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A NMR strategy to unambiguously distinguish nucleic acid hairpin and duplex conformations applied to a Xist RNA A-repeat

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ABSTRACT

All RNA sequences that fold into hairpins possess the intrinsic potential to form intermolecular duplexes because of their high self-complementarity. The thermodynamically more stable duplex conformation is favored under high salt conditions and at high RNA concentrations, posing a challenging problem for structural studies of small RNA hairpin conformations. We developed and applied a novel approach to unambiguously distinguish RNA hairpin and duplex conformations for the structural analysis of a Xist RNA A-repeat. Using a combination of a quantitative HNN-COSY experiment and an optimized double isotope-filtered NOESY experiment we could define the conformation of the 26-mer A-repeat RNA. In contrast to a previous secondary structure prediction of a double hairpin structure, the NMR data show that only the first predicted hairpin is formed, while the second predicted hairpin mediates dimerization of the A-repeat by duplex formation with a second A-repeat. The strategy employed here will be generally applicable to identify and quantify populations of hairpin and duplex conformations and to define RNA folding topology from inter- and intra-molecular base-pairing patterns.

INTRODUCTION

All RNA sequences that fold into hairpins possess the intrinsic potential to form intermolecular duplexes because of their high self-complementarity. The thermodynamically more stable duplex conformation is favored under high salt and high RNA concentrations. This is a challenging problem for structural studies: crystallization for X-ray crystallography often requires high salt conditions and NMR structural studies require sample concentrations in the millimolar range. Unambiguous identification of sample stoichiometry under experimental conditions is essential as early as possible in RNA structural studies to ensure that the RNA is present in a biologically relevant conformation.

Xist (X inactivation specific transcript) RNA is a large non-coding RNA essential for the initiation of X-inactivation in mammalian females (1). Early in embryonic development it is expressed from the X-chromosome that will be silenced and coats it in cis, which coincides with transcriptional shutdown through an unknown mechanism (2). The conserved so-called ‘A-repeats’ at the 5'-end of Xist are essential for its silencing function, while several other regions are redundantly responsible for chromosome association (3) (Figure 1A). In humans, the A-repeats are constituted of 7.5 copies of a 26 nt motif, connected by long U-rich linkers. A Mfold secondary structure prediction of a single A-repeat suggested a double hairpin structure where the two hairpins possibly stack on top of each other (3) (Figure 1B). As no structural information on the A-repeats is available, we started NMR studies on a single A-repeat (Figure 1C) with the goal to solve its atomic structure and to obtain molecular insight into X-Inactivation. The construct used in our study shown in Figure 1C is identical to the 5th human A-repeat, apart from switching the positions of G and C in the third G–C base-pair to facilitate chemical shift assignments. Previous studies have shown that altering the sequence of the stem in hairpin 1 does not influence Xist activity as long as base pairing is not disrupted (3). During our structural studies we encountered difficulties...
complete NMR assignments of the second predicted hairpin. Signals from this hairpin were broad, and sometimes doubled (data not shown), which indicated possible dynamics or sample heterogeneity, although native gel analysis of the 26-mer A-repeat RNA suggested a homogenous monomeric population (Supplementary Material). The strategy described in this article was essential to characterize and distinguish the intramolecular and intermolecular base pairs in monomeric/dimeric forms of RNA at sample conditions required for structural biology. The approach provided valuable insight into the possible architecture of the A-repeats.

NMR structures are determined from proton–proton distance restraints derived from nuclear Overhauser effects (NOEs), dihedral angle restraints derived from J-coupling constants and Residual Dipolar Couplings (4–8). Since chemical shifts and the pattern of NOEs in NMR spectra of hairpin and duplex species of a given nucleic acid are very similar, standard NMR techniques do not normally suffice to distinguish between the two. Other methods to determine the stoichiometry of nucleic acids include native gel electrophoresis and UV melting (9). However, these experiments are carried out at low concentrations. Hydrodynamic measurements such as ultracentrifugation, light scattering and NMR diffusion (10) are also available, but can be difficult to interpret for elongated molecules like RNA and for low molecular weight systems.

Several NMR methods have been developed to unambiguously distinguish between monomers and dimers of nucleic acids under NMR conditions (11–14). All these use an equimolar mixture of isotope labeled and unlabeled RNA/DNA. In such a mixture, different populations will exist for a RNA hairpin conformation with intramolecular base-pairing or for a duplex involving intermolecular base pairs. If the RNA folds into a hairpin, 50% of the molecules will be labeled and 50% will be unlabeled (Figure 2A). If the RNA adopts a duplex form, 25% of the molecules will be labeled, 25% will be unlabeled and 50% of the duplexes will consist of one labeled and one unlabeled strand (Figure 2B). Nucleic acid duplexes are detected based on differences in NMR parameters such as chemical shifts (11), cross-hydrogen bond $J_{NN}$ coupling constants (12,13) or NOEs (14).

The chemical shift based method (11) requires the introduction of a mutation that shifts a G imino signal to a characteristic frequency in the case of duplex formation. An obvious drawback of this method is that a change in the RNA sequence can possibly alter the monomer–dimer equilibrium.

The method using scalar couplings (12,13) is based on the HNN-COSY experiment (15), which relies on the transfer of magnetization across the hydrogen bond in nucleic acid base pairs using the two-bond $J_{NN}$ coupling. RNA duplexes are detected through a difference in intensities between cross- and diagonal peaks in HNN-COSY spectra, as in a duplex species with one labeled and one unlabeled strand magnetization transfer over the hydrogen bond cannot take place. We have extended this method and demonstrate that the molar fractions of hairpin and duplex species in a mixed population can be determined.

The NOE based method uses isotope editing/filtering techniques to distinguish intermolecular and intramolecular NOEs (14). NOEs arising from pairs of protons where one is bound to $^{15}$N and the other one bound to $^{14}$N can only arise from a duplex species. This is a complementary approach if an efficient J-based magnetization transfer across the base pair is not possible, for example in G–U pairs. We have employed simultaneous filters for protons attached to nitrogen and to carbon. Thereby, intermolecular NOEs involving imino-imino (in G–U base pairs) and imino–amino (in G–C base pairs) can be discriminated from NOEs involving imino protons and H2s in A–U base pairs.

Here we present the combination of HNN-COSY and NOE based methods to distinguish between monomeric
hairpin and duplex conformations of nucleic acids. The approach is demonstrated and was crucial for the determination of the architecture of the Xist RNA A-repeats in our structural studies.

MATERIALS AND METHODS

Sample preparation

$^{13}$C,$^{15}$N uniformly labeled and unlabeled r(GGCCGC [AUACG] GCCCUUCGG [AUACG] CUGC) A-repeat 26-mer RNA (consisting of both predicted A-repeat hairpins, Figure 1C) and r(GGCCGC [AUACG] GCCGCC) A-repeat 14-mer RNA (containing only the first hairpin, Figure 1D) was prepared by in vitro transcription with T7 RNA polymerase (EMBL Protein Expression and Purification Core Facility, Heidelberg, Germany) using synthetic DNA oligos (MWG Biotech/Eurofins MWG Operon, Ebersberg, Germany) as a template (16) and either $^{13}$C,$^{15}$N labeled (Silantes, Munich, Germany) or unlabeled (Sigma-Aldrich, Munich, Germany) NTPs. The positions of G and C in the fourth base pair of the first predicted hairpin were reversed to facilitate NMR assignment. 2'-O-methyl groups were incorporated into the two 5'-residues of the DNA template strand to reduce the amount of $n + 1$ transcription products (17).

The RNA was purified on preparative denaturing 20% (w/v) polyacrylamide (19:1 acrylamide: bisacrylamide) gels. Gel bands were visualized by UV shadowing using fluorescent thin layer chromatography plates (Whatman) and the band corresponding to the full-length product was excised and electroeluted (Schleicher & Schuell/Whatman, Dassel, Germany). The RNA was precipitated, resuspended and extensively washed in 1K MWCO concentrators (Pall, Dreieich, Germany) with 10 mM NaH$_2$PO$_4$/Na$_2$HPO$_4$ buffer (pH 6.0) of progressively decreasing NaCl concentration (1–0 M). Finally the RNA was desalted (PD-10, Amersham/GE Healthcare, Freiburg, Germany) and lyophilized.

NMR samples were prepared in 10 mM NaH$_2$PO$_4$/Na$_2$HPO$_4$ buffer (pH 6.0), 100 mM NaCl, 0.02% Azide, 0.02 mM EDTA in 95% H$_2$O, 5% D$_2$O. The RNA concentration of the $^{13}$C,$^{15}$N uniformly labeled sample for recording the HNN-COSY reference experiment was 0.8 mM. An equal amount of unlabeled RNA was added to this sample, mixed, lyophilized and resuspended in the same amount of H$_2$O/D$_2$O. Both the edited/filtered NOESY and HNN-COSY were run on this 1.6 mM sample. In addition, the HNN-COSY was repeated on the same sample diluted to a total RNA concentration of 0.8 mM. Just before the NMR measurements samples were heated to 95°C for 5 min followed by snap-cooling on ice with the rationale to trap the kinetically favored intramolecular monomeric hairpin conformation over a possible intermolecular dimer. The stoichiometry of the NMR samples was initially checked with native PAGE (Supplementary Material).

NMR Spectroscopy

HNN-COSY. NMR experiments were recorded at 5°C on a Bruker DRX600 spectrometer equipped with a cryoprobe. HNN-COSY spectra were recorded with the pulse sequence described by Dingley & Grzesiek (15) shown in Supplementary Figure 1. The delay $T$ for evolution of the $^{13}$J$_{NN}$ coupling was set to 20, 30 and 40 ms. One thousand and twenty-four complex points were collected for each complex data point. A 13C filter in which anti-phase magnetization is created for $^{13}$C/14N bound protons. The $^{15}$N 180° pulses are composite pulses of the form $90°/180°/90°$, $\psi = x$, $\gamma$, $x$ for distinguishing $^{15}$N/14N bound protons. Phase cycle: $\phi_1 = x$, $\gamma$, $x$ + TPPI; $\phi_2 = x$, $\gamma$, $-x$, $\phi_{180} = x$, $\gamma$, $-x$, $\gamma$, $x$; Water suppression is accomplished by WATERGATE (25) combined with water flip back (26). Gradients of 800 µs length were applied with 30% (g1) and 50% of maximal power (g2).

Isotope-edited/filtered NOESY. The $\omega_1$, $\omega_2$ double isotope-filtered NOESY experiment employed is shown in Figure 3. Compared to the experiment proposed by Aboul-ela et al. (14) for discrimination between intra- and intermolecular NOEs we have added a $^{13}$C filter in $\omega_1$. This allows the discrimination of intermolecular imino–imino NOEs in G–U base pairs, and the identification of intermolecular imino-H2 NOEs in A–U base pairs. Moreover, the $\omega_1$ filter is combined with semi-constant time chemical shift evolution (18,19) in $\tau_1$ to exploit the filter delay 2$\Delta$ simultaneously for chemical shift labeling and J-coupling evolution (20) for improved sensitivity. Editing/filtering for both $^{15}$N and $^{13}$C in $\omega_1$ is accomplished by a ‘jumping’ 180 pulse (Figure 3). The pulse is applied before or after a delay $\Delta$ and $\Delta'$, 1/2J$_{HN}$ and 1/2J$_{HC}$ in which anti-phase magnetization is created for H$_N$ and H$_C$ proton spins, respectively. If the pulse is applied at position (1) in Figure 3, the signals of $^{15}$N/$^{13}$C bound proton spins are effectively inverted and have an opposite phase compared to other spins. If it is applied at position (2) (Figure 3), evolution of heteronuclear
coupling is completely refocused during the semi-constant time evolution period, and $^{15}$N/$^{13}$C- and $^{14}$N/$^{12}$C-bound protons have the same phase. Transients are stored separately for the two positions of the jumping $180^\circ$ pulses in order to be able to separate the inter- and intramolecular NOEs. The sum of scans (1) and (2) selects $^{14}$N/$^{12}$C bound protons in $\omega_1$, its difference $^{15}$N/$^{13}$C bound protons. In $\omega_2$, editing is achieved by applying two consecutive 90 pulses with phase $\psi$ alternating between $x$ and $-x$ (21) and the two transients are stored separately. When both pulses have the same phase the signal of $^{15}$N bound protons is inverted. Again, the sum of scans with $\psi = x$ and $\psi = -x$ selects $^{14}$N bound protons in $\omega_2$, its difference $^{15}$N bound protons. A total of four different FIDs are recorded as described in Table 1, which results in (i) inverting signals of $^{15}$N/$^{13}$C bound protons in $\omega_1$ and $^{15}$N bound protons in $\omega_2$; (ii) inverting only $^{15}$N bound proton signals in $\omega_1$; (iii) inverting only $^{15}$N/$^{13}$C bound protons in $\omega_1$; and (iv) without any signal inversion. Linear combinations of these four FIDs yield four sub-spectra I–IV as shown in Table 2. The NOESY mixing time was set to 300 ms. The experiment was performed with the $^1$H carrier positioned at the $\text{H}_2\text{O}$ resonance and the $^{15}$N and $^{13}$C carriers both at 150 p.p.m. Pulse lengths for the $^{15}$N and $^{13}$C 180$^\circ$ pulses were 84 and 30 $\mu$s, respectively. For the $^{15}$N and $^{13}$C spins of interest in the G–C and A–U base pairs these pulses achieve better than 98% inversion. Therefore, incomplete inversion which could degrade the editing/filtering performance should not be an issue. One thousand and twenty-four complex points were collected in $t_1$ with a sweep width of 13.9 kHz, and 256 complex points were recorded in $t_1$ with a sweep width of 13.9 kHz. Thirty-two scans of four separate FIDs were collected for each complex point in $t_1$. The data was zero filled to 512 $\times$ 2K complex data points and apodized using Lorentz-to-Gauss and cosine functions in $t_2$ and $t_1$, respectively before Fourier transformation.

Spectra were processed with NMRPipe (22) and analyzed using NMRVIEW (23). Imino protons were assigned with a combination of HNN-COSY (15) and a sequential walk in 2D NOESY spectra (300 ms mixing time) (24) (Duszczyk et al., in preparation). All pulse sequences use the WATERGATE sequence (25) and water flipback (26) for water suppression.

### Quantitative analysis of HNN-COSY

A description of the magnetization transfer in the HNN-COSY pulse sequence is given in the Supplementary Material. The HNN-COSY spectrum shows cross and diagonal peaks at the chemical shifts of the $^{15}$N1 and $^{13}$N3 nuclei for each G–C, A–U, and some non-canonical base pairs. If both hydrogen bond acceptor and donor are isotope labeled, intensities of cross and diagonal peaks are proportional to $\sin^2(\pi h^2 J_{NN}T)$ and $\cos^2(\pi h^2 J_{NN}T)$ respectively.

Values of $h^2J_{NN}$ coupling constants were calculated from $I_{\text{cross}}/I_{\text{diagonal}}$ extracted from the spectra recorded on the fully labeled RNA samples with:

$$I_{\text{cross}}/I_{\text{diagonal}} = \tan^2(\pi h^2 J_{NN}T)$$

$I_{\text{cross}}/I_{\text{diagonal}}$ was determined as the amplitude ratio of the time domain oscillations using the time domain fitting routine nlinLS contained in the NMRPipe package (15,22). Errors in the intensity ratio and couplings were calculated based on the effect of random noise for the peak height.

### Table 1. Transfer amplitude factors for the signals in the four different FIDs of the double filtered NOESY experiment shown in Figure 3

| FID | Jumping pulse position | Phase $\psi$ | Amplitude factor | Overall amplitude factor |
|-----|------------------------|-------------|-----------------|-------------------------|
|     |                        | (1)         | (2)             |                         |
| $\omega_1$ | $^{13}$C, $^{15}$N       | $x$         | $-$             | $+$                     |
| $^{13}$C, $^{14}$N |                        | $x$         | $+$             | $+$                     |
| $^{15}$N |                        | $-x$        | $-$             | $+$                     |
| $^{12}$C, $^{14}$N |                        | $-x$        | $+$             | $+$                     |

### Table 2. NOEs observed in the four subspectra that result from linear combinations of the four different FIDs recorded

| Subspectrum | Linear combination of FIDs | NOEs involving protons bound to $\omega_1$ | NOEs involving protons bound to $\omega_2$ | NOEs observed |
|-------------|----------------------------|------------------------------------------|------------------------------------------|---------------|
| I           | $a + b + c + d$            | $^{12}$C, $^{14}$N                       | $^{12}$/$^{13}$C, $^{14}$N               | G–U, A–U      |
| II          | $a - b - c + d$            | $^{13}$C, $^{15}$N                       | $^{15}$N                                  |               |
| III         | $a - b + c - d$            | $^{13}$C, $^{15}$N                       | $^{12}$/$^{13}$C, $^{14}$N               |               |
| IV          | $a + b - c - d$            | $^{12}$C, $^{14}$N                       | $^{15}$N                                  |               |
estimated by nlinLS. No correction for an underestimation of 10–20% due to the finite excitation bandwidth of the $^{13}$N radio frequency pulses (15) was made.

Discrimination between intra- and intermolecular hydrogen bonds can be accomplished by comparing the relative intensities of cross and diagonal peaks in HNN-COSY spectra of a fully labeled and 50% labeled, 50% unlabeled RNA sample. If the RNA forms monomeric hairpins, the relative intensities of the diagonal and cross-peaks ($I_{\text{diagonal}}/I_{\text{cross}}$) will be equal in both samples, apart from an overall 50% loss of intensity as only the 50% labeled molecules contribute to the signal. In the case of duplex formation $I_{\text{diagonal}}/I_{\text{cross}}$ will increase as 50% of the RNA population of which one strand is labeled and the other unlabeled will only contribute to the diagonal peak intensity but not to the cross peak intensity.

The $^1$H and $^{15}$N chemical shifts of corresponding positions in hairpin and duplex conformations are usually degenerate. Therefore, if the interconversion between monomer and dimer species is either fast or slow on the time scale of the $h^2J_{NN}$ couplings and if there is no significant population of a non-hydrogen bonded form, molar fractions in a mixed population of monomers and dimers can be derived from the intensity ratio of cross and diagonal peaks in a 50% labeled, 50% unlabeled sample. The requirements are: (i) corresponding $h^2J_{NN}$ coupling constants are similar in the hairpin and duplex conformations; (ii) the size of the $h^2J_{NN}$ couplings is known (for example from measurements on a 100% labeled sample) and (iii) transverse $^{15}$N T2 relaxation times of the monomer and dimer species are known. Then, the molar fractions of the hairpin ($\chi_{\text{monomer}}$) and that of the duplex conformations ($\chi_{\text{duplex}} = 1 - \chi_{\text{monomer}}$) in a 50% labeled, 50% unlabeled sample contribute to the intensities of diagonal and cross peaks as follows:

$$I_{\text{diagonal}}/I_{\text{cross}} = \frac{\chi_{\text{monomer}} \cos^2(\pi h^2J_{NN} T) e^{-2T/T2(\text{monomer})} + (1 - \chi_{\text{monomer}}) (1 + 0.5 \sin^2(\pi h^2J_{NN} T) e^{-2T/T2(\text{duplex})})}{\chi_{\text{monomer}} \sin^2(\pi h^2J_{NN} T) e^{-2T/T2(\text{monomer})} + (1 - \chi_{\text{monomer}}) 0.5 \sin^2(\pi h^2J_{NN} T) e^{-2T/T2(\text{duplex})}}$$

Note, that the transverse $^{15}$N relaxation has only a very small effect on $I_{\text{diagonal}}/I_{\text{cross}}$. Moreover, in the case of a fully dimeric or fully monomeric conformation $I_{\text{diagonal}}/I_{\text{cross}}$ does not depend on the transverse relaxation time at all.

**RESULTS AND DISCUSSION**

**Xist RNA A-repeat 26-mer assignment**

Assignment of RNA usually starts with confirmation of the secondary structure by identifying Watson–Crick base pairs in A-form helical regions. In long mixing time NOESY spectra in H2O these cause a characteristic pattern of inter-base NOEs within base pairs and between consecutive stacked base pairs. With the latter a so-called ‘sequential walk’ through the stem is possible. Sequence specific assignment of the imino protons is facilitated with the HNN-COSY experiment where the imino proton shifts are correlated with base-specific chemical shifts of the nitrogens in the base pairs.

After recording these two experiments on our Xist 26-mer sample, imino assignment could be almost completed. However, we noticed an unusual downfield shifted signal, which did not show any imino–imino NOEs in the NOESY. In the HNN-COSY experiment this imino was correlated to a uridine N3 nitrogen and to a nitrogen at ~222 p.p.m., indicative of either adenine N1, N3 or purine N7 nitrogens (27). A correlation to adenine N1 would correspond to a canonical A–U base pair, but such a base pair was not expected in the predicted 26-mer double hairpin. We considered two possibilities: either that this imino was involved in a non-canonical base contact in the second predicted loop, or that it was involved in an intermolecular base pair in a duplex species, although native gel analysis of the 26-mer A-repeat RNA suggested a homogenous monomeric population (Supplementary Figure 2).

**Quantification of HNN-COSY**

To rule out that our Xist samples are dimerizing under NMR conditions, we used the approach based on the quantitative HNN-COSY experiment (Figure 4). The HNN-COSY spectrum recorded on a 100% $^{15}$N-labeled sample of the 26-mer is shown in Figure 4C. Figure 4D shows slices through the maxima of cross- and diagonal peaks for two base pairs (G2–C13 in the first predicted stem and the base pair involving the U20 imino) from HNN-COSY spectra recorded on a fully labeled (blue) and 50% labeled, 50% unlabeled (green) 26-mer RNA. Corresponding spectra were recorded on a shorter 14-mer Xist RNA sample containing only the first predicted hairpin (Figure 1D) to compare measurements in the first hairpin within the 26-mer sample to the ones in a small and stable hairpin that provided superior spectral quality (data not shown). Only non-overlapped signals were used to measure intensities. All base pairs in the 14-mer could be analyzed except G1–C14, which is not visible presumably due to end-fraying. Measured intensities for diagonal and cross peaks for the base pairs in the 14-mer and the 26-mer are given in Table 3. It is clear that for the 14-mer Xist RNA $I_{\text{diagonal}}/I_{\text{cross}}$ ratios are comparable within error for the 100% labeled sample and the 50% labeled, 50% unlabeled sample. This confirms a monomeric state of the 14-mer hairpin. The non-overlapped signals in the 26-mer show different results for base pairs from the first and second predicted hairpin. For G2–C13, G12–C3 and G4–C11 no significant difference between $I_{\text{diagonal}}/I_{\text{cross}}$ is seen between the two samples. In contrast, for the base pair involving the U20 imino proton, $I_{\text{diagonal}}/I_{\text{cross}}$ increases four-fold in the 50% labeled, 50% unlabeled sample. This unambiguously shows that U20 is involved in an intermolecular base pair. The most probable partner in this intermolecular base
Figure 4. Secondary structure assignment of the Xist RNA A-repeat (A) G–C, A–U and G–U base pairs with magnetization transfer in the HNN-COSY. (B) The predicted structure of the 26-mer A-repeat construct with sequential imino-imino NOE and HNN transfers as dotted and solid lines respectively, in magenta for the first predicted hairpin, in green for the second one (C, Top/middle) 600 MHz 1H, 15N HNN-COSY spectrum recorded on a 100% 13C15N labeled sample in 95:5 H2O:D2O at 278 K. (Top) Intra-base imino-N1 correlations leading to diagonal peaks for the guanosines in the stem are shown in black and are connected by solid lines to the inter-base imino-N3 cross peak correlations of the cytosines they are base paired to. (Middle) The correlation for the A–U base pair is shown. (Bottom) A 2D 1H NOESY spectrum, recorded on an unlabeled sample in H2O under the same conditions, connects the iminos by a sequential walk through the stem as indicated by a dotted line. Connectivities for the first predicted stem are shown in magenta, for the second predicted stem in green as shown in (B). Signals arising from the U20 imino proton are boxed in magenta. (D) Diagonal and cross peak intensities in a fully labeled (blue) and 50% labeled, 50% unlabeled (green) Xist A-repeat 26-mer sample are compared for the intramolecular base pair G2-C13 (boxed in black) and the intermolecular base pair U20-A21 (boxed in magenta). The spectra have been normalized with respect to the diagonal peak height, as experimental times of the spectra recorded on the fully labeled and 50% labeled samples were different.
Table 3. Experimental ratios between diagonal and cross peak intensities extracted from HNN-COSY spectra of fully labeled and 50% labeled, 50% unlabeled—Panel A: Xist RNA A-repeat 14-mer and Panel B: Xist RNA A-repeat 26-mer with calculated J-couplings for each base pair

| Base pair | 1_{\text{diagonal}}/I_{\text{cross}} | 1_{\text{diagonal}}/I_{\text{cross}}, 50% labeled 1.2 mM | 5_{\text{JNN}} coupling | 1_{\text{diagonal}}/I_{\text{cross}}, 50% labeled 0.8 mM each |
|-----------|-----------------|-----------------|----------------|-----------------|
| Panel A: 14-mer |
| C5-G10 | 1.6±0.1 | 5.3±0.1 | ND | 1.7±0.1 |
| G4-C11 | 1.4±0.1 | 5.6±0.1 | ND | 1.3±0.1 |
| C3-G12 | 1.4±0.1 | 5.6±0.1 | ND | 1.4±0.1 |
| G2-C13 | 1.9±0.1 | 5.0±0.1 | ND | 1.8±0.1 |
| Panel B: 26-mer |
| C5-G10 | 0.1/8.5 | 0.1/1.8 | ND | ovl. |
| G4-C11 | 0.1/8.5 | 0.1/1.8 | ND | ovl. |
| C3-G12 | 0.1/1.4 | 0.1/1.3 | ND | ovl. |
| G2-C13 | 0.1/1.3 | 0.1/1.3 | ND | ovl. |
| C16-G25 | ovl. | ovl. | ovl. | ovl. |
| G18-C23 | ovl. | ovl. | ovl. | ovl. |
| U20-A21 | 2.0±0.1 | 4.9±0.1 | 5.6±0.1 | 8.5±1/ noise |

For the 26-mer calculated 1_{\text{diagonal}}/I_{\text{cross}} values are given for spectra recorded on samples with a total RNA concentration of 1.6 mM and 0.8 mM. Intensities that could not be analyzed because of overlap are labeled with ‘ovl’ for ‘overlapped’ and their calculated 5_{\text{JNN}} couplings are labeled with ‘ND’ for ‘not determined’. 1_{\text{diagonal}}/I_{\text{cross}} for U20-A21 could not be determined for the sample with 0.4 mM labeled and unlabeled 26-mer each as the cross peak intensity was in the noise level. Errors in the intensity ratios and couplings were calculated based on the effect of random noise for the peak height estimated by the time-domain fitting routine nlinLS contained in the NMRPipe package.

pair is A21 in a second 26-mer molecule. Thus, A20-U21 is indeed a canonical Watson–Crick base pair, however, from an unexpected duplex RNA species. Although 1_{\text{diagonal}}/I_{\text{cross}} for the G18–C23, G10–C5 and G25–C16 base pairs could not be analyzed quantitatively due to spectral overlap, the signals involving G18–C23 and G10–C5 and G25–C16 in the second predicted hairpin show a significant decrease in cross peak intensity in the 50% labeled sample (data not shown), consistent with intermolecular base pairs in the second predicted hairpin.

Calculation of 5_{\text{JNN}} coupling constants from a 100% isotope labeled RNA yielded values between 4.9 and 5.6 Hz (Table 3), which are at the lower end of the range of reported values of 6–7 Hz for Watson–Crick base pairs (15,28,29). Repetition of the experiment on a different spectrometer and with different transfer times for the N–N transfer resulted in the same range of couplings. Smaller couplings are usually observed in nonlinear H-bond geometries or due to fraying at the interfaces with non-regular secondary structure elements (30). Presumably, apart from a potential underestimation of the couplings due to imperfections of the 15N radio frequency pulses, the size of the couplings in our system is reduced by fraying and/or other conformational dynamics in the small hairpins investigated.

Figure 5 shows the correlation of 1_{\text{diagonal}}/I_{\text{cross}} with the transfer time for full monomeric and full duplex conformations, as well as for the case where only 90% of the sample is either monomeric or dimeric. Transverse 15N T2 relaxation times were estimated to be 50 ms for the dimer and 70 ms for the monomer based on reported values for RNAs of similar size (31) and considering that the experiments were recorded at 5°C. For a transfer time T of 40 ms and a 5_{\text{JNN}} coupling of 5 Hz, 1_{\text{diagonal}}/I_{\text{cross}} is calculated to be 7.6 for a pure duplex and 1.9 for a pure hairpin conformation. Thus, a four-fold increase in the ratio is expected for a full duplex species upon mixing with unlabeled RNA. If a mixed population exists, the 1_{\text{diagonal}}/I_{\text{cross}} ratios shift closer together. Variations in the 15N T2 relaxation times also lead to an increase/decrease of 1_{\text{diagonal}}/I_{\text{cross}} for monomer/dimer species, but the effects are negligible (data not shown). By comparing the experimentally measured 1_{\text{diagonal}}/I_{\text{cross}} values with the simulated curves it is clear that all base pairs in the first hairpin of the 26-mer A-repeat RNA are consistent with a fully monomeric conformation, while the A–U base pair in the second predicted hairpin exists in a fully dimeric form.

With an A–U base pair originating from a dimeric conformation, two possibilities exist for the A-repeat topology. Dimerization could involve a kissing hairpin with two equivalent intermolecular A–U base pairs (Figure 6B), or correspond to a duplex conformation with full base pairing with the second part of another A-repeat (Figure 6C). To distinguish between these two conformations information on the state of the G–U base pair in the second predicted stem would be useful. In a kissing hairpin complex this base pair would be intramolecular while in a duplex this base pair would be intermolecular. Unfortunately, the analysis of J-couplings across hydrogen bonds is not applicable to G–U base pairs due to a lack of sizable J-couplings (Figure 4A).

Analysis of the double-isotope-filtered NOESY spectra

As a characteristic and strong NOE pattern is visible between the G and U imino protons in a G-U base pair we analyzed the 50% labeled, 50% unlabeled sample following the approach first proposed by Aboul-ela et al. (14) based on NOEs in base pairs with optimizations and extensions as described in the Materials and Methods section.

Linear combinations of the four FIDs a–d (Table 1) recorded in the double-filtered NOESY experiment yield
thick lines) and $^{15}$N T2 relaxation times of 70 and 50 ms for the monomer $I_{20}$–$A_{21}$ base pair is $h_{NN} = 4.9$ Hz, while the couplings for the G–C base pairs are 5.0–5.5 Hz.

The question remains if the dimerization seen in vitro under NMR conditions is significant in vivo, or if it is merely an artifact of the high concentrations used in the experimental conditions. In this respect, we note that the imino signal of $U_{20}$, which is involved in the intermolecular basepair, is visible in 1D NMR spectra even at concentrations as low as 50 $\mu$M. In addition, analytical ultracentrifugation data of a 0.1 mM sample show that the A-repeat 26-mer exists as a dimer in solution (Duszczek et al., in preparation). It should also be considered that in vivo there is a high local concentration of the A-repeats as they are connected by relatively short linkers and are localized to the X chromosome. Thus, it is possible that the second region of the A-repeat may function as a multimerization platform for several A-repeats, either within a single Xist RNA molecule, or between different ones. Further experiments to address these issues are underway.

The architecture of the Xist RNA-repeat

The combined data from the HNN-COSY and double-isotope-filtered NOESY experiments are in agreement with an architecture of the A-repeats as shown in Figure 6C with a fully dimerized second stem. The fact that the imino–imino NOEs for the G17–$U_{24}$ base pair in the spectra shown in Figure 6D and E have comparable intensities indicates that the 26-mer RNA exists in a fully dimeric form in solution. If an equilibrium would exist between the kissing hairpin and duplex forms shown in Figure 6B and C, the G–U imino–imino NOEs in Figure 6E should have lower intensities than those in Figure 6D.

CONCLUSION

We developed and applied a novel approach to unambiguously define the stoichiometry of the NMR sample for the structural analysis of a Xist RNA A-repeat. Using a combination of quantitative HNN-COSY and a novel double isotope-filtered NOESY experiment we could define the conformation of the 26-mer A-repeat RNA. We show that in a single 26-mer A-repeat only the first predicted hairpin is formed, while the second predicted stem-loop forms a RNA duplex and mediates dimerization of the 26-mer A-repeat. Our strategy will be generally applicable to identify and quantify populations of hairpin and duplex conformations in RNAs and to define RNA folding topology from inter- and intra-molecular base-pairing patterns.

SUPPLEMENTARY DATA

Supplementary data are available at NAR Online.
NOEs involving the intermolecular G17–U24 and U20–A21 base pairs are shown. The corresponding NOE transfers are indicated schematically for G–U and A–U base pairs in D and E, respectively. Zoomed-in views of the edited subspectra and (E) edited/filtered subspectra derived from the double filtered NOESY. In (E) NOEs between pairs of protons where one is attached to an isotope labeled heteroatom and the other one to an unlabeled one are observed. These NOEs can only originate from a duplex RNA species. The corresponding NOE transfers are indicated schematically for G–U and A–U base pairs in D and E, respectively. Zoomed-in views of the NOEs involving the intermolecular G17–U24 and U20–A21 base pairs are shown.

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