DNA-Catalyzed DNA Cleavage by a Radical Pathway with Well-Defined Products

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ABSTRACT: We describe an unprecedented DNA-catalyzed DNA cleavage process in which a radical-based reaction pathway cleanly results in excision of most atoms of a specific guanosine nucleoside. Two new deoxyribozymes (DNA enzymes) were identified by in vitro selection from N40 or N100 random pools initially seeking amide bond hydrolysis, although they both cleave simple single-stranded DNA oligonucleotides. Each deoxyribozyme generates both superoxide (O2•− or HOO•) and hydrogen peroxide (H2O2) and leads to the same set of products (3′-phosphoglycolate, 5′-phosphate, and base propenal) as formed by the natural product bleomycin, with product assignments by mass spectrometry and colorimetric assay. We infer the same mechanistic pathway, involving formation of the C4′ radical of the guanosine nucleoside that is subsequently excised. Consistent with a radical pathway, glutathione fully suppresses catalysis. Conversely, adding either superoxide or H2O2 from the outset strongly enhances catalysis. The mechanism of generation and involvement of superoxide and H2O2 by the deoxyribozymes is not yet defined. The deoxyribozymes do not require redox-active metal ions and function with a combination of Zn2+ and Mg2+, although including Mn2+ increases the activity, and Mn2+ alone also supports catalysis. In contrast to all of these observations, unrelated DNA-catalyzed radical DNA cleavage reactions require redox-active metals and lead to mixtures of products. This study reports an intriguing example of a well-defined, DNA-catalyzed, radical reaction process that cleaves single-stranded DNA and requires only redox-inactive metal ions.

INTRODUCTION

Many deoxyribozymes have been identified by in vitro selection1 for a growing reaction scope with biomolecular substrates.2 The first report3 of a deoxyribozyme was for nonhydrolytic RNA cleavage by mediating attack of a 2′-hydroxyl on an adjacent phosphodiester (the ribonuclease mechanism),4 and many RNA-cleaving deoxyribozymes were subsequently found.5 While seeking DNA-catalyzed amide bond hydrolysis,6 eventually with success,7 we initially identified deoxyribozymes that hydrolyze specific phosphodiester linkages of single-stranded DNA (ssDNA) substrates8 and RNA substrates.9 We and others also found deoxyribozymes that deglycosylate DNA, with subsequent elimination reactions that lead to strand scission.10 Other reported DNA-catalyzed cleavage reactions include oxidative DNA cleavage11,12 and phosphoramidate cleavage.13 Here we report a new, distinct type of DNA-catalyzed ssDNA cleavage. This cleavage process occurs by a superoxide/H2O2-dependent radical mechanism rather than an ionic mechanism and leads to well-defined reaction products that are identical to those formed by the action of the glycopeptide natural product bleomycin on double-stranded DNA (dsDNA).14 Unlike the reported examples of DNA-catalyzed oxidative DNA cleavage,11,12 the reactions described here do not require redox-active metal ion cofactors such as Cu2+ or Mn2+. Instead, only redox-inactive Zn2+ and Mg2+ are needed. This finding expands the scope of reactions that can be catalyzed by DNA to include a radical pathway that forms well-defined products without obligatory involvement of redox-active metal ions.

RESULTS

Identification of a New DNA-Cleaving Deoxyribozyme by in Vitro Selection. The two new deoxyribozymes in this study were found by in vitro selection, initially seeking
DNA-catalyzed amide hydrolysis. A successful in vitro selection experiment often depends upon implementing a suitable "capture reaction", which is highly selective for a functional group characteristic of the desired reaction product. Amide bond hydrolysis leads to a carboxylic acid functional group, which can be captured during each selection round using an amino-modified oligonucleotide and a suitable coupling agent such as the carbodiimide EDC (Figure 1).

During successful selection experiments for amide hydrolysis in which modified DNA nucleotides were incorporated into the catalytic DNA populations, we identified one additional and previously unreported deoxyribozyme, named RadDz3, which cleaves a simple DNA oligonucleotide substrate that lacks any amide bond at all (Figure 2A). From the PAGE analysis, the product is >85% of one species. RadDz3 was identified from an N40 random pool, i.e., DNA sequences with 40 random nucleotides. As originally identified, RadDz3 incorporated several 5-(hydroxymethyl)dU nucleotides, but these modifications were found to be dispensable for activity of the new RadDz3 deoxyribozyme reported here. The N40 selection experiment used unmodified DNA and led to the new RadDz6 deoxyribozyme. Both RadDz3 and RadDz6 lead to DNA cleavage, not amide hydrolysis.

Figure 1. In vitro selection process originally intended to capture the carboxylic acid product of DNA-catalyzed amide hydrolysis. The substrate presents its single amide bond embedded within two DNA oligonucleotide segments, which form Watson–Crick base pairs with the fixed-sequence 5′- and 3′-segments that flank the initially random region, which was either N40 or N100. The N40 selection experiment included modified nucleotides, although these modifications are dispensable for activity of the new RadDz3 deoxyribozyme reported here. The N100 selection experiment used unmodified DNA and led to the new RadDz6 deoxyribozyme. Both RadDz3 and RadDz6 lead to DNA cleavage, not amide hydrolysis.

Figure 2. Assays of the RadDz3 deoxyribozyme that cleaves ssDNA by a radical pathway. Incubation conditions: 70 mM HEPES, pH 7.5, combinations of 1 mM ZnCl2, 20 mM MnCl2, and 40 mM MgCl2 as indicated, and 150 mM NaCl at 37 °C. (A) PAGE analysis of single-turnover DNA cleavage by RadDz3, using 5′-32P-radiolabeled DNA substrate. Representative time points at t = 30 s, 5 h, and 24 h are shown. S = substrate; P = product. (B) Kinetic plots. For this data set, k_{obs} values (h⁻¹): Zn²⁺/Mn²⁺/Mg²⁺ 0.21, Zn²⁺/Mn²⁺ 0.22, Zn²⁺/Mg²⁺ 0.004, Mn²⁺/Mg²⁺ 0.17, Mn²⁺ 0.38. When Mn²⁺ was included at 1−300 μM along with 1 mM Zn²⁺ and 40 mM Mg²⁺, the DNA cleavage yield was the same as with Zn²⁺/Mg²⁺ in the absence of Mn²⁺ (data not shown), indicating that trace Mn²⁺ is not responsible for the Zn²⁺/Mg²⁺ reactivity. See the Experimental Section for quantitative analysis information on metal ion salts. (C) Determination of metal ion concentration dependence for Zn²⁺ (in the presence of 20 mM Mn²⁺) and Mn²⁺ alone. Data points with error bars were n = 2 (weighted averages; error bars by propagation from curve fit error); data points without error bars were n = 1. For Mn²⁺, squares were fit, and triangles were not fit. Apparent K_d = 2 ± 2 mM, Hill coefficient n = 2.8 ± 0.4, k_{max} = 0.10 ± 0.01 h⁻¹.
in 24 h, $k_{\text{obs}} = 0.0002 \text{ h}^{-1}$). The apparent $k_f$ value was $26 \pm 2$ mM for Mn$^{2+}$ (Figure 2C), with some inhibition above 100 mM. The Zn$^{2+}$ concentration dependence was assayed in the presence of Mn$^{2+}$, due to the low catalytic activity with Zn$^{2+}$ alone. Between 0.2 and 2 mM Zn$^{2+}$, $k_{\text{obs}}$ varied by ~2-fold, with inhibition above 2 mM (Figure 2C), as observed with several other deoxyribozymes and perhaps due to precipitation of Zn(OH)$_2$.8a,15 Assays under multiple-turnover conditions did not provide evidence of turnover by RadDz3 (data not shown).

**Establishing a Radical Pathway for RadDz3.** During the in vitro selection process, the survival of RadDz3 cannot be explained by DNA phosphodiester hydrolysis, which would not have led to a capturable product. We therefore considered alternative reaction pathways. RadDz3 was fully inhibited by either glutathione or catalase (Figure 3A), where glutathione quenches radical intermediates and catalase destroys $\text{H}_2\text{O}_2$, suggesting a radical pathway and involvement of H$^\cdot$O$^2$- . Conversely, adding $\text{H}_2\text{O}_2$ strongly enhanced the catalysis (Figure 3B).16 At 100 $\mu$M $\text{H}_2\text{O}_2$, $k_{\text{obs}} = 10^{-1}$ s$^{-1}$ (50-fold higher than without $\text{H}_2\text{O}_2$), with even higher $k_{\text{obs}}$ (not measurable by manual mixing methods) at 1 mM $\text{H}_2\text{O}_2$. Superoxide dismutase (SOD) did not suppress cleavage product formation, but this outcome is difficult to interpret because superoxide (O$_2^\cdot$ - or HOO$^\cdot$) could react further immediately after its formation and have no opportunity to be contacted by SOD; also, SOD produces $\text{H}_2\text{O}_2$ from superoxide. Indeed, separately adding potassium superoxide (KO$_2$) also substantially increased the catalysis (Figure 3C), suggesting a pathway in which RadDz3 sequentially forms superoxide and then $\text{H}_2\text{O}_2$.

This combination of observations led us to consider the bleomycin radical pathway (Figure 4), in which abstraction of a hydrogen atom from the C4′ position of a nucleoside in the presence of O$_2$ leads to a 3′-phosphoglycolate product. Because 3′-phosphoglycolate is a carboxylic acid, this DNA-catalyzed reaction product would be capturable during the in vitro selection process of Figure 1, explaining the survival of RadDz3. Of all 2′-deoxyribose hydrogen atom abstraction intermediates that can be formed from ssDNA, only the C4′ intermediate leads to a carboxylic acid.19 Glutathione is known to quench formation of bleomycin products, presumably by returning a hydrogen atom to the C4′ radical.20 The amide linkage in the originally used substrate would not necessarily be required, and indeed, RadDz3 functions well using an entirely DNA substrate, with 1.7-fold higher $k_{\text{obs}}$ than with the amide-containing substrate (data not shown).21

The RadDz3 products, formed either with or without inclusion of $\text{H}_2\text{O}_2$ in the reaction, were analyzed by MALDI mass spectrometry, revealing formation of the 3′-phosphoglycolate and 5′-phosphate products that result from excision of most of a particular guanosine nucleoside of the DNA substrate, located in the middle of the substrate (Figure 5). The small (MW = 205) base propenal fragment formed from the excised mononucleoside was also detected via a colorimetric assay (Figure 6), thereby accounting for all DNA substrate atoms.24 The mfold-predicted secondary structure of RadDz3 is shown in association with its DNA substrate in Figure 7.

**A Second, N$_{100}$ Deoxyribozyme That Cleaves DNA by the Same Radical Pathway.** In parallel with the above experiments, we performed a separate, new in vitro selection experiment using an N$_{100}$ random sequence population, considering that, in general, the length of the initially random region can be an important experimental variable.26 This selection experiment used the amide substrate of Figure 1 along with solely unmodified DNA and resulted in one new deoxyribozyme, named RadDz6, that cleaves a simple DNA oligonucleotide substrate. Under single-turnover conditions and in the absence of $\text{H}_2\text{O}_2$, RadDz6 has only 6% yield in 24

![Figure 3](image-url)
h, whereas, in the presence of 100 μM H₂O₂, kₐ is 1.0 h⁻¹ with 39% yield in 24 h; both kₐ and yield are even higher at 1 mM H₂O₂ (Figure 8A). As was observed for RadDz3, inclusion of KO₂ also increased the catalytic ability of RadDz6 (Figure 8B). Mn²⁺ supported the greatest activity by RadDz6, either alone or in combination with Zn²⁺ or Mg²⁺ (Figure 8C). Significantly, the combination of 1 mM Zn²⁺ and 40 mM Mg²⁺ in the absence of Mn²⁺ also supported substantial catalysis, demonstrating that RadDz6, like RadDz3, does not require a redox-active metal ion cofactor for its catalytic activity. As for RadDz3, glutathione quenches product formation (data not shown). Analysis of the RadDz6 DNA cleavage products by MALDI mass spectrometry revealed the same 3′-phosphoglycolate and 5′-hydroxyl products as formed by RadDz3 (Figure 8D), and the base propenal product was detected colorimetrically (Figure 8E). The combined data strongly suggest that RadDz3 and RadDz6 form their DNA cleavage products through the common Figure 4 mechanistic pathway.

**DISCUSSION**

In this study, we describe a new DNA-catalyzed reaction in which a deoxyribozyme, accompanied by redox-inactive metal ion cofactors, cleaves a single-stranded DNA substrate with excision of most atoms of a specific guanosine nucleoside. The biochemical and mass spectrometry data fully support a radical reaction pathway directly analogous to the O₂-dependent bleomycin pathway, in which creation of a C4′ radical at the guanosine leads to formation of 3′-phosphoglycolate and 5′-phosphate DNA termini, along with loss of the remainder of the guanosine nucleoside as the base propenal fragment. The present findings constitute the first report of DNA-catalyzed oxidative DNA cleavage that does not require redox-active metals such as Cu²⁺ or Mn²⁺, as both the RadDz3 and RadDz6 deoxyribozymes can form their DNA cleavage products using only redox-inactive Zn²⁺ and Mg²⁺ as the cofactors. In contrast, the Cu²⁺/Mn²⁺-dependent
Oxidatively cleaving deoxyribozymes do not lead to well-defined products, instead forming a variety of fragments via "oxidative destruction" of one or more nucleosides.11,12 The new DNA-catalyzed cleavage reaction appears to represent a new mechanistic class for nucleic acid enzymes, considering the combination of radical-based chemistry and selectivity in product formation, in contrast to the myriad of oxidative DNA cleavage products formed by other deoxyribozymes.11,12 The current mechanistic model for the RadDz3/RadDz6 reaction pathway has unresolved aspects. The homogeneity of the reaction products indicates that the C4′ radical is formed at a specific guanosine nucleoside of the substrate, presumably as directed by the deoxyribozyme, but the details of C4′ radical generation are undefined by the current data. Metal ion redox activity is not required, because each deoxyribozyme is active in the presence of Zn2+/Mg2+. Additional work is required to understand formation of the C4′ radical and the detailed roles of superoxide and H2O2 in the reaction pathway. No damage of the deoxyribozymes themselves was detected by postreaction isolation and piperidine treatment (data not shown), indicating no obligatory redox participation of any of the deoxyribozyme nucleotides.

EXPERIMENTAL SECTION

Oligonucleotides. DNA oligonucleotides (including deoxyribozymes) were obtained from Integrated DNA Technologies (Coralville, IA) or prepared by solid-phase synthesis on an ABI 394 instrument using reagents from Glen Research. All oligonucleotides were purified by 7 M urea denaturing PAGE with running buffer 1×TBE (89 mM each Tris and boric acid and 2 mM EDTA, pH 8.3), extracted from the polyacrylamide with TEN buffer (10 mM Tris, pH 8.0, 1 mM EDTA, 300 mM NaCl), and precipitated with ethanol as described previously.7 The DNA substrate sequence was 5′-GGATAATACGACTCACTATTTGAAGAGATGGCGACTTCG-3′, where most of the underlined G is excised by RadDz3 and RadDz6.

In Vitro Selection and Identification of RadDz3 and RadDz6 Deoxyribozymes. RadDz3 was identified in our recent study that sought DNA-catalyzed amide hydrolysis, specifically from the in vitro selection experiment in which the H2O2 modification was included.7 The incubation conditions used during identification of RadDz3 were 70 mM HEPES, pH 7.5, 1 mM ZnCl2, 20 mM MnCl2, 40 mM MgCl2, 150 mM NaCl, and [H2O2] as indicated at 37°C. The new DNA-catalyzed cleavage reaction appears to represent a new mechanistic class for nucleic acid enzymes, considering the combination of radical-based chemistry and selectivity in product formation, in contrast to the myriad of oxidative DNA cleavage products formed by other deoxyribozymes.11,12 The current mechanistic model for the RadDz3/RadDz6 reaction pathway has unresolved aspects. The homogeneity of the reaction products indicates that the C4′ radical is formed at a specific guanosine nucleoside of the substrate, presumably as directed by the deoxyribozyme, but the details of C4′ radical generation are undefined by the current data. Metal ion redox activity is not required, because each deoxyribozyme is active in the presence of Zn2+/Mg2+. Additional work is required to understand formation of the C4′ radical and the detailed roles of superoxide and H2O2 in the reaction pathway. No damage of the deoxyribozymes themselves was detected by postreaction isolation and piperidine treatment (data not shown), indicating no obligatory redox participation of any of the deoxyribozyme nucleotides.
MnCl₂, 40 mM MgCl₂, and 150 mM NaCl at 37 °C. The RadDz3 sequence is 5′-CGAAGTCGCGCTCTTCTGACGTGGCAATAGTGAGTCGTATTATA-3′ (74 nt), where the initially random N₁₀₀ region is underlined. RadDz6 was identified in a new in vitro selection experiment in which only unmodified DNA was used, along with an initially random N₁₀₀₀ region. This experiment used the amide-containing substrate from Figure 1 (same substrate as was used in ref 7) and our earlier selection procedure, which uses the same capture strategy as depicted in Figure 1. The same incubation conditions as in identification of RadDz3 were used. After eight selection rounds, the capture yield was 8%, compared to 60–70% capture yield for the capture standard reaction, and individual deoxyribozymes were cloned from round 8. The RadDz6 sequence is 5′-CAGAGTGGCAATAGTGAGTCGTATTATA-3′ (134 nt), where the initially random N₁₀₀₀ region is underlined. No homology is evident between the RadDz3 and RadDz6 sequences.

Single-Turnover Deoxyribozyme Assay Procedure. The DNA substrate was 5′-[^32]P-radiolabeled using T4-PAT and polynucleotide kinase. A 10 μL sample containing 5 pmol of DNA substrate (of which 0.2 pmol was 5′-[^32]P-radiolabeled) and 15 pmol of deoxyribozyme was annealed in 5 mM HEPES, pH 7.5, 15 mM NaCl, and 0.1 mM EDTA by heating at 95 °C for 3 min and cooling on ice for 5 min. The DNA-catalyzed DNA cleavage reaction was initiated by bringing the sample to 300 μM total volume containing 70 mM HEPES, pH 7.5, 1 mM ZnCl₂, 20 mM MnCl₂, 40 mM MgCl₂, 150 mM NaCl, and 100 μM H₂O₂. The sample was incubated at 37 °C for 2 h; longer incubation times could not be used due to malondialdehyde degradation. After addition of 500 μL of 50 mM thiorbituric acid in water, the sample was incubated at 95 °C for 20 min and diluted by addition of 200 μL of water to a total of 1 mL volume. The absorbance at 532 nm was measured in a quartz cuvette with 1 cm path length (Thermo Scientific NanoDrop; 600 nm baseline subtraction). A calibration curve was created using malondialdehyde, which was prepared as follows. A sample containing 100 μL of 50 mM aqueous malondialdehyde bis(dimethyl acetal) (1,1,3,3-tetramethoxypropane; Aldrich), 100 μL of 1 M HCl, and 800 μL of water was heated at 55 °C for 1 h, cooled to room temperature, and diluted by addition of 4 mL of water, providing a 10 mM solution of malondialdehyde. A 0.1 mM stock solution of malondialdehyde was prepared by diluting 100 μL of the 10 mM solution to 10 mL. The calibration curve was created using 50–2000 pmol of malondialdehyde (0.5–20 μL of the 0.1 mM stock solution) in the above DNA cleavage procedure. The slope of the calibration plot was 0.145 nmol⁻¹.

### REFERENCES


### COMMUNICATIONS

The authors declare no competing financial interest.

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### Notes

The authors declare no competing financial interest.
analogous products but with excision of most of the T nucleoside.


They found that when the glucose oxidase oxygen scavenging system was precluded by its scavenging system, the PCD enzyme itself, in the absence of oxygen, could not be added as usual to the deoxyribozyme. Finally, physical purging of O2 using argon had no measurable effect.


21 We made several unsuccessful attempts to exclude O2. Use of the glucose oxidase oxygen scavenging system was precluded by its direct production of H2O2, and catalase could not be added as usual for this system because the deoxyribozyme itself generates and requires H2O2. Use of the protocatechuic acid oxygen scavenging system was thwarted because the PCD enzyme itself, in the absence of its protocatechuic acid substrate, was found to inhibit the deoxyribozyme. Finally, physical purging of O2 using argon had no measurable effect.