > NA > NT ~NG), which can be reconciled by the deletion of T2.1 and/or T12. Positions 15 and 15.0 of 8–17NC were identical to 8–17NG, and therefore the same selectivity was observed at substrate position N18 in groups of junctions defined by the same 3'-nt (such as GG > AG > CG > UG, GT > AT > CT > UT and so on). The G3A4T5 stem utilized by 8–17NA is expected to modestly decrease the activity against NG and NT junctions, while having a relatively neutral effect for NA junctions and slightly beneficial effect for NC junctions. 8–17NC also utilized a G3A4T5 stem, but had a G–A mismatch between positions 3 and 11. The functional significance of this mismatch is still unclear, but it does reaffirm the notion that a stable stem is not an absolute prerequisite for robust activity (at least against NC junctions). 8–17NA and 8–17NC had single-nucleotide bulges/insertions at positions 10 and 8, respectively. These mutations are expected to cause only a modest decrease in the reactivity against NG junctions, and slightly increase the activity (by ~2-fold) against NA, NC and NT junctions.

In general, the observed selectivity and sequence variations in 8–17NA and 8–17NC are consistent with the functional significance assigned to specific positions during our systematic analysis of the catalytic core.

This provides some measure of confidence that the conclusions from this study are not artefacts of the chosen WT sequence. However, additional studies will be necessary to fully understand the interactions between different nucleotide positions and their functional consequences. High-resolution structural studies, including NMR spectroscopy and/or X-ray crystallography, would be particularly useful, and could benefit in turn from the many sequence variants characterized herein.
Prospects for faster 8–17 variants

Herein we have shown that the 8–17 deoxyribozyme has the ability to cleave all dinucleotide junctions, but exhibits a hierarchy of reactivity against groups of dinucleotide junctions according to the order NG > NA > NC > NT. This order was based on the fastest 8–17 variants (i.e. 8–17NG, 8–17NA and 8–17NC) that could be identified from a sample of ~75 different sequence variants, representing some of the most commonly observed mutations. In theory, however, there are still thousands of additional sequence permutations that conform to the sequence and secondary structure requirements of the 8–17 motif. Thus, it remains to be determined if this hierarchy reflects the true functional limitations of 8–17, or alternatively, a possible sampling artefact. The in vitro selection results reported by Cruz et al., provides evidence against the latter possibility, because the observed hierarchy of reactivity closely parallels the relative frequency with which 8–17 variants were isolated against each of the various junctions (34). Nevertheless, these selection results are not sufficient to completely discount the prospect of faster sequence variants. This is because the 43-nt random domain used by Cruz et al. is too large to sample all possible sequence permutations. In addition, this study used a constant and relatively permissive reaction time during each round of selection, which, as our laboratory has subsequently shown, may contribute to the random loss of deoxyribozymes, including proficient catalysts (41). Therefore, faster 8–17 variants for cleavage of the NA, NC and NT junctions are still plausible.

Practical implications, limitations and future directions

The three optimal 8–17 sequence variants described herein can collectively cleave 10 junctions with useful rates of $\geq 0.1 \text{ min}^{-1}$ (Figure 6). The number of junctions and/or the reaction rate can be further supplemented by other deoxyribozymes like the 10–23 (33) and bipartite (42). However, proficient deoxyribozymes for the cleavage of pyrimidine–pyrimidine junctions have yet to be reported. This study represents one step closer towards the goal of a complete set of practical deoxyribozymes for the site-specific cleavage of all 16 dinucleotide junctions. These enzymes would be of immediate benefit to researchers conducting structural/functional studies on RNA. Deoxyribozyme-mediated RNA cleavage offers several advantages over the alternative ribozyme (43) and RNase H (44) techniques including extra versatility, simplicity, stability and/or cost effectiveness.

This study also provides new insight into the distribution of RNA-cleaving deoxyribozymes in sequence space, and more importantly, how best to access them. For instance, several studies have suggested that longer random libraries (for in vitro selection) provide greater structural complexity (45–47), which in turn can be correlated with greater functional activity (48). We shared this view in the past, and therefore used the longest random-sequence domain (i.e. 80-nt) ever reported for the in vitro selection of RNA-cleaving deoxyribozymes (32). Contrary to expectations, however, the fastest deoxyribozymes that emerged were 8–17 variants. These results are complemented by the current study, in which we have demonstrated that all junctions can actually be cleaved by a deoxyribozyme as small and as simple as 8–17. These findings suggest that our goals may be better served from the complete sampling of sequence space afforded by a library containing a small random domain (N < 25), rather than from the increased structural complexity afforded by a library with a larger random domain. This hypothesis is currently under our investigation. Yarus and co-workers have previously investigated the relative merits of different random library sizes for the selection of isoleucine-binding RNA aptamers (49). Their results also suggest that the selection of certain functional motifs can benefit from smaller library sizes.

Although it is beyond the scope of the current study, further examination of the substrate and metal-ion dependency of these 8–17 deoxyribozymes may be the subject of a future report. In the meantime, it remains to be determined if the 8–17 variants described herein will be equally effective against all-RNA substrates, and whether the observed cleavage site versatility can be supported by different buffer conditions. However, it should be noted that 8–17 has been isolated from in vitro selection experiments using both chimeric (24,31,34) and all-RNA substrates (32,33). Furthermore, an 8–17 variant described by Lu and colleagues was only ~4-fold less active against an all-RNA substrate versus the corresponding chimeric version (24). Previous studies have also indicated that 8–17 can utilize a wide variety of divalent metal ions as cofactors, with transition metal ions generally providing greater activity than the alkaline earth metal ions (22,24).

CONCLUSIONS

Deoxyribozyme 8–17 represents an ideal model system to address theoretical questions about the functional limitations of nucleic acid-mediated catalysis. This RNA-cleaving deoxyribozyme also represents considerable practical interest because of its utility in many diverse applications. Herein, we conducted a comprehensive mutational study to understand the functional significance of 8–17 sequence variations on cleavage site selectivity. We report for the first time that the 8–17 motif has the unique ability to cleave all 16 dinucleotide junctions, with rate enhancements of at least 1000-fold over background. To the best of our knowledge, no comparable deoxyribozyme or ribozyme has been reported. Sequence space is densely populated by 8–17 sequence variants that differ widely in both reaction rate and cleavage site preference. We have provided evidence suggesting how each nucleotide position in the catalytic core might contribute to the observed cleavage site selectivity of a given 8–17 sequence. In particular, core positions 2.1 and 12 represent a primary determinant of selectivity at the dinucleotide 3' position, N13,1, while core positions 15 and 15.0 represent a primary determinant of selectivity at the dinucleotide 5' position, rN18.

The optimal 8–17 variants identified in this study can cleave 10 out of 16 junctions with rates $\geq 0.1 \text{ min}^{-1}$, and exhibit an overall hierarchy of reactivity towards groups
of related junctions according to the order NG \(\succ\) NA \(\succ\) NC \(\succ\) NT. However, our results also suggest that the identification of faster sequence variants against less reactive junctions may still be possible through focused in vitro selection efforts. The remarkable structural and functional plasticity demonstrated by this very simple deoxyribozyme, also serves to challenge the general perception that complexity is a prerequisite for greater function.

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