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Developments in solution-state NMR yield broader and deeper views of the dynamic ensembles of nucleic acids



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Abstract

Nucleic acids do not fold into a single conformation, and dynamic ensembles are needed to describe their propensities to cycle between different conformations when performing cellular functions. We review recent advances in solution-state nuclear magnetic resonance (NMR) methods and their integration with computational techniques that are improving the ability to probe the dynamic ensembles of DNA and RNA. These include computational approaches for predicting chemical shifts from structure and generating conformational libraries from sequence, measurements of exact nuclear Overhauser effects, development of new probes to study chemical exchange using relaxation dispersion, faster and more sensitive real-time NMR techniques, and new NMR approaches to tackle large nucleic acid assemblies. We discuss how these advances are leading to new mechanistic insights into gene expression and regulation.

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Introduction

Composed of only four chemically similar building block nucleotides, the three-dimensional (3D) structures of nucleic acids are nowhere near as diverse or complex as those of proteins with their 20 chemically diverse amino acids. Yet, nucleic acids are specifically recognized by the cellular machinery and participate in highly complex reactions, both as substrate and enzyme. During cellular processes, DNA and RNA cycle through different conformations each satisfying distinct requirements of

multistep biochemical reactions. These structural dynamics provide a means for amplifying specificity and functionality beyond that which can be attained based on a rigid 3D structure [1,2].

To deeply understand how nucleic acids fold and function at the atomic level, we need to reimagine their 3D structures to include a description regarding propensities to undergo conformational changes over timescales spanning 12 orders of magnitude in time, from picoseconds to seconds. The dynamic ensemble provides one such unified description [1,3,4].

In solution, a nucleic acid molecule will sample a vast number of different conformations. Some conformations may be highly populated (>10%), whereas others may be vanishingly low in abundance (<0.00001%). Some may form on the picosecond timescale, whereas others may form slowly on the second timescale. Some may differ markedly from the native state, whereas others may only differ by the relocation of a single proton. The Boltzmann-weighted distribution of interconverting conformations is referred to as a 'dynamic ensemble.' The dynamic ensemble provides a statistical mechanical description that can be used to rigorously compute the thermodynamic cost accompanying conformational transitions and also the rates with which they occur [1].

Because a continuous landscape comprising thousands of 3D structures needs to be specified along with the Boltzmann-weighted populations and kinetic rates of interconversion, determining dynamic ensembles continues to be an outstanding challenge in structural biology and biophysics. Many biologically important conformations exist in low abundance and/or are exceptionally short-lived, and are therefore difficult to detect experimentally. Over the past 5 years, significant progress has been made toward the development of experimental and computational approaches that provide different cross-sectional views in space and time of the dynamic ensembles formed by DNA and RNA. Here, we highlight recent advances in solution-state nuclear magnetic resonance (NMR) methods for ensemble determination. and when applicable, their integration with computational techniques. We also discuss how the characterization of dynamic ensembles is leading to a deeper and more quantitative understanding regarding biochemical processes mediated by nucleic acids.

Dynamic ensembles from ensemble-averaged NMR data

Approaches for ensemble determination

Many NMR observables including residual dipolar couplings (RDCs), the nuclear Overhauser effect (NOE), and chemical shifts are averaged over many conformations in the dynamic ensemble. In conventional NMR structure determination protocols, a single conformation is identified that satisfies the ensemble-averaged experimental data. A collection of such structures each satisfying the data within experimental uncertainty are often pooled together in a bundle, and such ensembles need to be distinguished from the statistical mechanical definition in terms of a Boltzmann distribution used throughout this review [1,3,4].

Three different approaches have been developed to interpret NMR data in terms of a Boltzmann distribution of nucleic acid conformations. In one approach, the NMR data are used to guide the assignment of Boltzmann weights to a library of conformations generated using computational methods, typically molecular dynamics (MD) simulations [5,6]. In a second approach, the data are applied as a restraint during MD simulations [7]. Finally, the NMR data can also be used in conventional structure refinement programs to explicitly solve for multiple conformations [8]. Although the experimental data are typically not used to encode timescale information [9], they are often sensitive to motions occurring over submillisecond timescales.

Dynamic ensembles that satisfy different types of NMR data are being reported for a small but steadily growing number of nucleic acid motifs, including apical loops [10-13], bulges [14-16], single-strands [17], duplexes [18], and a G4 quadruplex [19] (Figure 1a). Together, these ensembles are beginning to outline the conformations that dominate the landscape of nucleic acid ensembles.

Nucleotides in single-strand and loop regions frequently transition between a narrow distribution of stacked conformations and a broader distribution of unstacked extrahelical conformations [14,16,17,19] (Figure 1a). Extra-helical flipping can be accompanied by changes in sugar pucker and coaxial interhelical stacking [14,17] (Figure 1a). The ensembles can also include partially melted states [13] or non-native secondary structures [14] (Figure 1a). Dynamic ensembles of canonical DNA duplexes reveal sequence-specific variations particularly for backbone torsion angles ε and ζ [18] (Figure 1a).

Measurements of ensemble-averaged NMR data and their use in ensemble determination

Because determining a dynamic ensemble is an inherently underdetermined problem, much effort has been directed toward measuring new types of ensembleaveraged NMR data to improve the accuracy of the determined ensembles. A recent study underscored this point [20], showing that increasing the amount of input RDC data used to determine dynamic ensembles of HIV-1 TAR RNA results in ensembles that better predict the binding of small molecules using ensemblebased virtual screening.

NMR chemical shifts can be measured with great ease and abundance and carry rich information regarding the distribution of torsion angles, sugar puckers, hydrogenbonding, and ionization/tautomerization states of the nucleobases [21]. Recent advances in predicting chemical shifts from 3D structure are enabling their application in determining and assessing dynamic ensembles of nucleic acids.

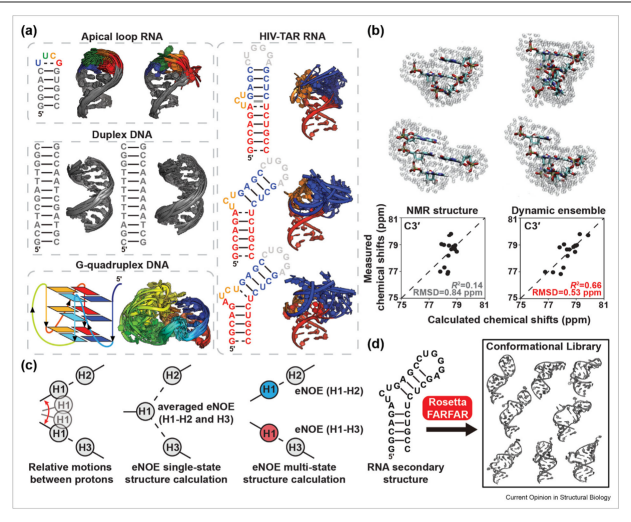
Frank et al. [16] developed an empirical approach for rapidly predicting ¹H and ¹³C chemical shifts based on a 3D RNA structure. This made it possible to apply the chemical shifts as restraints during an MD simulation. The approach was used to determine a dynamic ensemble for the U6 intramolecular stem-loop RNA, which is a key component of the spliceosome.

Empirical approaches are constrained to the available database of nucleic acid chemical shifts with known 3D structures and by difficulties in taking into account motional averaging contributions. Some of these limitations can be overcome using ab initio approaches for computing chemical shifts based on 3D structures. However, these calculations can be prohibitively timeconsuming for large biomolecules, and until recently, the agreement between predicted and measured chemical shifts was not sufficiently good for nucleic acids to allow quantitative applications.

To accelerate the *ab initio* calculations, Swails et al. [22] developed automated fragmentation quantum mechanics/molecular mechanics (AF-QM/MM), in which a nucleic acid is divided into small quantum fragments each of which is subjected to quantum mechanical calculation while treating the remaining parts of the molecule using molecular mechanics (Figure 1b). For four DNA duplexes [18] as well as the HIV-1 TAR RNA [14], quantitative agreement was obtained when averaging the predicted ¹H, ¹³C, and ¹⁵N chemical shifts over an ensemble of conformations determined using RDCs and MD simulations (Figure 1b) [14,23]. For both the DNA and RNA examples above, the agreement deteriorated considerably for the conventional single NMR structure, underscoring the importance of treating motional averaging when calculating chemical shifts, in addition to validating the accuracy of the calculations.

Ensemble-averaged interproton distances derived from quantitative measurements of exact NOEs (eNOEs) [24] have been applied to determine nucleic acid ensembles (Figure 1c). Relative to semiquantitative

Figure 1



Determining dynamic ensembles of nucleic acids using NMR and computational methods. (a) Shown are NMR-derived dynamic ensembles for RNA (UUCG apical loop [13] and HIV-TAR bulge [14]) and DNA (duplexes [18] and a G-quadruplex [19]). The figures were adapted from the original publication. (b) Representative fragments of an RNA hairpin structure used to compute chemical shift using the AF-QM/MM approach (top). The figure was adapted from Ref. [22]. Also shown is a comparison between measured and AF-QM/MM predicted ¹³C chemical shifts for a conventional NMR structure and an RDC-derived dynamic ensemble of a DNA duplex (bottom). The data shown in the figure was adapted from Ref. [23]. (c) The principle of eNOE. A mobile proton H1 samples positions that are close to both protons H2 and H3. The eNOE data are better explained by a two-state ensemble shown in blue and red relative to a model that assumes a single-state structure. The figure was adapted from Ref. [25]. (d) The Rosetta FARFAR structure prediction algorithm uses an RNA secondary structure to rapidly generate a conformation library, which can be used in conjunction with NMR data to determine a dynamic ensemble. The figure was adapted from Ref. [14].

standard NOEs, eNOE can be more readily interpreted quantitatively in terms of many conformational states in an ensemble. eNOEs have been used to determine ensembles for a 14-mer UUCG tetraloop either through multistate ensemble refinement [25] or *posteriori* refinement of an MD trajectory using a maximum entropy/Bayesian approach [26] establishing the robustness of these measurements in RNA ensemble determination.

Dynamic ensembles by combining NMR data with structure prediction

The accuracy of dynamic ensembles obtained with the aid of NMR data can be highly dependent on the

accuracy of the force field used to generate the starting conformational library or to run restrained MD simulations. Despite improvements in nucleic acid force fields [27], even for simple motifs, ensembles of nucleic acids generated using MD often poorly predict NMR data [10,14,17,19,23,28]. By examining what aspects of the MD-derived ensembles need to be altered to satisfy experimental NMR data, studies are beginning to pinpoint potential biases in nucleic acid force fields, including a tendency to form intercalated structures [17] and certain biases in sugar pucker [14]. These findings should help guide and accelerate future advances in force field development.

Recent studies have demonstrated the power of combining NMR data, including ¹H chemical shifts [29] and imino proton NOEs [30] with structure prediction algorithms when solving a 3D structure of an RNA. Shi et al. [14] showed the utility of combining RNA structure prediction algorithms with RDC and chemical shift measurements in determining dynamic ensembles of RNAs. Rather than using the time-consuming step of solving a 3D structure then subjecting the structure to MD simulations, Rosetta's Fragment Assembly of RNA with Full-Atom Refinement (FARFAR) RNA structure prediction algorithm [31] was used to rapidly and directly generate a conformational library for an RNA secondary structure [14] (Figure 1d). FARFAR [31] generates physically plausible conformations by stitching together fragments from a library of existing structures in the protein data bank [32], and it was shown to overcome some of the biases and inaccuracies in MD force fields as supported by comparisons to quantummechanical calculations of NMR chemical shifts [14].

Excited conformational states from relaxation dispersion

The dynamic ensembles of nucleic acids also include low abundance excited conformational states (ESs) that have exceptionally low populations (as low as 0.01%) and short lifetimes (as low as a few microseconds) [1,2]. Methods based on NMR relaxation dispersion (RD) are enabling the characterization of excited conformational states with unprecedented detail providing information regarding their structure, population, and kinetic rates of interconversion [33–35].

Recent developments in methods

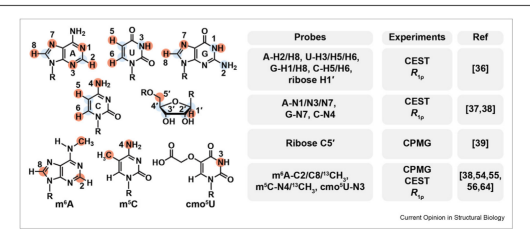
NMR techniques to characterize ESs come in four flavors optimized for different exchange rates ($k_{\rm ex} = 1 \, {\rm s}^{-1}$ to 10^5 s⁻¹) and/or type of nuclei all of which are now routinely applied to study nucleic acids. These include spin relaxation in the rotating frame (R_{10}) , Chemical Exchange Saturation Transfer (CEST), Carr-Purcell-Meiboom—Gill (CPMG), and ZZ-exchange experiments [33–35]. Although early studies focused on the protonated base (U/C-C5, T/U/C-C6, A/G-C8, and A-C2) and sugar (C1', C2', C3', and C4') carbons, and the base imino (G-N1 and T/U-N3) as well as amino nitrogen (G-N2 [33]), the list of RD probes now includes base and sugar protons (U-H3, G-H1, A-H2; A/G-H8, U/C-H5/H6; and ribose H1') [36], nonprotonated (G-N7 and A-N1/N7) [37] and amino nitrogen (C-N4) [38], as well as the ribose C5' [39] (Figure 2). Leveraging the practical advantages of CEST, a recent study broadened the timescale sensitivity of the experiment to fast exchange through the use of high radiofrequency fields [40].

Approaches to stabilize ESs using mutations or chemical modifications [41] and improvements in predicting chemical shifts based on secondary structures (Y Wang et al., unpublished) are making it possible to determine the structure and dynamic properties of nucleic acid ESs at atomic resolution [23]. Techniques have also been developed that rely on biasing the conformational equilibria through the use of mutations to assess the abundance of RNA ESs within the physiological cellular context [42,43].

Biological activities of ESs

Several recent examples highlight how the cellular machinery recognizes and acts on the ESs of RNAs that form by reshuffling base pairs in and around noncanonical motifs. For example, microRNAs (miRNA) downregulate gene expression by binding to the three prime untranslated regions (3'-UTRs) of mRNAs and activating the RNA-induced silencing complexes (RISCs), which promote mRNA degradation and translation repression [44].

Figure 2



A growing number of nuclei are targeted for relaxation dispersion measurements (CEST, CPMG, and R₁₀) in studies of nucleic acid ESs. The list of probes is growing steadily from an initial focus on protonated carbons, imino and amino (G-N2) nitrogens (highlighted using blue circles) to now include protons, non-protonated and amino (C-N4) nitrogens, ribose C5', and nuclei in modified bases including m⁶A, m⁵C, and cmo⁵U (highlighted using red circles). Also shown are the types of RD experiments performed for the various probes.

Using R_{10} RD, Baronti et al. [43] showed that the catalytically active RISC conformation acts on an ES formed by miR-34a:mRNA duplex. The ES features an elongated 8-mer seed with a coaxially stacked topology that better satisfies the substrate requirements for binding the active conformation of RISC (Figure 3a). Mutations to various mRNA targets that stabilize the ES of the miR-34a:mRNA duplex were shown to robustly increase the miR-34a mediated downregulation of the target genes in human cells.

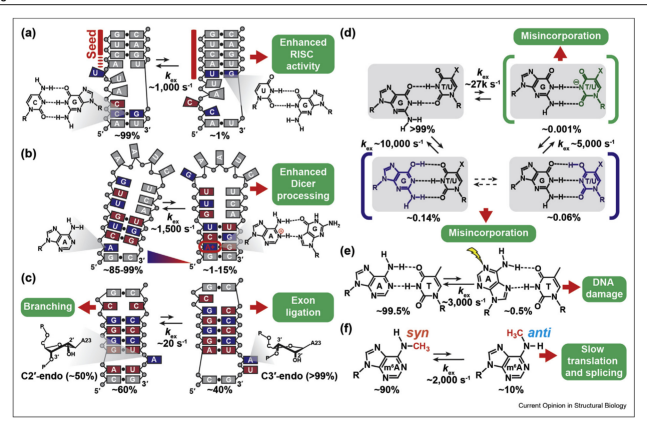
Using CEST, Baisden et al. [45] showed that the Dicer protein more efficiently recognizes and processes a pHdependent ES in pre-miR-21 that features a protonated $A^+(anti) \cdot G(syn)$ mismatch, and which also better satisfies its substrate requirements (Figure 3b). Mutants that stabilize the ES were shown to be more readily processed by Dicer relative to mutants that favor the

GS. It was proposed that pH-dependent ESs may provide a general mechanism for regulating miRNA biogenesis in response to environmental stimuli.

RNA ESs can also feature noncanonical sugar puckers [46]. Using ZZ-exchange, Plangger et al. [47] showed that the group II intron ribozyme exists as a dynamic equilibrium between two conformations in which a catalytic adenine residue adopts either ~50:50C2'endo:C3'-endo or >99% C3'-endo sugar pucker conformations. The equilibrium was proposed to regulate the switch between branching and exon ligation during splicing and reverse splicing (Figure 3c).

Combining RD with chemical shift perturbations and Bayesian inference, two ligand molecules were shown to recognize and bind an ES of HIV-1 TAR that has an exceptionally low population of <0.001%, highlighting

Figure 3



NMR reveals RNA and DNA ESs that play important roles in biochemical processes (a-c) RNA ESs form by reshuffling base pairs. (a) The miR-34a:mRNA duplex undergoes conformational exchange between the GS and the ES, which has an elongated 8-mer seed and altered RNA topology, leading to the formation of the active RISC complex. The figure was adapted from Ref. [43]. (b) Pre-miR-21 forms a pH-dependent protonated ES with an A⁺(anti)·G(syn) base pair (bp) that is more efficiently recognized and processed by Dicer. The figure was adapted from Ref. [45]. (c) Domain 6 of type II intron forms an ES with a bulge adenine that has a predominantly (>99%) C3' endo sugar pucker conformation, suitable for exon ligation during splicing and reverse splicing. The figure was adapted from Ref. [47]. (d) Wobble G T and G U mismatches form tautomer (blue) and anionic (green) Watson-Crick-like ES conformations that can evade fidelity checkpoints during DNA replication and cause spontaneous mutations. cmo⁵U increases the population of the anionic Watson-Crick ES conformation. The figure was adapted from Ref. [50]. (e) Watson-Crick A·T bps form Hoogsteen ES conformations, which increase the damage susceptibility of canonical duplex DNA. (f) The methylamino group of m⁶A forms an anti-ES needed for base pairing, thereby slowing down duplex hybridization and conformational transitions

how ligands with apparently weak binding affinity actually recognize exceptionally lowly populated conformations in the dynamic ensemble with high binding affinity [48]. However, the ESs of HIV-1 TAR and RRE stem IIB [49] were shown to disrupt key elements of structure needed for protein recognition and cellular activity making them interesting HIV-1 drug targets [42].

The cellular machinery can also act on the ESs of DNA. ES conformations of the G·T mismatch that satisfy the Watson-Crick geometry through tautomerization or ionization of the bases were shown to be recognized by high fidelity polymerases leading to copying errors during DNA replication [50] (Figure 3d). A kinetic mechanism based on the NMR-derived exchange kinetics was shown to predict copying errors across a range of high-fidelity polymerases, conditions, and modifications [50]. Damaging reagents were also shown to act on ES Hoogsteen base pairs, which expose the Watson-Crick face of nucleotide bases in duplex DNA to solvent [51,52] (Figure 3e).

Impact of epigenetic and post-transcriptional modifications on excited conformational states

Application of NMR RD is helping reveal how epigenetic and post-transcriptional modifications of DNA and RNA reshape the dynamic ensembles of nucleic acids providing new insights into their biological mechanisms [53]. Key to many of these studies has been the organic synthesis of ¹³C- and/or ¹⁵N-labeled phosphoramidites bearing these different modifications [38,54-56] (Figure 2).

The highly abundant epigenetic and epitranscriptomic modification m⁶A was shown to slow annealing of DNA and RNA duplexes by \sim 5- to 20-fold [55] through isomerization of the methylamino group [56], potentially explaining why the modification slows a variety of biochemical processes (Figure 3f). However, 5-formylcytosine (5fC), a semipermanent epigenetic modification, which also forms as an intermediate along the 5-methylcytosine (m³C) demethylation pathway, was shown to both slow DNA duplex annealing and to also accelerate duplex melting [57].

Uridine 5-oxyacetic acid (cmo⁵U), a naturally occurring tRNA modification, was shown to increase the population of an anionic Watson-Crick like G·U ES mismatch (Figure 3d), potentially explaining why the modification enhances the decoding of mRNAs containing G·U mismatches at the third wobble position [54]. By stabilizing paired nucleotides, 2'-O-methylation of the sugar moiety was shown to increase the abundance and prolong the lifetime of the TAR ESs [58]. Conversely, m⁵C was shown to minimally impact exchange kinetics between Watson-Crick and Hoogsteen conformations in duplex DNA [38].

Real-time NMR

Within cells, many RNAs function while they fold in a process referred to as 'co-transcriptional folding.' Many biochemical processes are coupled to co-transcriptional folding and require a kinetic description in terms of dynamic ensembles that evolve with time.

Time-resolved NMR experiments using photolabile protection groups are providing a window into the time evolution of biologically important meta-stable intermediates during co-transcriptional folding that plays important roles in the mechanisms of riboswitches [59,60]. Helmling et al. [60] showed that the formation of an antiterminator structure that ensures gene expression in the esoplasma florum 2'-deoxyguanosine (2'dG)-sensing riboswitch is slowed down below the rate of transcription when binding to its cognate ligand, thus mediating the switch into the functional OFF state (Figure 4a).

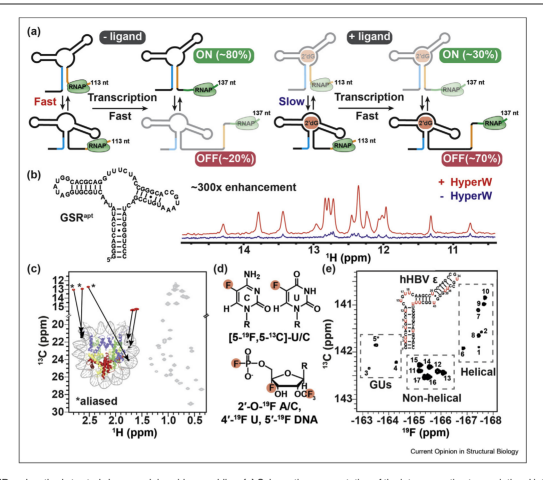
The scope of time-resolved NMR experiments to study RNA folding was recently dramatically improved through the application of the hyperpolarized water technique [61] to boost NMR sensitivity by 300-fold [62] (Figure 4b). With this approach, real-time NMR was successfully used to monitor RNA refolding with subsecond time resolution [62]. Time-resolved NMR experiments have also provided fresh insights into the folding landscape of DNA G-quadruplexes, revealing multiple folding pathways [63,64].

Applications to large molecular weight nucleic acids

Although most studies of nucleic acid dynamics have focused on relatively small motifs, recent advances push application to much larger molecular weight assemblies. Methyl-TROSY-based NMR techniques, which have been used with great success to probe the dynamics of large protein assemblies [2], have recently been successfully deployed to probe the dynamics of highmolecular-weight (as large as 200 kDa) DNA and DNA-protein complexes such as the nucleosome core particle [65]. The approach uses enzymatic methods to introduce m⁶A and m⁵C into DNA and then uses the methyl groups to probe dynamics using methyl-TROSY-based NMR experiments (Figure 4c). Developments in isotopic labeling strategies [66–69] as well as experiments for measuring RDCs in large molecular weight systems (up to 78 kDa [70]) are also making it possible to quantitatively measure NMR data in nucleic acid systems as large as 232 kDa [71].

Other exciting developments to probe the structural dynamics of nucleic acid include methods that introduce fluorine ¹⁹F into the bases [72,73], sugars [74–76], and the phosphate backbone [77] of nucleic acids (Figure 4d). These approaches take advantage of the high

Figure 4



Real-time NMR and methods to study large nucleic acid assemblies. (a) Schematic representation of the interconverting transcriptional intermediates of 2'dG-sensing riboswitch at transcript lengths 113 and 137 nt with and without ligand. Ligand binding slows the conformational exchange between the OFF and ON states below the transcription rate, triggering the OFF state function. The figure was adapted from Ref. [60]. (b) 1D imino spectra of the guanine-sensing riboswitch (GSR) aptamer domain (GSR^{apt}) of the *Bacillus subtilis xpt-pbuX* operon with (red) and without (blue) injection of hyperpolarized water (HyperW). The figure was adapted from Ref. [62]. (c) The 2D ¹³C – 1H HMQC spectrum of a nucleosome core particle (NCP) (shown is a cartoon representation of NCP, protein data bank [PDB] ID 6ESF), in which the 153-bp Widom DNA is highly deuterated with ¹³CH₃-methyl labeled at five sites indicated by black circles. The methyl group resonances belonging to the DNA and protein components are shown in red and gray, respectively, with the positions of the m⁶A methyls aliased (*). The figure was adapted from Ref. [65]. (d) Recently developed ¹⁹F labeling schemes for nucleic acids. (e) 2D ¹⁹F–¹³C TROSY spectra of 5-fluorouracil (red) modified human hepatitis B virus encapsidation signal epsilon (hHBV ε) element. The figure was adapted from Ref. [72].

sensitivity and broad chemical shift range of ¹⁹F nuclei as well as the favorable relaxation properties of the aromatic ¹⁹F—¹³C spin pair in 2D transverse relaxation-optimized spectroscopy experiments (Figure 4e) [72,73,78]. Coupled with advances in the synthesis of site-specifically labeled and/or modified nucleotides and phosphoramidites, and methods to incorporate them into large nucleic acids, these approaches should streamline dynamics studies of large nucleic acid assemblies.

Future perspective

Owing to continued advances in NMR methods, studies are beginning to view the vast landscape of

conformations in the dynamic ensembles of nucleic acids. Using the dynamic ensemble description, a deeper and more quantitative understanding of nucleic acid mechanisms is beginning to emerge. The continued development of computational methods to model nucleic acid structure and dynamics and their integration with data from NMR and other biophysical approaches not reviewed here will be necessary to increase the accuracy, throughput, and robustness of current ensemble determination methods. We can anticipate new insights into how nature amplifies the specificity and functional complexity of nucleic acids by evolving machinery capable of 'reading out' features of their dynamic ensembles.

Conflict of interest statement

HMA is an adviser to and holds an ownership interest in Nymirum Inc., an RNA-based drug discovery company. The other authors declare no competing interests.

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