In vitro selection of small RNA-cleaving deoxyribozymes that cleave pyrimidine–pyrimidine junctions

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ABSTRACT

Herein, we sought new or improved endoribonucleases based on catalytic DNA molecules known as deoxyribozymes. The current repertoire of RNA-cleaving deoxyribozymes can cleave nearly all of the 16 possible dinucleotide junctions with rates of at least 0.1/min, with the exception of pyrimidine–pyrimidine (pyr–pyr) junctions, which are cleaved 1–3 orders of magnitude slower. We conducted four separate in vitro selection experiments to target each pyr–pyr dinucleotide combination (i.e. CC, UC, CT and UT) within a chimeric RNA/DNA substrate. We used a library of DNA molecules containing only 20 random-sequence nucleotides, so that all possible sequence permutations could be sampled in each experiment. From a total of 245 clones, we identified 22 different sequence families, of which 21 represented novel deoxyribozyme motifs. The fastest deoxyribozymes exhibited $k_{\text{obs}}$ values (single-turnover, intermolecular format) of 0.12/min, 0.04/min, 0.13/min and 0.15/min against CC, UC, CT and UT junctions, respectively. These values represent a 6- to 8-fold improvement for CC and UC junctions, and a 1000- to 1600-fold improvement for CT and UT junctions, compared to the best rates reported previously under identical reaction conditions. The same deoxyribozymes exhibited ~1000-fold lower activity against all RNA substrates, but could potentially be improved through further in vitro evolution and engineering.

INTRODUCTION

Deoxyribozymes (or DNAzymes) are enzymes made from single-stranded DNA (1). Although they are yet to be found in nature, many different deoxyribozymes have been made in laboratories through a process known as in vitro selection (2). Enzymes made from DNA can offer several practical advantages over those made from protein or RNA, including simplicity/convenience, greater stability and lower production costs. RNA-cleaving deoxyribozymes represent a very important class (3), which has been used for a variety of purposes including (i) biosensors (4–7), (ii) DNA computational elements with Boolean logic functions (8,9), (iii) nanomachines (10), (iv) antiviral or gene control agents (11–13) and (v) the in vitro manipulation of RNA (14).

We are interested in understanding the functional limits of DNA-mediated catalysis, using RNA-cleaving deoxyribozymes as a model system. These efforts should expand their utility, and provide insight to better assess future applicative potential. One specific question that we have sought to answer is how many different dinucleotide junctions can be cleaved efficiently by RNA-cleaving deoxyribozymes?

This is an important question because the ability to efficiently cleave at any location in a given RNA molecule would be very useful for RNA structural and/or modification studies. Furthermore, improved cleavage site versatility would benefit applications involving long RNA molecules, which often possess higher order structures that can interfere with DNAzyme binding, and therefore limit the number and types of susceptible target sites (15).

We have previously established that all 16 dinucleotide junctions are in fact susceptible to deoxyribozyme-mediated cleavage (16), but efficient cleavage of all junctions has not been demonstrated. Existing RNA-cleaving deoxyribozymes such as the 10-23 (17), bipartite (18) and 8-17 (19) serve as a standard baseline, to which activity can be compared. These deoxyribozymes can achieve reaction rates from ~0.1 to 10/min for the cleavage of various purine–purine, purine–pyrimidine and/or pyrimidine–purine junctions (Table 1). However, comparable rates for cleavage of any of the four different pyrimidine–pyrimidine (pyr–pyr) junctions have not been reported. So far, only the 8-17 deoxyribozyme has confirmed...
activity against pyr–pyr junctions, with rates that are several orders of lower magnitude (19).

We have hypothesized that (i) efficient deoxyribozyme-mediated cleavage of all four pyr–pyr junctions is possible, and (ii) these deoxyribozymes can be isolated through a focused in vitro selection effort using a small (i.e. 20 nt) random library approach. These hypotheses are largely motivated by the recent discovery that 8-17 can cleave all 16 dinucleotide junctions (19), and different sequence variants can show different dinucleotide selectivities. Because only 4 of the 15 nt that comprise the 8-17 catalytic core are absolutely conserved, there may still be more proficient variants among the thousands of possible sequence permutations. A small random-sequence library would thereby ensure that all 8-17 sequence variations are represented in the initial pool. The identification of more efficient 8-17 sequence variants is an attractive prospect, because it would avoid the time-consuming characterization experiments that are often necessary before novel motifs can be used productively.

Toward this end, we conducted four separate in vitro selection experiments to target the four different types of chimeric (RNA/DNA) pyr–pyr junctions (i.e. CC, UC, CT and UT). Two additional selection experiments were also performed as controls, targeting AC and AT junctions, for which small and efficient RNA-cleaving deoxyribozymes are known to exist.

### MATERIALS AND METHODS

#### Oligonucleotides and reagents

Oligonucleotides were prepared by automated DNA synthesis using cyanoethylphosphoramidite 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. Nucleoside 5'-triphosphates, [γ-32P]ATP and [α-32P]dGTP were purchased from Perkin Elmer (Boston, MA, USA). T4 DNA ligase and T4 polynucleotide kinase (PNK) were purchased from MBI Fermentas (Burlington, ON, CAN). Tth DNA polymerase was purchased from Biotools (Madrid, Spain). All chemical reagents were purchased from Sigma (Oakville, ON, CAN).

#### In vitro selection procedure

The library design and in vitro selection procedure are illustrated in Figures 1A and 2, respectively.

**Step 1 and 2.** The full-length 73-nt library was constructed by splint-directed ligation of the 43-nt enzyme domain to the 30-nt substrate domain. Each selection experiment used the same enzyme domain, but different substrate domains and appropriate ligation splints (i.e. 22-nt DNA oligos complementary to adjacent regions in both the substrate and enzyme domains). A total of 1000 pmol of 5'-phosphorylated enzyme domain, 1100 pmol of ligation splint, 1200 pmol of substrate

### Table 1. Reaction rates of various RNA-cleaving DNAzymes against different dinucleotide junctions

| DNAzyme | Reference | Size (nt) | Dinucleotide junction | 10-23 | Bipartite | 8-17 | Various |
|---------|-----------|-----------|------------------------|-------|----------|------|---------|
| 17      | 17        | 15        | GG                     | 9.2   | N        |      | ND      |
|         | 18        | 20        | AG                     | 0.0001| 3.1      | 0.63 | N       |
|         |           |           | CG                     | 0.20  | N        |      | ND      |
|         | 19        | 15        | GA                     | 3.4   | N        |      | ND      |
|         | This work | 20        | AA                     | 1.4   | N        |      | ND      |
|         |           |           | CA                     | 1.7   | N        |      | ND      |
|         |           |           | UA                     | 0.47  | N        |      | ND      |
|         |           |           | GC                     | 0.36  | N        |      | ND      |
|         |           |           | AC                     | 0.11  | N        |      | ND      |
|         |           |           | CC                     | 0.12  | N        |      | ND      |
|         |           |           | UC                     | 0.04  | N        |      | ND      |
|         |           |           | GU/T                   | 0.037 | N        |      | ND      |
|         |           |           | AU/T                   | 0.0089| N        |      | ND      |
|         |           |           | CT                     | 0.13  | N        |      | ND      |
|         |           |           | UT                     | 0.15  | N        |      | ND      |

All $k_{obs}$ (min$^{−1}$) values were determined under single-turnover conditions in an intermolecular format. Reaction conditions for 10-23: 10 mM MgCl$_2$, pH 7.5, 37°C, all-RNA substrate. Reaction conditions for bipartite: 30 mM MgCl$_2$, pH 7.4, 23°C, all-RNA substrate. Reaction conditions for 8-17: 100 mM KCl, 400 mM NaCl, 7.5 mM MgCl$_2$, 7.5 mM MnCl$_2$, pH 7.0, 23°C, chimeric RNA/DNA substrate. The 8-17 rates are derived from several different 8-17 sequence variants. Reaction conditions for various DNAzymes from this work: 100 mM KCl, 400 mM NaCl, 7.5 mM MgCl$_2$, 7.5 mM MnCl$_2$, pH 7.0, 23°C, chimeric RNA/DNA substrate. Size refers to the length of the catalytic core. ND: no detectable cleavage activity. Blank squares indicate no reported cleavage activity.

#### Figure 1. Library design. (A) Sequences of libraries and PCR primers. Regions that differ in sequence are highlighted by gray boxes, which include the dinucleotide cleavage junction (denoted as rN–N in the sense primer) and the five terminal nucleotides of the substrate domain (denoted as NNNNN in the sense primer). It should be noted that six different sense primers were used to minimize PCR cross-contamination, although only one is shown for simplicity. N$_{20}$ refers to 20 random nucleotides, where N = A, G, C or T deoxyribo-nucleotides. Sp18 refers to an 18-atom hexa-ethyleneglycol spacer. (B) Two hypothetical secondary structure arrangements showing location of preengineered base-pairing interactions between substrate and enzyme domains. A 2-nt frameshift in base-pairing interactions on the 5'-side of the cleavage junction is possible. The structure on the right shows how an 8-17 motif might form.
domain and ~1 pmol of 5'-32P-labelled enzyme domain were combined, supplemented with DEPC-H2O to a final volume of 85 μl, and then heated at 90°C for 1 min. After cooling at RT for 10 min, 10 μl of 10× T4 DNA ligase buffer (manufacturer supplied) and 5 μl of T4 DNA ligase (5 Weiss units/μl) were added to initiate the reaction. The ligation mixture was incubated at RT for 2 h, then ethanol precipitated, and purified by 10% denaturing PAGE. The final yield of the 73-nt ligation product was ~600 pmol, which was used for the first round of selection. Therefore, on average ~300 copies of every possible sequence should have been represented in the initial pool.

**Step 3.** The RNA cleavage reaction was initiated by adding a 2× selection buffer (200 mM KCl, 800 mM NaCl, 15 mM MgCl2, 15 mM MnCl2, 100 mM HEPES pH 7.0 at 23°C) to an equal volume of DEPC-H2O in which the 73-nt library had been resuspended and preheated (90°C for 30 s, followed by 10 min cooling at RT), to a final concentration of ~1 μM. The reaction was incubated at RT for a designated period of time, quenched with the addition of EDTA (pH 8.0) to 30 mM, and ethanol precipitated. In rounds 1–4, a 100 min reaction time was used. In round 5, the population was divided into three subpopulations and subjected to 1, 10 and 100 min reaction times. In rounds 6–9, reaction times of 0.1, 1 and 10 min were used. From rounds 5 to 8, the 1 min subpopulation was used to seed the next round of selection.

**Step 4.** The 58-nt cleavage product was purified by 10% denaturing PAGE, excised from the gel and recovered by ethanol precipitation.

**Step 5.** The cleavage product was PCR amplified in a 100 μl reaction volume, containing 0.2 mM of each dNTP, 10 μl of 10× PCR buffer (Biotools), 0.5 μM each of sense and antisense primers, 5 U Tth DNA polymerase, 2 μl of 50× SYBR green (Invitrogen, Burlington, ON, CAN) and 20 μCi of [γ-32P]dGTP. Reactions were conducted on a Cepheid SmartCycler (Sunnyvale, CA, USA) and monitored in real time using SYBR green as the reporter. Each PCR cycle consisted of 94°C for 30 s, 50°C for 45 s and 72°C for 40 s. PCR amplification introduces the substrate domain, via the sense primer which contains the target ribonucleotide. The antisense primer contains an internal 18-atom hexa-ethyleneglycol spacer that blocks further DNA polymerization, leading to a dsDNA product of unequal length.

**Step 6.** The 73-nt sense strand was separated from the 88-nt antisense strand by 10% denaturing PAGE, excised from the gel and recovered by ethanol precipitation. Steps 3–6 were repeated for nine rounds.

**Cloning and sequencing of selected DNA populations**

Deoxyribozymes from the 100 min subpopulation of round 5, and the 10 min subpopulation of round 9 were cloned with the InstAclone PCR cloning kit (Fermentas). Approximately 60 clones from each selection experiment were chosen at random and sequenced by Functional Biosciences Inc. (Madison, WI, USA).

**Kinetic analyses**

All rates were determined under single turnover conditions, in which a large excess of deoxyribozyme (~1.7 μM) was used with a trace amount of 5'-32P-labeled substrate (~0.002 μM). Substrate and deoxyribozyme were heated together at 90°C for 30 s and allowed to cool at RT for ~10 min. An equal volume of 2× reaction buffer (200 mM KCl, 800 mM NaCl, 15 mM MgCl2, 15 mM MnCl2, 100 mM HEPES pH 7.0 at 23°C) was added to the deoxyribozyme/substrate mixture to initiate the reaction. 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. The cleavage products from a reaction timecourse...
strategy and the reproducibility of our results.

A measure of confidence in the effectiveness of our selection

known RNA-cleaving deoxyribozymes would provide a

8-17) are known to exist (Table 1). The recurrence of these

efficient RNA-cleaving deoxyribozymes (i.e. bipartite and

as controls, targeting AC and AT junctions, for which

two additional selection experiments were also performed

these junctions has not been demonstrated. However,

primary targets of interest, because efficient cleavage of

pyr–pyr junctions (i.e. CC, UC, CT and UT) were our

the surrounding deoxyribonucleotides (20). The four types

chimeric substrate is

diester bonds in an all-RNA substrate are equally suscep-

could be unambiguously targeted. Whereas the phospho-

all-RNA substrates, so that specific dinucleotide junctions

Figure 1A. The substrate domain contains a single ribonu-

were conducted in parallel using the library design illustrated in

Six independent in vitro selection experiments were con-

non-WT buffer . When necessary, slower reactions were followed

10 min were used. However, no appreciable cleavage

in Figure 2. In step 1, the sense primer (which serves as the

Library design

RESULTS

Metal ion dependency

Simple assays based on single-timepoint measurements were used to determine the relative activity of a given deoxyribozyme under various 'nonwild-type' buffer condi-

The last five nucleotides at the 3'-end of the substrate domain differed between experiments, in order to

were separated by denaturing 10% PAGE, and quanti-

tated using PhosphorImager and ImageQuant software. A

were used as described earlier; however, rates were approximated using the equation k_{obs} = \ln(1 - \text{fraction cleaved})/time. We have previously found that this method provides a reasonable estimation of the rate, when appropriate timepoints are chosen (i.e. prior to reac-

and upstream from the cleavage site, to provide some ver-

sequence of the substrate and enzyme domains according to

The percent cleavage product observed at each round of

The in vitro selection strategy

The in vitro selection scheme used in this study is depicted in

In vitro selection strategy

The percent cleavage product observed at each round of selection is illustrated in Figure 3. A total of nine rounds of selection were conducted in each experiment, and a distinct cleavage product (from 5% to 40%) was first observed by round 4. During the first four rounds of selec-

Selection progress

The percent cleavage product observed at each round of selection is illustrated in Figure 3. A total of nine rounds of selection were conducted in each experiment, and a distinct cleavage product (from 5% to 40%) was first observed by round 4. During the first four rounds of selection, a single reaction time of 100 min was used. In subsequent rounds, the selection stringency was increased by reducing the reaction time, in an effort to isolate the fast-

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were used to determine the relative activity of a given deoxyribozyme under various 'nonwild-type' buffer condi-

the same general reaction conditions and protocol were used as described earlier; however, rates were approximated using the equation k_{obs} = \ln(1 - \text{fraction cleaved})/time. We have previously found that this method provides a reasonable estimation of the rate, when appropriate timepoints are chosen (i.e. prior to reac-

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To determine the relative activity of a given deoxyribozyme under various 'nonwild-type' buffer condi-

RESULTS

Library design

were conducted in parallel using the library design illustrated in

Six independent in vitro selection experiments were con-

non-WT buffer /k_{WT} buffer. When necessary, slower reactions were followed

for at least 72 h. 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^{-17}/min.

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product was observed under the 0.1 min reaction time in any of the selection experiments. From rounds 5 to 9, the 1 min subpopulation in each selection experiment was used to seed (i.e. provide input molecules for) the next round of selection. By round 9 (or earlier), the composite population from each selection experiment could cleave its target dinucleotide junction with a rate of \( \frac{1}{C24} 0.01–0.1/min \).

All experiments were terminated at round 9 after no appreciable increase in activity was observed for several rounds.

**Deoxyribozyme sequence diversity and distribution**

Approximately 60 clones from each selection experiment were randomly chosen for sequence analysis. Approximately half of these clones were taken from round 5 (100 min subpopulation) and the other half were taken from the terminal round 9 (10 min subpopulation). Clones from the 10 min subpopulation of round 9 were sequenced instead of the 1 min subpopulation, in order to minimize the likelihood of background contaminating sequences due to the low level of cleavage observed in some populations under the latter selection time. Twenty-six different sequence families were identified in total, of which only 12 survived until round 9. The number of sequence families per population varied from 3 to 10, and several sequence families were identified in two or three different selection experiments. The frequency and distribution of clones are summarized in Figure 4. The sequences of all clones are available in Supplementary Figure 1.

**Recurrence of the 8-17 motif**

We identified 28 different 8-17 sequence variants from 34 clones and grouped them into one sequence family (S24) for simplicity. Based on previous characterization studies (16,19,21), we have defined an 8-17 motif by the presence of four absolutely conserved bases and 4–6 co-varying nucleotides that form a putative 2–3 bp stem. The 8-17 variants were observed in the AC, CC and CT populations, with the majority (22 sequence variants from 27 clones) found in the AC population. Only three 8-17 sequence variants survived until round 9, and all were found in the AC control selection experiment, consistent with expectations. Eight sequence variants were chosen for kinetic analysis in trans, under single-turnover conditions. Three sequence variants from the round 9 AC population exhibited comparable rates to the best NC-cleaving 8-17 variant (i.e. ‘8-17NC’) reported previously by our lab (19).

**Figure 3. In vitro selection progress.** The percent cleavage product is shown for each selection round. A 100 min selection time was used for the first four rounds of selection. In round 5, the population from each selection experiment was divided into subpopulations and subjected to a 1, 10 or 100 min selection time. In rounds 6–9, selection times of 0.1, 1 and 10 min were imposed. Less than 1% cleavage product was observed in the 0.1 min subpopulations. From rounds 5–9, the 1 min subpopulation from each selection experiment was used to seed each subsequent round of selection.
It should be noted that the exact 8-17NC sequence variant was not isolated herein, but this is not surprising since many sequence variants are known to be phenotypically equivalent. The activities of two 8-17 sequence variants from each of the CC and CT populations (round 5) were also determined. Interestingly, while the rates of three of these 8-17 variants were comparable to our previous report, one 8-17 variant (denoted as 8-17CT) actually exhibited a 1000-fold improvement with a rate of 0.13/min ($Y_{max} = 87\%$). The sequence and kinetic analysis of 8-17CT is shown in Figure 5. The 8-17CT contains a distorted stem (due to a mismatch in the central position), and a tetra-loop instead of a tri-loop, which are features also shared in common with 8-17NC described previously. The 8-17 variants have been shown to exhibit robust activity against groups of related dinucleotide junctions (e.g. NG group, where $N = G, A, C$ or U) rather than any single junction. Contrary to expectations, however, 8-17CT was quite selective (from ~25- to 100-fold) for a CT junction over the related GT, AT and UT junctions (Figure 5). The 8-17CT was also tested against GC, AC, CC and UC junctions, but showed no activity in 100 min (data not shown).

### Recurrence of the bipartite motif and identification of novel deoxyribozymes

Consistent with expectations, we identified the bipartite deoxyribozyme (S22) in both the AT and AC control selection experiments. It should be noted that the 10-23 motif was not reisolated in either control experiment, because it does not cleave chimeric substrates as proficiently as all-RNA substrates. The activity of 10-23 against chimeric substrates has not been previously reported; therefore, we tested two different 10-23 motifs optimized for cleavage of either AC or AU junctions (17), under our reaction conditions and with our chimeric AC and AT substrates. Only 2 and 8% cleavage product was observed in 100 min against the AC and AT junctions, respectively. These results indicate that the 10-23 deoxyribozyme would not be competitive under the imposed selection time pressure, and therefore unlikely to survive multiple rounds of selection.

The remaining 10 sequence families identified in round 9 represent novel deoxyribozymes that have not been previously reported. A representative sequence from each...
family is illustrated in Table 2. Secondary structure formation within the random-sequence domain of each family appears limited, with contiguous base-pairing up to 3 bp (Supplementary Figure 2). The preengineered substrate binding arms made for the facile separation of substrate and enzyme domains into an intermolecular format, which was further simplified with the removal of excess sequence elements peripheral to the core region. The random sequence domain also did not harbor any strong sequence complementarity to the substrate that might compete with the preengineered binding arms. Each deoxyribozyme sequence was confirmed to be active in trans under single-turnover conditions against their primary (i.e. most abundant) target dinucleotide junction (Table 2). The reaction rate ($k_{\text{obs}}$) varied from 0.007 to 0.15/min, with maximum cleavage yields ($Y_{\text{max}}$) between 74% and 90%. From among these new sequence families, deoxyribozymes S9, S4, S15 and S21 exhibited the fastest $k_{\text{obs}}$ values against CC (0.12/min), UC (0.04/min), CT (0.1/min) and UT (0.15/min), respectively. In comparison to the fastest rates previously reported and determined under identical reaction conditions (Table 1), S9 and S4 provide a 6- to 8-fold improvement, while S15 and S21 provide a 700- to 1600-fold improvement.

Metal ion dependency

The ‘wild-type’ (WT) reaction buffer used during in vitro selection contained several monovalent and divalent metal ions including 100 mM KCl, 400 mM NaCl, 7.5 mM MgCl$_2$, 7.5 mM MnCl$_2$, 50 mM HEPES pH 7.0 at 23°C. The underlined region of S22 is identical to the bipartite consensus sequence, with exceptions denoted in boldface. The sequence of the substrate (sub.) is also provided, written 3’ to 5’ in opposite orientation. YrH represents the dinucleotide cleavage junction, where Y = C or T, and H = A, C or U.

Cleavage activity against all-RNA substrates

The cleavage activity of each pyr–pyr cleaving deoxyribozyme from Table 2 was evaluated against all-RNA substrates, using the same reaction conditions and format described previously. Each substrate was identical to the substrate sequence shown in Table 2, except for the addition of two G residues on the 5’-end (to increase RNA transcript yield). Deoxyribozymes S9, 8-17CT, S4, S10-UC and S20-UT showed detectable but severely reduced (<1000-fold lower) activity against all-RNA substrates (Supplementary Figure 3). The remaining deoxyribozymes did not show detectable cleavage activity within a 72 h time period.

| Family | 5’ arm | N$_{20}$ region | 3’ arm | Junction | $k_{\text{obs}}$ (min$^{-1}$) | $Y_{\text{max}}$ (%) |
|--------|--------|-----------------|--------|----------|-----------------|-----------------|
| S3     | GAAAACCC--AAATCGAGGTTGGAGACCA--TCTTCG       | AC     | 0.03   | 75       |
| S22    | GAAAACCC--CGATAGGCGCTCGTGCA--TCTTCG       | AT     | 0.09   | 88       |
| S23    | GAAAACCC--GCTTTCTCCAGAAGGCGATAGG          | AT     | 0.05   | 83       |
| S9     | GAAAACCC--TACTGTCTTTACTGCGGCA--TCTTCG     | CC     | 0.12   | 90       |
| S13    | GAAAACCC--GGACAGAGAAGAGGGTTGA--TCTTCG     | CT     | 0.05   | 87       |
| S19    | GAAAACCC--TTTGGGAGGAAGGTGCGAA--TCTTCG    | CT     | 0.04   | 74       |
| S15    | GAAAACCC--AGATATCAAGAGTGAAGCGA--TCTTCG    | CT     | 0.10   | 85       |
| S10    | GAAAACCC--CGGGAGGTTTGCGTGAAGGAGCGG--TCTTCG | UC     | 0.005  | 85       |
| S4     | GAAAACCC--TGGGGTTGGTGGAAAGAGGG--TCTTCG   | UC     | 0.04   | 76       |
| S20    | GAAAACCC--TGGGAGGAGGATAGGGCGG--TCTTCG    | UC     | 0.007  | 89       |
| S21    | GAAAACCC--GGGGAGGTTGGTAGGAGCGA--TCTTCG    | UT     | 0.15   | 80       |
| Sub.   | 3’ CATTGCGGTGT---------YrH----------TAGAGAGAG 5’ |

The $k_{\text{obs}}$ values represent the average of at least two independent trials, which differed by <30%. Reactions were conducted under single-turnover conditions in 100 mM KCl, 400 mM NaCl, 7.5 mM MgCl$_2$, 7.5 mM MnCl$_2$, 50 mM HEPES pH 7.0 at 23°C. The underlined region of S22 is identical to the bipartite consensus sequence, with exceptions denoted in boldface. The sequence of the substrate (sub.) is also provided, written 3’ to 5’ in opposite orientation. YrH represents the dinucleotide cleavage junction, where Y = C or T, and H = A, C or U.
DISCUSSION

Herein, we have demonstrated that all four chimeric RNA/DNA, pyr–pyr junctions, can be efficiently cleaved by small deoxyribozymes. From a total of 245 deoxyribozyme clones isolated in four separate *in vitro* selection experiments, we identified 22 different sequence families that can cleave one or more pyr–pyr junctions. Twenty-one of these sequence families have not been previously described. The best deoxyribozymes (S9, S4, 8-17CT and S21) can collectively cleave all four pyr–pyr junctions with rates ranging from 0.04 to 0.15/min, which are 1–3 orders of magnitude faster than the best deoxyribozymes reported previously (Table 1). From among 115 clones isolated in two additional control experiments, targeting AC and AT junctions, we identified seven different deoxyribozyme sequence families including variants of the 8-17 and bipartite motifs. The remaining five sequence families represent novel motifs, but they do not provide any significant rate improvements over 8-17 or bipartite.

All of the deoxyribozymes isolated herein are 10- to 100-fold slower than other deoxyribozymes of comparable size (Table 1) that catalyze the cleavage of purine–purine junctions. Therefore, this study strengthens our previous suggestion that pyr–pyr junctions may be inherently more difficult to cleave (16,19). We have speculated that base-stacking interactions at the dinucleotide junction may play an important role in the cleavage mechanism of small deoxyribozymes, since purine bases can form stronger stacking interactions than pyrimidine bases (23–25).

This study also provides new insight into the functional versatility of the well known 8-17 deoxyribozyme. We have previously speculated that the structure of this deoxyribozyme might be flexible enough to support fast cleavage of all dinucleotide junctions, since only 4 of the 15 nucleotides that form its structural scaffold are catalytically essential. Because there are thousands of possible sequence variations (with unknown functional implications), a systematic and comprehensive mutational analysis would not be practical. However, by using a library containing only 20 random-sequence positions, we were able to sample every 20-nt sequence permutation in each selection experiment, including all 8-17 sequence variations. Consistent with expectations, we found 8-17 variants in the AC and CC populations that exhibited comparable rates to those described previously. No 8-17 variants were found in the AT, UC or UT.

Figure 6. Metal ion dependency of select deoxyribozymes. The sequence class and target dinucleotide junction is provided in each graph. Relative activity = $k_{\text{Mutant, buffer}}/k_{\text{WT, buffer}}$. WT buffer contains 100 mM KCl, 400 mM NaCl, 7.5 mM MgCl$_2$, 7.5 mM MnCl$_2$ (and 50 mM HEPES pH 7.0 at 23°C). Each metal ion was systematically removed individually, or in combination, as indicated above each graph. ‘Mn only’ refers to a buffer containing only 7.5 mM MnCl$_2$. ‘2M Na only’ refers to a buffer containing only 2 M NaCl.
selection experiments. However, we did identify one 8-17 variant (8-17CT) in the CT population that was ~1000-fold faster than the best 8-17 variant previously reported for the cleavage of this junction.

These results should be considered in the design of future in vitro selection experiments. A substantial effort has been expended to search for new and more useful RNA-cleaving deoxyribozymes, and in at least five independent studies (16,26–30), the recurrence of 8-17 has hampered this objective. Since UC and UT junctions are the least susceptible to 8-17-mediated cleavage, these dinucleotide junctions represent the best target cleavage sites to search for novel motifs, using larger random sequence domains. The likelihood of recurrence of other motifs described herein will depend on several factors (31), but in general, is expected to be lower due to the higher degree of sequence conservation in their putative catalytic cores (Supplementary Figure 1).

Limitations

It is important to recognize that the best possible solutions to a given selection experiment (typically presumed to be the most abundant sequences) might not always have the desired target phenotype. For instance, we did not always observe a direct correlation between the abundance of a given clone and its catalytic rate, as in the case of S20 and S21. Interestingly, S20 was about ~20-fold more abundant than S21, but ~20-fold slower than S21, contrary to expectations. This discrepancy may be due to the fact that rates were determined using a simplified intermolecular format (in which some fixed sequence regions were removed), and therefore may not reflect the activities of the full-length cis constructs that actually competed during selection. We tested this possibility, and determined that the activity of each full-length cis construct (S21 $k_{\text{obs}} = 0.056/min$, $Y_{\text{max}} = 46$%), S20 $k_{\text{obs}} = 0.036/min$, $Y_{\text{max}} = 22$%) differs from the activity determined by the simplified intermolecular format (Table 2). The difference in activity between the full-length cis constructs (~1.5-fold) is smaller than the difference observed between the simplified trans constructs (~20-fold); however, S21 still exhibits superior cleavage activity, which does not readily explain the discrepancy in their relative abundance.

Our previous study of in vitro selection population dynamics has suggested how secondary fitness constraints imposed unintentionally on the population, can unpredictably influence the relative abundance of different catalytic species (32). This scenario can enable sequences with non-target phenotypes (e.g. better PCR amplification efficiency than catalytic efficiency) to increase in abundance at the expense of those with target phenotypes. However, we have used several general strategies to try and mitigate the effects of this scenario including: (i) conducting control experiments under identical conditions; (ii) applying stringent time pressure to favor the target phenotype; (iii) performing extensive sequence analysis of each population at two different stages in the selection to minimize sampling artifacts and (iv) determining the rates of both major and minor sequence classes.

The practical utility of the deoxyribozymes described herein is currently limited by their inability to cleave all-RNA substrates as effectively as the chimeric substrates against which they were selected. Nevertheless, these catalytic DNA molecules can potentially serve as good starting points from which to evolve and/or engineer improved activity against pyr–pyr junctions within all-RNA substrates.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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