Inhibition of Viral Replication by Ribozyme: Mutational Analysis of the Site and Mechanism of Antiviral Activity

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A controlled mutational study was used to determine the site and mechanism of the antiviral action of ribozymes that inhibit Sindbis virus replication. A hairpin ribozyme targeting G575 of the Sindbis virus genomic RNA was designed and cloned into a minimized alphavirus ampiclon vector. Cells that were stably transfected with this construct expressed low levels of a constitutive transcript containing the ribozyme plus recognition sequences for Sindbis RNA replicase. Upon infection, the ribozyme transcript was amplified to high levels by the viral replicase, resulting in decreased viral production from infected ribozyme-expressing cells. Mutations were then introduced into the viral RNA target sequence to interfere with ribozyme binding, and compensatory changes were generated in the ribozyme recognition sequence. Single mutations in the virus or ribozyme decreased the efficacy of the ribozyme’s inhibition of viral replication, and compensatory mutations restored it. To confirm that ribozyme-catalyzed RNA cleavage was actually needed for inhibition, we performed tests with a cell line expressing an inactivated ribozyme and with a virus containing a single nucleotide target mutation that allowed the ribozyme to bind but blocked cleavage at the recognition site. The results show that most of the antiviral activity of ribozymes is due to ribozyme-catalyzed cleavage at the targeted RNA sequence, but some additional inhibition seems to occur through an antisense mechanism.

The use of nucleic acids to achieve the selective inhibition of gene expression has been broadly investigated by the use of antisense, triplex, ribozyme, and RNA interference strategies (for reviews, see references 1, 16, and 22). While each approach has met with success, they all face common challenges, including intracellular activity and selectivity, subcellular localization, target site accessibility, delivery or expression, metabolism, and toxicity. The development of effective nucleic acid inhibitors of gene expression has not proven to be as facile as some had anticipated. Even in cases in which significant reductions of the targeted gene product have been realized, most studies have not been designed to determine if the inhibition resulted from the intended mechanism and site of action.

Hammerhead and hairpin ribozymes are small, sequence-specific RNA endonucleases whose specificity can be readily manipulated while retaining catalytic efficiency. These ribozymes have been used in several viral inhibition studies (6–9, 11, 20, 32). In our laboratory, we have focused on expressing hairpin ribozymes that are engineered to inhibit the replication of Sindbis virus in cultured cells. We have shown that combinatorial target site selection methods are an effective means of identifying target sites within viral RNA transcripts (31) and that hairpin ribozymes can be effectively engineered by using rules derived from the results of in vitro genetic and biochemical studies (30). Hairpin ribozymes are efficient and highly selective RNA endonucleases in the test tube and within eukaryotic cells (3, 19, 24).

Sindbis virus is the type virus of the Alphavirus genus in the Togaviridae family (18, 23), which is a group of viruses with single-stranded, message-sense RNA genomes that includes human and veterinary pathogens, including the Eastern, Western, and Venezuelan equine encephalitis viruses. Sindbis virus has a wide host range spanning insect, avian, and many types of mammalian cells. Sindbis virus strain S.A.A.R86 (21), which was utilized in our study, contains an 11,663-nucleotide (nt) single-stranded 42S RNA genome that also serves as the mRNA encoding the nonstructural proteins (nSPs; first two-thirds of the genome) and as the template for minus-strand RNA synthesis. A 26S single-stranded subgenomic RNA is also produced, driven by the subgenomic promoter on the minus-strand RNA and encoding the viral structural proteins.

The structure-rich RNA genome is infectious, and the replication cycle of the virus takes place in the cytoplasm following a receptor-mediated infection. In a rapid course of events after infection, several hundred thousand genomic and subgenomic RNAs are generated, and viral particles are secreted from the infected cells at 4 to 6 h postinfection at a rate of ca. 2,000 PFU/cell every hour (23). Cell death usually occurs 24 to 36 h after infection due to virus-induced apoptosis (5).

Considering its structure-rich RNA, high level of viral RNA replication, and large amount of virus output, the Sindbis virus system presents a rigorous challenge for the development of antiviral ribozymes. Previously, we showed that engineered hairpin ribozymes expressed from a U6 (RNA polymerase III [Pol III]) promoter in clonal mammalian cell lines can effectively inhibit Sindbis virus replication (20). However, these experiments were not designed to determine the mechanism or site of antiviral action.

In this article, we report the results of a series of experiments in which we used a molecular genetic approach to determine the site and mechanism of antiviral activity. An analysis of the effects of compensatory mutations in ribozymes expressed from clonal cell lines and in the Sindbis virus RNA genome showed that the antiviral activity results from the action of the
ribozyme at the targeted site in the Sindbis virus genome. An examination of the effects of ribozyme and viral mutations that block catalysis without affecting the binding of the ribozyme to the viral target showed that the observed antiviral activity results from a combination of ribozyme-catalyzed RNA cleavage and antisense effects.

**MATERIALS AND METHODS**

**Preparation of RNA.** Oligonucleotide RNA substrates were synthesized on an Applied Biosystems 392 DNA-RNA synthesizer and deprotected by standard procedures (27). Ribozymes and long substrates were transcribed from PCR products generated for ribozyme expression vector construction (described below) and from PCR products corresponding to nucleotide positions 422 to 681 of the viral genome by the use of T7 RNA polymerase and were purified by electrophoresis through denaturing polyacrylamide gels.

**Screening of accessible ribozyme target sites in Sindbis virus genomic RNA.** A DNA fragment containing nucleotide positions 88 to 875 was amplified from a full-length cDNA plasmid of the Sindbis virus (SIN) S.A.A.Rö6 strain (ps13, a gift from R. E. Johnston, University of North Carolina, Chapel Hill, N.C.) (GenBank accession no. U38303). It was then transcribed into RNA, and the RNA was purified by electrophoresis through a denaturing polyacrylamide gel. Protocols for randomized ribozyme pool preparation, cleavage reactions with the randomized ribozyme pool, and the mapping of cleavage sites by primer extension were described previously (31). The primers employed in the primer extension mapping experiments annealed to the following regions of the SIN genome: at nt 286 to 305, 469 to 490, 661 to 681, and 854 to 875.

**Ribozyme expression vectors.** A ribozyme targeting the G575 site of the SIN genome was designed according to standard parameters for hairpin ribozymes, the randomized ribozyme pool, and the mapping of cleavage sites by primer extension were described previously (31). The primers employed in the primer extension mapping experiments annealed to the following regions of the SIN genome: at nt 286 to 305, 469 to 490, 661 to 681, and 854 to 875.

**In vitro ribozyme cleavage assays.** All ribozyme cleavage assays were performed under similar reaction conditions as described previously (31) (also see the figure legends). Total RNAs extracted from the ribozyme-expressing cells or from SIN-infected naive BHK-21 cells were employed as a source of expressed ribozymes or as viral RNA substrates, respectively, for some experiments. The cleavage products were resolved in denaturing polyacrylamide gels and quantified by radioanalytic imaging. Cleavage rates were calculated as described elsewhere (33).

**Ribozyme inhibition of Sindbis Virus.** Ten micrograms of an XbaI-linearized SIN cDNA plasmid was transcribed by Sp6 RNA polymerase in a 100-µl reaction by use of a high-yield capped RNA transcription kit (Ambion). The infectious viral RNAs were mixed with 3 x 10^6 exponentially growing BHK-21 cells (prewashed with 1× phosphate-buffered saline) in 0.5 ml of 1× phosphate-buffered saline. The cells were electroporated three times at 850 V with a capacitance of 25 µF in a cuvette with a 0.4-cm gap and then transferred to a 100-mm-diameter dish with 10 ml of warm growth medium. Supernatants were harvested at 30 to 35 h posttransfection, when most cells were dead. The supernatants were clarified by centrifugation at 12,000 × g for 10 min at 4°C, and aliquots were stored at −80°C. The concentrations of the viral stocks were determined by plaque assays using BHK-21 cells.

**Cell culture and transfection.** BHK-21 cells were obtained from the American Type Culture Collection and cultured in Dulbecco’s modification of Eagle’s medium (Mediatech Cellgro) containing 10% fetal bovine serum, 10% tryptose phosphate broth, 2 mM l-glutamine, 100 U of penicillin/ml, and 100 µg of streptomycin/ml. Cells and cell lines were maintained in tissue culture incubators at 37°C with 5% CO2. Cotransfections were performed with 6 x 10^5 exponentially growing BHK-21 cells during exponential growth phase and with 0.5 ml of deoxynucleoside triphosphate (dNTP) mix (a 375 µM concentration of each dNTP) when limited primer extension was performed, 5 mM DTT, 10 mM NaCl, and 6 mM MgCl2 and 5 to 7 U of avian myeloblastosis virus reverse transcriptase. The elongation step was completed in a 42 to 45°C water bath for 45 min. The reactions were terminated by the addition of an equal volume of formamide loading buffer containing 10 mM EDTA, denatured at 85 to 90°C for 1 min, and resolved in a sequencing gel. Gels were quantified by radioanalytic imaging. A special dNTP-dNTP mix was employed when direct RNA sequencing was performed. A 100 µM concentration of each ddNTP was included in addition to the regular dNTP mix, which allowed a partial termination of primer elongation and the formation of a sequencing ladder on a denaturing gel.
RESULTS

Combinatorial target site selection. To build upon previous experiments in which engineered hairpin ribozymes were used to inhibit Sindbis virus replication (20, 31), we made several changes which were designed to improve (i) the target site, (ii) the catalytic activity of the ribozyme under physiological conditions, and (iii) the expression levels and subcellular localization of the ribozyme. We chose to target the 5′/H11032 two-thirds of the Sindbis virus genome, which encodes nonstructural proteins that are essential for viral replication and contains sequences that are not contained within the highly expressed 26S subgenomic RNA.

A combinatorial library of hairpin ribozymes containing all possible substrate specificities (31) was transcribed and used to probe target site accessibility within the 800 nt at the 5′ end of the Sindbis virus genomic RNA. We identified 16 target sites within this sequence, each of which could be cleaved by one or more ribozyme variants within the combinatorial library with a moderate to high efficiency. Data for two of these sites, G575 and G586, are shown in Fig. 1B.

For the mutational analysis described here, we chose to develop ribozymes that cleave G575 of the genomic RNA. This target site permitted us to create mutations within the viral RNA in such a manner as to change the activity of the ribozyme without modifying the amino acid sequence of the encoded viral protein, nsP1. Although several other potential target sites identified by this screening showed stronger cleavage patterns (data not shown), their local sequences did not permit us to introduce proper silent mutations into the viral genome to address the specificity and mechanism of the hairpin ribozyme. The relatively modest activity of ribozyme 575 in vitro led us to construct a four-way junction (4WJ) version to reach a higher cleavage activity under intracellular conditions, as described below.

Ribozyme design and catalytic activity. Hairpin ribozymes targeted to G575 of the Sindbis virus genome were designed

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**FIG. 1.** Ribozymes and viral targets. (A) Secondary structure of ribozyme-substrate complex. Green, sequences contained in standard two-way helical junction ribozyme construct; yellow, additional sequences contained in four-way helical junction ribozyme construct; pink, sequences of targeted region of Sindbis virus genomic RNA. H1 to H6, helical elements of ribozyme-substrate complex. The ribozyme shown was designed by use of the results shown in panel B. Note that the identities of ribozyme nucleotides 16 and 48 have been changed from the standard sequence in order to prevent a predicted misfolding event. (B) Combinatorial screening for optimal ribozyme cleavage sites within viral genomic RNA. A library of hairpin ribozymes containing all possible substrate specificities (31) was incubated with Sindbis virus genomic RNA sequences, and cleavage sites were mapped by the extension of 5′-32P-labeled oligonucleotide primers with reverse transcriptase. Control lanes (lacking ribozyme and lacking magnesium) were used to identify reverse transcriptase stop sites that did not result from ribozyme-catalyzed RNA cleavage. Sequencing lanes used appropriate ddNTPs for sequence-specific chain termination. An autoradiogram of a 6% denaturing polyacrylamide gel is shown. (C) Time course of RNA cleavage with ribozymes targeted to the Sindbis genomic G575 target. Ribozymes were generated by in vitro transcription with T7 RNA polymerase and applied to cleave a 5′-32P-labeled oligoribonucleotide substrate in a buffer containing 2 mM MgCl2, 150 mM NaCl, 0.1 mM EDTA, and 50 mM HEPES (pH 7.5). (D) Cleavage of structured RNA substrates. The experiment was the same as that described for panel C, except that the substrate was a uniformly labeled 250-nt RNA generated by in vitro transcription under single-turnover conditions in a buffer containing 12 mM MgCl2 and 40 mM Tris-HCl (pH 7.6).
and transcribed for examinations of their catalytic activities in vitro. We generated both a minimal ribozyme, organized around a two-way helical junction (2WJ), and a 4WJ ribozyme in which two additional helices were introduced, one of which (helix 6) provided additional base pairing between the ribozyme and its substrate (Fig. 1A). To facilitate the dissociation of the 3'/H11032 cleavage product and thereby increase the extent of cleavage by reducing the ligation of the cleavage products, we destabilized helix 1 in the four-way junction ribozyme by reducing it from 6 to 3 bp (28). In addition, the C16-G48 base pair in helix 3 of the ribozyme was replaced with U16-A48 in order to avoid the predicted misfolding of the four-way junction construct.

Results of in vitro cleavage assays are shown in Fig. 1C and D. In a standard single-turnover reaction using long (ca. 250 nt) Sindbis virus-derived sequences conducted in cleavage buffer containing a higher concentration of magnesium ions (12 mM) than that available in the mammalian cytoplasm, the four-way junction ribozyme showed a higher rate and extent of cleavage than did the two-way junction ribozyme over the time interval examined. When an analogous experiment was conducted with an oligoribonucleotide substrate (Fig. 1D), the four-way junction ribozyme showed a higher rate and extent of cleavage than did the two-way junction ribozyme over the time interval examined. When an analogous experiment was conducted with an oligoribonucleotide substrate in a buffer with a more physiological magnesium ion concentration (2 mM), the catalytic advantage of the four-way junction ribozyme was even more pronounced. The enhanced activity of the four-way junction ribozyme 575 is likely due to more efficient folding into the active tertiary structure, which is stabilized by the stacking of helices 2 and 3 upon helices 6 and 5, respectively (10, 25, 26).

Because of its significantly higher activity under near-physiological ionic conditions, we selected the four-way junction form of ribozyme 575 for subsequent cellular expression and viral inhibition assays.

Viral genomic mutations and ribozyme mutations. For an analysis of the site and mechanism of antiviral activity in vivo, we designed a series of mutations within the antiviral ribozyme 575 and within the infectious viral genomic RNA, at and around the intended ribozyme target, nucleotide G575. Mutations in the viral target sequence were designed to generate synonymous codons so that the amino acid sequence of the nsP1 gene product was unchanged. To ensure that the ribozyme and viral target sequences interacted in the expected manner, we conducted in vitro cleavage assays with in vitro-transcribed ribozymes and oligoribonucleotide substrates (Fig. 2).

The ribozyme-substrate complex was destabilized by single base substitutions at the terminal base pair of helix 1 (H1) proximal to the cleavage site, formed by C578 of the viral genomic RNA G6C of the antiviral ribozyme 575. A C578G substitution in the viral target was inhibitory, and a G6C substitution in the ribozyme was strongly inhibitory (compare Fig. 2B and C to Fig. 2A), although the association of the two mismatched ribozyme-target combinations was apparently not affected, as shown by in vitro gel mobility shift assays (data not shown). The compensatory base pair substitution (G6C in the ribozyme and C578G in the viral RNA) significantly increased the cleavage activity with respect to each of the individual
FIG. 3. Virus-induced amplification of ribozyme transcripts. (A) Amplification scheme. pRSV, promoter recognized by RNA polymerase II; tRNAAsp, partial tRNA encoded by defective interfering viral RNA; 5'-CSE, cis-acting element of Sindbis virus genome; pSIN26S, promoter for 26S subgenomic RNA, recognized by Sindbis virus RNA replicase; 3'-UTR, untranslated region required for the transcription of 5'-strand RNA. (B) High-level expression of ribozyme amplicon upon infection with Sindbis virus. The results of limited primer extension assays, in which 5'-end-labeled oligonucleotide primers were extended by reverse transcriptase in a reaction containing three dNTPs and one ddNTP (ddTTP for ribozyme...
mutations, although the cleavage rate of the compensatory mutational combination was somewhat lower than that observed for the wild-type combination of ribozyme and substrate (Fig. 2D). These results are consistent with prior mutational studies on the hairpin ribozyme and its substrates.

Two other mutations provided important probes of ribozyme activity and target selectivity. First, we used mutations at ribozyme nucleotide G8, which is a component of the active site, where base substitutions are significantly inhibitory (13). The introduction of a G8A substitution into a mismatched G6C ribozyme caused a significant reduction in cleavage activity (Fig. 2E), although the incorporation of the four-way helical junction into these ribozyme constructs acted to partially suppress the inhibitory effects of this and other mutations (e.g., C578G). Second, a substitution of the cleavage site G (G575) nucleotide strongly inhibits cleavage because it functions to form a tertiary base pair with C25 of the ribozyme which is, in turn, important for positioning G8 within the active site (13–15). It is important that neither the G8A ribozyme mutation nor the G575A target site mutation inhibits the formation of the ribozyme-substrate complex (13, 14). Therefore, they were used as inactive controls for the present study in order to distinguish viral inhibition by ribozyme-catalyzed cleavage from antisense effects at the same site.

Cytoplasmic amplification of ribozymes upon Sindbis virus infection. Previous expression work in our lab focused on the use of the U6 snRNA promoter to direct the transcription of ribozymes by RNA Pol III. For this study, we explored the use of Sindbis virus infection to trigger the cytoplasmic amplification of antiviral ribozymes by the viral RNA replicase when the ribozyme was expressed constitutively at low levels as a Pol II transcript (Fig. 3A). To accomplish this, we cloned ribozyme 575 downstream of the 26S subgenomic promoter of a Sindbis virus amplicon derived from a naturally occurring defective interfering RNA, DI25 (2, 12). Note that this vector does not drive expression of the ribozyme sequences to form a tertiary base pair with C25 of the ribozyme which is, in turn, important for positioning G8 within the active site (13–15). It is important that neither the G8A ribozyme mutation nor the G575A target site mutation inhibits the formation of the ribozyme-substrate complex (13, 14). Therefore, they were used as inactive controls for the present study in order to distinguish viral inhibition by ribozyme-catalyzed cleavage from antisense effects at the same site.

Biochemical analysis of biological transcripts. To ensure that the cellular RNA transcripts containing ribozyme sequences possessed the expected cleavage activities, we used RNAs extracted from cell lines expressing ribozymes to cleave oligoribonucleotide substrates in vitro. The results confirmed that the ribozyme-expressing cell lines contained the expected endonuclease activities (Fig. 4A, lanes 1 and 2) and that RNAs extracted from a cell line expressing the G8A mutant ribozyme exhibited a marked reduction in activity (Fig. 4A, lane 3). In a complementary experiment, we used total RNAs extracted from virus-infected cells as substrates in cleavage experiments performed with ribozymes that were generated in vitro. These experiments showed the cleavage of viral RNA at the expected site when the antiviral ribozyme 575 was incubated with total RNAs extracted from cells infected with wild-type Sindbis virus (Fig. 4B, lane 4). As expected, no cleavage was observed with RNAs from cells infected with the uncleavable G575A mutant virus or when the ribozyme was omitted (Fig. 4B, lanes 5 and 6).

Mutational analysis of antiviral mechanism and target site. After the characterization of stable cell lines expressing each of the ribozyme variants (the antiviral ribozyme 575, the mismatched G6C ribozyme, and the reduced-activity G8A ribozyme), we used plaque assays to examine the time course of viral replication following infection by wild-type Sindbis virus or one of two engineered mutants (mismatched C578G and uncleavable G575A mutant viruses).

The abilities of ribozymes to inhibit the replication of wild-type Sindbis virus are shown in Fig. 5A to D. The expression of the antiviral ribozyme 575 resulted in an 8- to 10-fold inhibition of the formation of infectious viral particles from 12 to 36 h after infection (Fig. 5B and D), with the time course of virus production in cells that did not express the ribozyme being used as a reference. Smaller extents of inhibition were observed at the 5-h time point, when newly synthesized viruses were only beginning to be released, and at the 48-h time point, when ribozyme-expressing cells were thriving and releasing virus at a reduced rate, while infected naive cells had under-
FIG. 4. Biochemical analysis of biological transcripts. (A) Activity of ribozymes expressed within cells. Total cellular RNAs were extracted from clonal cell lines expressing ribozymes and used for in vitro cleavage of a 21-nt oligoribonucleotide substrate. In all cases, total RNAs were harvested from cells 16 h after infection with wild-type Sindbis virus at a multiplicity of infection of 0.5. M, markers. Lane 1, 5 μg of total RNA from cells expressing ribozyme 575; lane 2, 10 μg of total RNA from cells expressing ribozyme 575; lane 3, 10 μg of total RNA from cells expressing the G8A reduced-activity ribozyme. Total RNAs were incubated with 5'-end-labeled WT synthetic RNA (21 nt) in annealing buffer (150 mM NaCl, 0.1 mM EDTA, and 50 mM HEPES, pH 7.5) for two cycles of 95°C for 2 min and 37°C for 10 min in a PCR instrument. MgCl2 was added to 2 mM, and reactions were incubated for 4 h at 37°C. Reaction products were separated by electrophoresis through a denaturing 20% polyacrylamide gel. (B) Ribozyme cleavage of viral RNA synthesized within cells. BHK-21 cells were infected with either wild-type or noncleavable G575A mutant Sindbis virus at a multiplicity of infection of 0.3, and total cellular RNAs (including viral RNA) were extracted 24 h after infection. Ribozyme 575 (2 μM), generated by in vitro transcription and purified by denaturing gel electrophoresis, was incubated with the cellular RNA at 37°C for 2 h in a buffer containing 2 mM MgCl2, 150 mM NaCl, 0.1 mM EDTA, and 50 mM HEPES (pH 7.5). Cleavage products were identified by a primer extension assay, using a radiolabeled oligonucleotide primer complementary to the Sindbis virus RNA downstream of the cleavage site. Lane 4, wild-type Sindbis viral RNA incubated with ribozyme 575; lane 5, mutant Sindbis virus carrying the G575A mutation, which blocks ribozyme-catalyzed RNA cleavage at a step following the formation of the ribozyme-substrate complex; lane 6, wild-type Sindbis viral RNA incubated in the absence of ribozyme.

Studies in RNA biology have led to several novel strategies for the development of novel RNA-based agents that may have potential in therapeutics, including ribozymes, small interfering RNAs, and aptamers. Like other strategies for gene therapy, there are many potential difficulties for the development of a safe and practical therapy. We have been working systematically to identify and overcome the challenges necessary to develop ribozymes as antiviral agents and have shown that engineered hairpin ribozymes can be expressed within cells and successfully used to inhibit the replication of members of three diverse viral families, namely, human immunodeficiency virus type 1 (HIV-1), hepatitis B virus, and Sindbis virus (20, 29, 32).

Prominent among the challenges in gene therapy are the issues of the biological mechanism and the site of action of the therapies being developed. For example, our previous work...
with hairpin ribozymes to inhibit the replication of Sindbis virus is consistent with the intended mechanism, i.e., ribozyme-catalyzed cleavage of the targeted site within the viral RNA. However, these results do not rule out other plausible antiviral mechanisms, e.g., antisense inhibition, unintended cleavage of a cellular RNA whose product is essential for viral replication, or insertion of a ribozyme-encoding DNA within a host cell gene that is important for viral replication.

For this study, we used a genetic strategy to determine if the ribozyme was acting on the intended viral target site and to distinguish between viral inhibition due to ribozyme-catalyzed cleavage and that due to a general antisense mechanism involving binding of the ribozyme to the target site. Our approach has been to generate mutations based on our understanding of the behavior of the ribozyme-substrate complex in vitro and then to determine how these mutations affect antiviral activity in cell culture. To this end, we generated infectious virions with specifically designed mutations at and around the ribozyme’s intended target site within the viral genome and also constructed clonal cell lines expressing a series of ribozyme variants that would affect the site and mechanism through which the ribozyme would act.

In all cases, we chose single base mutations within the Sindbis virus genome. Because the cleavage site lies within an essential coding sequence, it was necessary to find a target site and to choose mutations that would have no effect on the amino acid sequence of the translation product and to demonstrate that viruses containing the mutations replicated with the same growth characteristics as the wild-type strain.

The present study establishes the site of antiviral action through two complementary experiments. First, we used a compensatory mutational strategy to test the effects of single base substitutions within helix 1 of both the ribozyme and the viral RNA target. Separately, these mutations each destabilized the ribozyme-substrate complex and diminished the antiviral activity of the ribozyme. Together, they restored full base pairing within the complex and were observed to restore the full antiviral effect. Second, we mutated the cleavage site guanosine (G575) at the intended viral target and observed a reduction in antiviral activity that resulted from the loss of ribozyme-catalyzed cleavage of viral RNA at position 575.

Our work also established that the mechanism of antiviral activity includes two components, (i) ribozyme-catalyzed cleavage at the target site and (ii) an antisense component involving the formation of an inactive ribozyme-substrate complex. The residual inhibition that was seen with two reduced-activity complexes, the G8A ribozyme mutation and the G575A mutant virus, was attributed to antisense inhibition, although the G8A mutation alone did not completely inactivate the catalytic activities of the four-way junction ribozymes used in our study. We believe that antisense inhibition of viral replication does not involve RNA-induced silencing complex-mediated RNA interference because all of the helices in the ribozyme-substrate complex are significantly shorter than those required for cleavage by the dicer endonuclease.

To ensure that the population of cells was homogeneous with respect to ribozyme activity, we chose to generate clonal cell lines that constitutively expressed hairpin ribozymes within...
a Pol II transcript that was amplified following Sindbis virus infection of the ribozyme-containing cell. The vector used, D125, is different from common alphanavirus replicon vectors, which encode viral nonstructural proteins that lead to cytopathology and cell death (17). In our system, the amplification of ribozyme-containing transcripts is triggered by Sindbis virus infection and is supported by the proteins encoded in the incoming viral genome. Previous studies (2, 12) and our controls clearly show that the amplicon per se does not inhibit viral replication.

This amplification system has two apparent advantages. First, ribozyme-containing transcripts are amplified to very high levels that are similar to the levels attained by the Sindbis virus genomic RNAs themselves. Second, the ribozymes and viral RNA targets are colocalized within the cytoplasm. However, there is also an apparent disadvantage in that early in the infection there may not be enough ribozymes present (prior to translation of the viral nonstructural proteins) to prevent infection of the cell. Our previous work has shown that the most successful ribozymes may be those that prevent the initiation of viral replication within a cell (20). Consequently, our present strategy may be improved through the generation of higher levels of ribozyme transcripts prior to viral infection and by an enhancement of ribozyme activity in the cytoplasm. In addition, Sindbis virus packaging cell lines (2) may be utilized to generate pseudoviruses that contain the ribozyme amplicon used for this study. With the wide host range of Sindbis virus, pseudoviruses may be a useful means to deliver ribozymes for therapeutic applications.

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