Volatility in mRNA secondary structure as a design principle for antisense

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ABSTRACT

Designing effective antisense sequences is a formidable problem. A method for predicting efficacious antisense holds the potential to provide fundamental insight into this biophysical process. More practically, such an understanding increases the chance of successful antisense design as well as saving considerable time, money and labor. The secondary structure of an mRNA molecule is believed to be in a constant state of flux, sampling several different suboptimal states. We hypothesized that particularly volatile regions might provide better accessibility for antisense targeting. A computational framework, GenAVERT was developed to evaluate this hypothesis. GenAVERT used UNAFold and RNAforester to generate and compare the predicted suboptimal structures of mRNA sequences. Subsequent analysis revealed regions that were particularly volatile in terms of intramolecular hydrogen bonding, and thus potentially superior antisense targets due to their high accessibility. Several mRNA sequences with known natural antisense target sites as well as artificial antisense target sites were evaluated. Upon comparison, antisense sequences predicted based upon the volatility hypothesis closely matched those of the naturally occurring antisense, as well as those artificial target sites that provided efficient down-regulation. These results suggest that this strategy may provide a powerful new approach to antisense design.

INTRODUCTION

The ability to manipulate gene expression is one of the most fundamental aspects of biotechnology. It has been accomplished through a variety of methods, including through the use of antisense nucleic acids (DNA and RNA). Since antisense is complementary to a target mRNA, the two strands may hybridize through hydrogen bonding. This double-stranded duplex may hinder ribosomal binding, block ribosomal migration or induce cleavage by an RNase (1,2). In this way, antisense has the potential to be used for numerous applications ranging from metabolic engineering to human gene therapy. Many antisense drugs are in clinical trial for the treatment of a wide variety of diseases, including cancer (3,4).

The process of selecting an antisense sequence that is able to effectively bind to a target mRNA and block protein synthesis is complex and governed by many factors. One of the most important factors is the secondary structure of the target mRNA, which is determined by intramolecular hydrogen bonding that helps to establish a more thermodynamically stable conformation (5). The accepted theory is that this secondary structure would be problematic for antisense-based down-regulation due to the majority of the target mRNA being paired to itself. This intramolecular bonding does not prevent translation because of the ribosome’s ability to unwind mRNA (6), but it greatly decreases accessibility for antisense binding.

There have been many attempts to try and accurately predict the efficacy of antisense sequences to save time, money and labor, all of which are wasted with brute force design and test methods of antisense synthesis. Some approaches involve searching an mRNA sequence for consensus sequences that are present in effective natural and artificial antisense and base their predictions on those motifs (7). Other methods offer the prediction of RNA–RNA interaction mechanisms and may suggest where the target would be in a given mRNA for a specified antisense or small-interfering RNA (siRNA) sequence (8,9). Still other strategies that focus on eukaryotic systems utilize large databases of known species-specific siRNA sequences and predict sequences based on that data. Finally, some methods focus mainly on predicting accessible sites on a target RNA (10,11) or fusing

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accessibility prediction with hybridization prediction (12–14). There is still much to learn about antisense prediction and the need for more effective strategies remains.

In recent years, the idea that an mRNA strand may not always take the form of a distinct fixed molecular structure has become much more prominent. It is believed that an mRNA molecule may actually be in a state of constant structural fluctuation, transitioning between different conformations near the minimum free energy (MFE) structure, particularly in an ever-changing cellular environment (15–17). Analyzing suboptimal mRNA structures with a thermodynamic stability comparable with that of the MFE structure may reveal that certain regions are more ‘volatile’ than others. Since these regions have the ability to change conformation without significantly altering the Gibbs free energy of the entire molecule, they may have more freedom to alter their hydrogen bonding. Therefore, these regions would likely be the most accessible targets for antisense binding because of their constant formation and breaking of intramolecular hydrogen bonds.

A computational framework, GenAVERT (http://www.rs labs.org), was developed to take advantage of this concept of structural fluctuation to predict the sites on a given strand of mRNA that are most likely to vary in structure within a defined range of free energy. These sites were hypothesized to be superior antisense targets. To test this idea, different types of antisense systems were examined. First, several naturally occurring antisense sequences from prokaryotes were analyzed by using GenAVERT. The analysis predicted that the most volatile regions of those mRNAs were essentially the same as those of the natural antisense target sites. Next, genes for which man-made antisense had been designed for down-regulation purposes were analyzed. The results of the best antisense compared favorably with GenAVERT predictions, indicating that those antisense exhibiting high levels of down-regulation targeted regions of relatively high volatility.

**MATERIALS AND METHODS**

**Natural and artificial antisense prediction**

The ability to predict natural antisense transcripts is a major part of developing a strategy for designing artificial antisense. As noted earlier, many strategies already utilize natural antisense as an indicator of a prediction system’s accuracy (8). Since these antisense expression systems have presumably evolved over millennia, they are thought to result in the most effective inhibitory duplexes possible for a given mRNA. Many of these antisense sequences have been experimentally tested to positively validate their efficiency in the down-regulation or silencing of their corresponding mRNAs.

Perhaps some of the most important and well-studied antisense expression systems are those of toxin–antitoxin systems (18). These systems generally consist of an mRNA that encodes for a ‘suicide protein’ that is extremely toxic to the host cell, as well as a *cis*-encoded antisense RNA that is transcribed from the same locus but in the opposite direction (as opposed to *trans*-encoded, where the antisense is encoded at a separate locus). The significance of these toxin–antitoxin systems is in the necessity for efficient inhibition of protein expression to avoid cell death. Translation levels must be brought to an extremely low level or blocked completely for a cell to continue to carry these suicide genes; therefore, their corresponding antisense inhibitors must be exceedingly effective. A successful antisense prediction system should be able to predict sequences similar to these RNA antitoxins after analyzing their correlating toxin-encoding mRNAs. The volatility hypothesis was evaluated using the following bacterial toxin–antitoxin systems (19): hok/sok (20), *pndA/pndB* (21), *hokC/sokC* (22), *gef/sof* (23), *hokA/sokA* (22) and *ldeA/rdLA* (24).

Apart from natural antisense prediction, the prediction of artificial antisense is perhaps just as enlightening. Examining artificial antisense systems that may exhibit a range of gene down-regulation levels provides another platform for measuring the efficacy of identifying volatile regions in mRNA secondary structure. These types of systems represent gene regulation beyond the scope of toxin–antitoxin systems and even those present in various types of bacteria. Three different systems were investigated, with mRNAs of varying lengths, one of which is polycistronic and upwards of 2000bp. The antisense from these systems demonstrated varying levels of down-regulation and their respective target sites were thus examined for volatility in an attempt to explain the variation in experimental efficiency.

**GenAVERT**

GenAVERT was developed to test the idea that structurally volatile regions of mRNA made more effective antisense targets. GenAVERT accomplished this objective by generating and comparing suboptimal secondary structures of a given mRNA sequence. Analysis of these comparisons revealed regions that were least ‘similar’ among the set of folds, indicating volatility in intramolecular hydrogen bonding and, according to the proposed hypothesis, accessibility for antisense binding. The program was written in Common Lisp (LispWorks, Cambridge, UK) and calls upon two external programs, UNAFold (http://mfold.rna.albany.edu/) and RNAforester (http://bibuserv.techfak.uni-bielefeld.de/rnaforester/), and a Perl script from the Vienna RNA Package.

GenAVERT functions simply by reading in an mRNA sequence with a given name and then generates a set of potential antisense sequences. Once GenAVERT receives its input, UNAFold is invoked. UNAFold uses the concept of nearest-neighbor thermodynamics to estimate how the bases of an RNA sequence will interact with each other to increase structural stability (25). It outputs the MFE secondary structure for the given mRNA sequence, as well as a set of suboptimal structures with slightly higher Gibbs free energies (26). Since the output from UNAFold is a set of ‘.ct’ files, each of which describes one structure, an external Perl script (Vienna RNA Package) (27) is called to convert each of these .ct files into Vienna bracket format, where periods represent
unpaired bases and open and close parentheses represent base pairs. For example, ‘…..((. . .))…..(( . .))…..’

Since it is believed, as previously mentioned, that an mRNA molecule is constantly fluctuating in its structure, it is assumed that the range of possible in vivo mRNA structures most likely consists of those that make up the suboptimal range of free energy. If these suboptimal folds are then compared based on structure, they should theoretically point out regions of the mRNA that change more often than others, yet still allow the entire structure’s free energy to remain relatively close to the MFE value. This in turn would indicate that these regions possess superior accessibility and act as likely antisense target sites. To accomplish this goal, the Vienna bracket structures are analysed with the external program, RNAforester.

RNAforester is a comparison program designed for phylogenetic analysis of different RNA molecules (28, 29). It takes both sequence and structure into account and compares only two structures at a time. As a result, the basic pairwise input would be two sequences and their corresponding Vienna bracket structures. It then proceeds to generate a homology file showing where the two sequences and structures are similar, using the common method of filling in regions that do not show homology in sequence and structure with ‘gaps.’ In the case of GenAVERT, RNAforester is called to compare the consecutive suboptimal folds that were generated from UNAFold. For example, assume that UNAFold outputs the MFE structure for a specific mRNA sequence as well as three suboptimal folds (1, 2 and 3), for a total of four structural folds. RNAforester would first be called to compare the MFE fold and suboptimal fold 1, then to compare suboptimal fold 1 with suboptimal fold 2, 2 with 3 etc. In this way, there is an artificial sense of transitioning from one structure to another because each file indicates which bases have changed their hydrogen bonding pattern between the two folds (note that all sequences used are that of the given mRNA sequence, while only the structures themselves are changing). Each of the structures is weighted equally. An example of this artificial transitioning is depicted in Figure 1. A more aesthetic method of viewing this transitioning is by simply using the program RNAmovies (15) where the only input required is the mRNA sequence and the set of structures in Vienna bracket format. Whenever GenAVERT is invoked, an RNAmovies input file is created for the user.

GenAVERT then searches all of these RNAforester homology files for any length of mRNA that has changed its conformation (altered hydrogen bonding) between two structures and includes them in a pool of possible volatile regions. It then searches through this pool of possible regions and enumerates the number of times that a certain base shows up. This count indicates the number of times that this base has changed position over the ‘structural transition’ from the MFE fold to the final suboptimal (and least energetically favorable) fold. The number of suboptimal structures is determined by the default settings of the UNAFold window parameter with all kept within 5% of the Gibbs free energy of the MFE structure (26). GenAVERT then uses these values to create a list of every base that meets or exceeds a certain level of volatility. Since this list is not always made of bases that are consecutively located in the mRNA sequence, it may be split into multiple individual volatile regions. The longest of these volatile regions is chosen as the most volatile and therefore the most accessible region on the mRNA strand. The reverse complement of this region is then generated as the antisense sequence that is most likely to down-regulate the expression of the target gene.

However, this first antisense sequence may not always be a practical choice and it may be necessary to continue to collect a set of potential antisense sequences. For example, consider a hypothetical set of five consecutive bases that alter in hydrogen bonding 20 times over the entire structural transition, with no other bases coming close to that level of volatility. As a result, an antisense sequence complementary to those five bases would be predicted as the optimal. However, such a short antisense is unlikely to be viable. As a result, the process is continued for bases that alter their bonding at the next highest level, for example, 15 times, 14 times, 12 times, etc., essentially providing a set of high ranking antisense possibilities of varying length. Therefore, GenAVERT continues to generate the optimal antisense sequence for each level of volatility until no significant difference between bases can be detected. A flowchart of the processes that make up GenAVERT is shown in Figure 2.

Some of the predicted results were considered too small to be viable. We therefore implemented a heuristic criterion to only select sequences that were at least 35 bp in length. Thus, although the program will generate antisense sequences less than 35 bp (see Supplementary Dataset 1), discussion is restricted to those sequence of 35 bp or greater as predicted by GenAVERT.

Comparison with Sfold
As a benchmark, the results from GenAVERT were then compared with that of a currently available program that has a similar goal of predicting inhibitory RNA sequences for prokaryotes. The program used for comparison was the Soligo partition of Sfold (http://sfold.wadsworth.org/cgi-bin/index.pl) (12–14). Sfold is designed to predict antisense sequences with the caveat that the user must declare a pre-determined length for the antisense sequence. Therefore, different values for the antisense length were used with Sfold to present a wider range of the program’s capabilities. Lengths of 35 bp, 50 bp and the exact target

![Figure 1. Transition between two possible structures of hok mRNA as predicted by UNAFold and displayed using the interpolating effects of RNAmovies. The large transitioning region (nucleotides 70-150) consists almost entirely of the sok target region (nucleotides 65-131).](image-url)
length were used. The authors have no knowledge of any other de novo antisense prediction software available at this time and none that requires as little input from the user as that of GenAVERT.

RESULTS

Natural antisense prediction

**hok/sok**
The hok (host killing) gene is located on plasmid R1 of *Escherichia coli* and encodes for a protein of 52 amino acids that is toxic to the host cell. A naturally occurring antisense transcript found at the same locus but encoded in the opposite direction, is denoted sok (suppression of killing). The overall function of the hok/sok expression system is for plasmid stabilization. Cells that lose the plasmid carrying the locus are killed due to translation of stable hok mRNA that remains behind in the cell, while the more quickly degraded sok RNA is unable to further inhibit protein synthesis (18,20,22).

When GenAVERT was used to evaluate hok mRNA, the highest ranking antisense sequence greater than 35 bp was the 70 bp antisense sequence shown in Figure 3. This sequence overlapped with the sok target region (sokT) by almost 93% and only had an overhang of 8 bases (Figure 3). All three of the other lengths of antisense predicted by Sfold were designed to target sites near the 3'-end of the mRNA, far from the naturally occurring target site. Table 1 summarizes these results, as well as all of the remaining results.

**pndA/pndB**
The pndA gene is located on plasmid R483 of *E. coli*. It is, in fact, a hok-homologue and has many of the same functional characteristics of the hok family genes, including a tac (translational activation) sequence, mok (method of killing) reading frame, as well as having a cis-encoded antisense denoted pndB. It also, unsurprisingly, encodes for a toxic protein (21).

Figure 4 displays the various antisense predictions, including antisense generated from GenAVERT that overlapped with ~87% of the pndB target region (pndBT) with an overhang of 11 bases. The Sfold antisense with the exact target site length, as well as the 50 bp length both overlapped with pndBT by only about 14.3%. The 35 bp antisense from Sfold was designed to target the complete opposite end of the mRNA, clearly illustrating the variability in Sfold predictions, even when antisense length is changed only by 15 bp.
The ldrA (long direct repeat A) gene is found on the E. coli K-12 genome (as opposed to being located on a plasmid like hok or pndA) along with its own cis-encoded antisense transcript known as rdlA (regulator detected in ldrA). ldrA is analogous (not homologous) to hok and is part of a different gene family. However, it also encodes for a toxic protein lethal to the host cell, consisting of only 35 amino acids.

rdlA was shown to effectively inhibit the translation of ldrA by Kawano et al. (24). Actual sequences were taken from the EcoCyc Database (http://www.ecocyc.org) (30) using the putative transcription start and end sites of the homologous and well-characterized ldrD gene.

GenAVERT predicted an antisense sequence of 95 bp, which overlaps with 92% of the 67 bp rdlA target region (rdlAT). However, it also has an overhang that is not complementary to rdlAT of about 34 bases. Despite this, when GenAVERT’s sequence is compared with those produced by Sfold, only one of the predicted Sfold sequences shared ~45% overlap with rdlAT, as shown in Supplementary Figure S1. Again, the 50 and 67 bp Sfold antisense strands target the 3' end, while the 35 bp strand targets the 5' end, showing the same variability as before.

hokC/sokC

The hokC gene (also homologous to hok) was found to be carried by E. coli ECOR24, and its transcript is believed to be ~330 nucleotides long. Its cis-encoded antisense was named sokC (22). After using GenAVERT to predict a potential antisense inhibitor for the hokC mRNA sequence, it was able to predict an antisense sequence complementary to 82% of the sokC target (sokT) sequence with an overhang of 28 bases. Sfold predictions were much more accurate in this case than in other examples with percent overlaps of 47%, 60% and 76% to sokT for 35 bp, 50 bp, and exact target lengths respectively (Supplementary Figure S2).

gef/sof

The gef (gene expression fatal) gene encodes for a 50 amino acid cell-killing protein and is found on the E. coli K-12 genome. It is essentially the same as hokC, with a cis-encoded antisense almost exactly that of sokC, denoted sof (suppression of fatality). Unlike hokC, however, there is an IS186 insertion sequence located downstream of the coding region, a sequence which is 1338 bp long (31) and is thought to disrupt the usual hokC fbi (fold-back inhibition) sequence and usual mRNA processing. However, despite the presence of this insertion, it has been shown that it is still active on a transcriptional level and is being regulated by sof RNA. The length of the gef mRNA in this case is thought to be about 644 nucleotides long and terminates within the...
insertion sequence, 200 bp downstream of the stop codon (22,23,32).

GenAVERT predicted an antisense sequence that was complementary to 77% of the sof target region (sof/T). However, it also targeted a portion of the mRNA outside of the sof/T region, as can be seen in Supplementary Figure S3. Comparing this data with the Sfold predictions revealed only one sequence that was predicted to overlap with 53% of the natural target. The other top scoring antisense sequences for 50 bp and for the exact target length did not overlap the natural target in any way. Again, the variability with small changes in sequence length is exhibited in this case.

\textit{hokA/sokA}  

Like \textit{pndA} and \textit{hokC}, \textit{hokA} is also a \textit{hok} homologue and displays the same characteristics but was found in \textit{E. coli} C instead of \textit{E. coli} K-12 or \textit{E. coli} ECOR24. Its cis-encoded transcript, \textit{sokA} regulates the expression of the toxic HokA protein in the usual manner (22).

It can be seen from Supplementary Figure S4 that all of the predicted antisense sequences included at least some complementary bases with the \textit{hokA} target region (\textit{hok/A}T). GenAVERT’s sequence contained a sequence that is complementary to all of \textit{hok/A}T (100%) with a short overhang of 13 bases. Sfold predicted a 35 bp region that exhibited 40% overlap with the target, whereas the antisense with the exact target length exhibited 75% overlap. However, even with the exact target length (52 bp) antisense showing such a high percent overlap, changing the antisense length by only 2 bp to 50 bp, radically shifted the sequence to display only 33% overlap.

As mentioned previously, Table 1 provides a summary of the results for all of the natural antisense prediction.

Antisense design applications

\textbf{Regulation of phosphotransbutyrylase and butyrate kinase in \textit{C. acetobutylicum}}

The \textit{adc} gene of \textit{C. acetobutylicum} ATCC 824 encodes for acetoacetate decarboxylase (AADC) and is a major component of the acetone formation pathway. The \textit{adc} mRNA is 859 bp long (33,34). Tummala \textit{et al.} (35) designed plasmids expressing three different antisense strands of varying lengths in an effort to try and alter acetone formation through the down-regulation of AADC. The first antisense RNA shared complementary base pairs with the first 38% of the \textit{adc} mRNA, the second shared complementary base pairs with the first 68% of the \textit{adc} mRNA, while the third shared complementary base pairs with 100% of the \textit{adc} mRNA. The percent down-regulation of AADC was reported to be greater than 80% for all three strains containing the three different antisense plasmids in both transitional and stationary growth phases. The level of AADC expression was found to be too low for quantification of down-regulation in the late exponential phase.

Figure 5 depicts the volatility profile for the \textit{adc} mRNA based on nucleotide position. The target for the first antisense sequence (38%) is from nucleotides 1 to 328, whereas the target for the second (68%) is from nucleotides 1 to 560. GenAVERT first predicts that the optimal antisense target region is from nucleotides 261 to 322. This 62 bp region is targeted by all three of the expressed antisense RNAs. The second best antisense target as predicted by GenAVERT targets nucleotides 252–488, again, a region with parts encompassed by all three. The next three subsequently scoring target sequences were predicted to be from nucleotides 252–504, 244–504 and 237–504, staying generally in the same volatile region on the mRNA (not depicted). The targeting of all three of the expressed antisense RNAs for this highly volatile region (>80% volatility within a 5% free energy range of the MFE structure) may be an explanation for their ability to induce efficient downregulation.

\textbf{Regulation of phosphotransbutyrylase and butyrate kinase in \textit{C. acetobutylicum}}

The \textit{ptb} and \textit{buk} genes of \textit{C. acetobutylicum} ATCC 824 encode for phosphotransbutyrylase (PTB) and butyrate kinase (BK), respectively. They are transcribed from the same operon and therefore the two genes are encoded for on one polycistronic mRNA that is 2128 bp long (36). PTB and BK are both essential parts of the butyrate production pathway and they were targeted in an effort to alter the primary metabolism, specifically, the solventogenesis pathways. Desai and Papoutsakis (37) designed two different plasmids, each expressing antisense RNAs that targeted either the \textit{ptb} region of the mRNA or the \textit{buk} region of the mRNA. Strains expressing the \textit{ptb} antisense resulted in about a 70% decrease in the peak level of PTB compared with the control, with peak levels of BK being ~80% less than that of the control. Strains expressing the \textit{buk} antisense resulted in about an 85% decrease in the peak level of BK compared with the control strain, while also showing about a 45% decrease in the peak level of PTB.

Figure 6 illustrates the volatility profile for the \textit{ptb-buk} mRNA, as well the first eight predicted volatile regions from GenAVERT. The most volatile region is 10 bp and falls within the \textit{ptb} target region (nucleotides 25–577) but is probably not viable. The region ranked as second is 65 bp and also falls within the \textit{ptb} target site. The third top scoring volatile region of 267 bp overlapped with the \textit{buk} target region (nucleotides 973–1018). The following three top scoring regions were all generally from nucleotides 1503–2059, indicating there was perhaps down-regulation potential by targeting sites nearer the 3'-end. Amazingly, volatile region number eight overlapped with almost the entire \textit{ptb} target region, encompassing nucleotides 5–568. It therefore seems likely that the two target regions on this polycistronic mRNA coincided with two highly volatile areas in the secondary structure, allowing for accessible binding and subsequent down-regulation of these two enzymes. When GenAVERT was restricted to searching for potential target sites in only the first 75% of the mRNA sequence, every single predicted site overlapped with the actual target sites (Supplementary Figure S5).

\textbf{Regulation of the \textit{\sigma}^{32} transcription factor in \textit{E. coli}}

The \textit{rpoH} gene of \textit{E. coli} encodes for the \textit{\sigma}^{32} transcription factor that is required for the transcription of specific
genes. Upregulation of $\sigma^{32}$ can be induced under multiple circumstances including ethanol shock, heat shock or the overexpression of recombinant proteins. Srivastava et al. (38) designed a plasmid to express an antisense RNA that would target rpoH mRNA under heat shock, ethanol shock and expression of organophosphorus hydrolase (OPH). However, it has been reported that under various conditions and with multiple strains there may be up to six possible rpoH mRNAs present in vivo due to different promoters under various regulation as noted in the EcoCyc Database (30). It is believed that the rpoHp1 (promoter 1 transcript) mRNA is present under most physiological conditions and is the primary transcript (39).

It was reported by Srivastava et al. (38) that under ethanol shock, control cultures showed a 10-fold increase in $\sigma^{32}$ expression, while cultures with induced antisense expression showed only an initial 3-fold increase, which then fell to a 2-fold increase. Likewise, the $\sigma^{32}$-regulated GroEL chaperone protein showed a significant decrease in expression during the first hour after ethanol shock when antisense was expressed. However, the GroEL expression levels were comparable in both antisense-expressing and control cultures beyond the 2 hour time point. Under heat shock conditions, GroEL levels dropped 30% in antisense-expressing cultures after the first 5 minutes. Finally, under conditions of OPH expression, OPH levels should theoretically have been higher
with antisense expression. However, it was instead observed that OPH was actually higher in the control cultures without antisense expression.

The volatility profile for \textit{rpoHp1} is shown in Figure 7 along with the first five predicted volatile regions from GenAVERT. It is clear from this profile, that the most conserved secondary structure of \textit{rpoHp1} is encompassed by the designated target site (nucleotides 90–369). GenAVERT ranks the top scoring volatile region as nucleotides 380–439, followed by the second ranked region of nucleotides 317–514. Regions ranked as fourth and fifth also show some overlap with the target region of ~52 and 73 bp, respectively. Volatility charts for \textit{rpoHp2}, \textit{rpoHp3}, \textit{rpoHp4}, \textit{rpoHp5} and \textit{rpoHp6} show the same trend, if not more pronounced conserved secondary structure in their respective target regions (Supplementary Figure S6 through Figure S10 for volatility profiles).

The level of regulation of \(\sigma^{32}\) that was detailed earlier using this target region indicates that there was perhaps a greater potential for superior down-regulation if the target had been chosen elsewhere, despite rationale of choosing the location of the Shine-Dalgarno sequence. This may be particularly true because of the presence of multiple mRNAs with very stable secondary structure in the target region. The yield of OPH could have perhaps been higher, while GroEL and \(\sigma^{32}\) levels could have been much lower under both heat shock and ethanol shock.

**DISCUSSION AND CONCLUSION**

The ability to determine accessible regions on a strand of mRNA for antisense binding has been the goal of many researchers for decades and continues to be a puzzling problem to this day. The proposed hypothesis that structurally volatile regions of mRNA make the best antisense targets may provide insight into how natural antisense transcripts evolved to be most effective in regulating the expression of their corresponding gene. It is particularly important to analyse \textit{cis}-encoded antisense systems because they most closely resemble artificial antisense sequences and reveal information about mechanisms through which completely complementary sequences are utilized in nature. This is opposed to the potentially less informative \textit{trans}-encoded antisense that often have unusual binding mechanisms or even multiple binding sites, many of which are difficult to predict \textit{in silico} (40).

All of the natural antisense systems described here are toxin–antitoxin systems, where an mRNA encodes for a host-killing protein and a shorter strand of antisense RNA that blocks the translation of this protein-encoding mRNA. This is a particularly interesting aspect of our findings because it is assumed that these antisense have evolved to be particularly effective, minimizing leaky translation so that a cell carrying such a ‘suicide protein’ gene will not only survive but will suffer the least amount of growth inhibition possible. The \textit{gef}/\textit{sof} example is of particular interest because even after the insertion sequence disrupted the gene and lengthened the mRNA sequence by more than 300 nucleotides, most of the volatile region was still overlapping with almost all of the \textit{sof} target sequence. It is surprising that this extreme lengthening did not alter the structural free energy calculations in such a way that another target region might be more volatile. The longer an RNA sequence is, the more options UNAFold has to predict suboptimal structures within the allowed deviance from the minimum free energy. Even if \textit{sof} did not initially evolve to regulate the significantly longer mRNA, the cells carrying the insertion may still have been able to survive because the volatile region on the mRNA remained relatively intact at the \textit{sof} target site. As a result, antisense continued to effectively bind to the same region.
The fact that the predicted sequences from GenAVERT most closely matched naturally occurring antisense when sequence length was restricted to 35 bp is very intriguing. In most cases, the antisense sequence length jumped from lengths of about 19, 32 or 34 bp up to lengths of 60, 70 or 90 bp at the next (and best) ranking. This indicates that there may be an underlying fundamental design aspect through evolution where these lengths may provide a greater antisense efficiency by maximizing specificity and thermodynamic hybridization capabilities while still minimizing antisense secondary structure.

In terms of benchmarks, Sfold does not recommend an antisense length and it simply defaults to a length of 20 and searches through the entirety of the given mRNA for accessibility and hybridization potential. The results show that the top scoring antisense sequences are clearly not predicted in the same way as GenAVERT’s, and their predicted target sites may be scattered all over an mRNA sequence. This is evident in all of the natural antisense examples except for that of hokC. GenAVERT was meant to overcome this problem and provide a better direction towards what the best antisense sequence would be without forcing a guess at a ‘preferred sequence length.’

In examining the mRNAs that had been previously targeted with artificial antisense, the volatility profiles provided extensive insight into their potential mechanisms of action. The most volatile regions on adc mRNA and ptb-buk matched up well with the antisense target sites, offering a possible explanation for their high level of down-regulation of the encoded proteins. The predicted antisense from GenAVERT for ptb-buk mRNA was particularly striking when only the first 75% of the mRNA was included as sequence search space. In this case, all of the top eight scoring antisense sequences encompassed or included the majority of the antisense target sites. In addition, after examining the set of rpoH mRNAs, it is apparent that the majority of the artificial antisense target sites on each of these sequences may have possessed a high level of conserved secondary structure and thus lack of volatility. The experimental results did indicate some protein down-regulation, which lines up with the fact that some shorter lengths of the target region exhibited high volatility. However, GenAVERT predicted regions of even higher volatility elsewhere, indicating that there was a much greater potential to inhibit $\sigma^{32}$ expression at those locations away from the target region. With the long lengths (800–2000 bp) of these mRNAs, it is apparent that GenAVERT has the potential to have an impact and be applicable in current research.

Since only bacterial examples have been examined, any benefits of using GenAVERT to design antisense sequences for eukaryotic mRNA remains unknown. Many eukaryotic mRNA sequences are extraordinarily long compared with the mRNA sequences of bacteria, and their structures may not, in fact, be accurately predicted by UNAFold. Likewise, making comparisons of sequences and structures that are thousands upon thousands of bases long with RNAforester exponentially increases the program’s runtime. However, the concept of volatility in mRNA secondary structure may still be a useful and applicable approach to blocking eukaryotic gene expression. This could perhaps be accomplished by breaking up long mRNA sequences into smaller overlapping parts and analyzing each individually for potential volatile regions. For the moment, 2500 bp may be a reasonable absolute limit for the sequence length that GenAVERT can handle because of runtime and the potential lack of accuracy in secondary structure prediction in RNAs longer than this.

Also, simply because GenAVERT predicts a certain volatile region on an mRNA sequence, does not mean that this entire region is the absolute optimal antisense binding site. Other factors may also play a significant role in antisense down-regulation. Thus, in the future, it may be helpful to incorporate hybridization algorithms to pinpoint a subsequence within a larger volatile region that would maximize hybridization. It has also been shown that perhaps different antisense sequences are more effective at different in vivo concentrations. As a result, it may be possible to search this volatile region for optimum hybridization while also utilizing equilibrium concentration data to identify the utmost effective antisense sequence.

Programs such as Ensemble_Calc (http://mfold.rna.albany.edu/?q=DINAMelt/Ensemble-calc) (41) may aid in this task.

Finally, since each individual base is crucial in the structural thermodynamic calculations and since so many different genes exhibit such intriguing patterns in volatility, it is almost certain that more antisense examples have yet to be found. By leveraging the idea that an mRNA molecule is not static, the hypothesis presented here may provide a new strategy in rational antisense design by predicting which sites on an mRNA strand are truly accessible for antisense targeting.

SUPPLEMENTARY DATA
Supplementary Data are available at NAR Online: Supplementary Figures 1–10 and Supplementary Dataset 1.

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