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Distance measurements between manganese(II) and nitroxide spin-labels by DEER determine a binding site of Mn²⁺ in the HP92 loop of ribosomal RNA[†]

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Mn²⁺ localization in hairpin 92 of the 23S ribosomal RNA (HP92) was obtained using W-band (95 GHz) DEER (double electron–electron resonance) distance measurements between the Mn²⁺ ion and nitroxide spin labels on the RNA. It was found to be preferably situated in the minor groove of the double strand region close to the HP92 loop.

The DEER (double electron-electron resonance) technique¹⁻³ has become very popular in recent years for obtaining nanometer scale distance restrains in structural studies of biomolecules in frozen solutions.⁴⁻⁷ The most common application of DEER is to measure distances between two nitroxide spin labels attached at specific points of interest in a biomolecule. It has also been successfully applied to determine the distances between other types of paramagnetic centers in biomolecules,8 such as Cu2+- $Cu^{2+9,10}$ $Gd^{3+}-Gd^{3+}$ 1^{1-13} $Mn^{2+}-Mn^{2+}$ 1^4 pairs of iron-sulfur clusters^{15,16} and trityl-trityl radicals.^{17,18} Biomolecular heterospin label distance measurements, including nitroxide-Cu²⁺,¹⁹⁻²¹ nitroxide-iron-sulfur cluster²² and nitroxide-Gd³⁺,^{23,24} have also been reported. A recent important alternative application of DEER, in combination with site directed spin labelling, has been locating metal ion binding sites by means of paramagnetic metal ionnitroxide distance measurements in a biomolecule, as has been demonstrated on Cu²⁺ binding sites.^{19,21}

Metal ions play a crucial role in RNA structure and function.²⁵ They stabilize its tertiary structure^{25–31} and are essential for catalysis in ribozymes.³¹ Several types of RNA-metal ion interactions, that can take place simultaneously, are recognized. One type involves poorly localized "diffuse ions", the charge of which

balance the negative charge of the phosphodiester backbone of the RNA.32 The second type involves ions with specific interactions at particular sites on the RNA molecule.³² These bind either through inner-sphere interactions involving direct coordination to the electronegative RNA functional groups, or via outersphere interactions mediated by water ligands.³¹ K⁺ and Mg²⁺ ions are usually considered the natural metal ion cofactors for nucleic acids in vivo. Mg2+, however, presents a challenge for optical and magnetic spectroscopic characterization. One way of overcoming this limitation is to substitute it with the paramagnetic Mn^{2+} ion, which has a similar radius and charge,33 and apply EPR spectroscopic methods.^{34,35} High resolution EPR methods that are typically used to probe Mn²⁺ binding sites in RNA yield local information, such as the hyperfine coupling with ¹⁴N from the nucleobases and phosphate ³¹P. While providing important structural information on the nature of the metal ion ligation, these methods, do not give direct information on the location of the binding site within the RNA.

In this communication we introduce high field (W-band, 95 GHz) Mn^{2+} -nitroxide DEER distance measurements for locating Mn^{2+} binding sites in RNA. We demonstrate this approach by locating a Mn^{2+} binding site in an RNA derived from hairpin 92 of the 23S ribosomal RNA (HP92). HP92 is a specific target to DbpA and YxiN RNA helicases (from *Escherichia coli* and *Bacillus subtilis* respectively). The feasibility of Mn^{2+} -nitroxide DEER measurements has been recently reported on a rigid model compound.³⁶

Fig. 1a shows the RNA construct used in this study and the spin labelling positions. Three HP92 RNA constructs were prepared: two singly labelled, (RNA(3) and RNA(31)), and one doubly labelled RNA (RNA(3,31)). Several methods exist for sitedirected spin labelling (SDSL) of nucleic acids,³⁷ but we chose post-synthetic labelling of 2'-amino groups with 4-isocyanto-TEMPO (Fig. 1b).³⁸ (see ESI[†] for details).

Measurements were carried out at W-band because of the high sensitivity that it features for Mn^{2+} , which is a half integer, high spin ion (S = 5/2). This arises from the reduced inhomogeneous broadening of the $|-1/2\rangle \rightarrow |1/2\rangle$ transition by the

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[†] Electronic supplementary information (ESI) available: Sample preparation, X-band EPR spectra, W-band nitroxide–nitroxide DEER details of DEER experimental conditions, Mn²⁺/nitroxide EDEPR spectra and DEER experimental details, DEER raw data, DEER data analysis using Tikhonov regularization. See DOI: 10.1039/c5cp01624j



Fig. 1 (a) The sequence and the secondary structure of the RNA construct studied. The spin labelling positions are indicated with circles. The putative location of the Mn^{2+} as determined by the DEER measurements is indicated by an arrow. (b) The RNA spin-labelling scheme.



Fig. 2 (a) DEER traces measured on RNA(3,31) after background removal along with the fit (red trace) obtained with the distance distribution shown in (b). The two upper DEER traces in (a) were shifted for clarity. Raw DEER data is presented in Fig. S3a (ESI†); data analysis was performed with DeerAnalysis.³ Sample composition was 166 μ M RNA/33 mM HEPES/66 mM NaCl/33% d₃-glycerol in D₂O.

zero field splitting (ZFS) at high fields.³⁹ First, DEER measurements were carried out on the doubly labelled RNA(3,31), without the addition Mn^{2+} or Mg^{2+} , to obtain structural restrains on the RNA itself and results are shown in Fig. 2. Measurements were carried out at three observer magnetic field positions to check for orientation selection (see Fig. S1, and ESI† for experimental details). All three measurements gave a similarly broad distance distribution, spanning a width of \sim 3 nm with a maximum in the range of 4.5–4.9 nm. In light of the width of the distribution and the lowest signal to noise ratio (SNR) of the " g_{zz} " trace, which gave the 4.5 nm maxima, we consider this range to be within experimental error and ignore orientation selection. The broad distance distribution is attributed to the nitroxide label local flexibility and high flexibility of the RNA. This is consistent with the room temperature X-band CW-EPR spectra (Fig. S1, ESI[†]) of RNA(3) and RNA(31) that are typical of highly mobile nitroxides.

 Mn^{2+} -nitroxide (Mn-NO) DEER measurements on Mn^{2+} /RNA(3) and Mn^{2+} /RNA(31) complexes (1:1 Mn^{2+} :RNA molar ratios) are shown in Fig. 3. Here the different spectroscopic properties of Mn^{2+} and the nitroxide spin label have to be carefully considered when the experiment is set up. EPR spectra of these samples and additional experimental details are presented in the ESI.†

In the Mn–NO DEER measurements, the observer pulses were set to the 3rd hyperfine line of the Mn²⁺ sextet, the pump pulse was set to either the maximum ($\Delta \nu = -120$ MHz) or more towards the g_{zz} position ($\Delta \nu = -65$ MHz) of the nitroxide



Fig. 3 DEER traces of Mn²⁺/RNA(31) (a) and Mn²⁺/RNA(3) (b) obtained for two $\Delta\nu$ values after background removal along with the fit (red trace) derived with the distance distribution shown in (c) and (d) respectively. The raw DEER traces are given in Fig. S3b, c (ESI†). All DEER measurements were carried out at 10 K and repetition time of 1 ms. Data analysis was done with DeerAnalysis.³ Sample compositions were 135 μ M RNA/135 μ M MnCl₂, 31 mM HEPES/62 mM NaCl/41% d₃-glycerol in D₂O.

spectrum (Fig. S2c, ESI[†]). By observing the Mn²⁺ signal we take advantage of its fast spin lattice relaxation rate that allows for fast signal averaging. This set up also minimizes the contribution of the nitroxide to the observed signal because of its saturation. In addition, the narrower nitroxide spectrum allows a higher modulation depth, namely more spins are affected by the pump pulse.²³ Fig. 3a, b present the DEER data of Mn²⁺/RNA(31) and the distance distribution obtained after fitting to a two Gaussians model. These measurements gave distance distribution with a narrow Gaussian at 2.3–2.4 nm superimposed on a much broader one with a width around 3.0 nm and a maximum at 2.5 nm. The data could not be satisfactory fitted with one Gaussian. Data analysis using Tikhonov regularization gave results consistent with this interpretation (see Fig. S4 and discussion in the ESI[‡] for details).

DEER data of $Mn^{2+}/RNA(3)$ are shown in Fig. 3c. Here the DEER traces could be satisfactorily fitted with a single Gaussian, with a broad distribution and a maximum at 3.6 nm (Fig. 3d). In neither constructs did we observe orientation selection similar to reported for W-band Gd³⁺–nitroxide DEER measurements.⁴⁰ In combining the NO–NO distances obtained from RNA(3,31) with those obtained from the Mn–NO results we assumed that addition of the equimolar amount of Mn^{2+} did not alter the structure of the RNA significantly.

W-band ENDOR (electron-nuclear double resonance) measurements of Mn^{2+} with non-labeled RNA revealed no ³¹P hyperfine couplings. This is unlike the single strand HP92, without the 11 base complementary strand, which showed a clear ³¹P coupling of ~9 MHz, typical of Mn^{2+} bound to phosphate of the backbone.⁴¹ In addition, X-band electron spin echo envelope modulation (ESEEM) measurements did not reveal any coupling to ¹⁴N, excluding the presence of a direct binding to an ¹⁴N of a nucleobase, a binding mode similar to one observed in the Hammerhead ribozyme with direct coordination to N7 nitrogen atoms of adenine and guanine.^{42–44} These findings suggest that the Mn^{2+} ion is bound to HP92 through an outer-sphere coordination. This mode of binding is similar to that reported for type 0 coordination of Mg^{2+} in the crystal structure of the large ribosomal subunit, which features hydrogen bonds involving six water molecules and zero direct contacts with RNA hydrophilic atoms.^{45,46}

The DEER results show that Mn²⁺ does bind to the HP92 constructs studied, although no 14N or 31P ENDOR signals were observed. The observed broad distance distribution could be due to either non-specific Mn2+ binding-sites on the RNA construct or flexibility of the RNA construct and/or the spin label. Both the X-band CW EPR spectrum of RNA(3) (Fig. S1, ESI[†]) and the broad nitroxide-to-nitroxide distance distribution obtained for RNA(3,31) (Fig. 2) indicate substantial flexibility of the label. On the other hand, the fact that we observe a superposition of a narrow and a broad distance distribution for $Mn^{2+}/RNA(31)$ indicates that the presence of non-specific sites cannot be excluded. The distance of 2.3-2.4 nm observed for RNA(31) suggests the presence of a specific Mn^{2+} binding site somewhere in the double stranded region of the 5' extension of the RNA construct. This is supported by the observed change in the coordination sphere of Mn²⁺ in HP92 upon binding of the 11-base complementary strand.⁴¹ The observation of a narrow distance distribution, that was distinct from the broad background, for RNA(31) construct, but not for RNA(3) construct, suggests that the broadening in the latter has significant contributions from the flexibility in the linker region between HP92 and the double stranded region of the 5' extension of the RNA construct.

The modulation depth observed, with the pump pulse set to the maximum of the nitroxide spectrum, was 1.2% for $Mn^{2+}/RNA(31)$ and ~3.0% for $Mn^{2+}/RNA(3)$. This is significantly lower than the ~5.5% observed for the nitroxide–nitroxide distance measurements, indicating that the binding affinity of the Mn^{2+} is not high and that there are free "diffuse" Mn^{2+} ions in solution. Measurements on a nitroxide–nitroxide model compound, under the same pump pulse parameters gave ~10% modulation depth, suggesting that the labeling efficiency was not 100%.^{47,48} Thus, the difference in modulation depth between the two constructs is attributed to differences in labeling efficiencies.

To substantiate the localization of the Mn^{2+} in the region of the 5' extension of the RNA construct we used a simple-minded model, taking the X-ray crystallography-derived RNA binding domain of YxiN bound to an extended fragment (nucleotides 2508–2580) of the 23S ribosome from *E. coli* (PDB: 3MOJ) as a template for the structure of the HP92^{49,50} and added to it the spin labels using MtsslWizard⁵¹ (details are given in the ESI†). Calculated DEER distance distributions were adapted from MtsslWizard's Distance mode taking into account all possible rotamers of the spin label, while keeping the RNA rigid.⁵¹ Fig. 4a highlights the spin label rotamers, yielding the shortest and longest distances to the Mn²⁺ position. In the case of the



Fig. 4 (a) The HP92 (grey) + 11 base complementary strand model (blue) and the Mn^{2+} (pink) location. Some of the nitroxide spin label rotamers are shown in green. Left are three rotamers in the 31 position, while on the right there are two rotamers in the 3 position. The lines highlight some nitroxide–nitroxide and Mn^{2+} –nitroxide distances. (b–d) The calculated distance distributions for the various constructs using this model (black) compared with the experimental ones (red).

spin label on 31, two distinct rotamer clouds were possible due to the curvature of the backbone, one internalized to the helix with a narrow distance distribution and a second broader cloud on the exterior of the helix curvature. Accordingly, Fig. 4a shows three rotamers at this position, two corresponding to the first cloud's shortest and longest distances and the third that corresponds to the second cloud's average. The distance for RNA(3,31), shown in Fig. 4b, yields a distance distribution with a maximum at 4.0 nm, which is shorter than the experimental result, ~4.9 nm. This suggests that in solution the structure of HP92 is on the average more extended. Moreover, the calculated distance distribution is significantly narrower than the experimentally derived distance distribution, which implies a large conformational flexibility of the HP92 construct in solution, not bound to YxiN.

While placing the Mn²⁺, it was taken as a hexaaqua complex, approximated as a sphere of diameter 1.0 nm (metal diameter + $2 \times$ bond length + $2 \times$ water's Van der Waals radius), with assumed position minimizing distortion of nucleotide bases and minimizing energy. It was placed in a position that would give the best agreement with the DEER data. According to this model, the Mn²⁺ is placed in a minor grove in the RNA twist shown in Fig. 4a (pink sphere). This position is also indicated as an arrow in Fig. 1a, to highlight the specific bases in the vicinity, one of which is guanine. Interestingly, outer-sphere hydrated Mg²⁺ ions are often found in the deep groove of A-form helices, forming hydrogen bonds to acceptors atoms of the guanine base.⁴⁶ This yields a most probable Mn²⁺nitroxide distance Mn²⁺/RNA(31) around 1.9 nm (Fig. 4c), which is rather close to the observed 2.3-2.4 nm. The modeled Mn²⁺/RNA(3) distance is 3.4 nm (Fig. 4d), also close to the experimentally observed distance of 3.6 nm. A reduced RNA backbone curvature, as suggested by the experimentally derived RNA(3,31) distance distribution, would generate a longer distance between the Mn²⁺ and nitroxide spin labels and a better

agreement. While this simplistic modelling suggest a reasonable location for the Mn^{2+} ion, in the future systematic molecular dynamics simulations should be carried out on the RNA, including the solvent and the Mn^{2+} ion to obtain the energy landscape of the Mn^{2+} potential coordination site and account for the multiple RNA conformations in solution, taking into account the experimentally derived distance distributions.

Conclusions

We have demonstrated that Mn^{2+} -nitroxide W-band DEER distance measurements can be added to the tool-box used for localizing Mn^{2+} binding-sites in nucleic acids, particularly when the Mn^{2+} (or Mg^{2+}) play a role in structure stabilization or catalysis. Our results show that HP92 with its 11 base complementary strand in solution is extended relative to the crystal structure (in the presence of YxiN) and highly flexible. We identified a specific binding site for outer-sphere coordinated Mn^{2+} in the minor groove of the double stand region close to the HP92 loop. In the future, the use of rigid spin labels⁵² would yield data where any observed flexibility could be directly traced to movements of the RNA, eliminating contributions from the spin label tether and will potentially allow for more precise localization of the binding sites.

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