Identification of inhibitors of ribozyme self-cleavage in mammalian cells via high-throughput screening of chemical libraries

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ABSTRACT

We have recently described an RNA-only gene regulation system for mammalian cells in which inhibition of self-cleavage of an mRNA carrying ribozyme sequences provides the basis for control of gene expression. An important proof of principle for that system was provided by demonstrating the ability of one specific small molecule inhibitor of RNA self-cleavage, toyocamycin, to control gene expression in vitro and vivo. Here, we describe the development of the high-throughput screening (HTS) assay that led to the identification of toyocamycin and other molecules capable of inhibiting RNA self-cleavage in mammalian cells. To identify small molecules that can serve as inhibitors of ribozyme self-cleavage, we established a cell-based assay in which expression of a luciferase (luc) reporter is controlled by ribozyme sequences, and screened 58,076 compounds for their ability to induce luciferase expression. Fifteen compounds able to inhibit ribozyme self-cleavage in cells were identified through this screen. The most potent of the inhibitors identified were toyocamycin and 5-fluorouridine (FUR), nucleoside analogs carrying modifications of the 7-position and 5-position of the purine or pyrimidine bases. Individually, these two compounds were able to induce gene expression of the ribozyme-controlled reporter; 365-fold and 110-fold, respectively. Studies of the mechanism of action of the ribozyme inhibitors indicate that the compounds must be incorporated into RNA in order to inhibit RNA self-cleavage.

Keywords: gene regulation; ribozyme self-cleavage; ribozyme inhibitors; toyocamycin; 5-fluorouridine

INTRODUCTION

The ability to “exogenously” control the expression of genes in mammalian cells has been a powerful tool of biomedical research (Gossen and Bujard 2002). Gene regulation technology has already played a critical role in efforts to understand the role of specific gene products in fundamental biological processes, in development, and in disease states. In the future, the technology offers the opportunity to have a major impact in new areas of research and medicine. To date, most commonly used gene regulation systems rely on the control of transcription (Gossen and Bujard 1992; Wang et al. 1994; Rivera et al. 1996; Suhr et al. 1998). More recently, the discovery of RNA interference and siRNAs has led to the development of novel strategies for controlling gene expression that rely on the targeting of RNA (Fire et al. 1998; McManus and Sharp 2002). In the hopes of extending the application and utility of gene regulation technology, we have been interested in the development of new methods for controlling gene expression that rely on the modulation of RNA self-cleavage rather than transcription. The basis for such a system is the incorporation of sequences encoding self-cleaving RNA motifs (i.e., ribozyme sequences) into a mammalian transcription unit, such that upon generation of the resulting mRNA, spontaneous ribozyme self-cleavage should lead to destruction of the mRNA and therefore no gene expression. Administration of agents capable of inhibiting RNA self-cleavage should result in preservation...
of the intact mRNA and therefore reinstates (i.e., “induce”) gene expression (Fig. 1A).

The two essential requirements of such an RNA-only gene regulation system are the availability of ribozyme sequences capable of efficient self-cleavage in mammalian cells when embedded in an mRNA transcript, and the availability of small molecules capable of inhibiting such self-cleavage in mammalian cells. With regard to the first requirement, we have recently tested a large number of natural and synthetic ribozyme sequences for their ability to function in mammalian cells when embedded in an mRNA transcript. We were able to identify and further develop a modified version of a hammerhead ribozyme, derived from *Schistosoma mansoni* (Sm1), that is capable of extremely efficient self-cleavage in mammalian cells (Yen et al. 2004). This ribozyme, termed N79, was found to function in a variety of commonly used mammalian cell lines and in primary cells in vivo. An important proof of principle for the use of inhibitors of RNA self-cleavage to control gene expression was provided by the demonstration of the ability of one specific small molecule inhibitor of RNA self-cleavage, toyoamycin, to control gene expression in vitro and vivo. Here, we describe the development of the high-throughput screening (HTS) assay that led to the identification of toyoamycin and other inhibitors of RNA self-cleavage. Rather than rely on in vitro screens of compounds, we chose to develop a mammalian cell-based screen in order to directly identify molecules capable of functioning within cells. Since a variety of aminoglycoside (Stage et al. 1995; Murray and Arnold 1996; Hermann and Westhof 1998; Tor et al. 1998; Jenne et al. 2001) and non-aminoglycoside antibiotics (Jenne et al. 2001) had been previously shown to be able to inhibit the self-cleavage of hammerhead ribozymes in the in vitro setting, we first screened such compounds in the cell-based assay. We then extended the studies to include the high-throughput screening of 58,076 compounds. We report here the results of those screening efforts and the characteristics (and in some cases, the mechanism of action) of the inhibitors that were identified.

RESULTS

Generation of reporter cell lines for cell-based screening

For screening purposes, we chose to generate a stable human (HEK-derived) cell line that carries an integrated mammalian expression vector in which a luc reporter gene’s expression is placed under the control of the CMV promoter and with two copies of the N79 ribozyme (Fig. 1B). Stable cell lines were generated by cotransfection of HEK 293 cells with the reporter construct, an expression vector containing the luciferase gene under the control of the CMV promoter, a ribozyme expression vector, and a vector coding for the N79 ribozyme. The stable cell lines were then screened for the presence of luciferase activity and for the ability to be induced by morpholino anti-sense oligonucleotides. The cell lines that showed the highest luciferase activity and were inducible by morpholino anti-sense oligonucleotides were selected for further analysis.

**FIGURE 1.** Strategy for identifying new ribozyme inhibitors via high-throughput screening. (A) Controlling gene expression via the modulation of RNA self-cleavage. When a cis-acting hammerhead ribozyme is embedded in the mRNA, self-cleavage leads to the destruction of the mRNA and results in the absence of gene expression. However, the administration of specific inhibitors of the ribozyme leads to the generation of intact mRNAs and results in protein expression. (B) Schematic representation of the N79 ribozymes and the reporter gene expression vector used for the cell-based screening assay. Stable cell clones that had low background luc expression and could be induced by morpholino anti-sense oligonucleotides as shown in the bioluminescent images were selected for the use in the screen. The black line identifies the sequence targeted by the anti-sense oligonucleotide. Nucleotide numbering follows standard nomenclature (Hertel et al. 1992).
vector encoding puromycin resistance, and subsequent selection in puromycin-containing medium. Approximately 10 clonal cell lines were generated in this way. To identify specific cell lines capable of responding to putative inhibitory molecules, all cell lines were tested for both their basal level of luc expression and for the extent of induction of luc expression achieved after transfection of the cells with an anti-sense morpholino oligonucleotide known to block the cleavage of N79 (Yen et al. 2004). Several cell lines tested in this way expressed very low basal levels of luc, yet were readily induced to express luc after administration of oligonucleotide. One of these cell lines, termed HEK79, showed a 20-fold induction of luc expression after oligonucleotide administration and was chosen for high-throughput screening (Fig. 1B).

High-throughput screening of compound libraries

As indicated above, several different antibiotic compounds have been previously shown to inhibit ribozyme activity in vitro. As a first step toward the identification of compounds capable of inhibiting ribozyme cleavage in mammalian cells, we tested a variety of such compounds (Table 1), using both the HEK79 cell line and cells transiently transfected with different N79 variants. None of the compounds tested revealed an appreciable inhibitory effect on ribozyme self-cleavage, when administered at concentrations of either 10 or 100 μM (data not shown).

Based on those results, we extended the screening effort to include 58,076 compounds. The full set of compounds included an annotated bioactive compound library (2036 compounds) (Root et al. 2003); a diversity library obtained from NINDS (1040 compounds) (Lunn et al. 2004); a collection of FDA-approved drugs (1000 compounds) (Rogers 2002); a combinatorial library from Comgenex (20,000 compounds); a library containing both natural products and synthetic compounds (TIC; 24,000 compounds) (Kelley et al. 2004); and a combinatorial library called the “World Diversity Set,” obtained from Specs, containing compounds chosen for geographic as well as chemical diversity (10,000 compounds). For this high-throughput screening effort, the HEK79 cells were seeded in a 384-well format and exposed to the different compounds at a final concentration of 4 μg/mL for 48 h, and luc activity was then measured using a plate reader. Molecules were considered positive if they induced luc expression in cells carrying functional N79 ribozymes but not in cells carrying inactive N79 ribozymes. Through this primary screening effort and subsequent analysis of similar analogs of positive molecules, 15 ribozyme inhibitors were identified. The maximal fold increase in luc expression achieved by each compound (Foldmax) and the concentration that yields 50% of maximal induction (EC50) are summarized in Table 2. All of the identified molecules were shown to possess comparable activity in a separately generated N79-carrying cell line to that observed in the HEK79 cell line used for the original screening (data not shown). Interestingly, none of the inhibitors that were identified were aminoglycosides, despite the presence of many such molecules in the libraries.

Characterization of positive compounds

Based on structural similarities, the identified molecules can be divided into three groups. The first group consists of adenosine analogs, including tubercidin (7-deaza-adenosine), tubercidin-monophosphate and -triphosphate, tubercidin-cyclic monophosphate, sangivamycin, 8-azaadenosine, tricyclic nucleoside, and toyocamycin. All members of this group contain at least a modified 7-position on the purine (Fig. 2A). Analogs that have modifications other than at the 7-position, such as 2-amino, 3-deaza, 6-chloro, 9-deaza, lacked an inhibitory effect. In addition, the identified compounds all contain a 2’OH group on the sugar base, indicating that a ribose is necessary for their inhibitory effects (Fig. 2A). Similar analogs that lack 2’OH or are missing the ribose entirely produced no appreciable inhibition (2’-deoxy-tubercidin, 2’-deoxy-toyocamycin) (Fig. 2B). Of all the compounds, toyocamycin (Aszalos et al. 1966) was found to be the most potent ribozyme inhibitor, resulting in up to 365-fold luc induction when applied to the stable cell line. The effectiveness of the remaining compounds in this group was <60-fold (Table 2). Based on its apparent higher effectiveness, a dose-response curve of toyocamycin on the stable cell line was generated. As shown in Figure 2C, the induction peaked rapidly around 1 μM and then dropped off sharply. This drop coincided with apparent cell toxicity observed by microscopy. In addition, a decrease of luc expression from cells carrying inactive ribozymes, presumably due to the toxicity effects of the compound, was observed when the concentration of toyocamycin reached 0.5 μM. The dose-limiting toxicity of toyocamycin thus excludes its use at higher concentration or for long-term applications.

### TABLE 1. Survey of ability of different antibiotics to inhibit hammerhead ribozyme function

<table>
<thead>
<tr>
<th>Antibiotic</th>
<th>Activity</th>
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<tr>
<td>Chlorotetracycline</td>
<td>Kanamycin</td>
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<tr>
<td>Demeclocycline</td>
<td>Neomycin B</td>
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<tr>
<td>Diminazene aceturate</td>
<td>Neomycin sulfate</td>
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<tr>
<td>Distamycin A</td>
<td>Netropsin</td>
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<tr>
<td>Doxorubicin</td>
<td>Paramomycin</td>
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<tr>
<td>5-epi-Sisomicin</td>
<td>Sisomicin sulfate</td>
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<tr>
<td>Gentamicin sulfate</td>
<td>Tetracyclin</td>
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<tr>
<td>G418</td>
<td>Tobramycin</td>
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<tr>
<td>Gramicidin D</td>
<td>Tuberactinomycin A</td>
</tr>
<tr>
<td>Hoechst 33258</td>
<td>Tuberactinomycin B</td>
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<td>Hoechst 33342</td>
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The second group of ribozyme inhibitors identified in the screen consists of uridine analogs, including 5-bromouridine, 5-fluorouridine (FUR), and 5-fluorouracil (5FU) (Fig. 3A). All contain a 5-position modification on the pyrimidine. Analogs that contain an iodo, a methyl, a hydroxyl, an ethyl, or an aza group on the 5-position lack an inhibitory effect. In the case of this group of inhibitors, FUR was the most potent. However, replacing the 2′OH group of FUR by either 2′deoxy or 2′O-methyl, or by inverting the stereo-center of the 2′-position (Fig. 3B), resulted in loss of inhibition, indicating that, as with adenosine analogs, a ribose is necessary for the inhibitory effects of uridine analogs. In contrast to toycamycin, the dose-response curve of FUR reached a plateau of 110-fold at 50 μM, and the plateau was maintained for a wide range of concentrations (Fig. 3C), suggesting that an efficient induction can be reached with a lower dosage before reaching severe toxicity. The related molecule 5FU (Longley et al. 2003), however, has a dose-response curve resembling that of toycamycin, reaching a peak of 65-fold followed by a quick drop-off (Fig. 3D). The effectiveness of 5FU was also significantly less, requiring a concentration of 500 μM to reach 50% of the maximal induction, as compared to the 6 μM required for FUR (see EC50 in Table 2). For both FUR and 5FU, a decrease of luc expression from cells carrying inactive ribozymes was observed when concentrations reached 800 μM.

The third group of ribozyme inhibitors belongs to the family of RNA-binding dyes, and includes Syto-83, ethidium bromide (homidium bromide), and acridine orange (Fig. 4A). These dyes were also far less potent inhibitors than the adenosine or uridine analogs (Table 2).

As mentioned earlier, the N79 ribozyme used in our cell-based assay was developed from Sm1 of S. mansoni (Ferbye et al. 1998), a human parasite that infects people in many areas of the world (Ross et al. 2002). It is intriguing that one of the drugs suggested in the past as potential treatments for Schistosoma infection was tubercidin (Jaffe et al. 1971; el Kouni 1991), the same molecule identified in our screen as an Sm1 ribozyme inhibitor. This prompted us to test other molecules with known anti-Schistosomal effects for their inhibitory potential. These included nebularine, dipryridamole, benzylacyclouridine, metrifonate, oxamniquine, praziquantel, and artemether. However, only nebularine (also an adenosine analog) (el Kouni and Cha 1987) showed an inhibitory effect on ribozyme self-cleavage, albeit minimal (Table 2). Interestingly, unlike most of the adenosine analogs found in our screen, nebularine contains a modification on the 6- but not the 7-position of the purine base (Fig. 2A).

Lastly, it should be noted that while there were numerous guanosine and cytidine analogs in the 58,076 compounds screened, none of them surfaced as ribozyme inhibitors. In particular, 7-deazaguanine and 7-deazaguanosine triphosphate (the purine counterparts of tubercidin) and 5F-cytosine (the pyrimidine counterpart of 5FU) (Fig. 4B) all showed no inhibitory effects when tested at similar concentrations that produced inhibition by adenosine analogs or uridine analogs. The fact that only adenosine and uridine analogs but no guanosine and cytidine analogs were found positive may provide an insight into the N79 structure–function relationship (see below).

### Mechanisms responsible for ribozyme inhibition: RNA incorporation vs. RNA binding

To examine the mechanisms responsible for ribozyme inhibition, we incubated the N79 ribozyme with either tubercidin or toycamycin in an in vitro catalytic reaction. Surprisingly, neither tubercidin (Fig. 5A, lanes 3–8) nor toycamycin (Fig. 5B, lanes 3–8) was able to inhibit ribozyme self-cleavage in vitro, as indicated by the unaltered cleavage percentage over a broad range of compound concentrations. The lack of direct inhibition by these molecules indicated that their mechanism of inhibition was not likely due to direct ribozyme–compound interactions. In light of our findings that inhibitory activity appeared to always be associated with a 2′OH group on the sugar base, we next asked whether covalent incorporation of these analogs into the ribozyme-containing RNA might be required for the inhibition of RNA self-cleavage. To test if covalent incorporation of identified inhibitory molecules can, indeed, lead
to cleavage inhibition, tubercidin-triphosphate was used in place of ATP in an in vitro transcription reaction to generate ribozyme RNA carrying substitutions in all adenine positions. As shown in Figure 5C, this substitution resulted in the complete loss of ribozyme self-cleavage (Fig. 5C, lanes 2–6), while the unsubstituted ribozyme cleaved efficiently under the same conditions (Fig. 5C, lane 7).

Similarly, FUR (Fig. 5D) and 5FU (Fig. 5E) exhibited no inhibitory effect when incubated directly with N79 ribozyme at concentrations that would normally produce pronounced induction in the cell-based assays. Our attempts to examine the effect of incorporating FUR into the N79 ribozyme were inconclusive due to the difficulty of generating full-length FUR-substituted N79 ribozyme by T7 polymerase. However, past studies using chemical synthesis to generate FUR-containing hammerhead ribozymes demonstrated that when incorporated at certain key positions of the ribozyme, FUR, indeed, impaired self-cleavage (Hammann et al. 2001).

**FIGURE 2.** Chemical structures of identified adenosine analog ribozyme inhibitors and related compounds. (A) Analogs capable of inhibiting ribozyme self-cleavage in cells contain at least a modified 7-position on the purine, with the exception of nebularine, which contains a 6-position modification. These compounds all have a 2′OH group on the sugar base. The differences between these compounds and the inactive adenosine are indicated by the yellow shade. (B) Examples of analogs that lacked an inhibitory effect. (C) Dose-response curve of toyocamycin in log scale. Error bars indicate 1 S.D. (standard deviation).

**FIGURE 3.** Chemical structures of identified uridine analog ribozyme inhibitors and related compounds. (A) Analogs capable of inhibiting ribozyme self-cleavage in cells contain a modified 5′-position on the pyrimidine. These compounds all have a 2′OH group on the sugar base. The differences between these compounds and the inactive uridine are indicated by the yellow shade. (B) Examples of analogs that lacked the inhibitory effect. (C) Dose-response curve of FUR in log scale. (D) Dose-response curve of 5FU in log scale. Error bars indicate 1 S.D.
Consistent with the mechanism of covalent incorporation of the compounds into the ribozyme-encoding cellular mRNA, it has been reported that tubercidin (Acs et al. 1964), toyocamycin (Suhadolnik et al. 1967), 5-FURidine (Glazer and Lloyd 1982), and 5-FU (Glazer and Lloyd 1982) are able to enter the ribonucleotide pool for RNA synthesis and can be found in the cellular RNAs of mammalian cells. This mechanism of inhibition of ribozyme function, dependent on inhibitor incorporation into RNA, may shed light on the basis for our identification of only adenosine analogs with a modified 7-position and uridine analogs with a modified 5-position as effective inhibitors and our inability to identify any analogous guanosine and cytidine analogs. While these findings might simply relate to the variable cell permeability of different molecules, or their ability to be recognized by RNA polymerases for incorporation, they may suggest that structural interactions involving the 7- and 5-positions of A and U bases, respectively, are particularly important for N79 ribozyme function in vivo. Although the 7-position of A does not form hydrogen bonds in the standard Watson-Crick base pairs, the X-ray structures of hammerhead ribozymes, indeed, showed its involvement in forming the nonstandard A-G base pairs in the conserved core (A9-G12, A13-G8; see Fig. 2 for the nucleotide numbering) (Fu and McLaughlin 1992; Pley et al. 1994; Scott et al. 1995). The role of the 5-position of U is less clear since it is not involved in hydrogen bonds of either the standard Watson-Crick or the G-U wobble base pairs. Nevertheless, it was shown that incorporating FUR at the U4 and U7 of the hammerhead core led to the impairment of self-cleavage (Hammann et al. 2001). Thus, modifying those A and U bases in the conserved core could lead to the graded loss of cleavage activity in vivo. Furthermore, the N79 ribozyme contains an unusually high percentage of A and U bases in its loops I and II. Substituting these bases could disrupt the critical interaction of the two loops required for the ribozyme to function in cells (De la Pena et al. 2003; Khvorova et al. 2003; Penedo et al. 2004; Yen et al. 2004).

Synergistic inhibition of ribozyme function by combined administration of toyocamycin and FUR

As toyocamycin (an adenosine analog) and FUR (a uridine analog) would occupy different positions if they were incorporated into the N79 ribozyme, we next asked whether combined administration of the two compounds might lead to improved inhibition of ribozyme function relative to addition of a single compound and perhaps define a less toxic regimen for achieving reasonable levels of induction of gene expression. To address this question, varying amounts of each compound were combined and added to HEK79 cells, and subsequently, the fold induction of luc expression was determined. As shown in Figure 6, administration of specific concentrations of the two compounds did, indeed, result in a more efficient inhibition of the ribozyme than was possible with either compound alone (e.g., 670-fold induction by a combination of the two compounds vs. 120-fold induction by FUR alone or 378-fold induction by toyocamycin alone). In addition, a variety of mixtures of the two compounds were able to induce gene expression between 200- and 600-fold (Fig. 6), in contrast to the narrow range of concentrations of toyocamycin for which induction can be achieved in the absence of significant toxicity (Fig. 2C). To document the practical use of this two-compound cocktail for the regulation of gene expression in vivo, we established an in vivo model in which an AAV vector encoding a luc reporter controlled by two copies of the N79 ribozyme (similar transcription cassette to that diagrammed in Fig. 1) was injected subcutaneously to the left ear of nude mice. To provide an internal control, all mice were also injected in the right ear with virus carrying the inactive N79. After 3 wk, these mice were imaged for basal luc gene expression (day 0), and immediately after imaging, a single pulse of cocktail was of mice, taken before and 2–3 d after cocktail injection, are shown in Figure 7. The images demonstrate that, as expected, the right ear carrying the inactive N79 showed little change in gene expression before and after cocktail treatment. In contrast, in the left ear carrying the functional N79, a pronounced induction was readily detectable after the treatment. Quantification of the photon output indicated that the induced gene expression reached 37-, 17-, and 13-fold in the three mice treated. The induction afforded by this single pulse injection method was somewhat variable from mouse to mouse, most likely due to the variability of cocktail volume delivered.
FIGURE 5. In vitro self-cleavage of N79 ribozyme in the presence of inhibitors and in vitro self-cleavage of derivatives of the N79 ribozyme containing analog substitutions. (A) Cleavage of N79 ribozyme in the presence of tubercidin. (Lane 1) RNA marker of 64 bases, (lane 2) without Mg$^{2+}$, (lanes 3–8) in the presence of increasing concentrations of tubercidin, (lane 9) in the presence of adenosine. The cleavage percentage remained unchanged from lanes 3 to 9. (B) Cleavage of N79 ribozyme in the presence of toyocamycin. Notations are as described in A. (C) Cleavage of N79 ribozyme in which all the adenosine positions were substituted by tubercidin. (Lane 1) RNA marker, (lanes 2–6) cleavage of substituted N79 ribozyme at various reaction times, (lane 7) cleavage of the unsubstituted N79 ribozyme. (D) Cleavage in the presence of FUR. Notations are as described in A. (E) Cleavage in the presence of 5FU. (Lane 1) Without 5FU, (lane 2) without 5FU and Mg$^{2+}$, (lanes 3–7) in the presence of increasing concentrations of 5FU, (lane 8) in the presence of uracil as the control. Conditions for all self-cleavage reactions: 50 mM Tris-HCl, 10 mM Mg$^{2+}$ (pH 7.5) for 20 min at 22°C, with the exception of E, which was carried out with 1 mM Mg$^{2+}$. 
DISCUSSION

The work described here represents the first cell-based screen aimed at identifying small molecules capable of modulating ribozyme self-cleavage in vivo. By screening 58,076 compounds, we were able to identify two small molecules, toyocamycin and FUR, that can potently inhibit ribozyme function in mammalian cells. It is of significant interest that the two most potent inhibitors of ribozyme function in cells that we identified appear to inhibit ribozyme self-cleavage via their covalent incorporation into the mRNA carrying the ribozyme sequences, as evidenced by the inability of these molecules to inhibit ribozyme self-cleavage in vitro and by the loss of self-cleavage activity when these molecules are incorporated. Based on the well-described interactions between specific aminoglycosides and other small molecules and RNA (Stage et al. 1995; Murray and Arnold 1996; Hermann and Westhof 1998; Tor et al. 1998; Jenne et al. 2001), we had expected that our extensive screen of compounds would have led to the identification of compounds that inhibit ribozyme self-cleavage by virtue of their direct binding to ribozyme-containing RNA. Our inability to identify such compounds may suggest that the complex structure of mRNA molecules and their associated proteins present in mammalian cells greatly restricts the possibilities for the types of interactions between small molecules and RNAs that can lead to the inhibition of RNA self-cleavage. The ability of morpholino oligonucleotides and several RNA-binding dyes to inhibit ribozyme activity in cells, however, suggests that appropriate interactions between specific small molecules and ribozymes that lead to inhibition of self-cleavage in vivo are indeed possible. We are hopeful that additional screening efforts will lead to the identification of new inhibitors of ribozyme function with improved toxicity profiles and more desirable pharmacokinetic properties.

We had already made use of one of the molecules identified in the current study, toyocamycin, to provide a critical “proof of principle” for the ability to regulate gene expression in mammalian cells both in vitro and in vivo via the modulation of RNA self-cleavage (Yen et al. 2004). As expected by the proposed mode of action of this inhibitor, we showed in that study that in the absence of toyocamycin, the ribozyme-controlled mRNA was not detectable in the nucleus or cytoplasm in cells carrying active ribozymes, consistent with the mechanism that ribozyme self-cleavage leads to rapid degradation of mRNAs. However, when these cells were treated with toyocamycin, the amount of luc mRNA increased to a level comparable to that of cells carrying inactive ribozyme, which is consistent with the mechanism that inhibition of ribozyme self-cleavage leads to the generation of intact mRNA in cells (Yen et al. 2004). In the present studies, we show that the combined administration of toyocamycin and FUR to cells harboring a ribozyme-regulatable expression construct leads to an improved induction of gene expression, relative to the use of toyocamycin alone. This combined regimen requires lower concentrations of the two compounds than must be used to achieve reasonable levels of inductions using either compound alone, and consequently results in a less toxic effect on the cells. While it is hopeful that the use of such a two-compound cocktail may enable more experimental applications of the ribozyme-based gene regulation system than are possible with toyocamycin alone, the identification
of significantly less toxic inhibitors of ribozyme function remains an important goal of future research.

In addition to the use of the ribozyme-based gene regulation system for the control of expression of gene products, the finding that one of the most commonly used chemotherapeutic agents, 5FU, can serve as a ribozyme inhibitor in vivo raises the interesting possibility that one novel application of RNA-only gene regulation systems might relate to the real-time “sensing” of the presence of specific compounds in vivo. In the specific case of 5FU, the ability of the molecule to induce expression of a reporter gene could provide an important means of monitoring 5FU uptake and retention by tumors and normal tissues, and/or facilitate efforts to develop more optimal routes of 5FU administration.

MATERIALS AND METHODS

High-throughput screening of compound libraries

ACL, NINDS, TIC, and Comgenex libraries were collected by Dr. Brent Stockwell, prepared as 4 mg/mL solutions in DMSO, and screened at a final concentration of 4 μg/mL. FDA2000 was collected by Dr. Steve Gullans (BWH), and each compound was prepared at 10 mM in DMSO and screened at 10 μM. The Specs library was prepared as 10 mg/mL and screened at 10 μg/mL.

HEK79 cells were seeded in white, opaque-bottom 384-well plates at a density of 8000 cells per well in medium with 10% serum and pen/strep using the Zymark dispenser. On the same day, cells were treated with compounds at a final concentration of 4 μg/mL and 0.1% DMSO, followed by an incubation period of 48 h at 37°C in a humidified incubator containing 5% CO₂. Cells were then lysed, and luciferin was added to the cells at a final concentration of 4 μg/mL; the amount of light output from each well was recorded on a Packer Fusion platerader. Induction of luc expression by a particular compound was calculated by subtracting the signal with instrument background and background from untreated cells. All screens were performed in triplicate repeats.

Cell culture assays for testing the effective concentration of compounds

Plasmid DNAs carrying a LacZ reporter under the control of two copies of the N79 ribozyme were transfected into subconfluent HEK293T cells using Fugene6 (Roche). The next day, cells were treated with compounds at various concentrations for 24 h. The level of β-galactosidase expression was then quantified by ONPG assay 24 h later. All experiments were performed at least in triplicate. Alternatively, HEK79 cells (stable cell lines containing a luc reporter under the control of two copies of N79 ribozymes; see Results) were seeded in white, opaque-bottom 96-well plates with a density of 40,000 cells per well, and treated with compounds the next day at various concentrations for 24 h. Luciferin was then added to the cells at a final concentration of 20 μg/mL, and the amount of light output from each well was recorded on a plate reader. All experiments were performed at least in triplicate. 5FU (from Sigma) was prepared as 100 mM stock in 100% DMSO; FUR (Sigma), as 10 mM stock in PBS; toyocamycin (Berry & Associates) and tubercidin (Sigma), as 10 mM stock in 10% DMSO.

In vitro transcription and catalytic reaction

DNA templates containing a T7 promoter and the ribozyme sequences were generated by PCR. 32P-labeled RNA was then transcribed by T7 RNA polymerase in the presence of 50 μM blocking anti-sense oligonucleotides using Ambion’s MEGashortscript kit. The anti-sense oligonucleotide sequence used was 5’-GATGCCAGTGAATCCAGGACGCAC-3’. In the absence of the anti-sense oligonucleotide, ~95% of the transcripts were cleaved during the 3-h in vitro transcription reactions. In the presence of the anti-sense oligonucleotide, the percentage of cleavage products dropped to 35%. To make ribozymes with incorporated tuberculosis, transcription was carried out by substituting ATP with tuberculosis triphosphate. Full-length RNAs were then purified by 10% denaturing polyacrylamide gel. To determine the cleavage efficiency, ribozyme was incubated in 50 mM Tris-HCl (pH 7.5) for 3 min at 95°C and slowly cooled to 23°C in 15 min. Self-cleavage reactions in the presence or the absence of inhibitory compounds were triggered by adding MgCl₂ to 1 mM or 10 mM at 23°C. The reaction was stopped by adding 5 volumes of stop solution (8 M urea, 50% formamide, 40 mM EDTA), and the cleaved fragments were separated by a 10% denaturing polyacrylamide gel. Cleavage percentage was then quantified by PhosphorImager. We did not detect any cleavage products in transcription reactions when tuberculosis triphosphate was used to replace ATP.

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