Secondary Structural Elements of the HCV X-region Involved in Viral Replication

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Abstract

Background and Aims: The noncoding regions in the 3'-untranslated region (UTR) of the hepatitis C virus (HCV) genome contain secondary structures that are important for replication. The aim of this study was to identify detailed conformational elements of the X-region involved in HCV replication. Methods: Ribonucleic acid (RNA) structural analogs X94, X12, and X12c were constructed to have identical conformation but 94%, 12%, and 0% sequence identity, respectively, to the X-region of HCV genotype 2a. Effects of structural analogs on replication of HCV genotypes 1b and 2a HCV RNA were studied by quantitative reverse transcriptase polymerase chain reaction. Results: In replicon BB7 cells, a constitutive replication model, HCV RNA levels decreased to 55%, 52%, 53%, and 54% after transfection with expression plasmids generating RNA structural analogs 5B-46, X-94, X-12, and X-12c, respectively (p<0.001 for all). In an HCV genotype 2a infection model, RNA analogs 5B-46, X-94, and X-12 in hepatic cells inhibited replication to 11%, 9%, and 12%, respectively. Because the X-12 analog was only 12% identical to the corresponding sequence of HCV genotype 2a, the sequence per se, or antisense effects were unlikely to be involved. Conclusions: The data suggest that conformation of secondary structures in 3'-UTR of HCV RNA genome is required for HCV replication. Stable expression of RNA analogs predicted to have identical stem-loop structures might inhibit HCV infection of hepatocytes in liver and may represent a novel approach to design anti-HCV agents.

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Keywords: HCV RNA genome; HCV X-region; Hepatitis C virus; Infection; Japanese Fulminant Hepatitis Virus; RNA secondary structure.

Abbreviations: 3' UTR, 3'-untranslated regions; cDNA, complementary deoxyribonucleic acid; CREs, cis-acting replication elements; CMV, cytomegalovirus; DMEM, Dulbecco's Modified Eagle Medium; FBS, fetal bovine serum; HCV, hepatitis C virus; JFH-1, Japanese Fulminant Hepatitis Virus-1; LDHA, lactate dehydrogenase A; MTT, 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide; NS, nonstructural; nt, nucleotide; PBS, phosphate buffered saline; qRT-PCR, quantitative reverse transcriptase polymerase chain reaction; RNA, ribonucleic acid; UTR, untranslated region.

Received: 02 February 2015; Revised: 26 February 2015; Accepted: 01 March 2015

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Introduction

Hepatitis C virus (HCV) is a ribonucleic acid (RNA) virus that causes chronic hepatitis and liver failure, worldwide. It consists of six different genotypes that are differentially distributed geographically. Success of treatment varies greatly depending on the genotype. The genome contains cis-acting replication elements (CREs) that are critical for HCV RNA replication and translation. RNA structural elements present in 5'- and 3'-untranslated regions (3'UTR) of the HCV genome interact with viral and cellular proteins to initiate and facilitate the replication and translation processes. In our previous studies, we showed that RNA secondary structure of the nonstructural (NS)5B coding region of the HCV genome was required for HCV RNA replication, and hence viral particle production. It has been shown that the X region in the 3'-UTR of the HCV RNA genome contains a highly conserved sequence. The latter have also been found to form stable secondary stem-loop structures that require physical contact between its RNA-dependent RNA polymerase for HCV replication. We hypothesized that RNA structural analogs resembling secondary stem-loop structure of the X region could be created that can compete with natural HCV genomic structure for binding to proteins and inhibit HCV replication. The aim of this study was to introduce RNA structural analogs resembling these CREs into human liver cells and to identify secondary structural elements of the HCV X-region involved in HCV replication and infection.

Materials and methods

Cell culture

In order to identify important secondary structural elements, the effects of HCV structural analogs on HCV replication in two model cell lines were studied: a constitutive replication model and an infection model. In addition, to evaluate whether differences in viral genotype could affect the possible interactions of structural analogs, viruses representing two different genotypes 1 and 2, were studied.

A genotype 1b BB7 constitutive replication system

Replicon cells, BB7, a cell culture system containing the HCV genotype 1b genome, were obtained from Apath (St. Louis, USA). Cells were maintained in Dulbecco's Modified Eagle Medium (DMEM) supplemented with antibiotic/antimycotic solution (Invitrogen, USA), 10% fetal bovine serum (FBS) and 0.5 mg/mL G418.
A Japanese Fulminant Hepatitis Virus-1 (JFH-1) HCV infection system

For JFH-1 HCV genotype 2a studies, Huh7.5 cells (human hepatoma cell line) (obtained from Dr. Charles Rice, Rockefeller University, NY, USA) were maintained in DMEM supplemented with antibiotic/antimycotic solution and 10% FBS. JFH-1 complementary deoxyribonucleic acid (cDNA), an HCV genotype 2a strain, from Dr. Takaji Wakita (National Institute of Infectious Diseases, Tokyo, Japan)\textsuperscript{22-24} was used to produce infectious HCV viral particles in Huh7.5.\textsuperscript{25,26} To make JFH-1 HCV stocks for infection, Huh 7.5 cells were infected with JFH-1 HCV. Media were collected 5 days post-infection and centrifuged at 1000 x g for 20 min to remove debris. JFH-1 HCV in media was concentrated with a centrifugal device (100K NMWL, Amicon ultra, Millipore, USA). Stocks were frozen at 
\( -80\, ^{\circ}\text{C} \) until further use.

Infectivity levels of viral stocks were checked by quantitation of JFH-1 HCV RNA in media 48 h post-infection of Huh7.5 cells with HCV genotype 2a specific primers (Table 1) using real time reverse transcriptase polymerase chain reaction (RT-PCR). The PCR conditions were: one cycle of 2 min at 50 \( ^{\circ}\text{C} \) and 10 min at 95 \( ^{\circ}\text{C} \) followed by 40 cycles of 15 sec at 95 \( ^{\circ}\text{C} \) and 1 min at 60 \( ^{\circ}\text{C} \).

Specificity of all designed primers was determined with PCR amplification and sequencing of amplified product (data not shown). Melt curve analysis was performed following each RT-PCR to identify the presence of primer dimers and analyze the specificity of the reaction.

RNA structural analogs

RNA structural analog X-94 was designed to be 100% identical to the (+) strand of X-region (9508-9605 nucleotides (nt)) on the 3' end of the HCV genotype 1b (94% identical to genotype 2a) genome (Fig. 1A). To determine whether stem-loop structures versus specific sequences of HCV of RNA were most important in HCV replication, base pair changes, described below, were made in stems (analog X-12) and loops (analog X-12c) of analog X-94 (Fig. 1B and 1D). Several software products are available for prediction of secondary structures of RNA based on thermodynamic parameters. Because of its high reliability and reproducibility, Mfold ver 3.2 was used in the current study.\textsuperscript{27-29} RNA structural analogs X-12 and X-12c were predicted to adopt stem-loop structure identical to analog X-94 (Fig. 1B and 1D). Table 3 shows the percent identity of the various RNA structural analogs with HCV genotypes 1b and 2a. To determine the minimal structure of the X-region that could inhibit HCV RNA replication, shorter RNA structural analogs X-12a and X-12b (Fig. 1C) were designed that were predicted to retain individual stem-loop structures of analog X-94.\textsuperscript{27-29} All analogs were named based on HCV genome region studied and sequence homology of the analogues relative to the JFH-1 HCV genotype 2a genome.

Cloning and expression

For expression studies, the X-94 sequence was subcloned into a pSilencer 4.1 cytomegalovirus (CMV) puro plasmid (Ambion, USA), as previously described.\textsuperscript{12} pSilencer 4.1 CMV puro plasmid enables high level expression of cloned hairpin short RNA templates. Expression vectors for RNA structural analogs X-12 and X-12c were constructed from a plasmid expressing the RNA analog X-94 using a QuikChange II Site-Directed Mutagenesis Kit (Agilent Technologies, Inc. USA) according to manufacturer’s instructions. Fig. 1B shows nucleotide replacements made in stems of analog X-94 resulting in analog X-12, and Fig. 1D illustrates nucleotide replacements made in loops of analog X-12 to construct analog X-12c. Shorter RNA structural analogs X-12a and X-12b (Fig. 1C) were constructed from a plasmid expressing RNA analog X-12 using a QuikChange II Site-Directed Mutagenesis Kit. These sequences were selected as non-overlapping fragments of X-12 that were individually predicted by Mfold to retain all the structural elements present in the parent X-12 analog. 5B-74 RNA analog, which was previously shown to inhibit viral genome replication\textsuperscript{12} was used as a positive control for inhibition of viral replication, and a plasmid expressing an unrelated sequence, HB, from hepatitis B virus was used as negative control. An RNA structural analog 5B-46 was predicted to adopt stem-loop structures identical to analog 5B-74 as determined by Mfold, and was constructed using a plasmid expressing RNA analog 5B-74 with QuikChange II Site-Directed Mutagenesis Kit. Sequences of each analog in pSilencer 4.1 CMV puro plasmid (Ambion) were verified with CMV puro primers (Table 1) as recommended by the manufacturer.

For BB7 HCV genotype 1b replicon studies, cells were plated in 6-well plates 2 days before transfection. Seventy-five percent confluent cells were transfected with various amounts of each plasmid to generate RNA structural analogs individually or in combinations using lipofectamine (Life Technologies) according to manufacturer’s instructions. In brief, lipofectamine and plasmid DNA were separately diluted in Opti-MEM I medium (Invitrogen) without serum. After 15 min incubation, cells were infected with JFH-1 HCV RV. Medium (Invitrogen) without serum. After 15 min incubation, JFH-1 HCV in media was concentrated with a centrifugal device (100K NMWL, Amicon ultra, Millipore, USA). Stocks were frozen at 
\( -80\, ^{\circ}\text{C} \) until further use.

Table 1. Sequences of primers used for quantification

| Primers          | Sequences                      |
|------------------|--------------------------------|
| LDHA FW          | 5’ TAATGAAGGACCTTGCGACATGAAC 3’|
| LDHA RV          | 5’ ACGGCTTTCCTTCCCTTGGCT 3’    |
| HCV1b FW         | 5’ CGTTCTCCAGCGAAGAGGCG 3’     |
| HCV1b RV         | 5’ CACTCGAAAGCACCCCTATCA 3’    |
| CMV puro FW      | 5’ AGGCGATTAGTTGGGTA 3’        |
| CMV puro RV      | 5’ CGTGAGGCCTGTAGCCTGTTGGT 3’  |
| JFH1 HCV FW      | 5’ TAGGAGGCCCCATGTTCAAC 3’     |
| JFH1 HCV RV      | 5’ CCCCTGCTTTCTGAGATGAC 3’     |

CMV, cytomegalovirus; FW, forward; HCV, hepatitis C virus; JFH-1, Japanese Fulminant Hepatitis Virus-1; LDHA, lactate dehydrogenase A; RV, reverse.
they were combined, incubated for 20 min at room temperature, and added to cells in varying concentrations. Cells were harvested 48 h post-transfection with Trizol (Invitrogen), and HCV RNA levels quantitated in the cell lysates.

For HCV genotype 2a infection studies, two models were used: transfection into cells with a pre-existing infection and transfection into cells before infection. For pre-existing JFH-1 HCV infection studies, 75% confluent Huh7.5 cells were infected with JFH-1 HCV for 8 h and then transfected with plasmids expressing RNA structural analogs with lipofectamine, as described above. In brief, cells were washed with phosphate buffered saline (PBS). Lipofectamine and plasmid DNA were separately diluted in Opti-MEM I medium without serum. After a 15 min incubation, they were combined, and incubated for 20 min at room temperature before being added to cells. Culture medium, 200 μL, was collected for quantification of JFH-1 HCV levels, and replaced with fresh 200 μL of cell culture medium at 0, 4, 8, 12, 24, 36, 48, and 72 h of transfection.

For before infection studies, 75% confluent Huh7.5 cells in 6-well plates were transfected with 16 μg of each plasmid expressing RNA structural analogs, as described above. Cells were then infected with JFH-1 HCV 48 h post-transfection. After 8 h of infection, cells were washed twice with PBS to

Fig. 1. A, a diagram of the Mfold ver3.2 computed secondary structure of analog X-94; B, a diagram of nt replacements made in the stems of analog X-94 to construct analog X-12 (secondary structure of X-12 was predicted identical to X-94); C, a diagram of analogs X-12a and X-12b, non-overlapping fragments of analog X-12; D, a diagram of nucleotide replacements made in loops of analog X-12 to construct analog X-12c (secondary structure of X-12c was predicted to be identical to X-94).

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remove input JFH-1 HCV. Culture medium, 200 μL, was collected for quantification of JFH-1 HCV levels and replaced with fresh 200 μL of culture medium at 0, 4, 8, 12, 24, 36, 48, and 72 h of infection.

**Quantitative RT-PCR (qRT-PCR)**

For BB7 HCV genotype 1b studies, whole cell RNA was isolated from replicon cell lysates with an RNAeasy kit (Qiagen, Germany) according to manufacturer’s instructions and treated with RNase free DNase (Invitrogen). cDNA was synthesized using 4 μg DNase treated RNA with SuperScript III First-Strand kit (Invitrogen) and quantified by real time RT-PCR with Power SYBR Green PCR Master Mix (Applied Biosystems, USA) using HCV genotype 1b specific primers (Table 1) according to manufacturer’s instructions. Human lactate dehydrogenase A (LDHA) mRNA levels were quantified in each sample to normalize HCV RNA levels using human LDHA specific primers (Table 1). Assays were done in quadruplicate, and results expressed as mean ± standard error of HCV RNA levels in cells transfected with analogs compared to untreated controls.

For JFH-1 HCV infection studies, viral RNA was extracted from 200 μL media collected from infected cells with a QIAamp Viral RNA kit (Qiagen, Germany) according to manufacturer’s instructions. cDNA was synthesized using 4 μg RNA with SuperScript III First-Strand kit (Invitrogen), and quantified by real time RT-PCR with Power SYBR Green PCR Master Mix (Applied Biosystems) using JFH-1 HCV RNA specific primers (Table 1) according to manufacturer’s instructions. Assays were repeated with three independent replicates, and results are expressed as means ± standard error of JFH-1 HCV RNA levels in media from infected cells transfected with analogs before or after infection compared to untreated controls.

**Results**

Table 2 shows the sequence of all RNA structural analogs. Analogs X-94 and X-12 were 100% and 59% identical to HCV genotype 1b, respectively, but were only 94% and 12% identical to the JFH-1 HCV genome, respectively. Analog X-12c was 50% identical to HCV genotype 1b, and 0% identical to the JFH-1 HCV genome.

To determine whether short RNA sequences predicted to fold into secondary structural analogs of the X region of HCV RNA genome could inhibit HCV replication, plasmids expressing RNA analogs X-94, X-12, and X-12c were transfected in replicon cells. Fig. 2 shows that the effects of transfection of plasmids expressing RNA structural analogs were dose-dependent and most effective at 16 μg of plasmid. Higher doses did not increase effects beyond those at 16 μg, and based on this information, 16 μg of each plasmid was used for transfection for subsequent experiments. 3-(4,5-dimethylthiazol-2-Yl)-2,5-diphenyltetrazolium bromide (MTT) assays showed no significant toxic effects due to DNA transfection (data not shown).

Fig. 3 shows that transfection of plasmids generating RNA analogs affected HCV RNA levels in replicon cells. After transfection of 5B-74, 5B-46, X-94, X-12, and X-12c at 16 μg, HCV RNA levels were decreased to 42%, 55%, 52%, 53%, and 54%, respectively, compared to levels of untreated controls. These differences were significant (p<0.001 for all). An unrelated control plasmid (HB) generating an HBV sequence had no significant effect under identical conditions. Furthermore, combinations of 5B-74 plus X-94 and 5B-46 plus X-12 administered at the same total dose, and under identical conditions, decreased HCV RNA levels to 21% and 30%, respectively, compared to untreated controls (Fig. 3). These levels of inhibition for the combinations were greater than that for any individual analog alone.

The X-region analog is 98 nt long. We wondered whether smaller structural elements could be produced that retained inhibitory effects. To test this hypothesis, two smaller analogs, X-12a and X-12b, were created that corresponded to nt 1-55 and nt 56-95 regions, respectively, of analog X-12 (Fig. 1C). Transfection with X-12a decreased HCV RNA to 58%, about the same as intact X-12 (p<0.001). However, the other fragment, X-12b, was much less effective, resulting in a level of only 78% and not significantly different from untreated control. A combination of X-12a plus X-12b inhibited levels to 60% of untreated controls, which was similar to the effects of X-12a alone (Fig. 4). The data suggested that the X-12a region was the portion of the X analog that was responsible for the majority of the inhibitory
activity. The X-12b region appeared to contribute little to X analog activity.

To determine whether RNA structural analogs were also effective in an infection model and against a different HCV genotype, a JFH-1 infection model was studied. Fig. 5 shows that the levels in untreated controls increased progressively, and by 12 h, exceeded the level of input HCV RNA levels by 6-fold (extreme left bars) at 72 h. In contrast, Huh7.5 cells transfected with analogs 5B-74, 5B-46, X-94, and X-12 and infected with JFH-1 48 h later, inhibited HCV RNA levels to 5.9%, 6.2%, 6.6%, and 1.8%, respectively, compared to untreated controls ($p < 0.001$). Even after 72 h, no cells treated with analogs had HCV RNA levels that exceeded more than 30% of input levels.

To determine whether RNA structural analogs could inhibit a pre-existing HCV infection, cells were infected with JFH-1 HCV for 8 h and then transfected with analogs HB, 5B-74, 5B-46, X-94, and X-12. Medium was tested for JFH-1 HCV RNA post-transfection at various time points. Fig. 6 shows that there was a progressive increase in JFH-1 HCV RNA levels with time in the media compared to uninfected controls. HCV RNA levels exceeded those of input virus at 12 h post-transfection and were four-fold higher by 72 h. However, 72 h after transfection with analogs 5B-74, 5B-46, X-94, and X-12; HCV RNA levels were 8.8%, 10.5%, 9.0%, and 11.6%, respectively, compared to untreated controls.

**Discussion**

Many previous studies, including our own, have shown that specific domains of the genomes of some RNA viruses are critical for viral translation and replication.\textsuperscript{30–32} The NS5B coding region of the HCV genome adopts a stem-loop structure that is involved in the replication of HCV. Expression of RNA structural analogs predicted to mimic the stem-loop structure identical to the NS5B region of the HCV genome was able to inhibit replication of HCV genotype 2a.\textsuperscript{12} The current study confirms previous reports that RNA secondary structure is important for HCV RNA replication.\textsuperscript{16,33} Conserved genomic RNA sequences have been shown to fold to adopt stem-loop motifs that interact with other RNA motifs and/or proteins required for translation and replication.\textsuperscript{10,31,34–37} Identification of such RNA sequences and determination of secondary structure formed by these sequences are challenging because structural motifs depend on various parameters, including host cellular microenvironments and the presence of other host and viral interacting molecules.\textsuperscript{29,38–40} Several types of software have been developed to predict the stable structures formed by RNA sequences based on thermodynamic parameters. The current data generated by Mfold software confirm our previous findings that structures predicted using these two-dimensional models do have substantial inhibitory activity against HCV replication. It is also clear that the actual molecules exist not in a two-dimensional but a three-dimensional state, and it is the latter that causes the inhibitory activity. Such structures cannot be predicted by the current software. Nevertheless, the data support the notion that two-dimensional structures are related to and can predict the activity of analogs in three-dimensions.

We intentionally studied models of two different HCV genotypes. The structural analogs were effective in models of both genotype 1 and 2 viruses, suggesting that because the design of the molecules was based on secondary rather than primary structure, the effects are more likely to be multigenotypic. This may be clinically relevant as it has already

| Analog   | HCV1b   | HCV 2a   |
|----------|---------|----------|
| X-94     | 100%    | 94%      |
| (dG=−46.70 kcal/mol) |        |          |
| X-12     | 59%     | 12%      |
| (dG=−47.40 kcal/mol) |        |          |
| X-12c    | 50%     | No significant identity |
| (dG=−47.50 kcal/mol) |        |          |

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**Table 3. Percent identity of RNA structural analogs with HCV genotypes 1b and 2a (dG: free energy)**

![Fig. 2. HCV RNA levels in replicon cells 48 h post-transfection with structural analogs.](image1)

![Fig. 3. HCV RNA levels in replicon cells 48 h post-transfection with structural analogs.](image2)
been demonstrated that some current direct acting antiviral agents vary in efficacy against HCV genotypes and even subtypes. Design of novel anti-HCV agents based on secondary structural considerations may offer a strategy to develop new agents that are independent of viral genotype or subtype.

For HCV genotype 1b studies, a replicon cell model with an integrated HCV genome was used to constitutively generate subgenomic HCV1b replicons in Huh7.5 cells. This system has been used extensively to determine the effects of various drugs and proteins on viral replication and infection. However, because HCV RNA replication in this model is constitutive, it is not a simulation of HCV infection. For this reason, we examined here the effects of structural analogs on JFH1, to provide a more realistic HCV infection model system. The results from the JFH-1 infection model systems offered insight into the differences in efficacy among structural analogs depending on whether they were introduced before or after viral infection. The JFH-1 HCV infection system has been used to determine the anti-HCV activity of several proteins and inhibitors (example: Raloxifene, NSC compounds) before or after infection of hepatic cells. Introduction of RNA structural analogs after viral infection resembles treatment strategies for hepatocytes already infected with HCV, while exposure of cells before viral infection represents a potential prophylactic approach.

Expression of the stem-loop structure of X-region in the 3'UTR of HCV genome (using RNA analogs X-94, X-12, and X-12c) was found to be effective against HCV replication, regardless of the sequence of RNA. Furthermore, we have identified a small portion, one of the stem-loop structures of the X-region, X-12a, as the smallest identified portion of the X-region analog that retains inhibitory activity. The other stem-loop structure, X-12b of approximately the same length; which also possesses natural HCV sequences, was virtually ineffective. These data confirmed that a specific structure, the stem-loop conformation, was involved, and that the observed inhibitory effects were not due a nonspecific interaction of HCV sequences. The studies on nucleotide base substitution in the X-region showing reduction in the identity to the natural HCV sequence to less than 50%, while retaining secondary structure, indicated that the observed inhibitory effects did not likely involve anti-sense mechanisms. These results are consistent with previous studies.

Conclusions

The data indicate that conformation of secondary structures in 3'UTR of HCV RNA genome is required for HCV replication. Stable expression of RNA analogs predicted to have identical
stem-loop structures, but sequences vastly different from HCV genomic RNA, might inhibit HCV infection of hepatocytes in liver, and may represent a novel approach to design anti-HCV agents.

Acknowledgments

We thank Dr. Charles Rice (Rockefeller University, NY, USA) for providing Huh7.5 cells, Dr. Takaji Wakita (National Institute of Infectious Diseases, Tokyo, Japan) for generous donation of HCV JFH-1 cDNA, and Amy Pallotti and Annilise Larosa for their secretarial assistance.

Conflict of interest

George Y. Wu serves on medical advisory boards for Gilead, Janssen, and Bristol-Myers Squibb.

Author contributions

Conceiving the study and contributing interpretation of data (GYW), designing and performing the research, as well as writing the manuscript (NG), contributing technical advice and review of manuscript (CHW).

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Characterization of the Terminal Regions of Hepatitis C Viral RNA: RNA structures involved in viral replication.