Structural basis for Diels-Alder ribozyme-catalyzed carbon-carbon bond formation

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The majority of structural efforts addressing RNA's catalytic function have focused on natural ribozymes, which catalyze phosphodiester transfer reactions. By contrast, little is known about how RNA catalyzes other types of chemical reactions. We report here the crystal structures of a ribozyme that catalyzes enantioselective carbon-carbon bond formation by the Diels-Alder reaction in the unbound state and in complex with a reaction product. The RNA adopts a λ -shaped nested pseudoknot architecture whose preformed hydrophobic pocket is precisely complementary in shape to the reaction product. RNA folding and product binding are dictated by extensive stacking and hydrogen bonding, whereas stereoselection is governed by the shape of the catalytic pocket. Catalysis is apparently achieved by a combination of proximity, complementarity and electronic effects. We observe structural parallels in the independently evolved catalytic pocket architectures for ribozyme- and antibody-catalyzed Diels-Alder carbon-carbon bond-forming reactions.

The discovery of the catalytic activity of RNA^{1,2} and the hypothesis of a prebiotic 'RNA world' have expanded the scope of enzymology to include other biopolymers than proteins. The currently known natural ribozymes catalyze only a narrow range of chemical reactions, namely the hydrolysis and transesterification of internucleotide bonds^{4,5}, and probably peptide bond formation⁶. However, in vitro selection and evolution have demonstrated that ribozymes are capable of accelerating a much broader reaction spectrum⁷. This finding and recent discoveries of metabolite-controlled RNA switches and ribozymes^{8,9} suggest that RNA might have performed an even broader range of activities in the preprotein realm, and that in vitro-selected ribozymes could be analogs of the missing links in the transition from an RNA world to modern protein-dominated life¹⁰. Whereas high-resolution structures and biochemical investigations of several natural ribozymes provide a basic understanding of how RNA carries out phosphodiester chemistry^{5,11}, little is known about how RNA catalyzes other reactions. To obtain a comprehensive picture of the catalytic abilities and limitations of ribozymes, it is thus important to expand structural and mechanistic investigations to *in vitro*–selected ribozymes^{12–14}. Such structural information can be especially valuable in the determination of the minimal RNA folds required for catalysis and, therefore, could be helpful in the investigation of the origin and evolution of natural ribozymes¹⁵.

Two examples describe the *in vitro* selection of ribozymes that accelerate the formation of carbon-carbon bonds by the Diels-Alder reaction 16,17 , a [4+2] cycloaddition reaction between a tethered diene and a

biotinylated dienophile. This reaction type, creating two carbon-carbon bonds and up to four new stereo centers in one step, has broad applications in organic chemistry for the stereospecific synthesis of complex molecules ¹⁸. In one case, a minimal 49-nucleotide ribozyme was rationally deduced from the sequences of the active RNA molecules ¹⁶, and this ribozyme was later found to accelerate the cycloaddition reaction free in solution (*in trans*) in a truly bimolecular fashion ¹⁹. The minimum ribozyme adopted a secondary fold composed of three helical segments, a central element defined by an asymmetric bubble containing opposing invariant UGCCA and AAUACU fragments, and a 5′-terminal GGAG sequence (red, **Fig. 1a**). The minimal ribozyme could furthermore be engineered to give catalytically active bipartite (**Fig. 1a**) and tripartite systems ^{16,20}. Catalysis proceeds with multiple turnover and high enantioselectivity, making this a unique ribozyme system ¹⁹.

A first insight into the principles of substrate recognition and the molecular determinants of stereoselectivity has been obtained by a study of the interactions of the ribozyme with a panel of 44 systematically varied substrate analogs²¹. The results suggest a major role for hydrophobic and van der Waals interactions, whereas hydrogen bonding and metal ion coordination seem less important for catalysis. Mutation analysis and probing identified interactions between nucleotides of the 5' end and the internal asymmetric bubble, suggesting a pseudoknot topology²⁰. In contrast to most known RNA aptamers²², the probing data indicate a preformed tertiary structure that shows no major changes upon substrate or product binding. These results lead to the proposal

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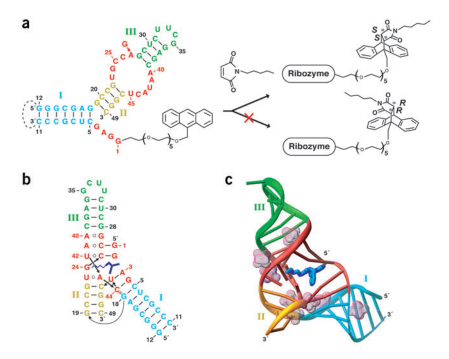


Figure 1 Secondary and tertiary folds of the Diels-Alder ribozyme. (a) Ribozyme secondary structure (left) with stems I-III in cyan, gold and green, respectively, and residues found invariant in the selection procedure in red. The same colors are used throughout this figure. The ribozyme consists of a 11-mer (G1-C11) HFG-linked with anthracene and a 38-mer (G12-C49). The dashed line indicates a loop present in the selected ribozyme and deleted in the bipartite system. The ribozyme-catalyzed Diels-Alder reaction between anthracene and N-pentyl maleimide results in formation of a single enantiomer of the complex (right). (b,c) Tertiary fold (b) and threedimensional topology (c) in the crystal structure of the ribozyme-product complex. The RNA bases and bound product (blue) are cylinders in **c**. The backbone is depicted with a ribbon, whereas the hydrated Mg²⁺ cations are in a mesh representation.

that the opposite sides of the asymmetric internal loop are most likely clamped by the four 5'-terminal nucleotides²⁰.

To obtain insights into the mechanism by which the Diels-Alder ribozyme catalyzes formation of carbon-carbon bonds, we determined the crystal structures of the ribozyme both in the unbound state and bound to the reaction product. These structures define underlying principles by which an RNA molecule carries out chemical reactions with small organic substrates and highlight the mechanism used to achieve stereoselectivity in catalysis.

RESULTS

Structure determination

Our structural efforts focused on the Diels-Alder ribozyme containing 38-mer and 11-mer RNA strands, with the shorter strand either unmodified or containing hexaethyleneglycol (HEG)-linked cycloaddition product (Fig. 1a). The product in complex with fully active A27G-substituted RNA gave the best crystals, and therefore initial efforts focused on determination of this structure. Because suitable heavy atom derivatives could not be identified using either soaking or halogen substitution approaches, we decided to rely entirely on the selenium (Se)-modification approach^{23,24}, which has been successfully applied for structure solution of a DNA duplex²⁵. Owing to the rapid radiation-induced decay of the crystals, six 2'-Se-methyl uridine (U_{Se}) and cytidine (C_{Se}) substitutions²⁴ were required for determination of the structure at a resolution of 3.0 Å by SAD (see Methods and Table 1). Next we determined the isomorphous crystal structure of the ribozyme-product complex lacking Se modifications and carrying adenine at position 27. These nearly identical structures provide details of the catalytic pocket, including bound product, surrounding RNA and coordinated divalent cations.

Overall topology and structural features

The molecule adopts a λ -shaped topology (**Fig. 1b,c**), with colinear stacking of stems II and III bridged by a zippered-up asymmetric bubble, whereas stem I is extended by two base pairs. Segments of the two arms of the tilted λ -shaped scaffold are positioned in close proximity, such

that the sugar-phosphate backbone of one of the strands of stem I is positioned within the major groove of stem II. The electron density centered about the catalytic pocket, built by the G1-G4 tetranucleotide and asymmetric bubble

segments, is shown in **Figure 2a**, and establishes that only the *S*,*S* stereoisomer of the Diels-Alder product can be readily traced into the map. In the structure of the product complex, we have found eight hydrated Mg²⁺ cations, which participate in stabilization of the RNA architecture and in crystal packing (**Fig. 1c**).

The 5'-G1-G2-A3-G4 segment plays a critical role in shaping both the RNA scaffold and the catalytic pocket (**Figs. 1b** and **2**). All four residues align through Watson-Crick base pairing with residues within the asymmetric bubble segment. The G1 and G2 residues form pairs with C26 and C25 respectively of one strand of the bubble, whereas the A3 and G4 residues form pairs with U45 and C44 respectively, of the opposite strand of the bubble. Notably, the 5'-GGAG segment bridges opposite strands of the internal bubble, thereby generating a complex nested pseudoknot topology.

The asymmetric bubble (residues 23–27 and 40–45) is zippered up through noncanonical pair formation between residues across its opposing strands. These involve a G/A27•A40 pair with purines aligned through their Watson-Crick edges (**Fig. 2b** and **Supplementary Fig. 1** online), and a reversed Hoogsteen U23•A43 pair thereby extending both stems III and II, respectively, by one step. The A41 residue aligns with the G1-C26 pair to form a three-base platform (**Fig. 2b** and **Supplementary Fig. 1** online), with potential for the formation of a C-H···O hydrogen bond. U42 participates in a U42•(G2-C25) base triple (**Fig. 3a**), where the alignment of U42 is dictated by formation of base-base and base-sugar hydrogen bonds. The unpaired G24 residue is sandwiched between U23 and U42 (**Fig. 3a** and **Supplementary Fig. 1** online), and forms three hydrogen bonds along its Watson-Crick edge with the phosphate oxygens of A43 and the carbonyl group of the maleimide ring of the bound product, while its 2' OH is hydrogen-bonded with U42.

All bases of the 5'-GGAG segment and all residues within the asymmetric internal bubble are involved in stacking interactions. The majority of the stacking is within individual strands but cross-strand stacking is found between U42 and G24 in the vicinity of the RNA catalytic pocket (**Supplementary Fig. 1** online). Extensive stacking is observed among the adjacent G27•A40 pair, A41•(G1-C26) three-base platform and U42•(G2-C25) base triple (**Fig. 2b**), between adjacent Watson-Crick

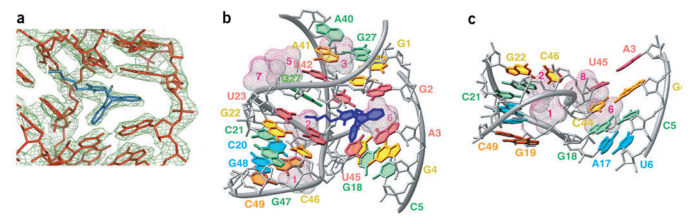


Figure 2 Details of the structure of the Diels-Alder ribozyme–product complex. (a) Final $2F_0 - F_c$ electron density map (green), contoured at 1 σ , for the cycloaddition product and surrounding RNA. The product (blue) and RNA (red), superimposed with the density, are from the refined model. (b) RNA tertiary structure surrounding the ribozyme catalytic pocket containing bound product. The product is blue, adjacent A3-U45 and U23•A43 base pairs and U42•(G2-C25) base triple are light red, G24 is green, with additional layers moving farther out from the pocket in gold (A41•(C26-G2), G22-C46 and C44-G4), light green (A40•G27, C21-G47 and G18-C5), cyan (C20-G48) and orange (G19-C49). The hydrated Mg²⁺ cations are in magenta-colored mesh. (c) Tertiary structure in the region of stems I and II. The RNA is slightly tilted compared with b to highlight the relative alignment of the stems. Individual base pairs and hydrated Mg²⁺ cations are shown as in b.

A3-U45 and Watson-Crick G4-C44 pairs (Fig. 3b), and between reversed Hoogsteen A43•U23 and Watson-Crick G22-C46 pairs (Fig. 3b), associated with the three stems that converge on the junction. Notably, the A43-C44-U45-C46 segment forms a sharp turn stabilized by two coordinated magnesium cations (Fig. 3b and Supplementary Fig. 1 online), with stacking of the first (A43) and last (C46) residues. The C44 residue, involved in turn formation, stacks on G18 (Fig. 3c). The G18 and G19 bases are splayed apart (Figs. 2c and 3c), thereby preventing stems I and II from adopting a colinear alignment, a feature recently observed in the structure of a group I intron²⁶. A disruption in stacking is also observed at the G24-C25 step (Fig. 3a and Supplementary Fig. 1 online), as G24 intercalates into the extended U42-A43 step. Finally, portions of stems I and II pack against each other, such that the backbone at the C44-U45 step of stem I is directed toward the major groove edge of stem II, with interactions mediated through bridging hydrated magnesium ions Mg1 and Mg2 (Supplementary Fig. 1 online).

Divalent cation-binding sites

Among the eight $\mathrm{Mg^{2+}}$ cations found in the structure, six participate in the stabilization of the RNA architecture (Mg1–Mg6) and two (Mg7 and Mg8) mediate contacts between RNA molecules in the crystal lattice. Mg2 seems to make the most important contribution to the structural scaffold because it makes multiple direct and water-mediated contacts

that bridge bases and sugar-phosphate backbones of six nucleotides situated beneath the product (**Fig. 3b**), and, together with Mg1, mediates the packing of helices I and II against each other (**Supplementary Fig. 1** online). Mg3 and Mg5 stabilize nonpaired G24 and the three-base platform and triple above the bound product (**Fig. 3a**), whereas Mg4 and Mg6 are located in the groove of helix I. There are no Mg²⁺ cations positioned in the immediate vicinity of the binding pocket that could potentially participate in the catalytic process. The divalent cation—dependence of the imino proton NMR spectra of the ribozyme—product complex has established that a minimum of two Mg²⁺ cations are required for RNA structure folding (**Supplementary Fig. 2** online). Although the direct evidence is missing, good candidates for such cations are Mg1 and Mg2 found in all Diels-Alder ribozyme structures reported in this article.

RNA-product interactions

The catalytic pocket is wedge-shaped (\sim 31° angle) and bracketed by the Watson-Crick A3-U45 pair, the noncanonical A43•U23 pair, and the U42•(G2-C25) triple (**Figs. 2b** and **3a,b**). Base G24 is positioned opposite to the opened edge of the pocket, and shares the space with the side chain projecting from the five-membered maleimide ring (**Figs. 2b** and **3a**). The cycloaddition product, which binds to the ribozyme with a dissociation constant in the 10 μ M range²¹, is positioned within the catalytic pocket with one of its aromatic rings sandwiched between purines

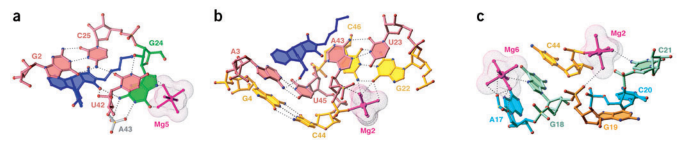
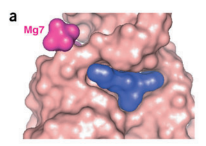
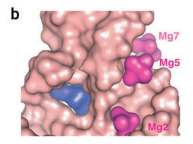


Figure 3 Structural elements of the Diels-Alder ribozyme–product complex. The view is rotated 180° along the vertical axis relative to the schematic in **Figure 1b**. The color code is the same as in **Figure 2**, unless specified otherwise. Oxygen, nitrogen and phosphorus atoms are, respectively, red, blue and yellow balls. Hydrogen bonds are dotted lines. Coordination bonds to Mg atom are magenta sticks. (a) RNA segment aligned above the product. The A43 backbone is silver. (b) RNA segment aligned below the product. (c) Structural motif highlighting splaying out of G18 and G19.





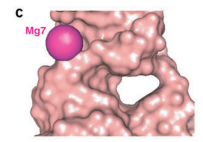
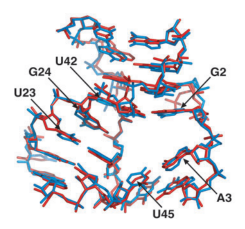


Figure 4 Surface representation of the catalytic pocket. (a,b) Front (a) and back (b) views of the catalytic pocket of the ribozyme–product complex. (c) Front view of the free ribozyme. Mg²⁺ ions are shown as hydrated cations in a and b for the 3.0-Å structure, and as balls in c for the 3.5-Å structure.

G2 and A3, and the other wedged between the base and sugar components of A43 and U45 (Figs. 2a and 3a,b). The HEG-linked bridgehead position is directed inward into the RNA scaffold, whereas the unmodified bridgehead position points outward toward solvent. The maleimide ring is stacked over C25, and oriented through a pair of hydrogen bonds made by one of its two carbonyl oxygens with the NH2 group of G24 and the 2' OH of U42 (Fig. 3a). The maleimide side chain runs inside a hydrophobic canyon created by bases C25 (top), A43 (bottom), and G24 (back) (Fig. 3a,b). The catalytic pocket, which is primarily formed by base edges with minimal contribution of the sugar-phosphate backbone, is accessible to the product from the front but not from the back (Fig. 4a,b). The 9' O atom (the one next to the bridgehead carbon) forms hydrogen bonds to the NH₂ of A3 (Fig. 3b). The hydrophobic pocket has room along the long edges of the cycloaddition product, but there is a snug fit along the short edges. The catalytic pocket contains not only a cavity to accommodate the cycloaddition product but also a directionspecific surface canyon to hold the maleimide side chain, thus allowing binding of only one product enantiomer.

Comparison of ribozyme structures

We have also determined the structure of the free Diels-Alder ribozyme at 3.5 Å, which is virtually identical to the structure of the product complex, indicative of a preformed architecture of the catalytic pocket (**Figs. 4c** and **5**). This conclusion is strongly supported by the imino proton NMR spectral patterns of the free ribozyme, its anthracene conjugate and product complex in Mg²⁺-containing solution, all of which exhibit resonance patterns associated with pairing of the G1-G2-A3-G4 segment (**Supplementary Fig. 2** online). This pairing probably generates the key interactions defining the geometry of the catalytic pocket. The imino proton NMR spectra of the samples with 38-mer plus 11-mer and the 38-mer plus 11-mer HEG-linked anthracene are very similar,



whereas the sample of 38-mer plus 11-mer HEG-linked product shows a better-resolved spectrum with a total of 20 imino protons. The unusually well-resolved spectrum for the product complex probably reflects both its compact scaffold and the slow exchange of imino protons lining the pocket, as a consequence of the positioning of the Diels-Alder reaction product within the catalytic cavity.

DISCUSSION

Architecture of the Diels-Alder ribozyme

This is the first report on the three-dimensional architecture of a ribozyme catalyzing a synthetic reaction between small organic molecules. The catalytic pocket is centered about a λ -shaped three-helical junction that is distinct from the reported topologies of natural ribozymes. In the hammerhead ribozyme²⁷, three helices converge toward a conserved sequence junction in a Y-shaped folding topology including a four-nucleotide turn element. The Varkud satellite ribozyme builds an active site using two three-way junctions²⁸. In the Diels-Alder ribozyme the three stems abut directly against the catalytic pocket, and are in part involved in the formation of an unusual nested pseudoknot. A double-pseudoknot architecture has been reported previously for the hepatitis delta helper virus (HDV) ribozyme²⁹, and both systems share common structural features, despite differences in pseudoknot topologies (Supplementary Fig. 3 online) and catalytic activities. Both ribozymes have preorganized active sites located in crevices, and both show a marked intertwining of secondary structure elements, leading to a densely packed, stable three-dimensional architecture. It is a notable coincidence that the HDV and Diels-Alder ribozymes, two of the fastest known ribozymes, both use nested pseudoknot architectures. The same pseudoknot topology is common to both the Diels-Alderase ribozyme and the putative fold in the regulatory region of the Escherichia coli α operon (Supplementary Fig. 3 online), which is specifically recognized and bound by ribosomal protein S4, leading to translational repression of the operon³⁰. Thus, the Diels-Alder pseudoknot topology may have relevance in a different biological context.

Correlation with biochemical data

The crystal structure is in excellent agreement with the observed structure-activity relations reported for ribozyme mutants²⁰. The bridging role of the G1-G2-A3-G4 segment correlates with the observation that essentially no single substitutions are tolerated within this sequence²⁰, whereas pairwise-complementary double substitutions lead to fully active mutants, at least for the A3-U45 and the G4-C44 base pairs. For

Figure 5 Superposition of the Diels-Alder ribozyme structures around the catalytic pocket. The RNA structures in stick representation are from the ribozyme–product complex (red) and the free ribozyme (blue).

G1 and G2, no compensatory double mutants were found, consistent with their involvement in the A41•(G1-C26) three-base platform and the U42•(G2-C25) base triple. The unpaired G24 residue is anchored in place by three specific hydrogen bonds involving its Watson-Crick edge; therefore, it is not surprising that its replacement by any other nucleotide results in loss of >90% of the catalytic activity²⁰.

Chemical probing has established that the N⁷ position of A43 is protected from modification, whereas the N¹ position of this nucleotide becomes more exposed upon tertiary structure formation²⁰. This is consistent with the observation of the important reversed Hoogsteen U23•A43 pair in the crystal structure, resulting in pairing of the N⁷ position along the Hoogsteen edge and exposure to solvent of the

 $\rm N^1$ position along the Watson-Crick edge. Related modification studies also established that the $\rm N^3$ positions of C25, C26 and C44 become increasingly protected from methylation upon tertiary structure formation, in agreement with formation of Watson-Crick G1-C26, G2-C25 and G4-C44 pairs.

Preformed catalytic pocket

The Diels-Alderase ribozyme forms a wedge-shaped catalytic pocket lined predominantly by bases and base pairs that are involved in extensive stacking interactions with neighboring residues. There are no backbone phosphates in the vicinity of the pocket, nor were divalent cations identified that could participate in the catalytic process. Sugar rings participate in defining the dimensions of the pocket, but not in hydrogen bonding with the product. The formation of a well-defined hydrophobic pocket, and its use to bind the reactants, is generally thought to be very difficult for RNA to achieve³¹.

The catalytic pocket architectures for the ribozyme in the free state and in the product complex are very similar. Independent chemical probing experiments support this fact, indicating that overall RNA folding is not changed upon addition of the anthracene substrate or the cycloaddition product²⁰. The observation of a preformed catalytic pocket for the Diels-Alder ribozyme contrasts with known examples of adaptive recognition found in ligand–RNA complexes^{22,32,33}.

The catalytic pocket must be able to accommodate both the diene and maleimide components of the reactants and the product. Previous studies have shown that substitutions along the longer edges of the diene ring are tolerated whereas substitutions along the shorter edges result in loss of catalytic activity²¹, consistent with the dimensions of the catalytic pocket in the crystal structure. The maleimide component is positioned within the walls of a canyon that seems to be wide enough to accommodate an unbranched alkyl or an aryl chain. Substitutions at the maleimide double bond positions result in total loss in activity²¹, as they would clash against the pocket-forming face of the tertiary G2-C25 base pair. Side chain reduction from pentyl down to ethyl results in decreased acceptance of the substrates, probably owing to reduced hydrophobic interactions with the walls of the canyon. By contrast, the open-ended nature of the canyon predicts that it should readily accommodate longer side chains, in agreement with experimental findings^{16,21}.

A proposed reaction mechanism

The Diels-Alder ribozyme structures, coupled with extensive chemical and biochemical experiments^{16,19–21}, suggest that the ribozyme should bind the substrates *in trans* in a precisely defined orientation within

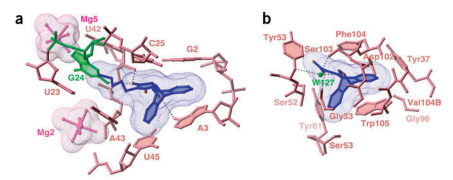


Figure 6 Comparison of the RNA and protein catalytic sites for the Diels-Alder reaction. The surface of the bound ligand is in mesh representation. (a) Catalytic pocket from the Diels-Alder ribozyme–product complex. (b) Catalytic pocket of the antibody catalyzing the retro-Diels-Alder reaction⁴¹ (PDB entry 1LO2). Anchoring of the ligand is achieved through direct and water-mediated (W127) hydrogen bonds.

the catalytic pocket, thereby facilitating the reaction by reducing translational and rotational degrees of freedom. The anthracene substrate could be bound by stacking interactions between G2 and the A3-U45 pair. The reaction requires that the maleimide substrate be stacked on top of the anthracene, and parallel to it, at a distance of ~3.5 Å. In this position, maleimide can form hydrogen bonds from its carbonyl oxygen to the exocyclic amino group of G24 and to the 2′ OH of U42. The N-alkyl side chain can be placed within the hydrophobic canyon (**Supplementary Fig. 4** online), which would allow the approach of the maleimide to the bound anthracene from only one direction, thereby providing the stereoselectivity of the reaction.

In the transition state of the reaction, computed using MOPAC (http:// www.cachesoftware.com/mopac/index.shtml) and published geometries of similar systems^{34,35}, two new single bonds are partly formed (bond lengths ~2.2 Å), and the anthracene ring system is bent out of planarity to yield an angle of ~154° (Supplementary Fig. 4 online), thereby further reducing the minimal unoccupied space underneath the bridgehead carbons seen in the structure of the ribozyme-product complex (Fig. 4a). Notably, the transition state has near-perfect shape complementarity with the catalytic pocket, a feature of prime importance in antibody catalysis of the Diels-Alder reactions^{36,37}. In addition to steric factors, both stacking and hydrogen bonding may contribute the energetics of the reaction. Stacking of the anthracene with nucleotides A3 and U45 could accelerate the reaction by increasing the diene's electron density, whereas hydrogen bonding of the maleimide would make it more electron-deficient, thereby increasing catalytic reactivity³⁸. Thus, it seems that ribozyme-based catalysis of Diels-Alder reactions reflects a combination of proximity, shape complementarity and energetic contributions to the catalytic process.

Comparison with protein-catalyzed reactions

The structural and mechanistic principles outlined above can be compared with the strategies used by antibodies catalyzing Diels-Alder and retro-Diels-Alder reactions. Several such antibodies have been crystallized in complex with either products or transition state analogs^{36,39–41}. Common features observed in these structures are the formation of hydrophobic cores lined predominantly by the side chains of tyrosine, phenylalanine, valine and tryptophan residues, and the stacking of one reactant over an aromatic residue, preferably tryptophan. Notably, the retro-Diels-Alderase catalytic antibody 10F11, the only antibody using anthracene as the diene⁴¹, forms a hydrophobic catalytic pocket with an overall shape similar to the Diels-Alder ribozyme pocket (**Fig. 6**). The anthracene ring is stacked over a tryptophan, similar to the posi-



tion of A3 in the ribozyme, with a few direct and water-mediated hydrogen bonds observed between heteroatoms on the bicyclic adduct and the amide backbone⁴¹.

The natural Diels-Alder enzyme also has a mainly hydrophobic catalytic pocket and interacts with the substrates by hydrogen bonding. Major differences, however, lie in the use of a bound Mg²⁺ cation, which directly coordinates to the carbonyl oxygens of the dienophile and participates in the catalytic mechanism, and in the generally lower importance of stacking in enzyme-substrate interactions^{42,43}.

Another Diels-Alder ribozyme requires both a covalent modification of the RNA and the presence of Cu²⁺ ions for catalytic activity, and apparently uses a different catalytic strategy^{17,44}. The structural prerequisites, however, are yet unknown.

The obvious similarities between antibody and RNA catalytic systems lead us to conclude that similar structural principles and catalytic mechanisms underlie macromolecule-mediated catalysis of Diels-Alder reactions. Notably, although RNA is equipped with a much less varied arsenal of functional groups than proteins, it has independently evolved matching criteria for generating catalytic pockets capable of facilitating carbon-carbon bond-forming reactions with comparable catalytic efficiency and enantioselectivity.

METHODS

RNA preparation. RNA (38-mer) was ordered from Dharmacon and CSS. Anthracene-HEG-linked 11-mer RNA was synthesized as described 16 . Synthesis of the stereochemically correct 11-mer with the HEG-linked Diels-Alder product was achieved by reaction between N-pentyl maleimide and anthracene-linked 11-mer annealed with 38-mer RNA 16 , and subsequent HPLC purification. RNAs containing Se modifications were chemically synthesized based on the 2'-O-TOM methodology, implementing $U_{\rm Se}$ and $C_{\rm Se}$ phosphoramidites according to published procedures 24 . The 5'-anthracene-HEG-linked 11-mer with $U_{\rm Se}$ at position 6 and $C_{\rm Se}$ at position 10 was converted into the product as described above. The 38-mer was prepared with $C_{\rm Se}$ at positions 15

and 20 and U_{Se} at positions 30 and 33. Details of preparation and purification of the RNAs are summarized in **Supplementary Methods** online.

Complex formation and crystallization. For annealing, 11-mer and 38-mer RNAs (200 μ M) were heated in 5 mM Tris-HCl, pH 8.0, 30 mM NaCl, 5 mM MgCl₂, at 95 °C for 2 min and cooled on ice. The RNA sample was combined with the same volume of the reservoir solution, and crystals were grown by hanging-drop vapor diffusion. Reservoir solutions were: (i) Se-modified ribozyme–product complex, 30% (v/v) PEG400, 10% (v/v) 2-methyl-2,4-pentadiol, 70 mM Na-HEPES, pH 7.5, 200 mM MgCl₂, 200 mM NaCl, 50 mM RbCl; (ii) unmodified ribozyme–product complex, 30% (v/v) PEG400, 100 mM Na-HEPES, pH 7.5, 200 MgCl₂, 0.5% (v/v) ethylacetate; (iii) free ribozyme, 30% (v/v) PEG400, 50 mM Na-cacodylate, pH 6.5, 80 mM Mg-acetate, 3% (v/v) 1,6-hexandiol, 50 mM NaCl, 15 mM CdCl₂. Crystals of the RNA-product complex grew to a maximal size of 0.4 × 0.1 × 0.02 mm in ~20 d at 4 °C. Free ribozyme crystals were grown at 20 °C.

Table 1 Data collection and refinement statistics

| | Product complex (Se and A27G) | Product complex | Free ribozyme |
|--|----------------------------------|-------------------------|---------------------|
| Data collection | | | |
| Space group | P2 ₁ | <i>P</i> 2 ₁ | P2 ₁ |
| Cell dimensions | | | |
| a, b, c (Å) | 77.87, 43.27, 79.93 | 78.28, 44.04, 80.00 | 77.69, 44.24, 79.63 |
| β (°) | 106.19 | 106.87 | 107.12 |
| Resolution (Å) | 30.0-3.0 (3.11-3.00) | 20.0-3.3 (3.42-3.30) | 20.0–3.5 (3.62–3.5) |
| R _{merge} | 8.4 (52.0) | 9.2 (16.2) | 15.4 (72.9) |
| Ι / σΙ | 11.2 (2.6) | 14.0 (8.0) | 8.1 (1.8) |
| Completeness (%) | 99.3 (98.2) | 97.7 (88.9) | 97.7 (98.2) |
| Redundancy | 3.6 (3.6) | 4.9 (4.4) | 3.8 (3.8) |
| Refinement | | | |
| Resolution (Å) | 20.0–3.0 | 20.0–3.3 | 20.0–3.5 |
| No. reflections | 10,481 | 7,909 | 6,563 |
| R _{work} / R _{free} | 0.209 / 0.234 | 0.228 / 0.262 | 0.286 / 0.324 |
| No. atoms | | | |
| RNA | 2,106 | 2,092 | 2,078 |
| Cycloaddition product | 56 | 56 | _ |
| Cations (Mg ²⁺ and Cd ²⁺) | 13 | 7 | 12 |
| Water | 83 | _ | - |
| B-factors ^a | | | |
| RNA | 25.0 | 23.9 | _ |
| Cycloaddition product | 26.0 | 23.8 | _ |
| Cations (Mg ²⁺ and Cd ²⁺) | 51.3 | 10.6 | _ |
| Water | 48.0 | _ | _ |
| B-Patterson / B-Wilson ^b | 42.4 / 37.9 | 46.3 / 25.8 | 70.4 / 91.2 |
| R.m.s deviations | | | |
| Bond lengths (Å) | 0.017 | 0.016 | 0.015 |
| Bond angles (°) | 1.539 | 1.480 | 1.612 |
| Estimated coordinate error ^c | 0.34 | 0.38 | 0.73 |
| Real-space R-factor ^d | 0.138 | 0.122 | 0.104 |

Values in parentheses are for the highest-resolution shell.

 $^{\circ}$ Owing to low resolution, individual B-factors were assigned to 40 Å $^{\circ}$ for all atoms during refinement. $^{\circ}$ Overall B-factor, estimated from Patterson origin peak, and Wilson B-factors were calculated using SFCHECK 49 . $^{\circ}$ CEstimated overall coordinate error based on maximum likelihood (ESU) was calculated with REFMAC 50 . 4 The mean real-space R-factor for composite simulated-annealing omit maps was calculated using CNS 51 .

Structure determination. X-ray data on the modified Diels-Alder ribozyme-product complex (containing six Se sites and A27G substitution) were collected at beamline X25 at the Brookhaven National Synchrotron Light Source (NSLS) from a single crystal cooled at 100 K. Data were processed with HKL2000 (HKL Research)⁴⁵. Owing to substantial radiation damage inflicted on the crystal, only the peak-wavelength data were used for phasing and all subsequent calculations.

Selenium sites were identified with SHELXD⁴⁶ run against all Bijvoet differences up to a resolution of 3.5 Å. The SAD phasing was carried out with SHARP⁴⁷, including the solvent flattening procedure with SOLOMON^{48,49}, assuming an optimized solvent content of 63% (**Supplementary Fig. 5** online). Anomalous phasing power was 1.08, and figure of merit after solvent flattening was 0.87. Data collection and refinement statistics are listed in **Table 1**. The initial RNA model was built using the SAD electron density map and TURBO-FRODO (http://afmb.cnrs-mrs.fr/rubrique113.html), and then refined with REFMAC⁵⁰ for the resolution range 20.0–3.0 Å. The noncrystallographic symmetry restraints were

applied between two molecules in the asymmetric unit. The Diels-Alder product and hydrated $\mathrm{Mg^{2+}}$ cations were added to the model based on analysis of the $2F_{\mathrm{o}}-F_{\mathrm{c}}$ and $F_{\mathrm{o}}-F_{\mathrm{c}}$ electron density maps (**Supplementary Fig. 5** online). The asymmetric unit contains two ribozymes, each bound to its Diels-Alder product, and 13 hydrated $\mathrm{Mg^{2+}}$ cations. The ribozymes are composed of 38-mer and 11-mer RNA molecules. The HEG linker is disordered.

The isomorphous crystal structures of the unmodified ribozyme–product complex and the free ribozyme were refined at a resolution of 3.3 and 3.5 Å, respectively, using the data collected at beamlines 14-BM of Argonne Advanced Photon Source and X25 of NSLS. Owing to limited resolution, no B-factor refinement was carried out for the free ribozyme structure.

NMR experiments. Imino proton NMR spectra were recorded using Varian and Bruker NMR spectrometers at 25 °C, using jump-and-return water suppression for detection (Supplementary Fig. 2 online).

Graphics. The figures were prepared with Ribbons (http://sgce.cbse.uab.edu/ribbons/) and PyMOL (DeLano Scientific; http://pymol.sourceforge.net/).

Coordinates. Coordinates for the Se-modified A27G-substituted Diels-Alder ribozyme–product complex at a resolution of 3.0 Å have been deposited in the Protein Data Bank (accession code 1YLS). The structures of the free ribozyme (3.5 Å) and unmodified ribozyme–product complex (3.3 Å) have also been deposited (accession codes 1YKQ and 1YKV, respectively).

Note: Supplementary information is available on the Nature Structural & Molecular Biology website.

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COMPETING INTERESTS STATEMENT

The authors declare that they have no competing financial interests.

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