Automatic detection of conserved RNA structure elements in complete RNA virus genomes

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ABSTRACT

We propose a new method for detecting conserved RNA secondary structures in a family of related RNA sequences. Our method is based on a combination of thermodynamic structure prediction and phylogenetic comparison. In contrast to purely phylogenetic methods, our algorithm can be used for small data sets of ∼10 sequences, efficiently exploiting the information contained in the sequence variability. The procedure constructs a prediction only for those parts of sequences that are consistent with a single conserved structure. Our implementation produces reasonable consensus structures without user interference. As an example we have analysed the complete HIV-1 and hepatitis C virus (HCV) genomes as well as the small segment of hantavirus. Our method confirms the known structures in HIV-1 and predicts previously unknown conserved RNA secondary structures in HCV.

INTRODUCTION

One of the major problems facing computational molecular biology is the fact that sequence information is available in far greater quantities than information about the three-dimensional structure of biopolymers. While the prediction of three-dimensional RNA structures from sequence data is unfeasible at present (see, however, for a promising approach), the prediction of secondary structure is in principle tractable even for large molecules. Functional secondary structures are conserved in evolution (see for instance and they represent a qualitatively important description of the molecules, as documented by their application to the interpretation of molecular evolution data.

Almost all RNA molecules and, consequently also almost all sub-sequences of a large RNA molecule form secondary structures. The presence of secondary structure in itself therefore does not indicate any functional significance. In this contribution we show that potentially functional RNA structures can be identified by a purely computational procedure that combines structure prediction and sequence comparison. RNA viruses are an ideal proving ground for testing such a method.

(i) Distant groups of RNA viruses have very little or no detectable sequence homology and often very different genomic organization. Thus we can test our approach on essentially independent data sets.

(ii) RNA viruses show an extremely high mutation rate, of the order of 10⁻⁵–10⁻³ mutations per nucleotide and replication. Due to this high mutation rate they form quasi-species, i.e. diffuse ‘clouds’ in sequence space (3), and their sequences evolve at a very high rate. In contrast, functional secondary structures are strongly conserved. Due to the high sequence variation, the application of classical methods of sequence analysis is, therefore, difficult or outright impossible. Indeed, except for the family Mononegavirales (negative-stranded RNA viruses), there is no accepted taxonomy above the genus level.

(iii) The high mutation rate of RNA viruses also explains their short genomes, of less than ∼20 000 nt (3). A large number of complete genomic sequences is available in databases. The non-coding regions are most likely functionally important, since the high selection pressure acting on viral replication rates makes ‘junk RNA’ very unlikely. So far, a number of relevant secondary structures have been determined that play a role during the various stages of the viral life cycle in a variety of different classes of viruses, for instance lentiviruses (4–6), RNA phages (7,8), flaviviruses (9), pestiviruses (10,11), picorna viruses (12–17), hepatitis C viruses (10,18) and hepatitis D virus (19).

Three unrelated groups of viruses, which contain a variety of human pathogens of global medical importance, will serve as examples (see Fig. 1 for details). HIV-1 is a highly complex retrovirus. Its genome is dense with information for coding of proteins and is biologically significant. RNA secondary structures. The latter play a role in both the entire genomic HIV-1 sequence and in the separate HIV-1 mRNAs, which are basically (combined) fragments of the entire genome. Flaviviridae are small enveloped particles with an unsegmented, plus-stranded RNA genome. This virus family contains the genera flavivirus (which includes the viruses causing Japanese encephalitis, dengue fever, yellow fever and tick-borne encephalitis) pestivirus, hepatitis C and the recently discovered hepatitis G viruses (see 20 for a recent summary). Hantaviruses are serologically related members of the family Bunyaviridae (21). They are enveloped viruses with a tripartite negative sense RNA genome. The three genome segments are called L, M and S, encoding the viral transcriptase, envelope glycoproteins and nucleocapsid protein respectively. In this contribution we shall be concerned only with the small (S) segment. Hantaviruses
have been implicated as aetiological agents for two acute diseases: hemorrhagic fever with renal syndrome (HFRS) and hantavirus pulmonary syndrome (HPS). Both diseases are carried by rodent vectors. The total length of the genomic sequences of HIV-1 and hepatitis C virus (HCV), of the order of 10,000 nt, makes experimental analysis of the secondary structure of full genomes unfeasible. For RNAs of this size, structure prediction based on thermodynamic constraints is the only approach that is available at present.

MATERIALS AND METHODS

RNA structure prediction

RNA secondary structures are predicted as minimum energy structures by means of dynamic programming techniques (22–25). An efficient implementation of this algorithm is part of the Vienna RNA Package (available at http://www.tbi.univie.ac.at/ivo/RNA/; 26). Complete HIV and HCV genomes were folded on CalTech’s Delta using the message-passing version of the minimum folding algorithm described by Hofacker et al. (5,27). This version uses energy parameters based on Freier et al. (28), Jaeger et al. (29) and He et al. (30), but ignores dangling ends. The parameters are identical to those in Michael Zuker’s mfold 2.2 with the exception that stacking energies involving GU pairs were taken from He et al. (30). All other foldings were performed using version 1.2 of the package, which uses an updated parameter set described in Walter et al. (31).

Sequence and structure comparison

While computation of the secondary structures is a straightforward (yet computationally demanding) task, their comparison is less obvious. A variety of combined alignment plus structure prediction procedures have been proposed (32–34). The problem with this approach is three-fold for our task. (i) The computational efforts become prohibitive for longer sequences: CPU time scales of the order of \( n^4 \) in the approximate algorithm (32) and \( n^{3m} \) in the exact version (33), where \( n \) is the length of the sequence and \( m \) the number of sequences, by far exceed the available resources. (ii) Viral sequences show a large variation in sequence similarity along the chain. Furthermore, we do not expect a conserved secondary structure for all parts of the sequence, even if there is a significant level of sequence conservation. Combined folding and alignment algorithms will, therefore, produce poor alignments in such cases. (iii) The use of a combined algorithm for predicting structure and alignment would not allow independent verification of the predicted structural elements. The possibility of verifying the predicted structures, however, is particularly important when dealing with the relatively sparse data sets that are available.

We start the comparison procedure with an alignment of the sequences that is obtained without any reference to the predicted structures. The multiple sequence alignments are calculated using CLUSTAL W (35). A good alignment is a prerequisite for the success of our method. We find, however, that regions with conserved structures tend to align well, at least locally. We did not find it necessary to improve the alignments based on visual inspection. A modification of the alignment taking into account already predicted structures might increase the number of compensatory mutations and possibly also the number of detected structural elements. However, it would compromise the use of the sequence data for verifying the predicted structures.

The sequence alignment is then used to produce an alignment of the secondary structures by introducing the appropriate gaps into the minimum energy foldings. Up to this point our procedure is essentially the same as Riesner’s ConStruct (36), although we...
Figure 2. Scheme of the secondary structure analysis of viral genomes. Sequences are aligned using a standard multiple alignment procedure. Secondary structures for each sequence are predicted and gaps are inserted in the sequence alignment. The resulting aligned structures can be represented as aligned mountain plots. From the aligned structures consistently predicted base pairs are identified. The alignment is used to identify compensatory mutations that support base pairs and inconsistent mutants that contradict pairs. This information is used to rank proposed base pairs by their credibility and to filter the original list of predicted pairs.

start from minimum energy structures instead of base pair probabilities. The evaluation of the structure alignment, however, is quite different from these approaches. (i) We do not assume a priori that there is a conserved secondary structure for all parts of the sequence. Hence, we cannot simply search for the secondary structure that maximizes the sum of the predicted base pairing probabilities. (ii) We explicitly use the sequence information contained in the multiple alignment to confirm or reject predicted base pairs. A flow diagram of our approach is shown in Figure 2.

A quick overview of the data is conveniently obtained from the mountain representation. In the mountain representation (37) a single secondary structure is represented on a two-dimensional graph, in which the x-coordinate is the position k of a nucleotide in the sequence and the y-coordinate the number m(k) of base pairs that enclose nucleotide k. The mountain representation allows for a straightforward comparison of secondary structures and inspired a convenient algorithm for structure-based alignments of secondary structures (37, 38). Mountain representation plots, such as the one in Figure 3, can be used to identify conserved sub-structures. The consensusmountain of a set of N sequences can be defined as

\[
m(k)1/N \sum_{s}^{N} m_s(k) = 1
\]

The quality of a consensus mountain can be assessed at each position by comparing the slopes \(q_s(k) = m_s(k) - m_s(k-1)\) of the different sequences (39). These one-dimensional representations, such as \(m(k)\), provide a global overview of the structure and can be used to guide a manual reconstruction of consensus secondary structure elements. This approach turned out to be rather tedious for the 3'-non-coding regions (NCRs) of flaviviruses with a chain length of only 200–300 nt and is certainly not feasible for the analysis of entire genomes. On the other hand, the data contained in these simplified one-dimensional representations are not detailed enough to allow for automatic reconstruction of conserved patterns.

Automatic detection of conserved structural elements

The starting point of a more detailed analysis is a list of all predicted base pairs. This list will in general not be a valid secondary structure, i.e. it will violate one or both of the following two conditions: (i) no nucleotide takes part in more than one base pair; (ii) base pairs never cross, i.e. there may not exist two base pairs \((i, j)\) and \((k, l)\) such that \(i < k < j < l\).

Therefore, we rank the individual base pairs by their `credibility' (see below). Then we go through the sorted list and weed out all
base pairs that violate conditions (i) or (ii). Clearly, the sorting procedure is of crucial importance. For each predicted base pair \((i,j)\) we store the nucleotides occurring in the corresponding positions in the sequence alignment. We shall call a sequence non-compatible with a base pair \((i,j)\) if the two nucleotides at positions \(i\) and \(j\) would form a non-standard base pair, such as CA or UU. A sequence is compatible with base pair \((i,j)\) if the two nucleotides form one of the following six combinations: GC, CG, AU, UA, GU or UG.

When different standard combinations are found for a particular base pair \((i,j)\) we may speak of consistent mutations. If we find combinations such as GC and CG or GU and UA, where both positions are mutated at once, we have compensatory mutations. The occurrence of consistent and, in particular, compensatory mutations strongly supports a predicted base pair, at least in the absence of non-consistent mutations.

From the frequencies \(f_{ij}\) with which \((i,j)\) is predicted in the sample of sequences we derive the pseudo-entropy

\[
S_{ij} = -\sum_{k} f_{ik} \ln f_{ik} - \sum_{k} f_{kj} \ln f_{kj} + f_{ij} \ln f_{ij}
\]

where \((i,k)\) and \((k,j)\) are the alternative predicted base pairs involving \(i\) and \(j\) respectively. The pseudo-entropy is a measure for the reliability with which \((i,j)\) is predicted.

We call a base pair \((i,j)\) symmetrical if \(j\) is the most frequently predicted pairing partner of \(i\) and if \(i\) is the most frequently predicted pairing partner of \(j\). Note that for each sequence position \(i\) there is at most one symmetrical base pair involving \(i\). A symmetrical base pair \((i,j)\) necessarily has a rather large value for \(f_{ij}\); in particular, it does not allow a large number of structural alternatives.

In a first preprocessing step we remove for each \(i\) all but the most frequent pair \((i,j)\) from the list of predicted base pairs. The list is then sorted according to the following hierarchical criteria [i.e. criterion (ii) is used only when two pairs are not distinguished by criterion (i), and so on]:

(i) the more sequences non-compatible with \((i,j)\), the less credible is the base pair;

(ii) symmetrical base pairs are more credible than other base pairs;

(iii) a base pair with more consistent mutations is more credible;

(iv) base pairs with smaller values of pseudo-entropy \(S_{ij}\) are more credible.

Note that criteria (i) and (iii) make direct use of the sequence information without reference to frequency of a base pair.

Scanning the sorted list from the top, we remove a base pair if it conflicts with a higher ranking one that has already been accepted. Finally, base pairs with \(f_{ij}\) below some threshold are removed. This ensures that structures are predicted only for regions in which we have a strong signal. The threshold value is a conservative estimate for the reliability of the secondary structure prediction (40). In this study a value of 0.3 gave good results. The final output can be displayed as a colour-coded dot plot (as shown in Figs 4 and 5) or as a colour-coded mountain plot (Fig. 7).

The virtue of this approach can be tested quite easily. In Figure 4 we compare the predicted consensus structure for two sets of 2-error mutants of the same wild-type sequence. Sequences in the first set were generated by randomly mutating two positions. Even this small amount of sequence heterogeneity leads to a quite diverse set of structures (although some sequences fold into the wild-type structure) and, hence, no unambiguous secondary structure is predicted.

A second set was generated in the same way, but this time only sequences folding into the same structure as the wild-type were accepted. Since, by construction, all sequences fold into the same structure, we obtain a perfect prediction for this set which is supported by a small number of compensatory mutations.

The example shows that even a small number of mutations will disrupt the secondary structure in the absence of selection pressure to conserve the structure. Our approach is capable of distinguishing conserved secondary structure elements from pieces of sequence with high degrees of homology but without conserved structural features.

Figure 3. Aligned mountain representations \(m(k)\) of the RNA secondary structure of 13 complete HCV genomes. Peaks and plateaux in the mountain representation correspond to hairpins and unpaired regions in the secondary structure. The folds were computed with CalTech’s Intel Delta, a distributed memory parallel computer with 512 nodes and roughly 12 Mbytes memory per node. The thick full line is the average mountain representation. In the lower part of the sequence we plot the variance of the slopes (scattered dots) and a running average (full green line). Deep minima of the green curve correspond to consistently predicted parts of the structure, such as the two regions labelled A and B.
Figure 4. An artificial example. Two samples of 2-error mutants of the yeast tRNA\textsuperscript{phe} sequence were subjected to our procedure. A square in row \(i\) and column \(j\) of the dot plot indicates a predicted pair \((i, j)\). Its size and colour indicates the frequency and 'credibility' of the base pair. The area of the square is proportional to the frequency \(f_{ij}\) with which \((i, j)\) is predicted. Colours indicate the number of consistent mutations: red 1, yellow 2 and green 3 different types of base pairs. These saturated colours indicate that there are only compatible sequences. Decreasing saturation of the colours indicates an increasing number of non-compatible sequences; i.e. sequences that cannot form \((i, j)\). If there are more than two non-compatible sequences the entry is not displayed. (Upper right triangle) The 29 sequences that fold into the wild-type cloverleaf structure of tRNAs lead to a perfect reconstruction of the secondary structure. Each helix is supported by at least one consistent mutation. Forty-seven of the 76 sequence positions are conserved. (Lower left triangle) A sample of 20 randomly generated 2-error mutants of the tRNA\textsuperscript{phe} sequence does not produce a reasonable prediction of the cloverleaf structure: one stack of the cloverleaf is not predicted at all and another stack does not conform to the wild-type structure. Each helix contains several inconsistent mutations, despite the fact that 43 of the 76 positions are conserved. The only acceptable signal in this data set are the top-most three pairs of the anticodon loop.

Table 1. Predictions of 5s RNAs

<table>
<thead>
<tr>
<th>Sample</th>
<th>(n)</th>
<th>Sequence identity (%)</th>
<th>Base pairs Predicted</th>
<th>'False'</th>
<th>Phylogenetic</th>
</tr>
</thead>
<tbody>
<tr>
<td>Halobacteriales</td>
<td>12</td>
<td>81.8</td>
<td>31</td>
<td>1(^a)</td>
<td>31...36</td>
</tr>
<tr>
<td>Methanomicrobiales</td>
<td>9</td>
<td>75.0</td>
<td>29</td>
<td>2(^a)</td>
<td>32...39</td>
</tr>
<tr>
<td>Halobacteriales + Methanomicrobiales</td>
<td>11</td>
<td>71.1</td>
<td>30</td>
<td>1(^a)</td>
<td>31...39</td>
</tr>
<tr>
<td>Methanobacteriales + Methanococciales</td>
<td>12</td>
<td>67.3</td>
<td>33</td>
<td>4(^a)</td>
<td>33...38</td>
</tr>
<tr>
<td>Eubacteria</td>
<td>15</td>
<td>60.9</td>
<td>16</td>
<td>0</td>
<td>26...33</td>
</tr>
<tr>
<td>Eubacteria(^b)</td>
<td>10</td>
<td>58.8</td>
<td>19</td>
<td>2(^a)</td>
<td>25...33</td>
</tr>
<tr>
<td>Archaea</td>
<td>10</td>
<td>58.3</td>
<td>23</td>
<td>0</td>
<td>32...46</td>
</tr>
<tr>
<td>Eubacteria + Archaea</td>
<td>10</td>
<td>52.8</td>
<td>21</td>
<td>0</td>
<td>26...45</td>
</tr>
</tbody>
</table>

\(^{a}\)No 'falsely' predicted base pair is in conflict with the phylogenetic structure.

\(^{b}\)Disjoint samples.
As a second example, we applied our method to samples from the Berlin RNA Databank \(^{(41)}\), which contains 5S RNA sequences and phylogenetic structures. To investigate the influence of sequence heterogeneity we randomly selected 9–15 sequences from different sub-groups (see Table 1). For a relatively homogeneous sample, such as the Halobacteriales, we obtain an almost perfect prediction with only one small helix of 2 bp missing. For heterogeneous samples such as a random selection of five Eubacteria and five Archaea we still find most of the correct structure without introducing any false positives. Occasionally we find base pairs not in the phylogenetic structure. In all such cases they only elongate a helix present in the phylogenetic structure.

The accuracy of the minimum energy predictions for these sequences varies widely; on average \(\sim 70\%\) of the phylogenetic pairs are present in the minimum energy structure. Note, however, that different pairs are predicted in different sequences; in fact, no pair was present in the minimum energy structures of all sequences in a sample and some minimum energy structures do not contain a single correct pair.

Sometimes well-predicted stacked regions are interrupted by individual ‘holes’ or show a single base pair with a few non-compatible sequences. While in many cases these features reflect structural variability or the existence of an internal loop, they can be attributed to non-standard base pairs, like GA or UU, that do not necessarily disrupt the helix \(^{(42)}\) in other cases. An example is shown in Figure 5.

**RESULTS**

As a first application of our method we have investigated the minimum energy folding of 13 complete HCV sequences, a sample of 13 complete HIV-1 sequences and the S segment of 19 strains of hantavirus (access codes are listed in the Appendix). Minimum free energy structures of the complete HIV and HCV genomes were obtained on CalTech’s Intel Delta. For details of the parallel computer implementation of the folding algorithm see Hofacker *et al.* \((5)\). The hantavirus sequences were folded using the serial version of the folding program, which uses a slightly
HIV1

13

2

0

more recent parameter set (31). Multiple sequence alignments were obtained using CLUSTAL W. Both the folding outputs and the multiple alignments were processed without further modification.

Hepatitis C sequences have chain lengths of ∼9500 nt. The main differences in length stem from the 3′-end of the genome, where a poly(U) region separates a 98 nt sequence from the rest of the genome (18,43). This so-called X-tail is not present in the published ‘complete’ genomes, with a single (very recent) exception (Genbank accession no. D85516). In this sequence we found no long range interactions involving the X-tail and, hence, no evidence for the panhandle structure postulated in figure 7B of Kolykhlov et al. (43). We find that the poly(U) region acts as a spacer causing the X-tail to fold as a separate domain. It is justified, therefore, to consider the main part of the genome [before the poly(U) region] and the X-tail separately.

The length of the CLUSTAL W alignment of the main part of the genome, up to the poly(U), is 9538. Insertions or deletions appear in 267 positions before the poly(U). 4919 positions are conserved; the mean pairwise identity of sequences is 80%.

The 5′-NCR has recently been studied using a combination of thermodynamic prediction and biochemical methods (10,44,45). Unfortunately, the sequences at the 5′-end are highly conserved (97% pairwise identity, 89% of the 342 positions conserved). As a consequence of the very small sequence variation, our approach is not much better than thermodynamic predictions on a single sequence in this case. We find most of hairpins appearing in the model of Brown et al. (10), but predict no longer range base pairs.

A similar situation is encountered for the X-tail: of the 98 nt only five show any sequence variability. There are only three different sequences of these 98 nt among the eight database entries currently available. Our data agree with the three stem–loops predicted in Blight and Rice (46) and Ito and Lai (47) based on chemical probing. It is also supported by two consistent mutations in the long helix at the very 3′-end.

We do, however, find convincing structural motifs within the coding region of the viral genome, two examples of which are shown in Figures 5 and 6. Although neither region has been investigated before, the large number of compensatory mutations clearly indicates that these structural motifs are conserved.

The minimum free energy structures of the 13 sequences contain a total of 23 186 bp. Preprocessing leaves only 2805 list entries, which is already slightly less than the ∼3050 bp predicted in the individual minimum energy structures. Of these, 432 entries are inconsistent with higher ranking entries in the sorted list, 572 entries are removed because there are more than two inconsistent sequences and the frequency $f_{ij}$ of 298 of the remaining base pairs is below the threshold value 0.3. This leaves us with 1503 pairs in the dot plots, a reduction of >50% from the original. Of these, 985 have only compatible sequences, 179 have a single incompatible sequence and 339 have two.

The output generated for large molecules will in general still contain a substantial number of base pairs that are at best doubtful, such as isolated base pairs and short helices with several inconsistent sequences. The truly promising structures are, however, easy to detect by visual inspection of the resulting dot plot.

As a second example, we have re-analysed a sample of HIV-1 sequences from an earlier study (5). The number of predicted conserved base pairs is similar to the HCV case (see Table 2 for details). In the following we discuss the automatically generated predictions for two well-understood secondary structure motifs, namely the trans-activating responsive element (TAR) and Rev response element (RRE), in some detail.

Table 2. Predicted secondary structure elements

<table>
<thead>
<tr>
<th></th>
<th>HCV</th>
<th>HIV1</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of sequences (N)</td>
<td>13</td>
<td>13</td>
</tr>
<tr>
<td>Minimum sequence</td>
<td>9400</td>
<td>9074</td>
</tr>
<tr>
<td>Maximum sequence length</td>
<td>9502</td>
<td>9292</td>
</tr>
<tr>
<td>Alignment length</td>
<td>9538</td>
<td>9535</td>
</tr>
<tr>
<td>Conserved positions</td>
<td>4919</td>
<td>4779</td>
</tr>
<tr>
<td>Average sequence identity (%)</td>
<td>80</td>
<td>83</td>
</tr>
<tr>
<td>Different base pairs</td>
<td>23 186</td>
<td>20 667</td>
</tr>
<tr>
<td>Credible base pairs</td>
<td>1503</td>
<td>1121</td>
</tr>
<tr>
<td>1 Consistent mutation</td>
<td>460</td>
<td>300</td>
</tr>
<tr>
<td>2 Consistent mutations</td>
<td>80</td>
<td>44</td>
</tr>
<tr>
<td>3 Consistent mutations</td>
<td>8</td>
<td>2</td>
</tr>
<tr>
<td>4 Consistent mutations</td>
<td>2</td>
<td>0</td>
</tr>
</tbody>
</table>

At the 5′-end of the viral HIV-1 RNA molecule resides the trans-activating responsive (TAR) element (48), which interacts with the regulatory Tat protein. Binding of the Tat protein to TAR is responsible for activation and/or elongation of transcription of the provirus (49,50). On the basis of biochemical analysis (4) and computer prediction of the 5′-end of the genome it is known that the TAR region in HIV-1 forms a single isolated stem–loop structure of ∼60 nt with ∼20 bp interrupted by two bulges.

This structure is indeed predicted in the minimum free energy structures of 11 of the 13 sequences analysed here. The consensus prediction (Fig. 7) is identical to the structure reported in the literature. The mean pairwise sequence identity of this region is 85%.
Figure 7. The TAR structure of HIV-1. Almost all predicted base pairs are consistent with all 13 sequences, most of them are predicted in at least 11 sequences. A large number of compensatory mutations supports the thermodynamic predictions. Our computed consensus structure (lower left) matches the structure determined by probing and phylogenetic reconstruction (4). We display here the consensus dot plot, the classical secondary structure and a mountain representation. The latter is a convenient alternative to dot plots for larger structural motifs. Base pairs are represented by slabs connecting the two sequence positions. The width and colour of a slab corresponds to size and colour of the corresponding dot plot entry.

The Rev response element (RRE) is an important conserved RNA structure that is located within the env gene. The interaction of RRE with the Rev protein reduces splicing and increases the transport of unspliced and single spliced transcripts to the cytoplasm, which is necessary for the formation of new virion particles (51).

A long stem–loop structure (I) separates the binding region from the rest of the RNA. The long stem–loop structure furthermore indicates that the structure is easily accessible. The consensus secondary structure of the RRE in HIV-1 is a multi-stem–loop structure consisting of five hairpins supported by a large stem structure (52; see Figs 8 and 9). An alternative structure of only four hairpins, in which hairpins III and IV of the consensus model merge to form one hairpin, has also been proposed (53,54); it matches the minimum energy structure for some sequences, e.g. HIVLAI (55). A comparison of minimum energy structures (5,27) shows that there appears to be a third structure in which hairpin III is relatively large and a few of the other hairpins have disappeared from the minimum free energy structure. A comprehensive analysis of the base pairing probabilities in the RRE shows that hairpins II, IV and V, as well as the basis of hairpin III, are not well defined, in the sense that they allow for different structures with comparable probabilities (56).

As a final example we consider the S segment of hantavirus. The ~1700 nt long S segment contains a single ORF encoding a nucleocapsid (N) protein. In contrast to other members of the family Bunyaviridae, there is no evidence for a second non-structural (NS) protein coded by the S segment. We used the 19 sequences listed in the Appendix, which have a mean pairwise identity of 63.9%. The only detected structural feature in this case is a 19 bp stem–loop structure formed by the 5′- and 3′-ends.

This panhandle structure is highly significant: All sequences are compatible with the structure and it is part of the minimum energy prediction in 16 of the 19 minus strands and 14 plus strands. There are two positions which show compensatory mutations (see Fig. 10). The panhandle structure was postulated in the 1980s for all Bunyaviridae (57,58).

DISCUSSION

We have presented a combination of secondary structure prediction based on thermodynamic criteria and sequence comparison that is capable of reliably identifying conserved structural features in a set of related RNA molecules. The method has been designed for routine investigations of large RNA molecules, such as complete viral genomes. Indeed, the procedure does not require any intervention: CLUSTAL W alignments and minimum energy structures [as obtained from the Vienna RNA Package or Zucker’s mfold (59)] can be used ‘as is’. Our program currently does not support the detection of pseudo-knots. However, the method is in principle suitable for this task (with minor modifications), provided the structure prediction algorithm allows for pseudo-knots (60,61).
Conserved secondary structures are likely to be functional, thus our method can be used to find functional secondary structures. Since our method emphasizes sequence variation, it complements other methods for finding functional RNA secondary structures based on thermodynamic prediction (for example 55,62).

We have applied this technique to complete genomes of three quite different species of RNA viruses: HIV-1, HCV and the small segment of hantavirus. In all cases we have been able to identify most of the known secondary structure features. In addition, we predict a large number of conserved structural elements which have not been described so far.

We have designed our approach in such a way that it does not predict a structure for all parts of a molecule; the filtering procedure outlined in Materials and Methods is designed in such a way that only base pairs that may occur in almost all sequences and that are predicted in a sizeable fraction of the sequences will be accepted. It is not surprising, therefore, that the predicted RRE structures in Figure 9 do not contain every single base pair of the published, experimentally supported structures. Rather, we obtain a subset of base pairs that is consistent with known features. This suggests that we are not producing a large number of false positives. The fact that no false positives were produced in the analysis of 5S RNA sequences supports this claim. On the other hand, we recover most of the structures described for both HIV-1 and HCV, as well as the panhandle structure of hantavirus and the main structural features of 5S RNA.

Since the success of the method depends on the availability of a good alignment, it works best for samples with moderate sequence heterogeneity (say 80% identity). However, alignment errors should at worst cause some structures to be missed, but are not likely to lead to falsely predicted structures. For heterogeneous samples of sequences that code for proteins good quality alignments might be obtained by first aligning the protein sequences of the translation products and translating back to nucleic acid sequences.

Extensive computer analysis of the RRE region of HIV-1 and HIV-2 has shown that the sequence alignment does not completely coincide with the alignment at the level of the secondary structure (63). This has two important implications: (i) methods that predict secondary structure of RNA on the basis of co-variation of positions within the sequence (2) cannot provide unambiguous answer here; (ii) the RRE has structural versatility. As a consequence, we obtain slightly different predictions for the conserved structure depending on the set of sequences used for the analysis. This structural versatility could also play a role in a single HIV clone. Using McCaskill's algorithm (64) for predicting the matrix of base pairing probabilities, we have indeed identified a spectrum of alternative structures for the RRE of HIVLAI in a previous communication (55). Similar features have been detected in the 3'-NCR of flaviviruses (39,65). These facts make it worthwhile to generalize the present approach to using base pairing probability matrices instead of minimum energy structures,
Figure 9. Consensus structures of the HIV-1 RRE region from a set of 13 sequences and from the 21 sequences reported in Hofacker et al. (5). The main hairpins are present in both predictions; the only difference is hairpin Ia which is supported by a single compensatory base pair in the larger data set. The predictions agree very well with an experimentally supported structure (52) that also contains Ia. The sequence in the Ia region is conserved in the smaller data set and purely thermodynamic considerations favour the short stack extending stack III in the right hand structure [this stem is called IIc in Mann et al. (53) and Zemmel et al. (54)]. Interestingly, earlier studies (5,55) indicate a substantial structural versatility in this region which may explain minor disagreements between different published structures (see for example 52–54).

Figure 10. Consensus structures for the minus strand of the S segment of hantavirus. The only consistently predicted structure is a panhandle formed by the 5′- and 3′-ends.
as in ConStruct (1), despite the substantial increase in required computer resources. Preliminary data indicate promising increases in the accuracy of predicted structures.

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38 ConStruct (1), despite the substantial increase in required computer resources. Preliminary data indicate promising increases in the accuracy of predicted structures.

APPENDIX

In this study we have used the following viral RNA sequences (Gembank accession nos are given in parentheses).

HIV-1: HIVANT70 (M31171, L20587), HIVBCSG3C (L20317), HIVCMAM (D10112, D00917), HIVD31 (X61240, 3835)}
X16109 U23487, HIVELI (K03454, X04414), HIVLAI (K02013), HIVMAL (K03456), HIVMVP5180 (L20571), HIVNDK (M27323), HIVOYI (M26727), HIVRF (M17451, M12508), HIVU455 (M62320) and HIVZ2Z6 (M22639).

HCV: complete genomes (except for the X-tail): HCU16362 (U16362), HCU45476 (U45476), HPCCGAA (M67463), HPCCGENOM (L02836), HPCCGS (D14853), HPCEGS (D17763), HPCHCJ1 (D10749), HPCIJ483 (D13558, D01217), HPCRJRNA (D14484, D01173), HPCJT (D11168, D01171), HPCK3A (D28917), HPCPP (D30613) and HPCRNX1 (D10934).

X-tail sequences (accession numbers only): D63922, D67091, D67092, D67093, D67094, D67095, D67096, D85516 (the last sequence is a complete genome including the X-tail).

Hantavirus sequences: AF004660, HNVNPSS, HNU37768, HMU32591, HUS29210, KUH35255, AF005727, PHU47136, PHVSESEG, PSU47135, PUUSNP, PUUSSVI83, PUUSSVIRRT, PUU82404, PUU22423, VRANICAS, RMU52136, HPSNUPR and TUVSS5302.

The comparison algorithm described is implemented as an ANSI C program alidot. It generates a text file with information on all predicted base pairs and a postscript file of the dot plot of the predicted conserved base pairs. Alternative representations, such as the aligned mountain plots, input files for XRRA (66) and post-processing of XRRA output is handled by a collection of perl scripts. This software is available upon request from the authors.