Structural basis for the biological relevance of the invariant apical stem in IRES-mediated translation

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ABSTRACT

RNA structure plays a fundamental role in internal initiation of translation. Picornavirus internal ribosome entry site (IRES) are long, efficient cis-acting elements that recruit the ribosome to internal mRNA sites. However, little is known about long-range constraints determining the IRES RNA structure. Here, we sought to investigate the functional and structural relevance of the invariant apical stem of a picornavirus IRES. Mutation of this apical stem revealed better performance of G:C compared with C:G base pairs, demonstrating that the secondary structure solely is not sufficient for IRES function. In turn, mutations designed to disrupt the stem abolished IRES activity. Lack of tolerance to accept genetic variability in the apical stem was supported by the presence of coupled covariations within the adjacent stem–loops. SHAPE structural analysis, gel mobility-shift and microarrays-based RNA accessibility revealed that the apical stem contributes to maintain IRES RNA structure through the generation of distant interactions between two adjacent stem–loops. Our results demonstrate that a highly interactive structure constrained by distant interactions involving invariant G:C base pairs plays a key role in maintaining the RNA conformation necessary for IRES-mediated translation.

INTRODUCTION

Translation initiation is a key step in the process of protein synthesis (1). In some RNA viruses, exemplified by picornaviruses and hepatitis C virus (HCV), and in a subset of cellular mRNAs which are translated during stress, internal ribosome entry site (IRES) elements drive translation initiation using a cap-independent mechanism (2,3). IRES elements differ in nucleotide sequence, RNA secondary structure and trans-acting factors requirement (2,4,5). This complexity compromises the general understanding of the mechanism of internal initiation, leading to the view that IRES elements may promote translation initiation using a large variety of mechanisms (6–8).

RNA structure plays a fundamental role in IRES-dependent translation initiation (4,9–11), as well as in other processes guided by RNA regulatory elements (12–14). This is illustrated by the fact that compensatory substitutions in base-paired regions tend to conserve the secondary structure during RNA evolution (2,15–17).

Foot-and-mouth disease virus (FMDV) is a picornavirus characterized by a high genetic variability (18). This feature provides crucial information regarding the tolerance to accept nucleotide substitutions all along the viral genome, and particularly, within the highly structured untranslated regions (UTRs) located at each end of the genome (19). The IRES element located at the 5′-UTR of FMDV is organized in structural domains, termed 2–5 in 5′- to 3′-end, which appear to have a division of functions (20). Domains 2, 4 and 5 determine the interaction with RNA binding proteins and

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various translation initiation factors (eIFs), with the exception of eIF4E (21,22). Domains 4 and 5 are responsible for the recruitment of eIF4G, eIF4B, eIF3 and other IRES binding factors (23,24). Remarkably, domains 4 and 5 do not possess IRES activity by themselves (25), indicating that interaction with these factors is necessary but not sufficient for IRES function.

In contrast to the large number of factors interacting with the distal domains, the central one, termed domain 3, interacts with a reduced number of host factors, such as PCBP2, EBP1, hnRNPK and DAZ-1 (26). This result could be interpreted as the consequence of having a large proportion of its nucleobases hidden within a compact RNA 3D structure. In fact, secondary structure determination of the FMDV IRES by RNA probing (9,25) revealed that the apical region of domain 3 contains four GC-rich stems that join at a cruciform structure. At the base of this domain, a long stem interrupted by internal bulges serves as a bridge to upstream and downstream IRES domains. Mutational analysis has shown that conserved motifs in the RNA structure of domain 3 play a crucial role in IRES-dependent translation (20,27–29). However, the 3D structure of this domain is still unknown.

Picornavirus IRES elements are remarkably long regions compared to other viral IRES, including those of HCV and dicistroviruses (2,30,31). Determining the structural organization of long IRES elements is the first step to begin to understand the efficient mechanism of internal initiation driven by these elements. Their long size and flexibility is well suited to structural analysis by selective 2'-hydroxyl acylation analyzed by primer extension (SHAPE) (32). Although SHAPE probing does not reach the atomic level resolution of NMR or X-ray structural analysis, it has the advantage over the latter approaches of allowing the analysis of long RNA molecules (33) and hence, it sheds light on RNA organization of the entire RNA regulatory element. Recently, we have performed a combined study involving SHAPE probing and microarray-based oligonucleotide accessibility of the FMDV IRES element (34) confirming its modular organization in structural domains.

In this study, we sought to investigate the involvement in IRES function of RNA structural motifs within domain 3 possibly constrained by distant interactions. To this end, we took advantage of the large variability of FMDV RNA 3 possibly constrained by distant interactions. To this end, we took advantage of the large variability of FMDV RNA sequences. To this end, we took advantage of the large variability of FMDV RNA sequences. To this end, we took advantage of the large variability of FMDV RNA sequences.
**Random expectation of covariation.** In order to determine which pairs had an MI value significantly different from random expectation, we generated 300 randomized sets by vertical shuffling all the columns of the alignment. This maintains the proportion of nucleotides for each column, but shuffles the co-occurrences in each row. The distribution of the MI values in these randomized alignments was used to calculate the corresponding z-scores. Variable pairs (VV) were therefore classified as covariant, if z-score > 2.96, or independent, otherwise. Moreover, each variable pair (VV) was further classified as conservative or non-conservative, according to whether the fraction of changes that preserved the canonical base pairing was >50% or <50%, respectively.

**RNA structure modeling**

PDB RNA structure models were performed using Mc-Fold (www.major.iris.ca/MC-Pipeline) to obtain the secondary structure in bracket format, followed of 3D MC-SYM generator to produce the PDB RNA structure. RNA structure was visualized using Swiss-PdbViewer. The first PDB structure obtained with the MC-SYM generator using the most stable energy RNA structure according to Mc-Fold was used to depict the model of each IRES sequence.

**IRES activity assays**

Relative IRES activity was quantified as the expression of luciferase normalized to that of chloramphenicol acetyltransferase (CAT) from bicistronic mRNAs as described (36) in transfected BHK-21 monolayers. Experiments were performed on triplicate wells and each experiment was repeated at least three times.

**In vitro transcription**

Plasmids were linearized to generate domain 3 or the full-length IRES using SmaI or XhoI, respectively. Transcription was performed for 1h at 37°C using 1000–3000 U of highly purified T7 RNA polymerase in the presence of 10–15 μg of linearized DNA template, 40 mM Tris–HCl, 50 mM DTT, 0.5 mM rNTPs, as described (34).

**SHAPE analysis**

RNA (0.5 pmols) was treated with N-methylisatoic anhydride (NMIA) as described (34,37). For primer extension, equal amounts of NMIA-treated and untreated RNAs (10 μl) were incubated with 0.5 μl of the appropriate antisense 5'-end 32P-labeled primer (5'-CTACGAAGCAACA GTG, 5'-CCCGGGTGTGGGTACC, 5'-GGAATGGGGA TCCTCGAGCTCAGGGGTC). Primer extension was conducted in a final volume of 15 μl containing reverse transcriptase (RT) buffer (50 mM Tris–HCl, pH 8.3, 3 mM MgCl2, 75 mM KCl, 8 mM DTT) and 1 mM of each dNTP. The mix was heated at 52°C for 1 min, prior to addition of 100 U of Superscript III RT (Invitrogen) and incubation at 52°C for 30 min. cDNA products were fractionated in 6% acrylamide, 7 M urea gels, in parallel to a sequence obtained with the same primer. For SHAPE data processing, the intensities of RT-stops were quantified as described (34). Data from three independent assays were used to calculate the mean (±SD) SHAPE reactivity.

**Gel-shift assays**

For RNA–RNA interactions, the uniformly [z-32P]-CTP labeled GNRA hairpin RNA (nucleotides 160–196) (38) was incubated with increasing concentrations of unlabeled domain 3 RNAs (50–1000 nM) in 50 mM sodium cacodylate, pH 7.5, 300 mM KCl, 10 mM MgCl2 (35,39). RNA–RNA complexes were allowed to form for 90 min at 37°C and immediately analyzed by electrophoresis in native acrylamide gels supplemented with 2.5 mM MgCl2 as described (38).

**Microarray hybridization and data analysis**

DNA oligonucleotides complementary to the IRES region of FMDV were described in (34). Mutant RNAs encompassing domain 3 were fluorescently labeled with Alexa 647 using the Ulysis 647 kit (Invitrogen). Microarrays were prehybridized and hybridized as described (34). Data were retrieved using the Genepix pro 6.0 software. The differential capacity of transcripts for antisense oligonucleotide hybridization was measured in three independent experiments as described (34).

**RESULTS**

**Identification of an invariant apical stem within the IRES element**

The alignment of nucleotide sequences belonging to 183 FMDV RNA isolates deposited in databases readily indicated an extensive degree of sequence heterogeneity across the IRES element (Figure 1a). In addition to regions that accumulated a large number of substitutions, we observed others in which coupled nucleotide covariation led to compensatory changes (Table 1) that maintained the RNA secondary structure and, interestingly, specific nucleotide tracts that were less tolerant to substitutions (Figure 1b).

The region that accommodated the higher number of changes was mapped to the single-stranded region at the 3'-end of domain 5 (Figure 1a), although the polypyrimidine tract is conserved (Supplementary Table S1). Similarly, other nucleotide variations located in loops within the IRES secondary structure (Figure 1b) corresponded to conserved motifs, as illustrated by the GNRA and the RAAA motifs in the apical region of domain 3 as well as the pyrimidine-rich motif in the apical loop of domain 2.

Analysis of covariation between pairs of positions with MI values significantly different from random expectation (z-score > 2.96) (Table 1) readily demonstrated the presence of paired regions in the IRES element (Figure 1b) that matched stems according to RNA probing (25,34). Covariation between variable positions within domain 3 occurs more frequently than between these and variable positions of the other domains (Supplementary...
Figure 1. Sequence variability of the IRES element in FMDV RNA. (a) The total number of changes found in 183 aligned IRES sequences is plotted against the nucleotide position. Location of the residues conforming the apical stem–loop, including the variable R residue of the RAAA motif, is indicated by a rectangle. (b) Sequence heterogeneity maintains the secondary structure of the FMDV IRES. Invariant nucleotides in the IRES secondary structure are marked in bold. Covariant and conservative nucleotide changes are depicted by light and dark violet squares, respectively. Independent substitutions are marked by green squares. Nucleotide positions are denoted by dots 10 nt apart. The most relevant motifs referred to in the text are indicated. Distribution of the IRES in domains (2–5) is indicated below each panel.
Table S2), reinforcing the idea of a modular organization in the IRES element. In this regard, covariation pairs for which any of the positions had 3 or less changes were discarded, since these could be fortuitous or due to sequencing errors.

We anticipate that the invariant regions may have been subjected to selection pressure to keep their primary sequence, because of their involvement in RNA–protein interactions or in maintaining the correct 3D RNA structure. Indeed, we observed that invariant regions tend to accumulate around nucleotides previously identified as the target site of RNA binding proteins involved in IRES activity. Thus, the upper stem–loop of domain 2 (nucleotides 45–68) and the stem at the base of domain 4 (nucleotides 306–320 and 396–416) (Figure 1b) provide the binding site for PTB and eIF4G, respectively (40,41).

| Ref. bp | Absolute number of base pairs | MI | Z-score |
|---------|-------------------------------|----|---------|
| C41U72  | 166 AU, 10 GU, 7 GC           |    |         |
| C45G71  | 179 CG, 4 UG                  |    |         |
| U47G70  | 162 UG, 21 CG                 |    |         |
| U48A298 | 178 UA, 3 CA, 1 AA, 1 UG      |    |         |
| G49C297 | 169 GC, 13 GU, 1AU            |    |         |
| U51A296 | 76 UA, 62 CC, 22 CU, 16 CA, 3 AC, 1UG, 1 CG, 1 AU, 1 UC |    |         |
| U52A294 | 171 UA, 11 CG, 1 UG           |    |         |
| G50A292 | 128 AU, 44 GC, 8 GU, 3 AC     |    |         |
| U62A282 | 176 UA, 6 CG, 1 UG            |    |         |
| C105G281 | 179 CG, 4 UG                 |    |         |
| U105U279 | 125 GC, 56 AU, 2 AC         |    |         |
| U111U277 | 135 UG, 44 UA, 2 CG, 2 UC    |    |         |
| G112U269 | 114 GU, 42 AU, 13 GC, 13 AC, 1 UU |    |         |
| A126C257 | 86 AU, 42 AC, 34 GU, 21 GC   |    |         |
| C127G256 | 169 CG, 7 UA, 7 UG           |    |         |
| A128A255 | 165 AU, 16 GU, 1 UG, 1UU      |    |         |
| G129C254 | 164 GC, 7 AU, 6 CU, 6 GU     |    |         |
| U135A250 | 144 UA, 23 CG, 12 UG, 1 AG, 1 AA, 1 CA |    |         |
| G136C247 | 148 UA, 30UG, 4 CG, 1 CA     |    |         |
| G137C246 | 174 GC, 6 GU, 3 AU           |    |         |
| C138G245 | 141 GC, 36 UG, 6 UA          |    |         |
| U139A190 | 164 UG, 17 CG, 2 UG           |    |         |
| A142U233 | 81 GC, 80 AU, 13 UA, 5 AC, 3 GU, 1 GA  |    |         |
| C144G222 | 128 UA, 54 CG, 1 UG          |    |         |
| C146G193 | 178 CG, 5 UG                  |    |         |
| U154A191 | 115 UA, 64 CG, 2 UG, 2 CA     |    |         |
| U155A184 | 134 UA, 48 UG, 1 CG           |    |         |
| C160G218 | 147 UA, 33 CG, 2 CA, 1 UG     |    |         |
| U162G224 | 166 UG, 17 UA                 |    |         |
| C173G360 | 147 CG, 38 UG                 |    |         |
| C175G353 | 135 CG, 48 UG                 |    |         |
| A180U395 | 172 AU, 4 GU, 3 AG, 3 GC, 1-U |    |         |
| C182G394 | 171 CG, 9 UG, 2 UA, 1-A       |    |         |
| U187A393 | 140 UA, 41 UG, 2 CG           |    |         |
| G188C392 | 123 GC, 54 GU, 6 AU           |    |         |
| G189U391 | 132 GC, 41 GU, 7 AU, 3 AC     |    |         |
| G192G390 | 175 GC, 8 GU                  |    |         |
| G195C388 | 119 GC, 43 AU, 21 GU          |    |         |
| C196G387 | 179 CG, 2 GC, 1 UG, 1-G       |    |         |
| G203C439 | 175 GC, 6 AU, 2 GU            |    |         |
| U212A437 | 127 UA, 54 CG, 2 UG           |    |         |

Ref. bp indicates the sequence of the base pair in the reference IRES sequence. MI indicates the probability of finding simultaneously a particular combination of nucleotide at a given position. Z-score > 2.96 indicates the pairs with MI significantly different from random changes.

The apical region of domain 3 contains three invariant regions (Figure 1b), encompassing nucleotides 140–150 (the 140 bulge), 195–205 (the apical stem–loop, that includes the apical stem and the RAAA motif), and 229–243 (the C-rich bulge). While the latter is a candidate sequence to interact with poly(rC) binding proteins and Ebp1 (26,42,43), proteins recognizing the other two invariant regions have not been identified. Furthermore, RNA probing carried out with substitution mutants of the conserved GNRA motif evidenced apical stem–loop reorganization leading to the disruption of the stem, despite the mutation being located in a distant sequence (34,38). These data, together with results of the covariation analysis, led us to focus our attention to the invariant sequence of the apical stem of domain 3 as a region contributing to an RNA structure crucial for IRES function.
The conserved apical stem uncovers a determinant of IRES activity

To determine the biological relevance of the invariant apical stem of domain 3 for IRES activity, we generated a set of mutants aimed at modifying both, the primary sequence and the secondary structure of this region (Figure 2a). The S1 mutant carrying a double substitution G195G196 to AA, designed to disrupt the canonical G:C base pairs of the wt IRES, led to a severe loss of IRES activity (1.8% with respect to the wt IRES) (Figure 2b). Similarly, a double substitution on the opposite arm of the stem, C204C205 to UU (mutant S2) also abrogated IRES activity (1.4%), suggesting that G:U base pairs were not formed or, conversely, they were not sufficient to stabilize this short stem.

To confirm that disruption of the G:C base pairs of this stem lead to IRES inactivity, we generated a mutant carrying the double substitution C204C205 to GG (S3). This mutant turned out to be severely defective (0.7%)
calculated from the reactivity of each nucleotide, relative to the most reactive nucleotide (set to 100%). The resulting color-coded profiles (Figure 3) revealed gross modifications of SHAPE reactivity in domain 3 of the mutants S1–S4 with respect to that of the wt RNA. This was particularly evident in residues 150–250 encompassing the apical region, which showed increased SHAPE reactivity. However, SHAPE reactivity was only slightly modified in domains 2, 4 or 5. Furthermore, changes in the SHAPE reactivity of domain 3 accumulated in similar positions when the transcripts encompassing either the entire IRES or the domain 3 alone (data not shown) were analyzed, suggesting that the RNA reorganization mainly affected the central domain.

The differences in SHAPE reactivity of the wt RNA (Supplementary Figure S1) with respect to each IRES mutant are depicted in Figure 4. Specifically concerning the S1 mutant RNA, there was a remarkable increase in SHAPE reactivity in residues 198–207, supporting the notion that nucleotides corresponding to the apical stem were not engaged in base pairs (Figure 4). Concomitant with this increase, a strong decrease of reactivity was observed in nucleotides 169–171 and 181, revealing a protection of the 170 bulge and the GNRA motif, respectively. Therefore, these results suggest a functional interaction between residues located in two stem–loops of the apical region, with a distinctive feature: an increase in SHAPE reactivity of the apical stem–loop was accompanied by a decrease in reactivity of nucleotides belonging to the GNRA hairpin, including the 170 bulge. These results are also in agreement with the RNA structure model of S1 IRES (Figure 2c) that not only depicts the mutated nucleotides 195–196, and residues belonging to 170 bulge are in the vicinity of nucleotides C204-G208.

We conclude that the invariant stem of the apical stem–loop plays a crucial role for IRES activity, and that it has a strong preference for G:C base pairs. Their substitution by C:G retained activity although this IRES mutant did not reach the same activity levels than the wt. These results are in strong agreement with the lack of genetic variability found in the apical stem–loop within the IRES sequence alignment.

RNA structural analysis reveals the specific need for G:C base-pairs for IRES function

Next, to establish a direct relation between RNA structure and activity of IRES mutants carrying mutations in the apical stem, we performed structural analysis by SHAPE, which allows the study of long, flexible RNA molecules in solution (32,33). RT extension products obtained with primers spaced along the RNA sequence allowed overlapping readings of the entire IRES region (34). For this, cDNA products derived from the untreated and NMIA-treated RNAs were resolved in denaturing acrylamide–urea gels in parallel to a sequence ladder obtained with the same primer. The intensity of each RT-stop band was normalized to that of the full-length product in the corresponding gel lane, and the background values of the untreated RNA were subtracted from the respective RT-stop intensity yielded by NMIA-treated RNA. In all cases, the mean of three independent assays was calculated from the reactivity of each nucleotide, relative
S3) impaired IRES activity. Conversely, compensatory mutations that preserved the secondary RNA structure (mutant S4) only partially restored IRES activity, since the wt IRES carrying the GC stem performed better than that of the S4 mutant IRES.

We conclude that the invariant bases that compose the apical stem of the IRES play a crucial role in determining the RNA structural organization affecting the adjacent GNRA hairpin that, in turn, has also a strong influence in the ability of this IRES element to drive internal initiation of translation.

![SHAPE reactivity of apical stem IRES mutants](image)

Figure 3. SHAPE reactivity of apical stem IRES mutants. Values of SHAPE reactivity at each individual nucleotide position correspond to the mean reactivity (±SD) of three independent assays. RNAs, treated with NMIA or untreated, were subjected to primer extension analysis conducted with 5'-end labeled primers. The intensity of each band was normalized to the full-length cDNA product detected in the corresponding gel lane after subtraction of the corresponding background RT-stop signal in the untreated RNA. Nucleotide positions are indicated on the x-axis. The SHAPE reactivity is depicted using color-coded bars. Position of the apical region is indicated by a broken line rectangle.

Long-distance interactions within domain 3 and local RNA accessibility depends on the sequence composition of the apical stem

We have previously identified long-range RNA–RNA interactions within domain 3, which are sequence specific, as well as Mg$^{2+}$ and RNA concentration dependent (38). The changes in RNA structure observed in the GNRA hairpin and the apical stem of S1–S4 mutants strongly suggested that the mutations introduced in the apical stem could generate different conformations of
domain 3, profoundly modifying long-range RNA–RNA interactions. To test the hypothesis that distant RNA–RNA interactions could be affected in the S1–S4 mutants, we determined by gel mobility-shift assays the capacity of a labeled transcript encompassing the GNRA hairpin (corresponding to nucleotides 160–196) to interact with unlabelled transcripts of domain 3 bearing either the wt or the mutated S1–S4 sequences (Figure 5a). The mutant RNAs responded in different manners (Figure 5b), with $K_d$ ranging between 112.3 nM (wt), 271.7 (S1), 234.3 (S2), 279.0 (S3) and 173.1 (S4). Strikingly, S1 RNA showed a reduced capacity to

Figure 4. Impact of the apical stem sequence composition on SHAPE reactivity differences. SHAPE difference plots of each mutant domain 3 relative to the wt RNA. Nucleotides with absolute changes in SHAPE reactivity >60% are depicted in black, while those between 30 and 60% are marked in dark grey. The secondary structure model of each IRES mutant, with nucleotides colored as in Figure 3 to reflect their mean SHAPE reactivity, is shown on the right. The reference mean SHAPE reactivity of the wt RNA is shown in Supplementary Figure S1.
establish long-range RNA interaction with the GNRA hairpin, although, at the same time, it has acquired the property to form two retarded complexes clearly distinguishable in long exposures (Figure 5a). This result may imply the presence of at least two conformations of this mutant RNA, which very weakly interact with the GNRA hairpin. S2 and S3 mutant RNAs also showed a reduced capacity to bind the labeled GNRA hairpin probe all along the concentration range tested. In turn, the S4 RNA that bears the compensatory mutation evidenced an intermediate capacity to interact with the GNRA hairpin probe (Figure 5b). Consistent with IRES activity and RNA structural analysis, the binding efficiency of the S4 domain 3 was greater than that of the other mutants, achieving values similar to the wt RNA only at the highest concentration (1000 nM) used in the assay.

The results observed with S1 mutant suggested that this IRES region could adopt different conformation depending on the sequence of the apical stem. Thus, to gain additional information about the local RNA structure of the IRES mutants, we measured the accessibility of domain 3 RNAs to a panel of customized overlapping antisense DNA oligonucleotides covering the entire IRES sequence printed on microarrays (34). RNA hybridization was conducted using fluorescent-labeled transcripts carrying the indicated mutations in the apical stem under the same ionic conditions and temperature used for SHAPE probing. The hybridization signal displayed by each mutant RNA revealed changes in RNA accessibility (Figure 6). Interestingly, S1 RNA exhibited an overall higher accessibility in the microarray-based assay than the wt RNA, demonstrating a less tight RNA structure that particularly affected residues complementary to oligonucleotides 134–148, 176–183 and 232–253. In contrast, changes in RNA accessibility of the S2, S3 and S4 mutants were concentrated in the region complementary to oligonucleotide 253. A decrease in accessibility to oligonucleotides 190 and 197, complementary to nucleotides 190–211, can be explained by the presence of two substitutions in the RNA sequence of these IRES mutants. These results strongly suggest that domain 3 consists of

![Figure 5. Mobility-shift binding assays of domain 3 RNAs with the GNRA hairpin. (a) Representative examples of gel-shift assays for each construct, wt and S1 to S4 mutant RNAs. Gel-shift assays were carried out using 32P-labeled GNRA hairpin (corresponding to nucleotides 160–196 of domain 3) (50 nM) as probe and the wild-type or mutant unlabeled domain 3 RNAs (0–1000 nM). The position of the retarded complex is depicted by an arrow. Note that a longer exposure is shown for S1 RNA to allow the detection of two weak retarded complexes. (b) The percentage of retarded probe calculated from duplicate assays was plotted against the unlabeled RNA concentration. Error bars, SD.](https://academic.oup.com/nar/article-abstract/39/19/8572/1177208)
a highly interactive structure in which its apical stem and the GNRA stem–loop are in close structural and functional connection.

In summary, our results reinforce the notion that the nucleotide composition of the apical stem performs a crucial role in maintaining the correct RNA conformation of domain 3. Using complementary approaches such as functional analysis, SHAPE reactivity, gel-shift and microarray-based RNA accessibility, we have established the relevance of the RNA structural organization for the function of the invariant stem of domain 3 in IRES-mediated translation.

**Figure 6.** Hybridization of IRES transcripts to antisense oligonucleotides printed on microarrays. Hybridization signal of the fluorescent-labeled domain 3 transcripts plotted against each oligonucleotide (mean ± SD) averaged from three independent assays. The array contains 14-nt long oligonucleotides (termed 1–449, by the position on the IRES sequence complementary to the 3'-end of the primer), overlapping each 7 nt within domain 3 (34). Changes in accessibility of mutant IRES to oligonucleotides 190 (S1, S4) and 197 (S2, S3, S4), depicted with white bars, are likely due to mismatches in the sequence of these RNAs with the sequence printed in the microarray. Position of the apical region is indicated by a broken line rectangle. Differences in accessibility are depicted in black (>0.4), grey (range 0.4–0.25) and light grey (<0.25).
The modular organization of picornavirus IRES is consistent with a distribution of functions among the different domains. The central domain of the IRES, termed domain 3, is a self-folding region since it adopts a very similar RNA structure organization when it is expressed alone or in the context of the entire IRES (34). Moreover, the local RNA structure of this domain plays a critical role during IRES-dependent translation (9,38). SHAPE reactivity data provided evidence for the existence of stem–loops whose structural conformation depends on distant interactions within this domain, involving residues of the GNRA motif (34). These results point to the existence of a structural element in the apical region of domain 3 that performs an essential role during internal initiation. In support of a general implication of this structural feature to picornavirus IRES performance, not only FMDV but also encephalomyocarditis virus (EMCV) IRES-dependent translation is impaired by nucleotide substitutions in the conserved GNRA motif (27,28). Thus, the RNA structure adopted by the apical region of domain 3 could constitute a signature of type II IRES elements, typically found in picornavirus genomes.

The genetic variability of the FMDV IRES analyzed in 183 isolates indicated certain tolerance to accept changes in the sequence of domain 3 (Figure 1b). Despite this overall variability, the GNRA and the RAAA motifs are fully conserved (Supplementary Table S1). We report here that the G:C base pairs that hold these motifs are also invariant. This suggested to us that they perform a key role in maintaining the functional IRES structure. Furthermore, the sequence of the internal 170 bulge within the GNRA hairpin tolerates substitutions, but not insertions or deletions (Supplementary Table S1).

We have shown that the G:C base pairs that conform the apical stem–loop have a primary relevance for IRES activity. Importantly, disruption of this stem occurred concomitantly with a structural rearrangement of the apical region, affecting two adjacent stem–loops (Figures 2c and 4). The results of SHAPE reactivity of wild-type and mutant IRES RNAs strongly support a direct effect of residues G195–G196 and C204–C205 in governing the RNA conformation of the active IRES element. Specifically, SHAPE reactivity, gel mobility-shift and RNA accessibility displayed by the S1 mutant (where the G195G196 were substituted by AA) supported a change in RNA conformation of the apical region affecting distant regions in the secondary structure of domain 3. The results derived from the study of S2 (CC substituted by UU) and S3 (CC changed to GG) RNAs demonstrated that the disruption of the apical stem induced the reorganization of the entire apical region of domain 3, including the GNRA hairpin. Unexpectedly, although the substitutions in S4 RNA partially restored IRES activity, the organization of the apical region was modified, as indicated by differences in SHAPE reactivity of the GNRA stem–loop and the apical stem–loop. Therefore, this result further confirmed the biological relevance of the GG:CC composition of the apical stem, in agreement with the lack of tolerance to accept sequence changes.

A common feature observed in the IRES mutants studied here is that an increase in reactivity of nucleotides belonging to the apical stem–loop (195–205) is coupled to a decrease in reactivity of nucleotides 169–171 and 181, that belong to the 170 bulge and the GNRA motif, respectively. This is specifically consistent with the reorganization predicted in the S1 RNA structure (Figure 2c). The implications of our mutational analysis suggest a cross-talk between distant nucleotides of domain 3, located at the apical stem and the GNRA hairpin, that share the property of being conserved at the structural level between picornavirus IRES classified as type II (2,43). Specifically, the observation that S1 mutant RNA severely affected the capacity to form distant interactions (Figure 5a), and that S4 mutant only partially restored IRES activity (Figure 2b) strongly suggests that the base pair composition of this stem determines the architecture of this IRES region. This line of evidence prompted us to consider the possibility of whether this reorganization is due to the disruption of distant interactions.

Changes in the efficiency and pattern of long-distant interactions within domain 3 have been demonstrated by gel-shift analysis of the GNRA hairpin with specific mutant IRES transcripts (Figure 5a and b) (38) and by RNA structure probing of mutants in the conserved GNRA motif (9,34). Previous studies have shown that the GNRA motif of FMDV, EMCV and poliovirus IRES adopts a tetraloop conformation at the tip of a stem–loop (9,44,45). GNRA tetraloops are frequently involved in tertiary interactions (46,47). In particular, it has been shown that GYRA tetraloops interact with GG:CC receptors when R corresponds to A (as in the GUAA sequence of the wt IRES sequence used here), whereas it can accept CU:AG receptors when the R is a G (48). The GNRA motif of the FMDV IRES shows a strong preference for GUAA, but it tolerates sequence variation to GUGA, GCAA, GAGA and GCQA (Supplementary Table S1) (34). Thus, covariation analysis of the IRES region could give us a hint about the location of the putative receptor. However, we could not detect signals of a GNRA receptor based solely on the genetic variability analysis. Although it can be argued that this could be due to insufficient number of analyzed samples, we cannot rule out the possibility of alternative binding sites or the existence of different RNA conformations as it has been observed in various RNAs (12,49).

The lack of variability of the apical stem, supported by the covariation observed in the adjacent stems, pointed toward its relevant contribution to IRES activity. Our results of mutational analysis of this invariant stem revealed a better performance of G:C than C:G base pairs, demonstrating that it is not only the secondary structure of this region, but its 3D RNA conformation is what determines efficient IRES activity. Furthermore, changes in IRES activity were accompanied by a structural reorganization, as revealed by SHAPE reactivity and gel mobility-shift interactions. A region effectively modified, in addition to the mutated bases, includes the 170 bulge, the GNRA motif, the stem 209, loop 140 and 238. Interestingly, genetic variability of the 170 bulge showed that whereas its sequence accepts certain
nucleotide substitution, insertions or deletions were not observed (Supplementary Table S1). It is worth noting that the overall RNA organization of the 170 bulge within the GNRA hairpin may resemble that of the HCV IRES domain II (50).

Importantly, we have found a modification of RNA structure affecting multiple nucleotides of the apical region in all mutants analyzed here. Additionally, we have shown that conformational changes in RNA structure play a pivotal role in IRES function. Taken together, our results demonstrate that the active conformation of domain 3 consists of an interactive structure constrained by distant interactions, in which two invariant G:C base pairs within the apical stem–loop play a key role in maintaining the correct RNA structure, likely interacting with the GNRA hairpin.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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REFERENCES

1. Sonenberg,N. and Hinnebusch,A.G. (2009) Regulation of translation initiation in eukaryotes: mechanisms and biological targets. Cell, 136, 731–745.
2. Martinez-Salas,E. (2008) The impact of RNA structure on picornavirus IRES activity. Trends Microbiol., 16, 230–237.
3. Spriggs,K.A., Bushell,M. and Willis,A.E. (2010) Translational regulation of gene expression during conditions of cell stress. Mol. Cell, 40, 228–237.
4. Costantino,D.A., Pang斯顿,J.S., Rambo,R.P. and Kieft,J.S. (2008) tRNA-mRNA mimicry drives translation initiation from a viral IRES. Nat. Struct. Mol. Biol., 15, 57–64.
5. Vallejos,M., Deforges,J., Plank,T.D., Leiteler,A., Ramdohr,P., Abraham,C.G., Valiente-Echeverria,F., Kieft,J.S., Sargueil,B. and Lopez-Lasra,M. (2011) Activity of the human immunodeficiency virus type 1 cell cycle-dependent internal ribosomal entry site is modulated by IRES trans-acting factors. Nucleic Acids Res., April 10; epub ahead of print; doi:10.1093/nar/gkr189.
6. Schuler,M., Connell,S.R., Lescoute,A., Giesebrecht,J., Dubrowski,M., Schroer,B., Mielcicki,T., Penczek,P.A., Westhof,E. and Spahn,C.M. (2006) Structure of the ribosome-bound cricket paralysis virus IRES RNA. Nat. Struct. Mol. Biol., 13, 1092–1096.
7. Terenin,I.M., Dmitriev,S.E., Andreev,D.E. and Shatsky,I.N. (2008) Eukaryotic translation initiation machinery can operate in a bacterial-like mode without eIF2. Nat. Struct. Mol. Biol., 15, 836–841.
8. Locker,N., Chamond,N. and Sargueil,B. (2011) A conserved structure within the HIV gag open reading frame that controls translation initiation directly recruits the 40S subunit and eIF3. Nucleic Acids Res., 39, 2367–2377.
9. Fernandez-Miragall,O. and Martinez-Salas,E. (2003) Structural organization of a viral IRES depends on the integrity of the GNRA motif. RNA, 9, 1333–1344.
10. Easton,L.E., Locker,N. and Lukavsky,P.J. (2009) Conserved functional domains and a novel tertiary interaction near the pseudoknot drive translational activity of hepatitis C virus and hepatitis C virus internal ribosome entry sites. Nucleic Acids Res., 37, 5537–5549.
11. Morris,M.J., Negishi,Y., Pazzini,C., Schonhoft,J.D. and Basu,S. (2010) An RNA G-quadruplex is essential for cap-independent translation initiation in human VEGF IRES. J. Am. Chem. Soc., 132, 17831–17839.
12. Chaban,S. and Woodson,S.A. (2008) Tertiary interactions determine the accuracy of RNA folding. J. Am. Chem. Soc., 130, 1296–1303.
13. Zuo,X., Wang,J., Yu,P., Eyster,D., Xu,H., Starich,M.R., Tiede,D.M., Simon,A.E., Kasprzak,W., Schwieters,C.D. et al. (2010) Solution structure of the cap-independent translational enhancer and ribosome-binding element in the 3′ UTR of turnip crinkle virus. Proc. Natl Acad. Sci. USA, 107, 1385–1390.
14. McPheeters,D.S., Cremona,N., Sander,S., Chen,H.M., Averbeck,N., Leatherwood,J. and Wise,J.A. (2009) A complex gene regulatory mechanism that operates at the nexus of multiple RNA processing decisions. Nat. Struct. Mol. Biol., 16, 255–264.
15. Gorodkin,J., Hofacker,I.L., Torarinsson,E., Yao,Z., Havgard,J.H. and Ruzzo,W.L. (2010) De novo prediction of structured RNAs from genomic sequences. Trends Biotechnol., 28, 9–19.
16. Berkhourt,B. and Das,A.T. (2009) Virus evolution as a tool to study HIV-1 biology. Methods Mol. Biol., 485, 436–451.
17. Smith,D.B., Mellor,J., Jarvis,L.M., Davidson,F., Kolberg,J., Urdea,M., Yap,P.L. and Simmonds,P. (1995) Variation of the hepatitis C virus 5′ non-coding region: implications for secondary structure, virus detection and typing. The International HCV Collaborative Study Group. J. Gen. Virol., 76 (Pt 7), 1749–1761.
18. Domingo,E., Escarmis,C., Martinez,M.A., Martinez-Salas,E. and Mateu,M.G. (1992) Foot-and-mouth disease virus populations are quasispecies. Curr. Top Microbiol. Immunol., 176, 33–47.
19. Serrano,P., Pulido,M.R., Saiz,M. and Martinez-Salas,E. (2006) The 3′ end of the foot-and-mouth disease virus genome establishes two distinct long-range RNA-RNA interactions with the 5′ end region. J. Gen. Virol., 87, 3013–3022.
20. Serrano,P., Ramajo,J. and Martinez-Salas,E. (2009) Rescue of internal initiation of translation by RNA complementation provides evidence for a distribution of functions between individual IRES domains. Virology, 388, 221–229.
21. Martinez-Salas,E., Pacheco,A., Serrano,P. and Fernandez,N. (2008) New insights into internal ribosome entry site elements relevant for viral gene expression. J. Gen. Virol., 89, 611–626.
22. Andreev,D.E., Fernandez-Miragall,O., Ramajo,J., Dmitriev,S.E., Terenin,I.M., Martinez-Salas,E. and Shatsky,I.N. (2007) Differential factor requirement to assemble translation initiation complexes at the alternative start codons of foot-and-mouth disease virus RNA. RNA, 13, 1366–1374.
23. Lopez de Quinto,S., Lafuente,E. and Martinez-Salas,E. (2001) IRES interaction with translation initiation factors: functional characterization of novel RNA contacts with eIF3, eIF4B, and eIF4G1. RNA, 7, 1213–1226.
24. Pacheco,A., Lopez de Quinto,S., Ramajo,J., Fernandez,N. and Martinez-Salas,E. (2009) A novel role for Gemin5 in mRNA translation. Nucleic Acids Res., 37, 582–590.

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25. Fernandez-Miragall, O., Lopez de Quinto, S. and Martinez-Salas, E. (2009) Relevance of RNA structure for the activity of picornavirus IRES elements. *Viruses*, 139, 172–182.

26. Pacheco, A., Reigadas, S. and Martinez-Salas, E. (2008) Riboproteomic analysis of polypeptides interacting with the internal ribosome-entry site element of foot-and-mouth disease viral RNA. *Proteomics*, 8, 4782–4790.

27. Lopez de Quinto, S. and Martinez-Salas, E. (1997) Conserved structural motifs located in distal loops of aphthovirus internal ribosome entry site domain 3 are required for internal initiation of translation. *J. Virol.*, 71, 4171–4175.

28. Robertson, M.E., Seamos, R.A. and Belsham, G.J. (1999) A selection system for functional internal ribosome entry site (IRES) elements: analysis of the requirement for a conserved GNRA tetraloop in the encephalomyocarditis virus IRES. *RNA*, 5, 1167–1179.

29. Martinez-Salas, E., Regalado, M.P. and Domingo, E. (1996) Identification of an essential region for internal initiation of translation in the aphthovirus internal ribosome entry site and implications for viral evolution. *J. Virol.*, 70, 992–998.

30. Lukavsky, P.J. (2009) Structure and function of HCV IRES domains. *Viruses*, 139, 166–171.

31. Filbin, M.E. and Kiefer, J.S. (2009) Toward a structural understanding of IRES RNA function. *Curr. Opin. Struct. Biol.*, 19, 267–276.

32. Merino, E.J., Wilkinson, K.A., Coughlan, J.L. and Weeks, K.M. (2005) RNA structure analysis at single nucleotide resolution by selective 2'-hydroxyl acylation and primer extension (SHAPE). *J. Am. Chem. Soc.*, 127, 4223–4231.

33. Watts, J.M., Dang, K.K., Görecki, R.J., Leonard, C.W., Bess, J.W. Jr, Swanstrom, R., Burch, C.L. and Weeks, K.M. (2009) Architecture and secondary structure of an entire HIV-1 RNA genome. *Nature*, 460, 711–716.

34. Fernandez-Miragall, O., Garcia-Sacristan, A., Ramajo, J., Briones, C. and Martinez-Salas, E. (2011) Structural analysis provides insights into the modular organization of picornavirus IRES. *Virology*, 409, 251–261.

35. Ramos, R. and Martinez-Salas, E. (1999) Long-range RNA interactions between structural domains of the aphthovirus internal ribosome entry site (IRES). *RNA*, 5, 1374–1383.

36. Martinez-Salas, E., Saiz, J.C., Davila, M., Belsham, G.J. and Domingo, E. (1993) A single nucleotide substitution in the internal ribosome entry site of foot-and-mouth disease virus leads to enhanced cap-independent translation in vivo. *J. Virol.*, 67, 3748–3755.

37. Wilkinson, K.A., Merino, E.J. and Weeks, K.M. (2006) Selective 2'-hydroxyl acylation analyzed by primer extension (SHAPE): quantitative RNA structure analysis at single nucleotide resolution. *Nat. Protoc.*, 1, 1610–1616.

38. Fernandez-Miragall, O., Ramos, R., Ramajo, J. and Martinez-Salas, E. (2006) Evidence of reciprocal tertiary interactions between conserved motifs involved in organizing RNA structure essential for internal initiation of translation. *RNA*, 12, 223–234.

39. Paullart, J.C., Skripkin, E., Ehresmann, B., Ehresmann, C. and Marquet, R. (1996) A loop-loop “kissing” complex is the essential part of the dimer linkage of genomic HIV-1 RNA. *Proc. Natl Acad. Sci. USA*, 93, 5572–5577.

40. Luz, N. and Beck, E. (1991) Interaction of a cellular 57-kilodalton protein with the internal translation initiation site of foot-and-mouth disease virus. *J. Virol.*, 65, 6486–6494.

41. Lopez de Quinto, S. and Martinez-Salas, E. (2000) Interaction of the eIF4G initiation factor with the aphthovirus IRES is essential for internal translation initiation in vivo. *RNA*, 6, 1380–1392.

42. Walter, B.L., Nguyen, J.H., Ehrenfeld, E. and Semler, B.L. (1999) Differential utilization of poly(rC) binding protein 2 in translation directed by picornavirus IRES elements. *RNA*, 5, 1570–1585.

43. Yu, Y., Abaeva, I.S., Marintchev, A., Pestova, T.V. and Hellen, C.U. (2011) Common conformational changes induced in type 2 picornavirus IRESs by cognate trans-acting factors. *Nucleic Acids Res.*, 39, 4851–4865.

44. Phelan, M., Banks, R.J., Conn, G. and Ramesh, V. (2004) NMR studies of the structure and Mg2+ binding properties of a conserved RNA motif of EMCV picornavirus IRES element. *Nucleic Acids Res.*, 32, 4715–4724.

45. Du, Z., Ulyanov, N.B., Yu, J., Andino, R. and James, T.L. (2004) NMR structures of loop B RNAs from the stem-loop IV domain of the enterovirus internal ribosome entry site: a single C to U substitution drastically changes the shape and flexibility of RNA. *Biochemistry*, 43, 5757–5771.

46. Correll, C.C. and Swinger, K. (2003) Common and distinctive features of GNRA tetraloops based on a GUAA tetraloop structure at 1.4 Å resolution. *RNA*, 9, 355–363.

47. Geary, C., Baudrey, S. and Jaeger, L. (2008) Comprehensive features of natural and in vitro selected GNRA tetraloop-binding receptors. *Nucleic Acids Res.*, 36, 1138–1152.

48. Geary, C., Chowros, A. and Jaeger, L. (2011) Promoting RNA helical stacking via A-minor junctions. *Nucleic Acids Res.*, 39, 1066–1080.

49. Gulyaev, A.P., Franch, T. and Gerdes, K. (2000) Coupled nucleotide covariations reveal dynamic RNA interaction patterns. *RNA*, 6, 1483–1491.

50. Lukavsky, P.J., Kim, I., Otto, G.A. and Puglisi, J.D. (2003) Structure of HCV IRES domain II determined by NMR. *Nat. Struct. Biol.*, 10, 1033–1038.