The ribose 2'-hydroxyl group in RNA is largely responsible for differences in structure, hydration, and thermodynamic stability relative to DNA.1 In addition to promoting the C3'-endo pucker of the ribose, the 2'OH group provides a scaffold for the water network in the minor groove of RNA2 and plays a role in RNA catalysis.3,4 In H2O solution at low temperature, the 2'OH protons in double-helical, A-form RNA can be observed by NMR, and OH orientations have been studied primarily on the basis of semiquantitative NOE analysis.1,5,6 Energetically, three different orientations are plausible,7 referred to as the O4', O3', and base domains. A very recent study by Fohrer et al.8 concludes that OH bond orientations alternate between the base and the O3' domain.

Here, we describe a novel 3D NMR experiment that simultaneously yields 13C-dispersed NOEs from ribose protons to 2'OH, as well as E.COSY-type 1J_C–OH couplings. Highly reproducible and narrowly clustered 1J_C1–2OH and 1J_C2–2OH, combined with quantitative evaluation of the NOEs to 2'OH, indicates that in our system all OH groups predominantly occupy the base region. The pulse sequence for measurement of 1J_C–OH is shown in Figure 1 and uses the IPAP approach10 to encode the 1J_C–OH couplings into the 2'OH resonance frequency difference in two 3D NOESY spectra, obtained from the sum and difference of an in-phase (IP) and an anti-phase (AP) type of spectrum (i.e., IP + AP and IP − AP). A description of the experiment in terms of product operator formalism is provided as Supporting Information. In brief, a constant-time 1H element (time points a–f) precedes the NOE mixing period, t_m, which is followed by WATERGATE11 detection. Depending on whether the dashed 180° 13C pulse at the midpoint of the 1H(−11C) rephasing interval is applied or not, the detected OH proton is either anti-phase or in-phase with respect to the ribose 13C on whose attached 1H the magnetization originated at time point a. In the usual IPAP manner,10 the sum and difference of the two spectra yield the individual 2'OH−{13C} doublet components, and coupling constants between the 2'OH proton and the ribose carbons can be extracted from their frequency difference. The measured splittings, in principle, can be reduced by 13C spin flips during t_m and detection,12 but not by solvent exchange of the 2'OH. Considering that the 13C T1 relaxation time exceeds the sum of t_m and the t_c detection period by more than an order of magnitude, such reduction is negligible, however. For highest S/N, the intensities of NOEs between the 2'OH and the ribose protons are obtained from the IP spectrum only; due to partial cancellation of the anti-phase 2'OH−{13C} doublet components, the AP spectrum is much weaker.

The favorable relaxation properties of multiple quantum coherence,13 applicable during the 25 ms 13C chemical shift labeling, and 1J_CC decoupling resulting from constant-time evolution result in good sensitivity and 13C resolution. Importantly, the ribose 1H chemical shift is recorded simultaneously during this 13C evolution period14 and its preceding and following δ delays, yielding a 31 ms constant-time sampling of ribose 1H evolution without incurring any additional relaxation losses. J_HH and multi-bond J_CC and J_CCC couplings remain active during these constant-time evolution periods, but appropriate correction factors are derived from the relative intensities of “diagonal” resonances observed for the various ribose protons (Supporting Information).

The experiment of Figure 1 was applied to a 1.0 mM solution of a uniformly 13C/15N-enriched oligomer (10 mM phosphate buffer, pH 6.8, 10 mM NaCl, 0.02 mM EDTA, 95% H2O/5% D2O), derived from helix-35 of E. coli 23S ribosomal RNA (Figure 2) at 6 °C and 800 MHz. The exchange rate constants of the 2'OH protons with bulk water, measured from a series of 2D NOESY experiments at 6 °C in which the 2'OH proton magnetization is pre-inverted, were found to range from 18 s−1 for C758 to 52 s−1 for G738 (Supporting Information). The IP 3D NOESY spectrum shows relatively strong NOEs from the intra-ribose H2' to 2'OH protons (Figure 2A). On average, somewhat weaker NOEs from 2'OH to H1' are observed, but NOEs from 2'OH to H3' and H4' are much weaker or absent (Supporting Information). Fast exchange between 2'OH protons and water in the loop region prevented measurement of their NOEs. The couplings between the 2'OH protons and C2' or C1' are readily measured from the frequency difference in the 2'OH dimension (F1) of the 3D constant-time HMQC−IPAP−NOESY spectrum (Figure 2). The coupling constants between the 2'OH protons and the ribose carbons are remarkably similar for the different riboses. For example, 2J_C2'–2OH values all fall in the −2.2 to −3.3 Hz, indicative of the high precision at which these couplings can be measured. Three-bond couplings to C1' are also very uniform and small (1.3 ± 0.7 Hz, Supporting Information) indicative of a gauche conformation.8 The very weak intensity of the H3' to 2'OH NOE prohibits precise measurement of 3J_C3'–2OH from the 3D constant-time HMQC−IPAP−NOESY HMBC spectrum. Instead, these values were measured from quantitative HMBC experiments16 (Supporting Information). Measured values are again quite uniform (4.2 ± 0.7 Hz) and much larger than 3J_C1–2OH (Table 2S), in agreement with the relative trend of these couplings recently reported by Fohrer et al.8 In comparison to this study, much less variation in our 3J_C–OH couplings is seen, which may in part derive...
from the unique ability of E.COSY experiments to accurately measure minute or even zero couplings that are not affected by the highly non-uniform OH solvent exchange rates (Table 1S) or may result from the difference in the system studied. The relative size of $J_{OC\text{-}OH}$ and $J_{OH\text{-}OH}$ suggests that the 2′OH groups in A-form RNA duplexes are oriented toward either the base domain (i.e., approximately trans to C3′ but gauche to C1′) or a small region of the O3′ domain (i.e., 2′OH–O2′–C2′–H2′ dihedral angle > 120°) and excludes significant occupation of the O4′ domain.64 The ambiguity between the base and O3′ domains can be resolved by quantitative analysis of the 2′OH NOEs, also available from the experiment of Figure 1.

The high resolution in the indirect 1H and 13C dimensions of the 3D constant-time HMQC–IPAP–NOESY spectrum strongly reduces overlap compared to 2D NOESY spectra and permits unambiguous assignment and analysis of the 2′OH NOE cross-peaks. In addition to the NOEs from H1′ to H2′ to 2′OH, discussed above, weak NOEs between 2′OH and the intra-nucleotide H4′ protons are observed, whereas for eight nucleotides the H3′–2′OH NOEs also are above the signal-to-noise threshold. Although NOE intensities to 2′OH are differentially affected by evolution of J couplings during the constant-time evolution period in the pulse scheme of Figure 1, correction factors can be derived from resolved “diagonal” 13C–1H correlations in the 3D spectrum, and within the A-form helix are found to be quite uniform for a given type of carbon (Supporting Information). Comparing these corrected NOE intensities to the H1′–2′OH NOE indicates that H2′–2′OH is 1.6–3.5-fold stronger, whereas H3′–2′OH and H4′–2′OH are weaker by factors in the ranges of 0.1–0.4 and 0.2–0.5, respectively (Table 3S). Comparison of these normalized NOE intensities with values expected as a function of the 2′OH–O2′–C2′–H2′ dihedral angle (Supporting Information) indicates that the base domain is the only one where all the experimental NOEs are compatible with the calculated values, although low level occupation (≤20%) of the other domains cannot be excluded. These NOE-derived results resolve the conformational ambiguity associated with the observation of very small $J_{OC\text{-}OH}$ values and relatively large $J_{OH\text{-}OH}$ couplings. The presence of 2′OH groups predominately in the base region is perfectly compatible with the oxygen electron density seen for H-bond accepting water molecules in the minor groove of an A-form RNA helix, observed by X-ray crystallography.2

The 3D constant-time HMQC–IPAP–NOESY experiment takes advantage of overlapping, simultaneous 1H and 13C evolution periods and offers remarkably well-resolved 3D NOE spectra that permit measurement of J couplings through the E.COSY principle. The experiment can be readily adapted for the measurement of other NOEs and J couplings in both nucleic acids and proteins.

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Supporting Information Available: Four tables with 2′OH solvent exchange rate constants, $J_{OC\text{-}OH}$ coupling constants, and normalized experimental NOEs from HMQC– and HSQC–NOESY spectra; two figures for the projected planes (C2′/C3′/C4′) of the HMQC– and HSQC–NOESY; one figure for the pulse sequence and spectra of quantitative HMBC; one figure displaying interproton distances and corresponding NOE ratios calculated as a function of the 2′OH–O2′–C2′–H2′ dihedral angle; product operator formalism analysis and detailed parameters of the pulse scheme of Figure 1; derivation of NOE correction factors for the effect of passive couplings. This material is available free of charge via the Internet at http://pubs.acs.org.

References


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