Structural characterization of NORAD reveals a stabilizing role of spacers and two new repeat units

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Long non-coding RNAs (lncRNAs) can perform a variety of key cellular functions by interacting with proteins and other RNAs. Recent studies have shown that the functions of lncRNAs are largely mediated by their structures. However, our structural knowledge for most lncRNAs is limited to sequence-based computational predictions. Non-coding RNA activated by DNA damage (NORAD) is an atypical lncRNA due to its abundant expression and high sequence conservation. NORAD regulates genomic stability by interacting with proteins and microRNAs. Previous sequence-based characterization has identified a modular organization of NORAD composed of several NORAD repeat units (NRUs). These units comprise the protein-binding elements and are separated by regular spacers. Here, we experimentally determine for the first time the secondary structure of NORAD using the nextPARS approach. Our results suggest that the spacer regions provide structural stability to NRUs. Furthermore, we uncover two previously unreported NRUs, and determine the core structural motifs conserved across NRUs. Overall, these findings will help to elucidate the function and evolution of NORAD.

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1. Introduction

Long non-coding RNAs (lncRNAs) constitute a heterogeneous family of RNA transcripts defined by their length (>200 nucleotides (nt)), and their lack of protein-coding potential. Most lncRNAs are expressed at low levels and in a tissue- or cell type-specific manner, and are poorly conserved both at the sequence and exon–intron structure levels [1,2]. Over the last years, an increasing number of lncRNAs have been associated with a variety of cellular and developmental functions [3-5], and have been shown to be altered in various human diseases [6]. However, the functions and structures of the vast majority of lncRNAs, remain poorly understood.

LncRNAs can form complex secondary structures, which tend to be more conserved than primary sequences and are thought to mediate their molecular functions [7,8]. However, controversy remains as to whether secondary structures of lncRNAs are evolutionarily conserved [9]. Although molecular probing techniques allow assessing RNA structures folded in vitro, the large size of many lncRNAs poses a significant challenge. As a result, only a few lncRNA structures have been experimentally characterized in vitro [9], including those of HOTAIR [10], RepA [11], Xist [12,13], ncSRA [14], COOLAIR [15] and lincRNAp21 [14,16].

Many sequences, including riboswitches, mRNAs and lncRNAs [9], are expected to fold into a variety of competing structures that can coexist. To account for this structural diversity, several methods have been recently developed that allow studying alternative RNA conformations using experimental data [17-21].

The non-coding RNA activated by DNA damage (NORAD) is a 5.3 kilobases (kb) lncRNA that is widely expressed and conserved across mammals. Previous sequence analyses of this lncRNA have described the presence of several repeated elements - NORAD repeat units or NRUs - that are predicted to fold into similar secondary structures and tend to be conserved [22,23]. NORAD maintains genomic stability by sequestering RNA-binding PUMILIO (PUM) proteins (PUM1 and PUM2), thereby preventing them from interacting with their targets [24]. PUM proteins act by repressing the translation of specific target mRNAs by binding to PUMILIO response elements (PRE) present in their 3’ untranslated region (3’UTR) [25;25,26]. In the absence of NORAD, PUM proteins drive chromosomal instability caused by DNA repair, hyperactively repressing mitotic, and DNA replication factors [22]. NORAD...
interacts with RBMX and controls the ability of RBMX to assemble a ribonucleoprotein complex (NARC1), important to NORAD function. Although it was initially reported that NORAD regulates genome stability through NORAD:RBMX interaction [27], it was recently shown that PUM binding with NORAD is required for genome stability whereas NORAD:RBMX interaction is not essential for this function [24]. Moreover, it was shown that deletion of the NORAD mouse ortholog resulted in genome instability and premature aging [28].

In addition, NORAD acts as a competitive endogenous RNA (ceRNA) and regulates cancer progression by sponging MicroRNAs (miRNAs) [29,30]. miRNAs are small regulatory RNAs of about ~20 nt in length that play critical regulatory roles in eukaryotic cells where they mediate post-transcriptional silencing of specific genes [31-33].

Despite the increasing relevance and known functions of NORAD, the experimental characterization of its structure is still lacking. To shed new light on the relationships between structure, function and evolution of NORAD, we here characterized its secondary structure using experimental probing with the NextPARS approach [34]. In addition, we assessed how changes in sequence context and shifts in temperature impact the structure of NRUs and other relevant NORAD motifs. Our results provide evidence that the spacer sequences provide structural stability to NRUs and that the structure around conserved SAM68-binding sites is more robust to thermal shifts than other positions in the NORAD sequence. Finally, we define the core structural motifs conserved across NRUs, and uncover the presence of two previously unreported NRUs.

2. Materials and methods

2.1. Sample preparation

The structure of several NORAD fragments was experimentally probed using NextPARS (PMID: 29358234) at three different temperatures (23 °C, 37 °C and 55 °C) by spiking the fragments of interest into fungal total or polyA+ RNA samples.

The probed NORAD fragments were the following: three RNA fragments covering the full-length NORAD lncRNA (called NORAD#1, NORAD#2 and NORAD#3, overlapping 1903, 1862 and 1614 bases, respectively), and 11 NORAD repetitive units previously described [23] with their surrounding regions (NRU#1, NRU#2, NRU#3, NRU#4, NRU#6 NRU#7, NRU#8, NRU#9, NRU#10, NRU#11 and NRU#12, overlapping 179 nt each) [Supplementary Table S1]. To produce these fragments, NORAD was cloned into the pUC57 vector by GenScript Biotech (Netherlands). Once the plasmid preparation was obtained, PCRs were performed to amplify and linearize the different NORAD fragments using the Phusion High-Fidelity DNA Polymerase mix (#2021, DongSheng Biotech) and a 0.4 μM final concentration of each primer (forward and reverse) in a total volume of 40 μl.

Supplementary Table S1 shows primer sequences and amplicon sizes and Supplementary Table S2 specifies PCR conditions used per each fragment. PCR amplicons were purified using a QiAquick PCR purification kit following the manufacturer’s instructions (Qiagen). Quality controls with NanoDrop 1000 Spectrophotometer (Thermo Scientific) and Agilent 2100 Bioanalyzer with the RNA 6000 Pico LabChip® Kit (Agilent) were performed before and after the digestion, to confirm that samples were not digested with the initial denaturation or over-digested after the enzymatic treatment.

2.2. Secondary structure probing with NextPARS

We used our previously implemented NextPARS technique [34] to probe the changes in the RNA structures at different temperatures, adapting the original protocol when necessary. We tested three different temperatures (23 °C, 37 °C and 55 °C), which were chosen based on a previous work [35], with the aim to obtain a broad picture of the RNA structure transitions due to structure perturbations caused by thermal shifts. Briefly, 2 μg of PolyA+ RNA or total RNA were mixed with 20 ng of each NORAD RNA fragment and were brought to a final volume of 80 μl with nuclease-free water. Samples were first denatured at 90 °C for 2 min and immediately cooled down to 4 °C on ice for 2 min. Then, 10 μl of ice-cold 10X RNA structure buffer (Ambion) were added to the samples and mixed by pipetting up and down several times. When probing at 23 °C, RNA samples were subsequently brought from 4 °C to 23 °C, in about 20 min (1 °C per min). For samples probed at 37 °C and 55 °C, the temperature was brought from 4 °C to 30 °C (1 min per each °C), samples were incubated 5 min at 30 °C, and then brought to 37 °C and 55 °C, respectively, for enzymatic digestion. Finally, 10 μl of the corresponding dilutions of RNase V1 (Ambion), S1 nuclease (Fermentas), or nuclease-free water were added to the V1-digested, S1-digested and undigested samples, respectively. The enzyme concentration used was different depending on the probing temperature, to compensate for the effect of higher temperatures to prevent over-digestion of the RNA samples and according to previous studies [35]. Thus, the final reaction contained 0.03 U, 0.015 U or 0.0075 U of RNase V1, and 200 U, 100 U or 50 U of S1 nuclease for samples probed at 23 °C, 37 °C or 55 °C, respectively. After mixing by pipetting, samples were incubated at each corresponding temperature for 15 min more. The RNAs were purified using RNeasy MiniElute Cleanup kit following the manufacturer’s instructions (Qiagen). Quality controls with NanoDrop 1000 Spectrophotometer (Thermo Scientific) and Agilent 2100 Bioanalyzer with the RNA 6000 Pico LabChip® Kit (Agilent) were performed before and after the digestion, to confirm that samples were not digested with the initial denaturation or over-digested after the enzymatic treatment.

2.3. Library preparation and sequencing.

We used the TruSeq Small RNA Sample Preparation Kit (Illumina) to prepare the libraries for further sequencing. We initially performed a phosphatase treatment incubating 16 μl of the digested samples for 30 min at 37 °C and 5 min at 65 °C mixed with 2.5 μl of 10X phosphatase buffer, 2.5 μl of nuclease-free water, 1 μl of RNase inhibitor and 3 μl of Antarctic phosphatase (New England Biolabs Inc.). We then performed a kinase treatment adding 4 μl of T4 Polynucleotide Kinase (PNK, New England BioLabs Inc.), 5 μl of 10X PNK buffer, 10 μl of ATP 10 mM, 1 μl of RNase inhibitor and nuclease-free water up to a total volume of 50 μl, and incubating the samples 1 h at 37 °C. Samples were then purified using RNeasy MiniElute Cleanup kit following manufacturer’s instructions (Qiagen) with a 10 μl RNase-free water final elution step. Samples were concentrated using a centrifugal evaporator SpeedVac™ to a final volume of 5 μl and we started the TruSeq Small RNA Sample Preparation Kit (Illumina) protocol according to manufacturer’s instructions. After the final purification of the cDNA libraries, we performed quality control of each library using Agilent 2100 bioanalyzer with the DNA 1000 Kit (Agilent). Libraries were sequenced...
in single-reads with read lengths of 50 nucleotides in Illumina HiSeq2500 sequencers at the Genomics Unit of the CRG.

2.4. Computation of the nextPARS scores

The computation of the nextPARS scores was obtained following the protocol in [34,36]. Briefly, we removed adapters with cutadapt and mapped the reads using STAR [37]. Then, we parse the mapped reads and retrieve the 5′-end position in the NORAD transcript. This results in a digestion profile that indicates the number of cuts per position of the transcript. To obtain the scores from nextPARS probing experiments we used as an input the digestion profile. The output contains a structural profile of the NORAD transcript (single-or double-stranded state), with a score for each nucleotide that ranges from −1.0 (highest preference for single strand) to 1.0 (highest preference for double-strand). The code to obtain the structural profile from nextPARS experiments is available on github at https://github.com/Gabaldonlab/nextPARS_docker.

2.5. RNA secondary structure prediction and visualization

Using the nextPARS2SHAPE v1.0 script (https://github.com/Gabaldonlab/MutiFolds/blob/master/scripts/nextPARS2SHAPE.py), we converted the nextPARS score to SHAPE-like normalized reactivities that were used for the secondary structure prediction. Then, to obtain the secondary structure of NORAD and NRUs, we use the Fold software (Version 6.2) from RNAstructure package [38], using pseudo energy restraints. Residues for which there was no nextPARS data were assigned a reactivity of −999, as suggested by the Fold manual. RNA structures plots were constructed using the command-line VARNA (Version 3–93) [39] using the structure from the dot-bracket files generated with Fold.

The consensus structure models have been generated using the RNAalifold software (Version 2.4.15). We obtained similar results with TurboFold, LocARNA and Foldalign software. R2R software then was used for the visualization of the consensus structure (supplementary data) [40].

2.6. Identification of repetitive units in NORAD

To identify new repetitive elements in NORAD we first performed a multiple sequence alignment with T-Coffee (Version_11.00.8&be486) [41], with the slow_pair method for global pairwise alignments. The sequence from the core regions (189 nt) of the 12 previously described NRUs (NRU#1 - NRU#12) was the T-Coffee input. Then, a covariance model (CM) was built using Infernal (Version 1.1.4) [42]. To construct the CM we use cmbuild with the alignment file generated with T-Coffee of the 12 NRUs sequences as the input, with n=10 parameter to ignore the secondary structure annotation. Then, we calibrated the CM for E-values with cmlcalibrate program with default parameters. Next, cmsearch program was used to search for the calibrated CM against NORAD full sequence. Finally, the 14 sequences below the default E-value cutoff (0.01) were retrieved, as they are considered to be reliable enough to be included.

2.7. Prediction of multiple folding conformations, guided by nextPARS data.

Rsample (Version 6.2) [20] was used to calculate the partition function for the NRUs from NORAD sequence to determine the multiple conformations. To perform a complete Rsample calculation, we performed the following steps. First, we converted the nextPARS score to SHAPE-like reactivities with nextPARS2SHAPE v1.0 script and then we ran Rsample (with default parameters) using the SHAPE-like constraint to produce a Partition Save File (PSF). After that, we ran stochastic, using this PSF file as input, and we created a CT file with Boltzmann ensemble of 1,000 structures. Finally, using the R script from Rsample (RsampleCluster.R) we use the CT file to calculate the optimal number of clusters and their centroids to model the NRUs sequences that populate multiple structures, including the relative probabilities of those structures. To evaluate conserved structural elements within the ensemble of structures of the NRUs, we used the Beagle (BEar Alignment Global and Local) software [43]. We performed an all-vs-all alignment for all the NRUs to identify common secondary structures and compared them with the most stable structure.

2.8. Statistical methods

Statistics were performed using the non-parametric Wilcoxon rank-sum and Kruskal-Wallis tests because the data are not normally distributed. The similarity between the same residues in NRU segments folded independently or within each piece and larger NORAD fragments was compared using the nextPARS scores to measure Pearson’s correlation (P < 0.05 was considered significant). The variability at different temperatures in positions of UAAA motifs was measured by Pearson’s correlation, using nextPARS data from isolated NRUs and NRUs in the context of larger NORAD fragments in combination with PhastCons score or Phylocp score at different conservation levels. The Gardner-Altman estimation plots were produced using the python version of dabest [44].

2.9. Deposited data

Raw sequencing data of nextPARS experiments were deposited in the Sequence Read Archive under the BioProject ID PRJNA714002.

3. Results

3.1. Secondary structure of NORAD and identification of subdomains

Resolving the structure of NORAD in vitro is exceptionally challenging because of the large size of this transcript. To circumvent this problem, we synthesized, cloned, and in vitro transcribed (see Materials and Methods) three overlapping NORAD fragments [Fig. 1A], and subsequently subjected them to enzymatic probing using the nextPARS method [34]. This approach renders single-nucleotide resolution measurements of in vitro secondary structure, represented by a structural profile (single- or double-stranded state). Providing a score for each residue, ranging from −1.0 (highest preference for single strand) to 1.0 (highest preference for double-strand). We converted the nextPARS score to SHAPE-like reactivity and used this to provide constraints for secondary structure prediction [Fig. 1B]. Our results show that most of the NRUs form independently-folded structures (NRU#2, NRU#3, NRU#5, NRU#7, NRU#8, NRU#9, NRU#11 and NRU#12) but some of them seem to interact with each other or with other NORAD regions [Fig. 1B].

To analyze NRUs structures in detail, and to assess the effect of the NORAD sequence context in NRU folding, we designed non-overlapping 179 nt long fragments covering 11 of the 12 previously described NRUs [23] and performed nextPARS experiments [Fig. 1A]. To identify structural subdomains in NORAD, we followed a strategy previously used with HOTAIR [10,45]. In this approach, the full-length lncRNA profile is compared to the profiles of individual RNA segments to identify potential independently-folded sub-domains. Comparing the nextPARS profile of the NRUs with
the same sequence in the context of the full NORAD molecule, shows a high correlation, with a p-value < 0.01 [Supplementary Figure S1]. Pearson’s correlation coefficient values ($q$) are expected to be higher for fragments that do not break elements of the secondary structure observed in the full-length molecule. This was the case for the eleven NRUs analyzed ($q > 0.8$). Correlations values were particularly high ($q > 0.9$) for some of the NRUs (NRU#1, NRU#4, NRU#8, NRU#10 and NRU#12), suggesting that these regions form more stable local structures [Supplementary Figure S1].

3.2. Spacer sequences provide structural stability to NRUs.

To evaluate the robustness and stability of NORAD secondary structure, we performed nextPARS experiments at three different temperatures: 23°C, 37°C and 55°C to study the alterations prompted by these temperature shifts as a means of structural perturbation [Fig. 2A]. We compared the nextPARS scores obtained in equivalent residues when probed in NRU-specific fragments or in larger NORAD fragments, and when probed at different temperatures in the context of the two molecular sizes. On average, the correlation between secondary structures was higher across varying temperatures than when NRUs structures were compared between those folded in isolated fragments with larger NORAD fragments [Fig. 2A,B]. These results suggest that the molecular context provided by the surrounding NORAD spacer sequences has a larger influence on the structure of NRU than sharp shifts in temperature.

Then, we measured the stability of NRUs by averaging the standard deviation of nextPARS scores at different temperatures [Fig. 2C]. Importantly, we observed that structural variability across temperatures was higher in NRUs fragments than larger NORAD fragments, supporting a stabilizing role of the molecular context when NRUs are embedded within larger NORAD fragments [Fig. 2C]. To ensure that the observed structural variability differences are not due to the effect of sequence length (i.e. longer molecules having higher stabilities), we use the information of nextPARS experiments of ten RNA molecules that were spiked with NORAD samples, with variable size [34], and measured the
structural stability at different temperatures. Our results did not detect any statistically significant correlation between stability and sequence length [Supplementary Figure S2], suggesting our results cannot simply be attributed to differences in length between NRU-specific, and larger NORAD fragments.

3.3. Conserved SAM68-binding sites in NORAD are particularly robust to structural perturbation by temperature shift.

SAM68 (KHDRBS1) is an RNA-binding protein (RBP) that performs several functions in the nucleus and cytoplasm [46]. It is
required for efficient recruitment of PUMILIO2 protein (PUM2) to NORAD and the regulation of Pum activity by NORAD [47-49]. SAM68 recognizes UAAA motifs [50], which in NORAD are present in 16 motif pairs separated by linkers of 15–35 bases. These motifs show higher evolutionary conservation as compared to the rest of the NORAD sequence [47]. When comparing nextPARS score variability at different temperatures for NRU-specific fragments in positions of UAAA motifs to other positions of NRUs, we observed that the motif sites were less variable as compared to the remaining residues in the NRUs (p-value = 0.010148) [Fig. 3A]. To further investigate the association between sequence conservation and structural flexibility in UAAA motifs, we measured the correlation between the evolutionary conservation as estimated by the PhyloP score (based on the 100-way vertebrate alignment) and the variability of structural profiles in the position of all UAAA motifs. First, we measured the structural variability of the UAAA motifs using the nextPARS probing data from NRU-specific fragments. We found no significant correlation between the level of conservation and the amount of structural variability [Fig. 3B]. Then, we calculated the same correlation measure but using the nextPARS data from the larger NORAD fragments and observed a significant negative correlation between sequence conservation and structural variability within UAAA sites- the higher the sequence conservation, the lower the structural variability at these positions - [Fig. 3B]. We obtained similar results using PhastCons score or Phylop score at different conservation levels [Fig. 3B]. These results suggest that SAM68-binding sites (UAAA motif) are robust to structural perturbation by temperature shift.

3.4. Accounting for structural diversity enhances evolutionary and functional comparisons.

We next set out to investigate other NORAD conserved elements beyond the known protein binding sites. We used our experimental data to study the two previously predicted hairpin structures present in the NRUs core, as such conserved secondary structure motifs are rarely detectable in IncRNAs [23]. We produced a multiple sequence alignment to derive a consensus secondary structure from the eight NRUs displaying a clear conserved long stem-loop hairpin [47] (NRU#1, NRU#3–8 and NRU#10). In the predicted consensus structure [Fig. 4A], we observed a short stem-loop structure comprising four paired bases and a variable loop followed by a second large stem-loop structure with nine base pairs.

Many sequences, including mRNA, IncRNAs and riboswitches, are expected to fold in multiple, coexisting structures [9]. To investigate this in the case of NORAD, we used Rsample [20] in combination with nextPARS data to predict the presence of alternative RNA structures for NRUs, and studied these conformations. NRU#2 was one of the four NRUs excluded from the analysis of the consensus structure, as this unit did not contain most of the repeat motifs compared to the rest of the NRUs [23]. When we predicted the structure of NRU#2, we observed that it was folded in two alternative conformations. The major conformation (inferred to represent 84.1% of the molecules in the mix) was predicted to form a large stem-loop after a U-rich stretch, and a small stem-loop before that U-rich [Fig. 4B]. These detected structure motifs...
were similar to those detected in some of the characterized NRUs (NRU#1, NRU#3–8 and NRU#10) [23]. In contrast, in the minor conformation (15.9%) or in the computationally predicted conformation, we could not detect the combination of sequence and structure motifs present in the other NRUs that were previously characterized [Fig. 4 B and Supplementary Fig. S3 A]. This indicates that considering the possibility of competing structural conformations enhances our ability to detect structural similarity otherwise missed by standard approaches based on a single stable structure.

Then, we searched for shared structurally related regions among the different NRUs, considering alternative conformations. For that purpose, we used the Beagle software [43] to perform an all versus all comparison of the most stable structure in all NRUs fragments. Then, we did the same all versus all comparisons but using the complete ensemble of alternative structures. We observed that the structural similarity (z-score) was higher for the ensemble of structures, as compared with the most stable one (p-value = 0.00095) [Fig. 4C], meaning again that using the alternative structures that may coexist within NORAD could be more accurate than using only the most stable one.

3.5. Discovery of two previously uncharacterized NRUs using covariance models.

Previous comparison of human NORAD sequences revealed that this lncRNA could be decomposed into at least 12 repeating units of ~300 nt each [47]. When we studied the relative positions of the core sequence of the NRUs throughout NORAD, we noticed that consecutive units were separated by a sequence spacers of similar length (between ~270 nt and ~340 nt), except for NRU#10 and NRU#11, which were separated by ~900 nt [Fig. 5A]. To investigate the presence of novel NRUs, we used infernal [42] to derive a model for NRUs (see Materials and Methods). Using this model on NORAD, we identified the 12 known NRUs, as expected, and we also detected the presence of two additional NRUs that we termed NRU#10a and NRU#10b. Interestingly, these units appear
Fig. 5. Discovery of two previously uncharacterized NRUs. A. Overview of the relative positions of the core sequence of the NRUs along with NORAD. The core positions of previously characterized NRUs are marked with grey bars, and the two new NRUs (NRU#10a and NRU#10b) are marked with green bars. The nextPARS score derived from the larger NORAD fragments experiment is shown, by an average window of 10nt. SAM68-binding sites are marked with brown lines. PhastCons score (based on the 100-way vertebrate alignment) is shown at the bottom.

B. On the top, secondary structures of NRU#10a and NRU#10b. On the bottom the sequence of NRU#10a and NRU#10b and the PhyloP score (based on the 100-way vertebrate alignment). Known structural and sequence motifs are marked in the sequence and in the structure. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)
within the unusually long intervening region between NRU#10 and NRU#11 and are separated by the approximately same distance as the rest of the NRUs [Fig. 5A]. Multiple sequence alignment of the core sequences of the known and newly identified NRUs showed a high degree of sequence similarity [Supplementary figure S4A]. Furthermore, we noticed a common sequence motif located at ~50 nt downstream of the end of the core of NRU#10a and other NRUs. [Supplementary figure S4B]. Moreover, in the new NRUs, we could identify most of the sequence and structure motifs typical for known NRUs, including the presence of SAM68-binding sites within the NRU or nearby [Fig. 5B] [47].

Finally, we assessed whether the structure of the newly discovered NRUs was similar to the previously characterized ones, aligning them to the previously mentioned alignment of the eight structurally conserved NRUs (NRU#1, NRU#3–8 and NRU#10). We were able to obtain a new consensus structure with the same short stem-loop structure. Still, the large consensus stem-loop was smaller compared with the consensus structure without the new NRUs [Supplementary Figure S5A], suggesting that part of the consensus structural similarity to the canonical consensus seems larger than that of NRU#2 and NRU#9. Based on these results, we conclude that the two newly identified NRU correspond to bona-fide functional elements in NORAD, and should be considered in future research.

4. Discussion

NORAD is a critical regulator of genome stability. Moreover, human NORAD is dysregulated in various types of cancer and could be an excellent diagnostic or therapeutic biomarker in tumour cells [30]. Since IncRNAs have no protein-coding potential, their structure may be key for exploring their functionality. Despite this, there have been no studies of the in vitro structurome of NORAD. To fill this important gap, we performed nextPARS experiments that allowed us to study temperature-dependent or context-induced structural changes in NORAD at a single-nucleotide resolution. Moreover, we experimentally determined NORAD's in vitro secondary structure and defined the core structural motifs conserved across NRUs. We identified two previously uncharacterized NRUs within an unusually long linker region. The addition of the two newly discovered NRUs results in all core units distributed along the whole NORAD sequence and placed at very regular distances. A possible explanation for this could be that the hairpin elements in the NRUs interact with each other or that they help to position the protein-binding sites at an advantageous distance from each other. The NRU#10b is well conserved with a similar PhastCons score than the previously characterized NRUs. However, we noticed that the NRU#10a was less conserved in their sequence compared to NRU#10b and the rest of the NRUs. Moreover, in contrast to the case of NRU#10b, we could not identify a putative SAM68-binding site sequence near the NRU#10a core. A plausible explanation for the poor sequence conservation could be that NRU#10a has been recently acquired during evolution.

The repetitive and modular nature of NORAD sequence allowed us to study several fragments at once and simplify the task of studying the structure of this long RNA transcript in its entirety. Our results indicate a stabilizing role of spacer sequences between NRUs, with their effect on the structure being larger than sharp shifts in temperature. How spacers stabilize NRUs is unclear. One possibility is that they favor interactions with other NRUs by placing them at some optimal distance, which would explain the regularity in spacer lengths. Alternatively, they may interact with some parts of the NRU structure, providing some additional stability. An RNA could fold in different structures inside the cell and many conformations for the same RNA coexist in equilibrium. Both for isolated NRUs and larger NORAD fragments experiments we observed that many sites exhibit clear signals for both single- and double-strand specific enzymes (nextPARS score near zero), which may result from noise, but also from the actual coexistence of alternative secondary structures during the experiment. Taking this into account, we estimated and compared the structural conformations of two NRUs and we detected structure motifs that were omitted when using the more stable structure. We revealed that these motifs were similar to the ones detected in some of the known NRUs. Moreover, we showed the importance of considering the ensemble of structures to compare structural elements in NRUs. Furthermore, a similar approach could be used to look for shared structural elements across species to study the evolution of NORAD.

NORAD contains several SAM68-binding sites (UAAG motif) and the interaction between NORAD and the RNA-binding protein SAM68 is required for NORAD function. We compared the nextPARS data at different temperatures to evaluate the stability of SAM68-binding sites motif secondary structure. We revealed that positions of UAAG motifs were more robust to thermal shifts than other positions in the NRUs sequence. Interestingly, we observed a correlation between conservation and flexibility in UAAG motifs but only when we use nextPARS data from larger NRAD fragments experiments and not when using data from NRU-specific fragments.

For certain short RNAs, the in vitro folding landscape can recapitulate the in vivo one effectively, while for long RNAs, the structures sometimes differ in vitro versus in vivo [51,52]. These differences are mainly due to interactions with other molecules. In vivo structure probing may provide more accurate information on biologically relevant RNA structures [53,54]. In this work, we use only information from in vitro experiments, so we lack direct information on which regions interact with other molecules. But we think that one future perspective could be combining these two experimental methods to obtain the full picture of the RNA structurome. Nonetheless, both in vitro and in vivo approaches have limitations [55]. Furthermore, NORAD has sponge potential on several miRNAs and some of them were validated in different studies. Another future perspective could be to study the secondary structure of miRNA binding sites. We hypothesize that NORAD structure could modulate miRNA cleavage by facilitating or preventing accessibility of the miRNA to its binding site.

Taken together, our analyses contribute important novel insights into the structural organization of NORAD. Future studies should focus on determining how the different structural elements of NORAD interact with each other and with other molecules, and how different mutations may affect these interactions and the functions mediated by them. Only by understanding the complex relationships between, sequence, structure, and function will we be able to comprehend the roles of NORAD in the normal functioning of the cell and in diseases such as cancer. Eventually, this may open the door to design novel therapeutic strategies in the fight against cancer.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.csbj.2021.05.045.

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