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Structural Determination of the 5' Untranslated Regions of IRE-containing mRNAs

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Structural Determination of the 5' Untranslated Regions of IRE-containing mRNAs

By

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Dr. Bin Wang, Thesis adviser

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ABSTRACT

The expression of ferritin and amyloid precursor protein (APP) is post-transcriptionally regulated by iron-regulating proteins via binding to a stem-loop structure known as an iron-responsive element in the 5’-untranslated region (5’UTR) of ferritin and APP mRNAs. In this study, we used atomic force microscopy (AFM) to visualize the conformation of the 5’UTRs of ferritin heavy chain (Ferritin-H), ferritin light chain (Ferritin-L), and APP mRNA transcripts from human and mouse, and determined the secondary RNA structures using selective 2’-hydroxyl acylation analyzed by primer extension (SHAPE). The AFM imaging did not provide high resolution structural information about these RNAs, whereas the SHAPE procedure successfully interrogated the secondary RNA structures at single nucleotide resolution. To our knowledge, this is the first time that the secondary structures of the entire 5’UTRs of these RNA molecules have been experimentally mapped. This study paves the way for the further investigation of RNA-ligand interactions in these RNA molecules.
ACKNOWLEDGEMENTS

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CHAPTER 1

Introduction

1.1. Iron storage and ferritin

Iron is an essential element required for normal cell growth and proliferation. Iron participates in the cellular processes of aerobic metabolism and enzymatic reactions, and is necessary for the synthesis of myelins, the development of neuronal dendritic trees, and the signal transduction of neurotransmitters in the brain. For the proper function of organs, it is crucial to maintain iron homeostasis. Iron deficiency (also called hypoferremia) results in anemia, whereas iron overload (i.e., the build-up of excess iron in the body) induces hemochromatosis.

The ubiquitous iron storage protein, ferritin, is responsible for both intracellular iron storage in a nontoxic state, and the controllable release of iron when necessary. Mammalian ferritin forms a spherical protein cage that contains 24 subunits, each made of a 4-helix bundle (see Figure 1). There are channels on the protein shell to allow iron exchange and proton transfer. The 24 subunits of the ferritin protein are composed of two functionally-distinct species encoded by two different genes. The first species, ferritin heavy chain (Ferritin-H), has an apparent molecular weight of 21,000 Da; the second, ferritin light chain (Ferritin-L), has an apparent molecular weight of 19,000 Da. The ratio of Ferritin-H to Ferritin-L in a ferritin protein varies in different tissues.
Figure 1. The structure of the mammalian ferritin protein. (A) Each subunit folds into a 4-helix bundle, and (B) the 24 subunits pack to form a spherical shell for iron storage. (Adapted, with permission, from Reference 4)

The deposition of iron into the ferritin cavity involves iron and oxygen chemistry. Ferrous ions (Fe$^{2+}$) react with oxygen (O$_2$) and/or peroxide (H$_2$O$_2$) to form a ferricoxohydroxide core, which is similar to the mineral ferrihydrite (reactions shown below). Ferritin-H catalyzes the oxidation of Fe$^{2+}$; whereas Ferritin-L promotes the nucleation of ferricoxohydroxide in order to store iron.$^{6,7}$

\[
2\text{Fe}^{2+} + \text{O}_2 + 4\text{H}_2\text{O} \rightarrow 2\text{Fe(O)OH}_{\text{core}} + \text{H}_2\text{O}_2 + 4\text{H}^+
\]

\[
2\text{Fe}^{2+} + \text{H}_2\text{O}_2 + 2\text{H}_2\text{O} \rightarrow 2\text{Fe(O)OH}_{\text{core}} + 4\text{H}^+
\]
Excess free (i.e., unbound) iron is cytotoxic because it can react with reactive oxygen species such as H$_2$O$_2$ to form highly toxic hydroxyl free radicals (the Fenton chemistry reaction is shown below). Free radicals can attack other stable molecules and steal their electrons. The attacked molecules then become free radicals themselves, thus setting off a chain reaction. Free radicals can damage DNA, RNA, proteins, and lipids, and cause the breakdown of cell membranes. Therefore, the synthesis of intracellular ferritin must be regulated in order to accommodate the need to store excess iron.

$$\text{Fe}^{2+} + \text{H}_2\text{O}_2 \rightarrow \text{Fe}^{3+} + \text{OH}^- + \cdot\text{OH} \quad \text{(Fenton reaction)}$$

1.2. Iron-responsive elements in the 5’-untranslated region of ferritin mRNA

Ferritin expression is post-transcriptionally regulated by two iron-regulating proteins (IRP1 and IRP2) via binding to a stem-loop structure known as an iron-responsive element (IRE) in the 5’-untranslated region (5’UTR) of the ferritin messenger RNA (mRNA). When the intracellular free iron content is low, IRPs specifically bind to ferritin IRE RNA, thus inhibiting ribosome binding and the corresponding ferritin translation. When the iron concentration is high, IRP-IRE binding is inhibited, thus allowing the synthesis of additional ferritin proteins for iron storage.

IREs are conserved RNA motifs that contain an approximately 30-nucleotide-long sequence that forms a hairpin stem-loop structure. The apical loop of the hairpin includes a
highly conserved CAGUGN sequence, where $N$ can be $A$, $C$, or $U$, but not $G$. Ferritin-H and Ferritin-L mRNAs both have an IRE in their 5’UTRs.

1.3. Iron accumulation and neurodegenerative diseases

Studies have demonstrated that severe iron accumulation in the brain is correlated to neurodegenerative disorders such as Alzheimer’s disease and Parkinson’s disease.\textsuperscript{13-19} Alzheimer’s disease is the most common progressive human neurodegenerative disorder, and is the fifth leading cause of death in adults aged 65 or older.\textsuperscript{20} This brain disorder was first described by German physician Alois Alzheimer in 1906.\textsuperscript{21} People with Alzheimer’s disease undergo memory decline and thinking and behavioral changes, followed by the loss of self-care ability.\textsuperscript{22} There are 5.3 million Americans suffering from Alzheimer’s disease, which imposes a heavy burden on those individual’s families and the health care system more generally.\textsuperscript{23}

One of the main physiological characteristics of Alzheimer’s disease is the formation of extraneuronal senile plaques containing aggregated amyloid $\beta$ peptide (A$\beta$), a 40-42 amino acid product derived from the serial cleavage of the amyloid precursor protein (APP) by $\beta$-secretase and $\gamma$-secretase.\textsuperscript{24-29} The accumulation of A$\beta$ interrupts synaptic transmission and alters synaptic plasticity. APP is a large type-1 transmembrane protein that is expressed in most cell types and is concentrated in the synapses of neurons.\textsuperscript{25} Beta-secretase, also known as $\beta$-site APP-cleaving enzyme 1 (BACE1), is a type-1 transmembrane protein that has aspartyl protease activity.\textsuperscript{30,31} BACE1 extracellularly cleaves APP at the N-terminus.\textsuperscript{32}
The C-terminal fragment of APP, which contains 99 amino acids (C99), remains bound to the membrane. Gamma-secretase, a protein complex consisting of presenilins, nicastrin, anterior pharynx-defective phenotype-1, and PS enhancer-2, further cleaves C99 within the transmembrane domain to release an extracellular, hydrophobic Aβ peptide (see Figure 2). It has been discovered that the deposition of Aβ in the brain is largely due to increased APP/BACE1 expression and activity. In addition, increased oxidative stress due to misregulated iron homeostasis also plays an important role in the pathogenesis of Alzheimer’s disease. Elevated iron concentrations have been detected in hippocampus, basal nucleus of meynert, senile plaques, and neurofibrillary tangles in the brain of Alzheimer’s patients.

**Figure 2.** An overview of the cleavage of APP by β-secretase and γ-secretase. (Adapted, with permission, from Reference 24)
Parkinson’s disease is the second most common human neurodegenerative disorder, with symptoms that include resting tremors, difficulty balancing, moving and talking, stiffness of limbs, and dementia. The etiology of Parkinson’s disease remains unclear. However, several factors play pivotal roles, including the elevated expression level of the presynaptic protein α-synuclein (SNCA), and the elevated iron concentration. The aggregated SNCA is the main component of Lewy bodies, abnormal protein clusters found in the brains of Parkinson’s patients. SNCA duplication and triplication as well as genetic variability in the promoter and 3’ untranslated region (3’UTR) have been reported to associate with familial Parkinson’s disease.

Elevated iron concentrations in the substantia nigra pars compacta have been implicated in the development of idiopathic Parkinson’s disease. An individual ferritin molecule can bind up to 4500 iron atoms. However, ferritin in the substantia nigra of Parkinson’s patients was reported to contain more iron than ferritin in the substantia nigra of healthy people. The overloaded ferritin was still unable to take up enough iron to properly regulate iron levels in the brains of Parkinson’s patients. Furthermore, research shows that the ferritin level in Parkinson’s patients was independent of iron level, i.e., the increased iron content observed in the brains of Parkinson’s patients was not accompanied by an increased ferritin level. The absence of ferritin up-regulation might be due to the increased binding activity of IRP to the IRE of ferritin RNA in the presence of nitric oxide. Other mechanisms may also be involved.
1.4. IRE in the 5’UTRs of APP and SNCA mRNAs

In 2002, an IRE-type structure was found in the 5’UTR of the APP mRNA. In 2007, an IRE-like stem-loop structure was found in the 5’UTR of the SNCA mRNA. These IRE-type sequences interact with IRPs to control the corresponding protein synthesis. The mechanism is similar to that of the iron-dependent translational regulation of Ferritin-H and Ferritin-L synthesis via IREs in their 5’UTRs. Tens of thousands of compounds have been screened to identify ligands that can bind to the 5’UTR of the human APP mRNA, which would allow the treatment of Alzheimer’s disease by decreasing the production of APP and the corresponding Aβ; fifteen compounds were identified as inhibiting APP translation by interacting with the 5’UTR of APP mRNA. The fifteen molecules are: iron chelators desferrioxamine (DFO), iodochlorhydroxyquin (clioquinol), M30, HLA20, and VK28 (varinel), copper chelator tetrathiomolybdate, lead and mercury chelator dimercaptopropanol, naturally occurring iron and copper chelators (-)-epigallocatechin-3-gallate (EGCG) and curcumin, bi-functional metal chelator XH-I, macrolide antibiotics Azithromycin and Erythromycin, selective serotonin reuptake inhibitor and chelator Paroxetine, anticholinesterase Phenserine, and antioxidant N-acetyl cysteine.

Some of the molecules mentioned above showed clinical effectiveness in slowing AD dementia. However, the underlying mechanism(s) has not been determined. For example, what conformation does APP mRNA 5’UTR adopt after the binding of a small molecule? Which nucleotides do the ligands bind to in the IRE? Do the ligands also bind to other regions in the 5’UTR of APP mRNA, such as the interleukin-1 responsive acute box?
If the molecules mentioned above have the capability to reduce APP expression, can they bind to the IRE in human SNCA mRNA 5’UTR to inhibit SNCA synthesis? Do they have the capability to inhibit Ferritin-H and Ferritin-L translation as well? It is an advantage if these molecules can down-regulate APP and SNCA expression, but certainly a disadvantage if they down-regulate the expression of ferritin.

1.5. Objective of the study

In order to substantially improve the understanding of ligand-RNA binding mechanisms and to shed light on the design of novel molecules targeted to specific RNA motifs, a complete elucidation of the RNA conformation before, during, and after ligand binding is necessary. The objective of this study is to determine the conformations of the 5’UTRs of human and mouse Ferritin-H, Ferritin-L, and APP mRNA transcripts in the absence of the ligand. The long-term goal of this study is to speed up the discovery of potential therapeutic agents for the treatment of neurodegenerative diseases.

1.6. Techniques used in the study

1.6.1. Atomic force microscopy

For this project, two cutting-edge technologies were used to study the structural characteristics of RNA. First, atomic force microscopy (AFM) was applied to visualize the conformation of RNA transcripts. AFM was invented in 1986, and the technique soon
became a potent tool for biological research. The principle of AFM is based on the mechanical interaction that occurs between a sample surface and a cantilevered sharp silicon tip (5-10 nm radius at the point of the tip), when a small, constant force is applied. A piezoelectric tube scanner is used to scan the tip over the sample surface. The tip-sample interaction is detected by monitoring the deflection of the cantilever using a photodiode detector (see Figure 3). AFM enables the study of the three-dimensional topography of a sample surface with a resolution at the Angstrom level. The simple sample preparation procedure for AFM imaging makes it much more convenient than conventional imaging techniques such as X-ray crystallography and electron microscopy.

AFM can be operated both in air and in a liquid solution, and no sample coating or staining is required; therefore, the conformation of biological macromolecules (e.g., DNA, RNA, and proteins) adsorbed onto a smooth mica surface can be visualized under quasi-native (i.e., close to physiological) conditions. Jaeger’s research group has recently used AFM to visualize the static RNA nanoarchitectures they designed, including tectoRNA and kissing-loop RNA. Samori’s research group applied AFM to study the secondary structure of Turnip Yellow Mosaic Virus RNA. Gamarnik and colleagues used AFM to investigate the long-range RNA interactions that circularize the 5’- and 3’-end of the dengue virus.
**1.6.2. Selective 2’-hydroxyl acylation analyzed by primer extension**

In addition to AFM visualization, the SHAPE (Selective 2’-Hydroxyl Acylation analyzed by Primer Extension) technique was used to determine RNA secondary structures. Invented in 2005, the theoretical basis of SHAPE chemistry is that the local nucleotide flexibility of an RNA can be monitored by treating the RNA with 2’-hydroxyl-reactive electrophiles such as N-methylisatoic anhydride (NMIA) or 1-methyl-7-nitroisatoic anhydride (1M7), which selectively and covalently modify flexible nucleotides (i.e., single-
stranded RNA nucleotides) at the 2’-ribose position.\textsuperscript{78-80} Treating RNA with the NMIA/1M7 reagent (usually \( \leq 10 \) mM) results in modifications approximately once in every several hundred nucleotides. The detection of the modified RNA nucleotides is based on primer extension to convert RNA to its complementary DNA (cDNA) by reverse transcriptase. The presence of a 2’-\( O \)-adduct causes the reverse transcription to stop exactly one nucleotide prior to the modified position, whereas RNA without NMIA/1M7 treatment produces a full-length cDNA.\textsuperscript{78} The amplified cDNA fragments are analyzed by conventional slab gel electrophoresis (for radio-labeled DNA fragments) or capillary gel electrophoresis (for fluorescently-labeled DNA fragments) to provide RNA structural information.\textsuperscript{81,82} Compared to traditional RNA structure-mapping techniques such as enzymatic cleavage and chemical modification, which are time consuming and evaluate only a subset of nucleotides in an RNA, SHAPE chemistry is a rapid, high-throughput technology with single nucleotide resolution.

Figure 4 shows an example of SHAPE data obtained from capillary gel electrophoresis. The SHAPE reactivity of each nucleotide of yeast tRNA\textsuperscript{Asp} can be determined by subtracting the control peak area from the NMIA peak area and the subsequent normalization. Peaks with high amplitudes correspond to the conformationally flexible (i.e., single-stranded) nucleotides, whereas peaks with low amplitudes correspond to the conformationally-constrained (i.e., base-paired or tertiary interaction-involved) positions. SHAPE data are applicable to constrain RNA structural prediction algorithms and can dramatically improve the predication accuracy of the RNA secondary structures.
Figure 4. An example of SHAPE data obtained from capillary gel electrophoresis. Left: SHAPE raw data from *Saccharomyces cerevisiae* tRNA<sub>Asp</sub> transcript obtained by capillary gel electrophoresis. The blue trace is +NMIA sample; the green trace is -NMIA negative control; black and red traces are two dideoxy ladders (ddC and ddT, respectively). The x-axis indicates the number of data points collected. The y-axis enumerates the fluorescence units. Right: Individual nucleotide reactivity based on SHAPE data. Red-coded nucleotides correspond to 60-100% of SHAPE reactivity; yellow-coded nucleotides correspond to 25-60% of SHAPE reactivity; green-coded nucleotides correspond to 10-25% of SHAPE reactivity; black-coded nucleotides correspond to 0-10% of SHAPE reactivity.

Figure 5 shows an example of the use of SHAPE chemistry coupled with an autoradiography/slab gel electrophoresis detection system to study magnesium-induced conformational changes in yeast tRNA<sub>Asp</sub>. Wang and colleagues used 1M7 instead of NMIA to run SHAPE experiments because the reactivity of 1M7 is not significantly modulated by ions in solution. As expected, SHAPE reactivity exactly recapitulates the native structure of tRNA<sub>Asp</sub>. Wang et al. explored Mg<sup>2+</sup>-dependent structural changes over a
wide range of ion concentrations (20 to 0 mM, Figure 5A, 19 left-most lanes). Smooth transitions for almost every nucleotide in tRNA<sub>Asp</sub> were observed. A well-defined model for the final non-native state was developed based on the SHAPE reactivity information (Figure 5C). Removing Mg<sup>2+</sup> destabilizes tertiary interactions in the variable loop and thereby allows this region to form stable, non-native base pairs with nucleotides in the D-loop.81
Figure 5. Mg$^{2+}$-dependent unfolding of tRNA$^{Asp}$. (A) tRNA$^{Asp}$ modification upon removing Mg$^{2+}$ as visualized by RNA SHAPE chemistry. Experiments were performed as a function of [MgCl$_2$] in the presence (+) and absence (–) of 1M7 reagent. (B) Mg$^{2+}$-induced
structural transitions, illustrated for instructive nucleotide positions.  (C) Structural model for the unfolding of tRNA$^{\text{Asp}}$ in the absence of Mg$^{2+}$.  Mg$_{1/2}$ values are listed for nucleotides that show greater than a two-fold reactivity change over the titration.  Nucleotides exhibiting 2-fold or larger changes in reactivity as compared to the native state are colored red (increase) and green (decrease); nucleotides showing <2-fold changes are colored blue (unreactive, SHAPE reactivity $\leq$0.3), orange (moderately reactive, $0.3<\text{SHAPE reactivity}<0.7$), and red outline (highly reactive, SHAPE reactivity $\geq0.7$).  (Adapted, with permission, from Reference 81)

In this study, we used AFM to visualize the conformation of the 5’UTRs of Ferritin-H, Ferritin-L, and APP mRNA transcripts from two species (human and mouse), and determined the secondary RNA structures using SHAPE analysis coupled with a fluorescence detection system.  To our knowledge, this is the first time that the secondary structures of the 5’UTRs of APP mRNA in human and mouse have been experimentally mapped.  This study paves the way for the further investigation of RNA-ligand interactions in these RNA molecules.
CHAPTER 2

Experimental Methods

2.1. DNA

*DNA of human Ferritin-H 5’UTR:* The 200-nucleotide-long DNA template (5’-
ACAAGCGACCGCAGGGCCAGACGTTCTTCGCCGAGAGTCGTCGGGGTTTCCTG
CTTCAACAGTGCTTTTGACGGAACCCGGCGCTCGTTCCCCCACCCCCCGCCGGCCCGCC
CATAGCCAGCCCTCCGTCACCTCTTTCAACCACCCCTCGGACTGCCCCAAGCCGCCCCCC
CGCCGCGCGCTCCAGCGCCGCAGCCGAGCCGACCGGCACCGCCCGCAG-3’), forward primer that
included a T7 promoter sequence (5’-
TAATACGACTCACTATAGGATAAGAGACGCGACGTTCTTCGCCGAGAGTCGTCGGGGTTTCCTG
CTTCAACAGTGCTTTTGACGGAACCCGGCGCTCGTTCCCCCACCCCCCGCCGGCCCGCC
CATAGCCAGCCCTCCGTCACCTCTTTCAACCACCCCTCGGACTGCCCCAAGCCGCCCCCC
CGCCGCGCGCTCCAGCGCCGCAGCCGAGCCGACCGGCACCGCCCGCAG-3’),
and reverse primer (5’-
GGCGGCGACTAAGGAGAGGCGGCGGCGGCGGCGGTGGCTGCGCGGCGCTG-3’)
were obtained from Integrated DNA Technologies, Inc. (IDT, San Diego, CA) and were used
to amplify the 254-base pair Ferritin-H 5’UTR. The Polymerase Chain Reaction (PCR)
mixture (1.0 ml total volume) contained 20 mM Tris (pH 8.4), 10% dimethyl sulfoxide
(DMSO), 50 mM KCl, 2.5 mM MgCl₂, 0.2 mM of each dNTP (dATP, dGTP, dCTP, and
dTTP), 0.5 µM each of forward and reverse primer, 5 pM of DNA template, and 0.025 U/µl
Taq DNA polymerase. The PCR product was inspected by ethidium bromide-stained
agarose gel electrophoresis and purified by ethanol precipitation.
**DNA of human Ferritin-H short 5’UTR:** To obtain a PCR ready cDNA, Human Brain Total RNA (Ambion, Austin, TX) was reverse-transcribed using a high capacity cDNA reverse transcription kit from Applied Biosystems (Foster City, CA). A 224-base pair segment of DNA that encodes the Ferritin-H short 5’UTR was amplified using the resultant cDNA as a template, along with a forward primer containing the T7 promoter (5’- TAATACGACTCACTATAGGATAAGAGACCACAAGCGACCCG-3’) and reverse primer (5’-GCGGTGGCTGCGCGGCGCTGGAG-3’), both of which were synthesized at IDT. The PCR reaction was performed using 0.2 mL, but was otherwise as described above. The product was inspected by ethidium bromide-stained agarose gel electrophoresis and purified using the QIAquick PCR purification kit (Qiagen, Valencia, CA).

**DNA of human Ferritin-L 5’UTR:** FirstChoice® PCR-ready Human Brain cDNA (Ambion, Austin, TX) (2 ng) was used as a template to amplify the 234 base-pair Human Ferritin-L 5’UTR. Both the forward primer, which contained a T7 promoter sequence (5’- TAATACGACTCACTATAGGGCAGTTCGGCGGTCCCGCGGGTCTGTCTCT-3’), and the reverse primer (5’-GAATCTGGGAGCTCATGGTTGGTTGGC-3’) were synthesized at IDT. A 0.2 ml PCR reaction was performed as described above. The product was inspected by ethidium bromide-stained agarose gel electrophoresis and purified using the QIAquick PCR purification kit (Qiagen, Valencia, CA).

**DNA of human APP 5’UTR:** FirstChoice® PCR-ready Human Brain cDNA (Ambion, Austin, TX) (2 ng) was used as a template to amplify the 180-base pair human APP 5’UTR. Both the forward primer, which contained a T7 promoter sequence (5’-
TAATACGACTCATACTAGGAGTTTCCTCGGCAGCGGTAGGCGAG-3’), and the reverse primer (5’-AAACCGGGCAGCATCGCGACC-3’) were obtained from IDT. A 0.2 ml PCR was performed as described above. The product was inspected by ethidium bromide-stained agarose gel electrophoresis and purified using the QIAquick PCR purification kit (Qiagen, Valencia, CA).

**DNA of mouse Ferritin-H 5’UTR:** FirstChoice® PCR-ready Mouse Brain cDNA (Ambion, Austin, TX) (2 ng) was used as a template to amplify the 186-base pair mouse Ferritin-H 5’UTR. Both the forward primer, which contained a T7 promoter sequence (5’-TAATACGACTCACTATAGGAGCAGACGTTCTCAGGCTCGCC-3’), and the reverse primer (5’-GGTGCGGGCGGGGCGAGGCGG-3’) were obtained from IDT. A 0.2 ml PCR reaction was performed as described above. The product was inspected by ethidium bromide-stained agarose gel electrophoresis and purified using the QIAquick PCR purification kit (Qiagen, Valencia, CA).

**DNA of mouse Ferritin-L 5’UTR:** Mouse Brain Total RNA (Ambion, Austin, TX) was reverse-transcribed using a high capacity cDNA reverse transcription kit from Applied Biosystems (Foster City, CA). The resultant cDNA product was used as a template to amplify the 224-base pair mouse Ferritin-L 5’UTR. Both the forward primer (5’-TAATACGACTCACTATAGGAGCAGCGCCTTGAGGAGGCTCGCC-3’) and reverse primer (5’-GGCTGATCCGGAGTAGGAGCTAAC-3’) were obtained from IDT. A 0.2 ml PCR reaction was performed as described above. The product was inspected by ethidium bromide-stained agarose gel electrophoresis and purified using the QIAquick PCR purification kit (Qiagen, Valencia, CA).
bromide-stained agarose gel electrophoresis and purified using the QIAquick PCR purification kit (Qiagen, Valencia, CA).

**DNA of mouse APP 5’UTR:** The reverse-transcribed mouse brain cDNA product described above was used as a template to amplify the 164-base pair mouse APP 5’UTR. Both the forward primer (5’-TAATACGACTCACTATAGGGTTTTCTCGGGCGGAGGCAGA-3’) and reverse primer (5’-CGTGATCCTGCGTGGGCCACCGAGT-3’) were synthesized at IDT. A 0.2 ml PCR reaction was performed as described above. The product was inspected by ethidium bromide-stained agarose gel electrophoresis and purified using the QIAquick PCR purification kit (Qiagen, Valencia, CA).

### 2.2. RNA

RNA from the human Ferritin-H 5’UTR was transcribed from 1.0 ml of a solution containing 50 µg of PCR-generated DNA, 40 mM Tris (pH 7.8), 20 mM NaCl, 6 mM MgCl₂, 2 mM spermidine HCl, 10 mM DTT, 1 mM of each NTP (ATP, GTP, CTP, and UTP), 0.1 U/µl SUPERaseIn™ RNase inhibitor, and 2 U/µl T7 RNA polymerase; incubation was at 37 °C for 4-5 hours. RNA transcripts were treated with TURBO™ DNase (Ambion, Austin, TX) to destroy any remaining DNA. The DNase was then deactivated by adding ethylenediaminetetraacetic acid (EDTA) to a final concentration of 15 mM, followed by heating at 75 °C for 15 minutes. The RNA transcripts were concentrated by ethanol precipitation, purified by 8% polyacrylamide gel electrophoresis (PAGE), and recovered by
passive elution overnight at 4 °C in a solution containing 0.5 M sodium acetate and 1 mM EDTA.

RNAs from the human Ferritin-H short 5’UTR, human Ferritin-L 5’UTR, human APP 5’UTR, mouse Ferritin-H 5’UTR, mouse Ferritin-L 5’UTR, and mouse APP 5’UTR were transcribed from 0.2 ml of a solution having the same composition as described above, except that only 5 µg of the PCR-generated DNA was used here. The RNA transcripts were purified using a MEGAclear™ Kit (Ambion, Austin, TX) instead of PAGE due to the relatively low yields. The purity and integrity of the seven RNA samples were tested by running a RNA 6000 nano-chip on an Agilent 2100 Bioanalyzer (Agilent, Santa Clara, CA).

2.3. AFM

A 3.5 µl aliquot of an approximately 1 µM RNA sample in TE buffer (10 mM Tris at pH 8.0 and 1 mM EDTA at pH 8.0) was combined with 1.5 µl of folding buffer (final concentration, 100 mM Tris-HCl at pH 8.0, 100 mM NaCl, and 6 mM MgCl₂) and incubated at 37 °C for 20-30 minutes. The 5.0 µl of solution was deposited on freshly cleaved mica, and the RNA was allowed to adsorb to the mica surface for 5 minutes. The non-adsorbed RNA was removed from the mica surface by washing with nuclease-free water, after which the sample was dried with a stream of nitrogen. All images were collected in air using a Thermomicroscopes Explorer AFM (Veeco, CA) with a Tube or Tripod scanner. Images were obtained while using either Micromash DP15/HiRes-W/AIBS tips (nominal radius of tip curvature ≈ 1 nm, Al reflective side coating, cantilever length = 230 µm, width = 35 µm,
resonant frequency = 325 kHz) or NSC15/AIBS tips (nominal radius of tip curvature < 10 nm, Al reflective side coating, cantilever length = 125 µm, width = 35 µm, resonant frequency = 325 kHz). Thermomicroscopes Scanning Probe Microscopy Lab Analysis software was then used to process and analyze the images for leveling and topography measurements.

2.4. SHAPE

SHAPE includes three major procedures: chemical modification of flexible RNA nucleotides, reverse transcription to identify modified RNA nucleotides, and finally data analysis and the construction of a model of the RNA secondary structure. The 8 detailed steps of SHAPE are itemized below.

1) Folding: The RNA was heated at 95°C for 2 minutes, and then cooled on ice to eliminate multimeric forms. The RNA began in 0.5× TE buffer (pH 8.0), folding buffer (final concentration, 100 mM Tris-HCl at pH 8.0, 100 mM NaCl, and 6 mM MgCl₂) was added to the RNA solution. This new solution was incubated at 37°C for 20 minutes. After incubation, half of the solution was transferred into a second tube.

2) 1M7 modification: 1M7 in DMSO was added to the RNA solution (the first tube) at a final concentration of 6 mM, and the solution was allowed to continue incubating for 2 minutes (greater than five 1M7 half-lives). A control experiment was run in parallel where 1M7 was omitted and only DMSO was added (the second tube).
3) An ethanol precipitation was then conducted to recover RNA.

4) Primer Extension: A different color-coded fluorophore-labeled DNA primer (human Ferritin-H 5’UTR primer: 5’-GACTAAGGAGAGG-3’; human Ferritin-H short 5’UTR primer: 5’-GCGGTGGCTGCGGCGCTGGGAG-3’; human Ferritin-L 5’UTR primer: 5’-GAATCTGGAGCTCATGGTT-3’; human APP 5’UTR primer: 5’-AAACCGGACGATCGAGAC-3’; mouse Ferritin-H 5’UTR primer: 5’-GGTGCGCCGCGGGCAG-3’; mouse Ferritin-L 5’UTR primer: 5’-GGCTGATCCGGAGTGGAGCATAAC-3’; and mouse APP 5’UTR primer: 5’-CGTGATCCGGAGTGGAGCACA-3’) was added to the (+) 1M7 and (-) 1M7 reactions. The two samples were incubated at 65°C for 6-10 minutes followed by 35°C for 15 minutes to allow primer annealing to the 3’ end of the RNA. Reverse transcription buffer (including final concentrations of 50 mM Tris-HCl at pH 8.3, 75 mM KCl, 3mM MgCl₂, 5 mM DTT, 0.5 mM dATP, 0.5 mM dCTP, 0.5 mM dGTP, and 0.5 mM dTTP) was added and heated to 52°C, then reverse transcriptase (Superscript III, Invitrogen, Carlsbad, CA) was added to the solution and allowed to incubate at 52°C for 15 minutes for primer extension. Two dideoxy sequencing reactions were run in parallel in order to identify peaks in the (+) and (-) 1M7 reactions.

5) Recovery of cDNA: Ethanol precipitation was used to recover cDNA, which was then resuspended in highly deionized formamide.
6) cDNA fragment analysis: The suspended cDNA was loaded onto the Applied Biosystems 310 Genetic Analyzer (ABI 310, Applied Biosystems, Foster City, CA). This system uses capillary gel electrophoresis coupled with a fluorescence detection system to determine the size of the cDNA fragments.

7) SHAPE data analysis: Data collected from ABI 310 underwent baseline adjustment, matrixing to remove signal overlap, and mobility shift adjustment using ShapeFinder software. Integrated SHAPE reactivity at each nucleotide position was normalized to a scale from 0 to ~2.

8) Model Construction: The normalized SHAPE reactivities were incorporated into RNAstructure 4.6 software, which uses both nearest-neighbor free energy parameters and SHAPE data as pseudo-energy parameters to develop a secondary structure prediction.
CHAPTER 3

Results

3.1. The 5’UTR of human Ferritin-H mRNA

The sequence of the 237-nucleotide-long human Ferritin-H mRNA 5’UTR is as follows:

5’-GG AUAAGAGACC ACAAGCGACC CGCAGGGCCA GACGUUCUUC
GCCGAGAGUC GUCGGGGUUU CCUGCUUCAA CAGUGCUUUGG ACGGAACCCG
GCGCUCGUUC CCCACCCCGG CCGGCGGCCC AUAGCCAGCC CUCUCUCGCC
UCUUCACCCGC ACCCUCGGAC UGCCCCAAGG CCCCCGCCGC CGCUCCAGCG
CCGCGCAGCC ACCGCGCGGC CCGGCCGCCUC UCCUUAGUCG CGGCC-3’

3.1.1. AFM images

An AFM image of the negative control is shown in Figure 6. The negative control was made by depositing 3.5 µl of TE buffer and 1.5 µl of folding buffer (see Experimental Methods), without RNA, onto freshly cleaved mica and allowing it to sit for 5 minutes, followed by washing with nuclease-free water, and drying with a stream of nitrogen. Figure 6A shows a two-dimensional image of the mica surface; Figure 6B is a three-dimensional view of the sample surface. The surface of the sample composed of mica with the negative
control solution deposited onto it is almost featureless, with no apparent blobs. The surface roughness is similar to that of the original mica surface.

Figure 6. AFM images of the mica surface with the negative control solution deposited onto it. (A) The two-dimensional view of the sample surface, and (B) the three-dimensional view of the sample surface.
An AFM image of a sample composed of human Ferritin-H mRNA 5’UTR deposited onto a mica surface is shown in Figure 7. The 237-nucleotide-long RNA molecule folds into a complex three-dimensional structure that appears as a blob on the mica surface. The heights of three blobs were measured using the “line analysis” tool in the Thermomicroscopes Scanning Probe Microscopy Lab Analysis software, with the maximum height of a blob recorded as its height (see Figure 8A). The heights of the eleven blobs were measured as: 0.81 nm, 0.55 nm, 0.35 nm, 0.38 nm, 0.31 nm, 0.21 nm, 0.33 nm, 0.22 nm, 0.23 nm, 0.13 nm, and 0.12 nm (average ± standard deviation = 0.33 ± 0.20 nm).

The volume of a blob was determined by drawing four lines on the blob (see Figure 8B, where only three lines are demonstrated), where line 1 was drawn parallel to the AFM tip’s scanning direction (horizontal blue line); line 2 was drawn at an angle that is 135 degrees clockwise to line 1 (green line); line 3 was drawn at an angle that is 90 degrees to line 2 (red line); and line 4 was drawn at 90 degrees to line 1 and 45 degrees to lines 2 and 3 (vertical line, not shown). The length of each line was measured at half the maximum height of the blob (see Figure 8C), a measurement called Full Width at Half Maximum (FWHM). The line with the maximum FWHM value is identified as the Major Axis, and the line with minimum FWHM value is identified as the Minor Axis. The blob was then treated as an ellipsoid, and the volume was determined using the formula: Volume = \(\frac{4}{3} \pi \times \text{FWHM}_{\text{Major Axis}} \times \text{FWHM}_{\text{Minor Axis}} \times \text{Height of the blob}\). The volumes of the eleven blobs were 1235 nm\(^3\), 823 nm\(^3\), 433 nm\(^3\), 313 nm\(^3\), 220 nm\(^3\), 103 nm\(^3\), 277 nm\(^3\), 112 nm\(^3\), 173 nm\(^3\), 97.0 nm\(^3\), and 131 nm\(^3\), respectively (average ± standard deviation = 356 ± 360 nm\(^3\)). There are several
possible explanations for the differences in height and volume of the blobs, such as 1) some
blobs on the mica surface were RNA aggregates; 2) some blobs were degraded RNA
molecules; and 3) RNA molecules adopted different conformations on the mica surface.
Figure 7. AFM images of a sample composed of the human Ferritin-H mRNA 5’UTR deposited onto a mica surface. (A) The two-dimensional view of the sample surface, and (B) the three-dimensional view of the sample surface.
Figure 8. Illustration of line analysis from the AFM image of the human Ferritin-H mRNA 5’ UTR. (A) Illustration of the height measurement of a RNA blob, (B) determination of the Major Axis and the Minor Axis for an RNA blob at half the maximum height, and (C) illustration of how to measure the full width of a blob at half the maximum height (FWHM).

3.1.2. SHAPE data

The human Ferritin-H mRNA 5’UTR transcripts were folded in a solution containing 6 mM Mg$^{2+}$ and subjected to the SHAPE procedure. The SHAPE experiments were repeated at least twice. The raw data from the ABI genetic analyzer underwent a series of
modifications in order to remove signal overlap and migration differences due to the cDNA fragments being labeled with different colored fluorophores. Figure 9 illustrates one set of SHAPE electropherograms for the human Ferritin-H mRNA 5’UTR.

After alignment and integration to obtain the Gaussian-fitted peak area at each nucleotide position for both the (+) 1M7 and (-) 1M7 traces, the background (-) 1M7 peak area was subtracted from the corresponding (+) 1M7 peak area. The absolute SHAPE reactivities obtained were normalized by excluding the 2% most reactive fluorescent intensities, and then dividing by the average of the remaining 8% most highly reactive positions. The normalized SHAPE reactivities were classified into three categories. Nucleotides with high SHAPE reactivities (≥0.7) are expected to be single-stranded; nucleotides with low SHAPE reactivities (<0.3) are unreactive, and are expected to be base-paired; and nucleotides with intermediate SHAPE reactivities (between 0.3 and 0.7) are difficult to judge. Nucleotides in this last category may be single-stranded or could be involved in base-pairing/tertiary interactions. Figure 10 presents the averaged SHAPE reactivities from two sets of SHAPE data for the human Ferritin-H mRNA 5’UTR.

Normalized SHAPE reactivity values (0 to ~2) were incorporated into the RNAstructure program as a pseudo free energy change term, so as to constrain the prediction of the secondary structure of the RNA. The predicted structure with the lowest free energy for the human Ferritin-H mRNA 5’UTR is shown in Figure 11. The human Ferritin-H mRNA 5’UTR shows a high degree of secondary structure, with numerous stem-loops, which is expected due to the high GC content in the RNA sequence. Interestingly, the
predicted structure does not contain a single stem-loop IRE where expected at nucleotide positions 61-88, as suggested by the presence of an apical loop that contains a CAGUGC sequence. In order to form the IRE stem-loop, nucleotides 68-72 should pair with nucleotides 83-79. However, in Figure 11, nucleotides 68-72 show medium to high SHAPE reactivities and are single-stranded, whereas nucleotides 79-82 pair with nucleotides 125, 123, 122, and 121, respectively. Based on the structure prediction, nucleotides at the 5’-end of the human Ferritin-H mRNA 5’UTR (nucleotides 2-10) form base pairs with nucleotides at the 3’-end of the RNA; this base-pairing may disrupt the secondary structure of the other regions in this RNA. We thus deleted 30 nucleotides from the 3’-end of the RNA, and synthesized an RNA named “human Ferritin-H mRNA short 5’UTR”.
Figure 9. SHAPE data for the 5’UTR of the human Ferritin-H mRNA transcript. The raw output from the ABI 310 genetic analyzer underwent baseline adjustment, matrixing, and mobility shift adjustment. Fluorescence-labeled DNA primer was annealed to the 3’-end of this 237-nucleotide RNA. The blue trace is the +1M7 sample; the black trace is the -1M7 negative control; green and red traces are two dideoxy ladders (ddG and ddT, respectively). The x-axis indicates the number of data points collected. The y-axis (not shown) enumerates the fluorescence units.
Figure 10. Single nucleotide resolution of normalized SHAPE reactivities for the human Ferritin-H mRNA 5' UTR. Error bars represent standard deviations calculated from two independent experiments.
Figure 11. The predicted secondary structure of the 5’UTR of human Ferritin-H mRNA. Nucleotides are colored black (unreactive, SHAPE reactivity < 0.3), or orange (moderately reactive, 0.3 ≤ SHAPE reactivity < 0.7), red (highly reactive, SHAPE reactivity ≥0.7), or grey (no data).
3.2. Human Ferritin-H mRNA short 5’UTR

The sequence of the 207-nucleotide-long human Ferritin-H mRNA short 5’UTR is as follows:

5’-GG AUAAGAGACC ACAAGCGACC CGCAGGGCCA GACGUUCUUC GCCAGAGAGUC GUCGGGGUUU CCUGCUUCAA CAGUGCUUGG ACGGAACCG GCGCUCGUUC CCCACCCCGG CCGGCCGCCC AUAGCCAGCC CUCCGUCACC UCUUCACCGC ACCCUCGGAC UGCCCCAAGG CCCCCGCCGC CGCUCCAGCG CCGGCAGGCC ACCGC-3’

3.2.1. AFM images

An AFM image of the human Ferritin-H mRNA short 5’UTR is shown in Figure 12. The 207-nucleotide-long RNA molecule folds into a complex three-dimensional structure that appears as a blob on the mica surface. The heights of eight blobs were measured as: 1.12 nm, 0.85 nm, 0.79 nm, 0.60 nm, 0.59 nm, 0.50 nm, 0.34 nm, and 0.41 nm (average ± standard deviation = 0.65 ± 0.26 nm). The volumes of the same eight blobs were 2403 nm³, 2001 nm³, 1451 nm³, 941 nm³, 694 nm³, 982 nm³, 471 nm³, and 357 nm³, respectively (average ± standard deviation = 1162 ± 732 nm³). The averaged volume of the blobs for the human Ferritin-H mRNA short 5’UTR (1162 ± 732 nm³) doubles that of the human Ferritin-H 5’UTR (356 ± 360 nm³), which demonstrates that no correlation has been observed between volume and RNA sequence length in this study.
Figure 12. AFM images of a sample composed of the human Ferritin-H mRNA short 5’UTR deposited onto a mica surface. (A) The two-dimensional view of the sample surface, and (B) the three-dimensional view of the sample surface.
3.2.2. SHAPE data

The human Ferritin-H mRNA short 5’UTR transcripts were folded in a solution containing 6 mM Mg\(^{2+}\) and subjected to the SHAPE procedure. The SHAPE experiments were repeated at least twice. Figure 13 illustrates one set of SHAPE electropherograms of this RNA. Figure 14 presents the averaged SHAPE reactivities from two SHAPE experiments. The predicted structure with the lowest free energy for the human Ferritin-H mRNA short 5’UTR is shown in Figure 15.

Human Ferritin-H mRNA short 5’UTR forms the correct IRE stem-loop structure at nucleotide positions 61-88, including a 73CAGUGC78 apical loop. It is known that C73 pairs with G77 within the loop, which is required for efficient IRP protein binding.\(^{12}\) Figure 16 shows that G77 is unreactive (i.e., low SHAPE reactivity) due to its involvement in the base pairing. The 65UGC67 and C84 nucleotides form a bulge in the RNA, which is a conserved feature for the ferritin mRNAs. The G66-C84 base pair is also required for IRP binding. The crystal structure of the IRP1-ferritin IRE complex resolved by Volz’s research group has shown that the L-shaped IRP1 interacts with the ferritin IRE stem-loop at two positions, the apical loop and the bulge.\(^{85,86}\)

SHAPE data from the human ferritin-H mRNA short 5’UTR demonstrate that the left side of the stem (68UUCAA72) of the IRE is more reactive than the right side of the stem (83AGGUU79), which means that the left side may be adopting a conformation that brings it toward the outside of the RNA, where it would be more accessible to 1M7.
Figure 13. SHAPE data for the 5’ UTR of the human Ferritin-H mRNA short transcript. The raw output from the ABI 310 genetic analyzer underwent baseline adjustment, matrixing, and mobility shift adjustment. Fluorescence-labeled DNA primer was annealed to the 3’-end of this 207-nucleotide RNA. The blue trace is the +1M7 sample; the black trace is the -1M7 negative control; green and red traces are two dideoxy ladders (ddG and ddA, respectively). The x-axis indicates the number of data points collected. The y-axis (not shown) enumerates the fluorescence units.
Figure 14. Single nucleotide resolution of normalized SHAPE reactivities for the human Ferritin-H mRNA short 5’UTR. Error bars represent standard deviations calculated from two independent experiments.
Figure 15. The predicted secondary structure of the 5’UTR of the human Ferritin-H mRNA short transcript. Nucleotides are colored black (unreactive, SHAPE reactivity <0.3), orange (moderately reactive, 0.3 ≤ SHAPE reactivity <0.7), red (highly reactive, SHAPE reactivity ≥0.7), or grey (no data).
3.3. Mouse Ferritin-H mRNA 5’UTR

The sequence of the 169-nucleotide-long mouse Ferritin-H mRNA 5’UTR is as follows:

5’-GG CAGACGUUCU CGCCAGAGU CGCCGCGGUU UCCUGCUUCA
ACAGUGCUUG AACGGAACCC GGUGCUCGAC CCCUCCGACC CCCGCCCGGC
GCUUUCGAGCC UGAGCCCUUU GCAACUUCGU CGUUCCGCCG CUCCAGCGUC
GCCAGCCGC CUGCCCGCGC CGCCACC-3’

3.3.1. AFM images

An AFM image of the mouse Ferritin-H mRNA 5’UTR is shown in Figure 16. The 169-nucleotide-long RNA molecule folds into a complex three-dimensional structure that appears as a blob on the mica surface. The heights of five blobs were measured as: 0.73 nm, 0.59 nm, 0.53 nm, 0.48 nm, and 0.30 nm (average ± standard deviation = 0.53 ± 0.16 nm). The volumes of the same five blobs were 1136 nm³, 851 nm³, 555 nm³, 468 nm³, and 680 nm³, respectively (average ± standard deviation = 738 ± 265 nm³).
Figure 16. AFM images of a sample composed of the mouse Ferritin-H mRNA 5'UTR deposited onto a mica surface. (A) The two-dimensional view of the sample surface, and (B) the three-dimensional view of the sample surface.
3.3.2. SHAPE data

The mouse Ferritin-H mRNA 5’UTR transcripts were folded in a solution containing 6 mM Mg$^{2+}$ and subjected to the SHAPE procedure. The SHAPE experiments were repeated at least twice. Figure 17 illustrates one set of SHAPE electropherograms of this RNA. Figure 18 presents the averaged SHAPE reactivities from two SHAPE experiments. The predicted structure with the lowest free energy for the mouse Ferritin-H mRNA 5’UTR is shown in Figure 19.

Mouse Ferritin-H mRNA 5’UTR forms the correct IRE stem-loop structure at nucleotide positions 29-61, including a 44CAGUGC49 apical loop. The 36UGC38 and C55 nucleotides form a bulge in the RNA. SHAPE data from the mouse ferritin-H mRNA 5’UTR demonstrate that nucleotides in the apical loop and the bulge of the RNA are reactive; nucleotides in the stem of the IRE are unreactive.
Figure 17. SHAPE data for the 5’UTR of the mouse Ferritin-H mRNA transcript. The raw output from the ABI 310 genetic analyzer underwent baseline adjustment, matrixing, and mobility shift adjustment. Fluorescence-labeled DNA primer was annealed to the 3’-end of this 169-nucleotide RNA. The blue trace is the +1M7 sample; the black trace is the -1M7 negative control; green and red traces are two di deoxy ladders (ddG and ddA, respectively). The x-axis indicates the number of data points collected. The y-axis (not shown) enumerates the fluorescence units.
Figure 18. Single nucleotide resolution of normalized SHAPE reactivities for the mouse Ferritin-H mRNA 5′ UTR. Error bars represent standard deviations calculated from two independent experiments.
Figure 19. The predicted secondary structure of the 5’UTR of the mouse Ferritin-H mRNA transcript. Nucleotides are colored black (unreactive, SHAPE reactivity <0.3), orange (moderately reactive, 0.3 ≤ SHAPE reactivity <0.7), red (highly reactive, SHAPE reactivity ≥0.7), or grey (no data).
3.4. Human Ferritin-L mRNA 5'UTR

The sequence of the 217-nucleotide-long RNA, which contains the 201-nucleotide-long human Ferritin-L mRNA 5’UTR plus 16 nucleotides that extend into the coding region of the human Ferritin-L mRNA, is shown below.

5’-GG GCAGUUCGGC GGUCGCCGCGG GUCUGUCUCU UGCUUCAACA GUGUUUGGAC GGAACAGAUC CGGGGACUCU CUUCCAGCCU CCGACCAGCCC UCCGAUUUCC UCUCGGCUUG CAACCUCGGG GACCAUCUUC UCGGCCAUCU CCUGCUUCUG GGACCUGGCA GCACCGUUUU UGUGGUUAGC UCCUUCUUGC CAACCAACCA UGAGCUCCCA GAUUC-3’

3.4.1. AFM images

An AFM image of the human Ferritin-L mRNA 5’UTR is shown in Figure 20. The RNA molecule folds into a complex three-dimensional structure that appears as a blob on the mica surface. The heights of fourteen blobs were measured as: 1.22 nm, 1.35 nm, 1.33 nm, 0.55 nm, 1.23 nm, 0.81 nm, 0.77 nm, 0.93 nm, 0.88 nm, 0.83 nm, 0.94 nm, 1.35 nm, 1.19 nm, and 0.81 nm (average ± standard deviation = 1.01 ± 0.26 nm). The volumes of the same fourteen blobs were 3355 nm³, 4407 nm³, 5845 nm³, 943 nm³, 2711 nm³, 768 nm³, 1241 nm³, 1671 nm³, 1404 nm³, 1463 nm³, 2780 nm³, 1384 nm³, 3212 nm³, and 1241 nm³, respectively (average ± standard deviation = 2316 ± 1481 nm³).
Figure 20. AFM images of a sample composed of the human Ferritin-L mRNA 5’UTR deposited onto a mica surface. (A) The two-dimensional view of the sample surface, and (B) the three-dimensional view of the sample surface.
3.4.2. SHAPE data

The human Ferritin-L mRNA 5’UTR transcripts were folded in a solution containing 6 mM Mg$^{2+}$ and subjected to the SHAPE procedure. The SHAPE experiments were repeated at least twice. The predicted structure with the lowest free energy for the human Ferritin-L mRNA 5’UTR is shown in Figure 21. Human Ferritin-L mRNA 5’UTR forms the correct IRE stem-loop structure at nucleotide positions 27-57, including a 41CAGUGU46 apical loop. The 33UGC35 and C52 nucleotides form a bulge in the RNA.

**Figure 21.** The predicted secondary structure of the 5’UTR of the human Ferritin-L mRNA transcript. Nucleotides are colored black (unreactive, SHAPE reactivity < 0.3), orange (moderately reactive, 0.3 ≤ SHAPE reactivity < 0.7), red (highly reactive, SHAPE reactivity ≥ 0.7), or grey (no data).
3.5. Mouse Ferritin-L mRNA 5'UTR

The sequence of the 207-nucleotide-long mouse Ferritin-L mRNA 5'UTR is as follows:

5'-GA GCAGCGCCUU GGAGGUCCCG UGGAUCUGUG UCUUGCUUCA ACAGUGUUUG AACGGAACAG ACCCGGGGAA UCCACUGUA CUCGCUUCCA GCCGCCUUUA CAAGUCUCUC CAGUCGCAGC CUCCGGGACC AUCUCCUCGC UGCCUUCAGC UCCUAGGACC AGUCUGCACC GUCUCUUCGC GGUUAGCUCC UACUCCGGAU CAGCC-3'

3.5.1. AFM images

An AFM image of the mouse Ferritin-L mRNA 5'UTR is shown in Figure 22. The RNA molecule folds into a complex three-dimensional structure that appears as a blob on the mica surface. The heights of fourteen blobs were measured as: 0.78 nm, 0.92 nm, 1.04 nm, 0.64 nm, 0.83 nm, 0.65 nm, 0.90 nm, 0.87 nm, 1.02 nm, 0.96 nm, 0.61 nm, 0.81 nm, 0.90 nm, and 0.70 nm (average ± standard deviation = 0.83 ± 0.14 nm). The volumes of the same fourteen blobs were 1103 nm$^3$, 1183 nm$^3$, 1022 nm$^3$, 782 nm$^3$, 828 nm$^3$, 709 nm$^3$, 952 nm$^3$, 1040 nm$^3$, 933 nm$^3$, 1220 nm$^3$, 486 nm$^3$, 749 nm$^3$, 1365 nm$^3$, and 619 nm$^3$, respectively (average ± standard deviation = 928 ± 247 nm$^3$).
Figure 22. AFM images of a sample composed of the mouse Ferritin-L mRNA 5’UTR deposited onto a mica surface. (A) The two-dimensional view of the sample surface, and (B) the three-dimensional view of the sample surface.
The mouse Ferritin-L mRNA 5’UTR transcripts were folded in a solution containing 6 mM Mg$^{2+}$ and subjected to the SHAPE procedure. The SHAPE experiments were repeated at least twice. The predicted structure with the lowest free energy for the mouse Ferritin-L mRNA 5’UTR is shown in Figure 23. Mouse Ferritin-L mRNA 5’UTR forms the correct IRE stem-loop structure at nucleotide positions 30-60, including a 44CAGUGU49 apical loop. The 36UGC38 and C55 nucleotides form a bulge in the RNA.

Figure 23. The predicted secondary structure of the 5’UTR of the mouse Ferritin-L mRNA transcript. Nucleotides are colored black (unreactive, SHAPE reactivity <0.3), orange (moderately reactive, 0.3 ≤ SHAPE reactivity <0.7), red (highly reactive, SHAPE reactivity ≥0.7), or grey (no data).
3.6. Human APP mRNA 5’UTR

The sequence of the 163-nucleotide-long RNA, which contains the 149-nucleotide-long human APP mRNA 5’UTR plus 14 nucleotides that extend into the coding region of the human APP mRNA, is shown below.

5’-GG AGUUUCCUCG GCAGCGGUAG GCGAGAGCAC GCGGAGGAGCGUGCGCGGGG GCCCCGGGAG ACGGCGGCGG UGGCGGGCGG GCAGAGCAA GGACGCGGCG GAUCCCACUC GCACAGCAGC GCACUCGGUG CCCCGCGCAG GGUCGCG AUGCUGCCCCG GUUU-3’

3.6.1. AFM images

An AFM image of the human APP mRNA 5’UTR is shown in Figure 24. The RNA molecule folds into a complex three-dimensional structure which appears as a blob on the mica surface. The heights of three blobs were measured as: 0.37 nm, 0.40 nm, and 0.35 nm (average ± standard deviation = 0.37 ± 0.03 nm). The volumes of the same three blobs were 476 nm³, 546 nm³, and 408 nm³, respectively (average ± standard deviation = 477 ± 69 nm³).
Figure 24. AFM images of a sample composed of the human APP mRNA 5’ UTR deposited onto a mica surface. (A) The two-dimensional view of the sample surface, and (B) the three-dimensional view of the sample surface.
3.6.2. SHAPE data

The human APP mRNA 5’UTR transcripts were folded in a solution containing 6 mM Mg$^{2+}$ and subjected to the SHAPE procedure. The SHAPE experiments were repeated at least twice. Figure 25 illustrates one set of SHAPE electropherograms of this RNA. Figure 26 presents the averaged SHAPE reactivities from two SHAPE experiments. The predicted structure with the lowest free energy for the human APP mRNA 5’UTR is shown in Figure 27.

The human APP mRNA 5’UTR contains an IRE stem-loop structure that is different from the IREs in the 5’UTRs of the Ferritin-H and Ferritin-L mRNAs; therefore, it is named IRE-Type II. Instead of having a 6-nucleotide apical loop comprised of CAGUGN (N can be A, C, or U, but not G), as present in the Ferritin IREs, the human APP IRE contains a 13-nucleotide loop comprised of nucleotides 83GGCAGAGCAAGGA95 (see Figure 27). The 85CAGA88 sequence in the apical loop is called an “amyloid” CAGA box. The mammal CAGA box is a Smad protein binding site for Smad3 and Smad4.87 The “amyloid” CAGA box within the IRE region of the human APP mRNA 5’UTR is believed to provide additional regulation of the synthesis of the human APP protein and to facilitate the production of Aβ.87

In addition to the difference in the apical loop, nucleotides 79CGCG82 in the human APP mRNA 5’UTR pair with nucleotides 99GC96 to form a 4-base-pair stem, whereas
IREs in the ferritin mRNA 5’UTRs have a conserved 5-base-pair stem. In addition, instead of having a UGC-C type of bulge, the human APP IRE contains a 78G-G100 bulge.

In the human APP mRNA 5’UTR, most nucleotides within positions 101-123 are reactive, which means that these nucleotides may adopt a conformation that makes them more accessible to 1M7. Interestingly, this RNA region is located within the interleukin-1 responsive element (called the acute box) domain, where interleukin-1, a cytokine released early in the acute phase of the immune response, binds in order to regulate and stimulate APP protein synthesis.

![Figure 25. SHAPE data for the 5’UTR of the human APP mRNA transcript. The raw output from the ABI 310 genetic analyzer underwent baseline adjustment, matrixing, and mobility shift adjustment. Fluorescence-labeled DNA primer was annealed to the 3’-end of this 163-nucleotide RNA. The blue trace is the +1M7 sample; the black trace is the -1M7 negative control; green and red traces are two dideoxy ladders (ddG and ddT, respectively).](image-url)
The x-axis indicates the number of data points collected. The y-axis (not shown) enumerates the fluorescence units.

**Figure 26.** Single nucleotide resolution of normalized SHAPE reactivities for the human APP mRNA 5’ UTR. Error bars represent standard deviations calculated from two independent experiments.
Figure 27. The predicted secondary structure of the 5’U TR of the human APP mRNA transcript. Nucleotides are colored black (unreactive, SHAPE reactivity <0.3), orange (moderately reactive, 0.3 ≤ SHAPE reactivity <0.7), red (highly reactive, SHAPE reactivity ≥0.7), or grey (no data).
3.7. Mouse APP mRNA 5’UTR

The sequence of the 147-nucleotide-long mouse APP mRNA 5’UTR is as follows:

5’-GG GUUUCCUCGG CGGCYGGGAGG CGAGAGCACC GGGAGCAGAG
CGAGCGCGGG GCCACCGGAG ACGGCGGCGG CGGCGCGGAC ACAGCCAGGG
CGCGGCCGGAU CUUCCACUCG CACACGGAGC ACUCGGUGGC CCACGCAGGA
UCACG-3’

3.7.1. AFM images

An AFM image of the mouse APP mRNA 5’UTR is shown in Figure 28. The RNA molecule folds into a complex three-dimensional structure that appears as a blob on the mica surface. The heights of five blobs were measured as: 2.60 nm, 2.90 nm, 2.50 nm, 2.90 nm, and 1.90 nm (average ± standard deviation = 2.56 ± 0.41 nm). The volumes of the same five blobs were 2509 nm³, 1881 nm³, 1054 nm³, 2006 nm³, and 1162 nm³, respectively (average ± standard deviation = 1722 ± 609 nm³).
Figure 28. AFM images of a sample composed of the mouse APP mRNA 5’UTR deposited onto a mica surface. (A) The two-dimensional view of the sample surface, and (B) the three-dimensional view of the sample surface.
3.7.2. SHAPE data

The mouse APP mRNA 5’UTR transcripts were folded in a solution containing 6 mM Mg$^{2+}$ and subjected to the SHAPE procedure. The SHAPE experiments were repeated at least twice. Figure 29 illustrates one set of SHAPE electropherograms of this RNA. Figure 30 presents the averaged SHAPE reactivities from two SHAPE experiments. The predicted structure with the lowest free energy for the mouse APP mRNA 5’UTR is shown in Figure 31.

The IRE-like stem-loop structure in the mouse APP mRNA 5’UTR contains a 9-nucleotide apical loop comprised of nucleotides 77GCGGACACA85, which does not contain an “amyloid” CAGA sequence. Nucleotides 74GGC76 pair with nucleotides 88CCG86 to form a 3-base-pair stem (see Figure 31). Rather than having the UGC-C type of bulge present in the ferritin IREs, the predicted mouse APP IRE structure contains a 6-nucleotide internal loop.
Figure 29. SHAPE data for the 5’UTR of the mouse APP mRNA transcript. The raw output from the ABI 310 genetic analyzer underwent baseline adjustment, matrixing, and mobility shift adjustment. Fluorescence-labeled DNA primer was annealed to the 3’-end of this 147-nucleotide RNA. The blue trace is the +1M7 sample; the black trace is the -1M7 negative control; green and red traces are two dideoxy ladders (ddG and ddA, respectively). The x-axis indicates the number of data points collected. The y-axis (not shown) enumerates the fluorescence units.
Figure 30. Single nucleotide resolution of normalized SHAPE reactivities for the mouse APP mRNA 5’ UTR. Error bars represent standard deviations calculated from two independent experiments.
Figure 31. The predicted secondary structure of the 5’UTR of the mouse APP mRNA transcript. Nucleotides are colored black (unreactive, SHAPE reactivity <0.3), orange (moderately reactive, 0.3 ≤ SHAPE reactivity <0.7), red (highly reactive, SHAPE reactivity ≥0.7), or grey (no data).
CHAPTER 4

Discussion

The AFM study of RNA conformation was not very successful. The length of the seven RNA transcripts investigated was between 140-240 nucleotides. Due to the relatively short length of the RNA molecules and their complex three-dimensional folding, AFM imaging only shows them as blobs. The difference in volume and height of the blobs representing these RNA molecules is somewhat random (see Table 1); no correlation has been observed between volume or height and RNA sequence length in this study. The day-to-day variation of the AFM instrument and tip conditions are considered to be important factors relative to these results.

In addition to the differences between RNA molecules of different types, the volume and height of the blobs for the same RNA molecule deposited onto the same mica surface show large variations (see standard deviations, SD, in Table 1). There are several possible explanations for the differences in height and volume of the blobs for the same RNA molecule, including 1) some blobs on the mica surface were RNA aggregates; 2) some blobs were degraded RNA molecules; and 3) the RNA molecules adopted different conformations on the mica surface. Overall, this study shows that AFM imaging is not an efficient way to visualize relatively short RNA molecules unless there are some distinct features present. For example, Jaeger’s research group visualized the dimerization of a 230-nucleotide RNA
molecule via “kissing-loop” interaction; the AFM images showed double or triple blobs caused by these “kissing-loop” structures.\textsuperscript{75}

**Table 1.** Summary of the averaged height and volume of the blobs in AFM images for each RNA molecule investigated (“nts” = nucleotides).

<table>
<thead>
<tr>
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</thead>
<tbody>
<tr>
<td>Height of the blob in nm (Average ± SD)</td>
<td>0.33±0.20</td>
<td>0.65±0.26</td>
<td>0.53±0.16</td>
<td>1.01±0.26</td>
<td>0.83±0.14</td>
<td>0.37±0.03</td>
<td>2.56±0.41</td>
</tr>
<tr>
<td>Volume of the blob in nm(^3) (Average ± SD)</td>
<td>356±360</td>
<td>1162±732</td>
<td>738±265</td>
<td>2316±1481</td>
<td>928±247</td>
<td>477±69</td>
<td>1722±609</td>
</tr>
</tbody>
</table>

SHAPE chemistry was used to map the secondary structures of the seven RNA transcripts at single nucleotide resolution. To our knowledge, this is the first time that the secondary structures of the 5’UTRs of APP mRNA from human and mouse have been experimentally mapped. Several research groups have used traditional X-ray crystallization techniques or enzymatic cleavage and/or chemical modification approaches to determine the conformation of the IRE region (30-50 nucleotides) in the Ferritin-H and Ferritin-L mRNA 5’UTR, in the presence and absence of small molecule or protein ligands, but none of them
determined the structure of the entire 5′UTR of these RNA molecules. This study paves the way for the further investigation of RNA-ligand interactions in the 5′UTR of IRE-containing mRNAs.

The seven RNA molecules show a high degree of secondary structure, with numerous stem-loops structures. The Human Ferritin-H mRNA short 5′UTR, mouse Ferritin-H mRNA 5′UTR, human Ferritin-L mRNA 5′UTR, and mouse Ferritin-L mRNA 5′UTR form the expected IRE stem-loop structure, which includes a CAGUGN apical loop (N can be A, C, or U, but not G). The first nucleotide C pairs with the fifth nucleotide G within the loop for efficient IRP protein binding. In addition to the apical loop, these IRE structures in the 5′UTR of the Ferritin mRNAs also contain a conserved 5-base-pair stem, followed by a UGC-C type of bulge. The G-C base pair in the bulge is required for IRP binding.

The APP mRNA 5′UTR in human and mouse contains an IRE stem-loop structure that is different from the IREs in the 5′UTRs of the Ferritin-H and Ferritin-L mRNAs. Instead of having a 6-nucleotide apical loop comprised of CAGUGN, where N can be A, C, or U, but not G, the human APP IRE contains a 13-nucleotide loop comprised of the nucleotides GGCAGAGCAAGGA. The mouse APP IRE contains a 9-nucleotide apical loop comprised of the nucleotides GCGGACACA. The human APP IRE contains a 4-base-pair stem, followed by a G-G bulge. The mouse APP IRE contains a 3-base-pair stem, followed by a 6-nucleotide internal loop.
In addition to the IRE stem-loop, the 5’UTR of the Ferritin-H, Ferritin-L, and APP mRNAs also contain an IL-1β-dependent acute box.\textsuperscript{92-95} The acute boxes in the Ferritin mRNA 5’UTRs are located at the 3’-end of the 5’UTR. For example, the core region of the acute box in the human Ferritin-H mRNA 5’UTR includes nucleotides 191CGCCGCGCAGCCACCGCCGCCGCCG215, where little structural information was obtained. This occurred because the fluorescently labeled DNA primer was annealed to the 3’-end of the RNA molecule for the SHAPE procedure. During primer extension, the large primer peaks mask the structural information of the neighboring (i.e., the 3’-end) nucleotides.

The core region of the acute box in the human Ferritin-L mRNA 5’UTR is located at nucleotides U111 through G136, which contains a single-stranded region and a stem-loop structure. The core region of the acute box in the human APP mRNA 5’UTR is located at positions 101-125, where most nucleotides are reactive.

The SHAPE technique was successfully used to map the secondary structures of the seven RNA molecules in the absence of the ligand. The next step of our study is to determine the secondary structures of the RNA molecules in the presence of different ligands at various concentrations. A slab gel electrophoresis system will be used instead of capillary gel electrophoresis to separate and size radio-labeled cDNAs. This will allow us to run 20-30 SHAPE reactions in parallel on one gel, and to visualize the gradual structural transition undergone by the RNA. The transition mid-point will be determined and the ligand binding site will be predicted.
CHAPTER 5

Conclusions

In order to substantially improve the understanding of ligand-RNA binding mechanisms and to shed light on the design of novel molecules targeted to specific RNA motifs, a complete elucidation of the RNA conformation before, during, and after ligand binding is necessary. Tens of thousands of compounds have been screened to identify ligands that can bind to the 5’UTR of the human APP mRNA, which would allow the treatment of Alzheimer’s disease by decreasing the production of APP and the corresponding Aβ; fifteen compounds were identified as inhibiting APP translation by interacting with the 5’UTR of APP mRNA, which includes an IRE stem-loop structure. However, the underlying mechanism(s) of these RNA-ligand interactions has not been determined. For example, what conformation does APP mRNA 5’UTR adopt after the binding of a small molecule? Which nucleotides do the ligands bind to in the IRE? Do the ligands also bind to other regions in the 5’UTR of the human APP mRNA, such as the interleukin-1-responsive acute box? If the molecules mentioned above have the capability to reduce APP expression, can they bind to the IRE region in the 5’UTR of the human Ferritin-H and Ferritin-L mRNA to inhibit the translation of these intracellular iron storage proteins as well? It is an advantage if the ligands can down-regulate APP expression, but certainly a disadvantage if they down-regulate the expression of ferritin.
The objective of this study was to determine the conformations of the 5’UTRs of Ferritin-H, Ferritin-L, and APP mRNA transcripts from two species (human and mouse) in the absence of the ligand. We used AFM to visualize the conformation of these RNA molecules, and determined the secondary RNA structures using SHAPE analysis coupled with a fluorescence detection system. To our knowledge, this is the first time that the secondary structures of the 5’UTRs of APP mRNA in human and mouse have been experimentally mapped. The AFM imaging did not provide high resolution structural information about these RNAs, whereas the SHAPE procedure successfully interrogated the secondary RNA structures at single nucleotide resolution. This study paves the way for the further investigation of RNA-ligand interactions in these RNA molecules.
References


63. Bandyopadhyay, S., Ni, J., Ruggiero, A., Walshe, K., Rogers, M.S., Chattopadhyay, N., Glicksman, M.A. & Rogers, J.T. A high-throughput drug screen targeted to the


