



Rapid access to RNA resonances by proton-detected solid-state NMR at >100 kHz MAS†

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Fast (>100 kHz) magic angle spinning solid-state NMR allows combining high-sensitive proton detection with the absence of an intrinsic molecular weight limit. Using this technique we observe for the first time narrow ¹H RNA resonances and assign nucleotide spin systems with only 200 μg of uniformly ¹³C, ¹⁵N-labelled RNA.

The discovery of the three-dimensional structure of ribonucleic acids (RNA), either in isolation or in complex with proteins, is essential to explain their numerous functions in cells. The structure of RNA is best studied by Nuclear Magnetic Resonance (NMR) spectroscopy, which can cope with the inherent flexibility of the polymer. However, unlike proteins, RNA consists of building blocks of low chemical diversity, resulting in poor chemical shifts dispersion. Consequently, NMR analysis of RNAs >40–50 nucleotides requires laborious selective isotopic labeling. Furthermore, the sensitivity and the resolution of solution NMR decrease with the molecular mass, posing a practical limit of about 60–80 nucleotides.

Magic-angle spinning (MAS) solid-state NMR spectroscopy¹ does not have an intrinsic molecular weight limit and holds the potential of being applicable to RNAs of any size. Recently, MAS NMR was employed for the first time to determine *de novo* the structure of a RNA in a non-diffracting solid preparation, the 26mer Box C/D RNA in complex with the protein L7Ae from *Pyrococcus furiosus* (Pf).^{2,3} The strategy comprised of ¹³C- and ¹⁵N-detected experiments for both resonance assignment and measurement of structural restraints. The low sensitivity

of ¹³C, ¹⁵N nuclei rendered high-dimensional NMR methods impractical, even when using >6–8 mg of RNA.^{2,3} The severe overlap of the resonances in two-dimensional spectra was lifted using single (A, U, C, G) nucleotide-type selective labeling for the assignment of nucleotide spin systems, and double (AU, CG, *etc.*) nucleotide-type selective labeling for sequential assignment and collection of structural restraints. Despite successful, this strategy necessitated the measurement of several samples, resulting in a substantial commitment of laboratory and instrument times. ¹H-detection provides the highest sensitivity; nonetheless, it has long been a unique mark of solution NMR spectroscopy, because of the extreme line broadening caused in solids by the dense network of strong ¹H–¹H dipolar couplings. ¹H-detection at moderate (24 kHz) MAS rates was shown to be practical with perdeuteration and partial reintroduction of exchangeable protons.^{4–6} However, in RNA this labeling scheme limits the NMR active probes to labile imino and amino ¹Hs, and prevents efficient resonance assignment strategies.^{5,7,8} Faster MAS rates are necessary to lift the requirement of proton dilution and at the same time ensure sensitivity and resolution. Recently, ¹H detection was used on a uniformly labeled 23mer DIS HIV-1 RNA to observe base pairs through ¹⁵N–¹⁵N proton assisted recoupling (PAR); however, at the MAS rate of 40 kHz, the proton resolution (500 Hz) was insufficient to enable site-specific assignments.⁹

The availability of hardware that is capable to reach fast (60–100 kHz) spinning rates has been critical for the development of ¹H-detected MAS NMR experiments for proteins, allowing faster backbone and side-chain assignments, as well as structural and dynamics studies.^{10–12} Despite its lower proton density when compared to proteins, RNA is a challenge for effective averaging of dipolar interactions by MAS, due to the spatial clustering of protons and the intrinsic flexibility. Here we show that MAS rates exceeding 100 kHz open an avenue for the study of fully protonated RNA at high resolution and sensitivity. 2D ¹H-detected fingerprint spectra can be acquired in minutes to hours using as little as 200 μg of RNA. This enables the exploitation of high-dimensional NMR techniques, which allow

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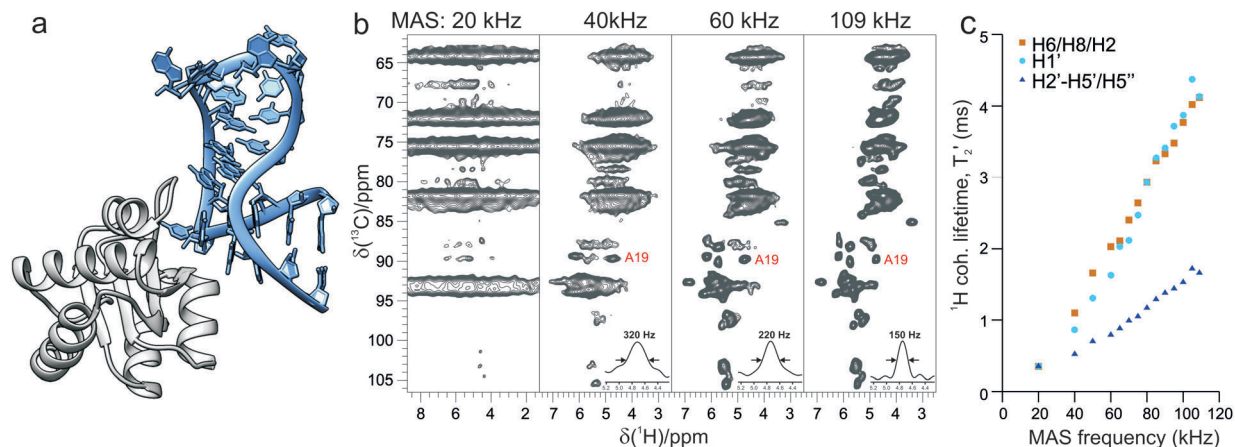


Fig. 1 (a) Structural model of the 26mer Box C/D RNA (light blue) bound to the L7Ae protein (grey) from *Pf*, established by alignment of the previously determined MAS NMR structure (PDB 2N0R)³ of the RNA from *Pf* with a homologous RNA sequence in a larger complex (PDB 3NMU). (b) Ribose and base C5–H5 regions of ¹³C–¹H CP-HSQC spectra recorded on a 800 MHz spectrometer at MAS rates of 20, 40, 60 and 109 kHz, characteristic for commercial 3.2, 1.9, 1.3 and 0.7 mm MAS probes. 1D traces of a representative resonance with the corresponding proton line width are shown in the bottom right corner. (c) Bulk coherence lifetimes (T_2') of base H6, H8 and H2 (brown squares), ribose H1' (cyan circles) and other ribose H2'–H5'' protons (blue triangles) measured on a 800 MHz spectrometer and MAS frequencies from 20 to 109 kHz.

the identification of ¹H, ¹⁵N and ¹³C resonances of nucleotide spin-systems in the 26mer Box C/D RNA from a single, uniformly ¹³C, ¹⁵N-labelled sample.

Fig. 1 shows the remarkable effect on ¹H resolution and sensitivity observed for the Box C/D RNA upon increase of the magic-angle spinning rates. 2D dipolar-based ¹³C–¹H cross-polarization (CP)-HSQC experiments,¹³ tailored for ribose resonances on a 800 MHz spectrometer, improve from a featureless spectrum at 20 kHz to a resolved fingerprint at 110 kHz MAS. ¹H line widths decrease substantially due to effective averaging of dipolar ¹H–¹H interactions, which is reflected by a nearly linear increase of the ¹H coherence lifetime (T_2') with MAS rates up to 110 kHz (Fig. 1c). In this forefront spinning regime, anomeric ribose H1' and base H6/H8/H2 resonances reach above 4 ms of bulk T_2' ; this corresponds to an 80 Hz homogeneous contribution to the ¹H line width of 130–170 Hz. At comparable MAS conditions, such long coherence lifetimes have been reported only for proteins in microcrystalline preparations featuring low dynamics.¹⁴ The resolution of ribose H2'–H5'/H5'' resonances is lower, as these protons are involved in a stronger network of dipolar couplings; however, they feature better chemical shift dispersion than their carbon counterparts.^{15,16} Seven C5–H5 correlations are visible out of the nine pyrimidines (Fig. S1a, ESI[†]).

Fig. 2 shows a magnification of the two anomeric and base ¹³C–¹H fingerprints recorded on a 1 GHz spectrometer. Analogously to solution NMR, non-helical regions ($\delta(C1')$ < 90 ppm, Fig. 2b) display the best dispersion of C1'–H1' resonances. 2D ¹³C–¹H correlations of base resonances are more overlapped (Fig. 2c), with only two bases showing distinct ¹³C chemical shifts and line widths of 150 Hz (¹H) and 120 Hz (¹³C). In the C2–H2 region (Fig. S1b, ESI[†]), five resonances can be resolved out of the nine As. The crowded appearance of the spectrum indicates a broader line width of most H8/H6 (purine/pyrimidine) with respect to the ribose ¹Hs. We hypothesized

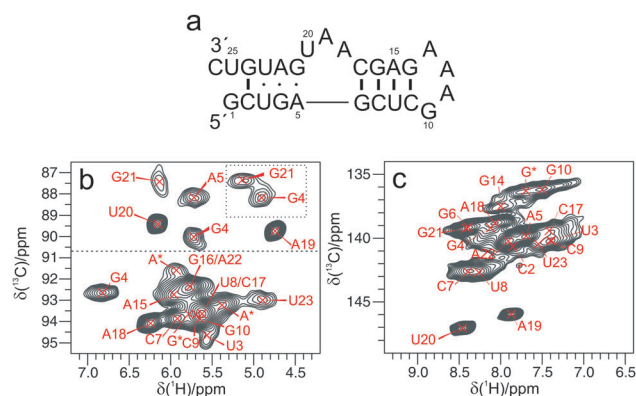


Fig. 2 (a) Sequence of the *Pf* 26mer Box C/D RNA. (b and c) Expansions of the ribose C1'–H1' and base C6/8–H6/8 regions of ¹³C–¹H CP-HSQC spectra recorded on a 1 GHz spectrometer and 111 kHz MAS. In (b) two resonances in the dotted box correspond to C4'–H4' correlations, and the horizontal dashed line at approx. 90 ppm ¹³C separates C1'–H1' resonances of nucleotides located in helical and non-helical regions. The signals are labeled according to the ¹H assignment determined in this work and the previously reported sequence specific ¹³C, ¹⁵N assignment.² The asterisk indicates resonances of non-assigned nucleotides.

that the broadening of the base resonances could stem from incomplete suppression of the dipolar interactions with either (i) the hydrogens of water molecules entering the major groove¹⁷ or (ii) the close-by H2's in helical regions. To test these two hypotheses we acquired CP-HSQC spectra on two additional samples: (i) L7Ae-Box C/D RNA complex dissolved in 90%/10% D₂O/H₂O rather than 100% H₂O. For this sample the H6/H8 line widths should not suffer from the vicinity to ¹Hs of the bound water; (ii) the L7Ae-Box C/D RNA complex with ²H, ¹²C, ¹⁴N-C, U, A and ¹H, ¹³C, ¹⁵N-G, dissolved in 90%/10% D₂O/H₂O. For this sample the H8 line width of the ¹³C, ¹⁵N-Gs should not suffer from the vicinity with the H2's of the preceding nucleotides. The fingerprint in the base region is



almost identical for all samples (Fig. S1, ESI[†]), with a slight improvement of the line width in deuterated buffer. Thus, we conclude that the largest contribution to line broadening must be due to local structural heterogeneity that mostly affects the aromatic ¹H shifts.

In the following, we demonstrate that, despite the limited chemical shift dispersion of RNA, 14 spin systems can be established out of 19 nucleotides in the structured regions of the 26mer Box C/D RNA (excluding the loop and the ends) without resorting to nucleotide-specific labeling. This is possible due to efficient magnetization transfers at MAS > 100 kHz and high-dimensional experiments.

Leveraging the high-sensitivity of the 2D (H)CH spectrum, we implemented an (H)NCH experiment. This is comparable to the (H)NCAHA experiment for proteins,^{13,18} which we adapted to the topology of ¹H, ¹³C and ¹⁵N spins in RNA and to the distribution of ¹³C chemical shifts. The experiment starts with a long-range ¹H-¹⁵N transfer, followed by a band specific CP step tuned to either ribose or aromatic ¹³C offsets, and yields either ribose-specific N1/N9-C1'-H1' or base-specific N1-C6-H6/N9-C8-H8 correlations (Fig. 3a and b). The ¹⁵N chemical shifts allow distinguishing pyrimidine from purine correlations (Fig. 3c and d) as well as associating C1'-H1' with C6-H6/C8-H8 resonances. These correlations are similar to those established in solution NMR, with the advantage that the dipolar-based long-range ¹H-¹⁵N and the one-bond ¹⁵N-¹³C

transfers used here maintain high efficiencies even for large RNAs, due to the sufficiently long $T_{1\rho}$ relaxation times at fast MAS (>12 and 50 ms for ribose ¹H and ¹³C, respectively; Fig. S3, ESI[†]). The ribose-specific (H)NCH spectrum shows excellent sensitivity (Fig. 3e-g); in less than 12 hours, resonances could be correlated for 17 out of the 19 previously assigned² spin systems (10 N9-C1'-H1' and 7 N1-C1'-H1' spin sets belonging to the region 3-10 and 15-23, with only nucleotides G14 and G24 missing, Table S4, ESI[†]). In the base-specific (H)NCH spectrum, the sensitivity of the transfer is affected by the multiple competitive coherence transfer pathways from the N1/N9 spins towards C2/C4 and C6/C8, as previously observed in CNC transfer schemes.² Nonetheless, we obtained correlations for 19 nucleotides in 26 hours.

At fast MAS rates, the long ¹³C $T_{1\rho}$ relaxation times enable efficient scalar transfers between bonded ¹³C atoms. These transfers allow establishing entire ribose spin systems in an efficient manner, as scalar ¹³C transfers are easier to interpret than those based on proton-driven spin diffusion (PDS) at slower MAS rates.¹⁹ Scalar transfers can be triggered by inserting into the (H)CH module a rotor-synchronized, low-power TOBSY mixing,^{20,21} or even less power-demanding schemes, such as WALTZ-16.²² In the resulting HCCH-TOCSY^{18,23} experiment, long mixing times (25 ms) are used to spread magnetization to all ribose ¹³C spins. The information content can be maximized if both ¹H and ¹³C shifts of starting and ending CH groups on

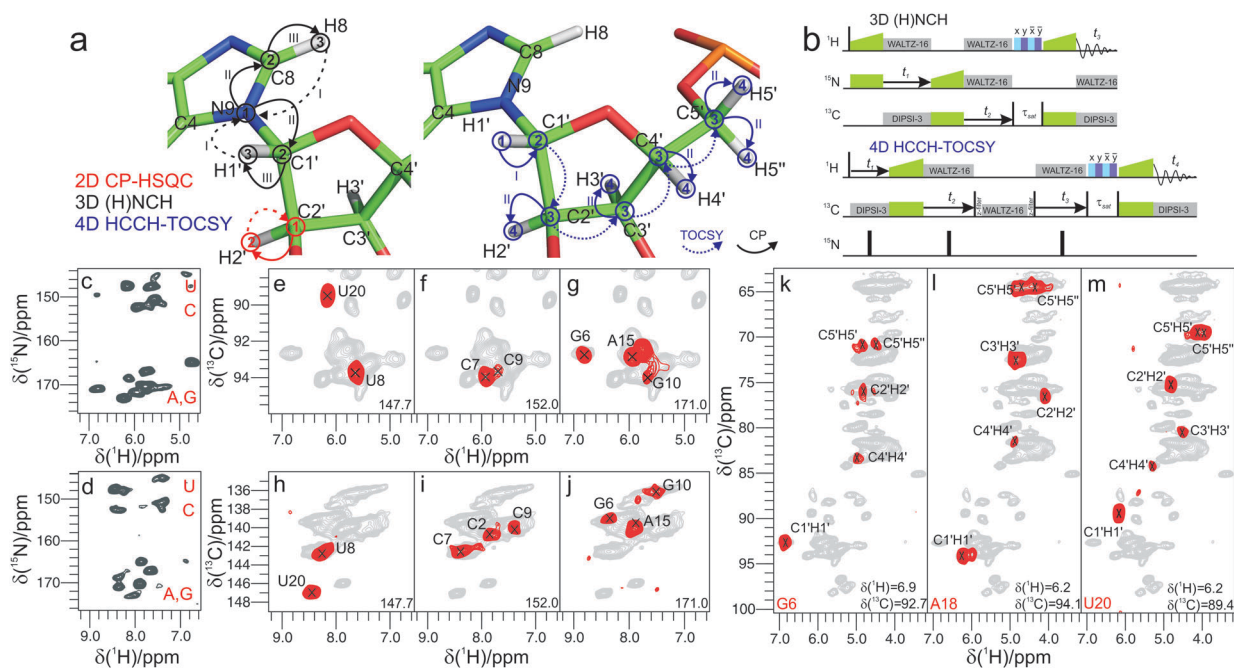


Fig. 3 (a and b) Magnetization transfer and radiofrequency irradiation schemes: 2D ¹³C-¹H CP-HSQC (color-coded in red), 3D (H)NCH (in black), and 4D HCCH-TOCSY (in blue). Arabic numbers in (a) indicate spectral dimensions (t_1 - t_4) in the pulse schemes (b); roman numbers indicate the CP transfers. (c and d) 2D ¹⁵N(t_1)-¹H projections of 3D (H)NCH experiments tuned to either ribose or base resonances. (e-j) Representative ¹³C-¹H cross-sections from the (H)NCH spectra showing either ribose N1/N9-C1'-H1' (e-g) or base N1-C6-H6 or N9-C8-H8 correlations (h-j), at the ¹⁵N frequency indicated in each panel. (k-m) Assignment of ribose spin systems with the 4D HCCH-TOCSY experiment. Representative 2D planes from the 4D experiment showing the spin systems of G6 (k), A18 (l), U20 (m). ¹H and ¹³C frequencies are indicated in each panel. For reference, in panels (e-m) the red contours of either (H)NCH or HCCH-TOCSY spectra are overlaid onto 2D CP-HSQC spectra (in grey). (H)NCH spectra were recorded on a 1 GHz spectrometer at 111 kHz MAS, while the HCCH-TOCSY spectrum was recorded on a 800 MHz spectrometer at MAS frequency of 100 kHz.



the mixing pathway are recorded in a 4D dataset, using non-uniform sampling (NUS) to ensure spectral resolution.^{24–28} 2D cross-sections from a 4D HCCH-TOCSY acquired for 3 days with exponentially relaxation-weighted random NUS,^{29,30} are shown in Fig. 3k–m. For most residues, the separation of the H1' and C1' resonances translates into fully resolved planes displaying the entire ribose spin systems. For few nucleotides in helical regions (U8/G10/C17), low dispersion of the C1' and H1' chemical shifts prevents the separation of their spin systems. To lift this ambiguity, we designed a 4D (H)NCCH-TOCSY experiment, which uses the N1/N9 dimension to further separate overlapped ribose spin systems (Fig. S4, ESI†). Even with moderate sampling of the ¹⁵N dimension, the experiment, which ran for 4.5 days, allows easy separation of purines and pyrimidines, as well as of spin systems within each subclass (U8 and C17, Fig. S4c–e, ESI†). All in all, we could unambiguously assign 14 out of 19 ribose spin systems in the structured part of the Box C/D RNA (Table S4, ESI†). The cross peak intensity of a few nucleotides was either low (G10, Fig. S4d, ESI†), or weak beyond detection (C2, C9, G14, A15 and G24). These nucleotides are mostly located in the junction or ends of secondary structure elements, where increased internal motions impair transfer efficiency during the two long CP steps required in ¹⁵N-resolved multidimensional spectra (Fig. S4a and b, ESI†).

In summary, we demonstrate that ¹H-detection at fast MAS rates can be applied to nucleic acids and allows the assignment of ¹H, ¹³C and ¹⁵N resonances in nucleotide spin systems with only 200 µg of uniformly ¹³C, ¹⁵N labeled RNA. In comparison to ¹³C-detected experiments, ¹H-detection bypasses the need of nucleotide-type selectively labeled samples, resulting in a fast and cost-efficient route to site-specific studies. The strategy we devised is similar to that used in solution NMR, but has the advantage of being applicable to large RNAs or RNA–protein complexes. We are currently developing this strategy towards the sequential assignment of RNA resonances and the collection of structural restraints based on ¹H-detection. The RNA–protein complex used in this study was in microcrystalline form. However, all building blocks of the pulse sequences have been shown to perform also on non-crystalline formulations.^{18,31–33} Given the difficulty of obtaining good-diffracting crystals for large RNA–protein complexes, we anticipate that these results will strengthen the role of MAS NMR in RNA structural biology, and more generally provide momentum in the atomic-level description of structure and dynamics of RNA.

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Conflicts of interest

There are no conflicts to declare.

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