An RNA dynamic ensemble at atomic resolution

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Abstract: Solving dynamic ensembles of biomolecules at atomic resolution is a major challenge in structural biology because it requires information which far exceeds that available from experimental measurements. We addressed the data gap in RNA ensemble determination by using structure prediction tools that leverage the growing database of RNA structures to generate a conformational library. Library refinement with NMR residual dipolar couplings enabled determination of an atomic-resolution ensemble for HIV-1 TAR as confirmed by quantum-mechanical calculations of NMR chemical shifts, comparison to a crystal structure of a substate, and atomic mutagenesis. The ensemble reveals how bulge residues cooperatively flip out and undergo sugar repuckering to allow the adjoining helices to stack. The generality of this approach will make determination of atomic-resolution RNA ensembles routine.
Main Text:

Biomolecules do not fold into a single 3D structure but rather form dynamic ensembles of many inter-converting conformations (1). Knowledge of dynamic ensembles is key for understanding how biomolecules fold and function, and for rationally manipulating their activities in drug discovery and synthetic biology (2-4). However, determining dynamic ensembles at atomic resolution remains an outstanding challenge in structural biology and biophysics, particularly for highly flexible ribonucleic acids (RNAs) (4-7).

The information required to specify the position of all atoms in thousands of conformations in an ensemble far exceeds the information content of experimental measurements. NMR spectroscopy is a rich source of ensemble-averaged measurements, and in combination with computational modelling, has been applied with success to determine ensembles of proteins at atomic resolution (8-10). In contrast, fewer atomic-level experimental measurements are typically available for characterizing nucleic acids ensembles. Moreover, while computational modeling methods such as molecular dynamics (MD) simulations are needed to address the experimental data gap (11), nucleic acid force fields remain underdeveloped relative to proteins, and MD simulations of RNAs often poorly predict experimental data even for simple motifs (12, 13). Consequently, relative to proteins, there is a greater danger of over-fitting an RNA ensemble, and assessing ensemble accuracy is more difficult.

To address the data gap, as well as dramatically simplify and accelerate RNA ensemble determination, we took advantage of structure prediction tools that leverage the growing database of RNA structures to directly generate a conformational library from a secondary structure that broadly samples energetically favorable 3D conformations. We used Fragment Assembly of RNA with Full-Atom Refinement (FARFAR) (14), given its high performance in extensive tests of blind
prediction of 3D RNA structure (15). We then determined RNA ensembles by using NMR residual dipolar coupling (RDC) data (16-18) to guide selection of conformers from the FARFAR library (7, 13, 19).

We tested our approach on the transactivation response element (TAR) (Fig. 1A) from HIV-1 (20, 21), which has served as a model system for bulge motifs. Bulges are one of the most common RNA secondary structural elements that act as dynamic joints connecting helical elements, enabling their relative orientation to change adaptively during folding and function (13, 20-22). We used FARFAR to directly generate a conformational library \( N = 10,000 \) from an input TAR secondary structure only constraining Watson-Crick base pairs (bps) inferred by NMR, and assuming an idealized A-form geometry (23) for these bps while predicting the structure for all remaining nucleotides (Methods). We then tested and optimized the FARFAR-library using a rich dataset of four independent RDCs (~8 RDCs per nucleotide) previously measured for four TAR molecules variably elongated to modulate alignment relative to the NMR magnetic field (7, 13).

Intriguingly, the FARFAR-library showed better agreement (RMSD 8.0 versus 8.6 Hz) with the RDCs (Fig. 1B) as compared to a previously reported (13) TAR library (Anton-MD) generated by subjecting an experimentally determined NOE-based NMR structure of TAR (PDB ID 1ANR) (20) to MD simulations with the CHARMM36 force field (24) (Fig. 1C). An optimized ensemble with \( N = 20 \) conformers (fig. S1) was generated (Methods) by using the RDC agreement to guide selection of conformers from the FARFAR-library (Fig. 1D) (13, 19). The optimized FARFAR-NMR ensemble also better predicted the RDCs (RMSD 3.1 Hz) relative to the optimized Anton-MD-NMR ensemble (RMSD 3.6 Hz) obtained using a similar procedure and the Anton-MD pool (Fig. 1E and fig. S1). Cross validation (13, 25) showed that the improved RDC
agreement was not due to over-fitting (fig. S2). Similar RDC agreement (Fig. 1F) was obtained when using FARFAR-NMR to generate ensembles for TAR in the presence of Mg$^{2+}$, and three additional TAR mutants (22) containing one (U1-TAR), two (U2-TAR), and seven (U7-TAR) bulge nucleotides in the presence and absence of Mg$^{2+}$ (fig. S3 to S4, and Supplementary Text), establishing the generality of the approach.

The improved agreement observed with the FARFAR-NMR ensemble was surprising given that the global inter-helical distribution (26) of the Anton-MD-NMR TAR ensemble has been independently validated via X-ray scattering interferometry (XSI) (27). Indeed, the FARFAR-NMR and Anton-MD-NMR ensembles show comparable agreement with the RDCs measured for helical bps (Fig. 1, D and E) and the two ensembles sample similar inter-helical orientational distributions (Fig. 1G and fig. S5).

Rather, the improved agreement was primarily driven by the RDCs measured in the locally more flexible bulge residues. For the FARFAR-NMR ensemble, the RDC RMSD (1.8 Hz) for bulge residues (Fig. 1D) was within experimental error (~2 Hz), but was substantially higher (3.7 Hz) for the Anton-MD-NMR ensemble (Fig. 1E). The bulge RDC RMSD could not be improved by running MD simulations using different force fields (Fig. 1H), indicating that the improved conformational sampling in the FARFAR-library makes it possible to surpass the accuracy with which the TAR bulge could be described using conventional MD simulations. This improved performance is particularly noteworthy when considering that determining a high-resolution structure and running MD can take several months and often years whereas the FARFAR-library is generated within 24 hours running on 100 cores in parallel.

To further evaluate the accuracy of the FARFAR-NMR TAR ensemble, we substantially expanded the breadth and depth of atomic-level experimental data that can be brought to bear when
evaluating the accuracy of RNA ensembles by predicting ensemble-averaged $^1$H, $^{13}$C, and $^{15}$N chemical shifts (~15 chemical shifts per nucleotide) using quantum-mechanical AF-QM/MM calculations (28, 29). Although rich in structural information that is complimentary to that obtained from RDCs, the agreement between measured chemical shifts and values predicted from crystal structures of nucleic acids has traditionally been poor (28, 30). We recently showed in studies of DNA duplexes that at least some of this disagreement originates from neglecting ensemble averaging when predicting $^{13}$C chemical shifts (29). This revelation and the improved FARFAR-NMR TAR ensemble led us to test this approach on flexible RNAs, and to extend its reach by incorporating $^1$H and $^{15}$N shifts in addition to $^{13}$C chemical shifts (Fig. 2A).

Remarkably, good agreement was observed between the measured $^1$H, $^{13}$C, and $^{15}$N base and sugar chemical shifts and values back-calculated for the FARFAR-library (fig. S6 to S7 and Supplementary Text). The agreement improved substantially for the FARFAR-NMR ensemble following RDC optimization for 48% of the atom-types analyzed, while the agreement was unaffected for the remaining atom-types (Fig. 2B and fig. S6 to S7). The agreement deteriorated substantially for any one conformer member of the ensemble, underscoring the critical importance of ensemble-averaging (fig. S8). The agreement with an independent set of measurements not used in ensemble determination that is highly sensitive to various structural features of the base, sugar, and backbone strongly suggests that FARFAR-NMR describes the TAR ensemble with atomic accuracy (< 2 Å, Methods). In sharp contrast, the agreement was much weaker for the Anton-MD-NMR ensemble relative to FARFAR-NMR for 90% of the atom-types, particularly for sugar chemical shifts, some of which show no apparent correlation even following RDC optimization (Fig. 2, C and D, and fig. S6 to S7). These results establish the utility of chemical
shifts in RNA ensemble determination and also show that the improved accuracy of the FARFAR-NMR ensemble relative to Anton-MD-NMR is even greater than alluded to by the RDC data.

We compared the FARFAR-NMR and Anton-MD-NMR ensembles (Fig. 3A) to better understand the features responsible for the more accurate ensemble description of the TAR bulge. Approximately 75% of the conformers in the FARFAR-NMR ensemble have the canonical UCU bulge, while the remaining 25% have a non-canonical AUC bulge that forms through a single nucleotide register shift (fig. S9). In contrast, conformers in the Anton-MD-NMR ensemble sample a broader set of junction topologies some of which diverge from the NMR-derived Watson-Crick pairing (fig. S9 to S10 and Supplementary Text), and this could contribute to the poor agreement observed with the imino $^{15}$N/$^1$H chemical shifts (Fig. 2C and fig. S6 to S7).

In addition, despite having a narrower range of junction topologies, the sugar pucker distribution (defined by angle $\delta$) for bulge residues in the FARFAR-NMR sample a broader range (Fig. 3, B and C), are substantially enriched in the non-canonical C2′-endo conformation (Fig. 3D) relative to the Anton-MD-NMR ensemble, and are in better agreement with the NMR-derived (31) sugar pucker (Fig. 3E and fig. S11). Moreover, ~15% of the conformers in the FARFAR-NMR ensemble had all three bulge residues simultaneously in the C2′-endo conformation but none did in the Anton-MD-NMR ensemble (Fig. 3F). Excluding all conformers containing C2′-endo sugar puckers in U23, C24, U25, A22 or U40 from the FARFAR-library diminished the RDC agreement to a level similar to the Anton-MD derived ensembles again confirming that the improved agreement is not due to over-fitting of the data (fig. S12). A similar behavior was observed for the backbone torsion angle $\gamma$ (Fig. 3B), which shows greater sampling of non-gauche+ angles in the FARFAR relative to the Anton-derived ensembles (Fig. 3C and fig. S11 to S12). This can account
for the better agreement observed for sugar chemical shifts (31) in the FARFAR-NMR versus Anton-MD-NMR ensembles (Fig. 2, B to D).

What causes conformers in the FARFAR-NMR ensemble to more greatly sample non-canonical sugar-backbone conformations at the bulge relative to conformers in the Anton-MD-NMR ensemble? Unpaired pyrimidine RNA nucleotides unconstrained by other interactions are enriched in the C2′-endo conformation (C2′-endo:C3′-endo is 40:60) (32) and C3′-endo becomes the predominant sugar-pucker when the nucleotides form bps or stack intra-helically (31). Indeed, ~80% of the residues with C2′-endo sugar puckers were extra-helical in the FARFAR-NMR ensemble (Fig. 4A and fig. S9). Moreover, for linear inter-helical conformations, there was a strong preference to have all three bulge residues simultaneously flip out (Fig. 4B) and adopt conformations enriched in the C2′-endo sugar pucker (Fig. 4C); the bulge residues flip out to allow the two helices to coaxially stack (Movie S1 and fig. S9). Such coaxial conformations have previously been hypothesized to exist within the TAR ensemble based on the Mg2+ dependence of the inter-helical ensemble (22) and a crystal structure of Ca2+ bound TAR (33).

Strikingly, two of the coaxial conformations in the FARFAR-NMR ensemble superimpose with the TAR crystal structure with a heavy-atom RMSD of 1.3-1.4 Å (Fig. 4D). Thus, the TAR crystal structure captures substates of the ensemble in solution. In sharp contrast, none of the conformations in the Anton-MD-NMR ensemble had the two helices coaxially stacked, and not a single conformer had all three bulge residues flipped out and having the C2′-endo sugar pucker (Fig. 4, A to C and fig. S9). The FARFAR-NMR ensemble (Fig. 4A and Movie S1) further shows that all three bulge residues simultaneously flip out with high cooperativity (~2.3 kcal/mol, see methods) most likely because this then permits favorable coaxial stacking of the helices (Fig. 4A, fig. S9, and Movie S1). This can explain prior results (34) showing that a TAR mutation that
promotes inter-helical stacking results in a predominantly coaxially stacked conformation in which all three bulge residues are flipped out.

Finally, we put key atomic features of the FARFAR-NMR ensemble to a test by rationally redistributing the conformer populations using atomic mutagenesis. The ensemble shows that the kinked unstacked conformations are enriched in bulge residues that have the C3′-endo sugar pucker relative to the coaxial stacked conformations (Fig. 4C). To test this feature of the ensemble, we incorporated 2′-O-Methyl (Nm) modifications (Fig. 4E) at U23 or C24 to bias the sugar pucker at these positions toward C3′-endo by ~0.2 kcal/mol (35). Indeed, methylating either U23 or C24 resulted in chemical shift perturbations in 2D NMR spectra of TAR (Fig. 4F and fig. S13) throughout the bulge and neighboring residues that are specifically directed towards the chemical shifts of the unstacked conformation (22), as expected for a cooperative redistribution in favor of the kinked conformation.

In conclusion, FARFAR-NMR lays the foundation for a new paradigm for RNA ensemble determination by combining reliably measurable NMR data with 3D structure prediction. The approach can immediately be applied to render many existing NMR structures of RNAs that were determined using conventional approaches into dynamic ensembles, and their accuracies can be tested using the available chemical shift data. The approach is general, rapid, and can also incorporate other sources of experimental data. Given its ease of implementation and higher throughput, FARFAR-NMR has the potential to unleash the ensemble description of RNAs to all corners of biology, from which a deeper and broader understanding of folding and function will undoubtedly emerge.
References and Notes:


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Supplementary Materials

Materials and Methods

Supplementary Text

Figures S1-S16
Tables S1-S5
Movie S1
Fig. 1. Using FARFAR-NMR to determine a TAR ensemble. (A) Secondary structure of TAR. (B-E) Comparison between measured and predicted TAR RDCs for the (B) FARFAR-library ($N = 10,000$), (C) Anton-MD library ($N = 10,000$), (D) FARFAR-NMR ($N = 20$) and (E) Anton-MD-NMR ($N = 20$). Values in parentheses denote the RDC RMSD. RDCs are color-coded according to the structural elements in Fig. 1A. (F) RDC RMSD for ensembles of TAR variants with ensemble size $N$ obtained using FARFAR-NMR. (G) Structural overlay of the TAR ensembles ($N = 20$) (Methods). $|\beta|$ is the absolute magnitude of the bend angle. (H) Comparison of RDC RMSD over all (blue) or bulge (red) residues for ensembles generated using different MD force fields (ff99 (36), CHARMM36 (24), ff99bsc0xOL3 (37) and modified ff14 DESRES (38)) and FARFAR.
Fig. 2. Evaluating TAR ensembles using chemical shifts. (A) Chemical structures of the sugar and base moieties with chemical shift probes used to test ensemble accuracy highlighted in green. (B-C) Comparison of measured and predicted $^{13}$C/$^{15}$N chemical shifts for (B) FARFAR-NMR (N was not certified by peer review) is the author/funder. All rights reserved. No reuse allowed without permission.
= 20) (C) Anton-MD-NMR (N = 20) by AF-QM/MM (Methods). Values are color-coded according to the structural elements in Fig. 1A. Chemical shifts for central Watson-Crick bps within A-form helices (C19-G43, A20-U42, G21-C41, A27-U38, G28-C37) are denoted using open circles. A correction was applied to the predicted chemical shifts (Methods) as described previously (29). For ¹H chemical shifts see fig. S7. (D) Comparison of RMSD (left) and R² (right) between measured and predicted ¹³C/¹⁵N chemical shifts for flexible residues (U23, C24, U25, A22-U40, G26-C39, C29-G36, G18-C44) for FARP-NMR (red) and Anton-MD-NMR (blue).
Fig. 3. FARFAR-NMR ensemble more broadly samples non-canonical sugar-backbone torsion angles relative to Anton-MD-NMR. (A) Overlay of the dynamic ensembles of the TAR bulge. (B) RNA backbone torsion angles exhibiting different and similar distributions between Anton-MD-NMR and FARFAR-NMR are colored red and green, respectively (Methods). (C) 2D density maps of \( \delta \) versus \( \gamma \) comparing Anton-MD-NMR and FARFAR-NMR ensembles (\( N = 2,000 \)) for
bulge residues as well as A22 and U40. The bin width is 20°. (D) Structure of the ribose moiety in C3'-endo and C2'-endo conformations. (E) Population of C2'-endo pucker at bulge residues as well as A22 and U40 in the FARFAR-library (N = 10,000, red open), FARFAR-NMR (N = 2,000, red fill), Anton-MD library (N = 10,000, blue open) and Anton-MD-NMR (N = 2,000, blue fill). Experimental estimates of the C2'-endo population based on $^{13}$C chemical shifts are indicated above the bars (Methods). (F) The population of conformers in the ensemble as a function of the number of C2'-endo bulge residues for FARFAR-NMR (red, N = 2,000) and Anton-MD-NMR (blue, N = 2,000).
**Fig. 4.** Cooperative extra-helical flipping and sugar repuckering of bulge residues are coupled to coaxial stacking. (A) Overlay of conformers showing motions in bulge residues for linear (|$\beta_h$| < 45°), intermediate bend (45° < |$\beta_h$| < 70°) and kinked (|$\beta_h$| > 70°) inter-helical conformations in the FARFAR-NMR and Anton-MD-NMR ensembles. (B-C) The fractional populations of
conformers with (B) extra-helical and (C) C2′-endo bulge residues (color-coded) as a function of bending angle in the FARFAR-NMR and Anton-MD-NMR ensembles. F and A denote FARFAR-NMR and Anton-MD-NMR, respectively. (D) Comparison of two coaxially stacked FARFAR-NMR conformers with the crystal structure of Ca²⁺ bound TAR (PDB: 397D) (33). (E) Nm shifts sugar pucker equilibrium towards C3′-endo. (F) Overlay of 2D [¹³C, ¹H] NMR HSQC spectra of the aromatic spins for TAR-Nm-C24 without Mg²⁺ (blue, inducing unstacking), TAR without Mg²⁺ (cyan) and TAR with Mg²⁺ (red, inducing stacking).