A trade-off for covalent and intercalation binding modes: a case study for Copper (II) ions and singly modified DNA nucleoside

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Selective binding to nucleic acids and, more generally, to biopolymers, very often requires at a minimum the presence of specific functionalities and precise spatial arrangement. DNA can fold into defined 3D structures upon binding to metal centers and/or lanthanides. Binding efficiency can be boosted by modified nucleosides incorporated into DNA sequences. In this work the high selectivity of modified nucleosides towards copper (II) ions, when used in the monomeric form, is unexpectedly and drastically reduced upon being covalently attached to the DNA sequence in single-site scenario. Surprisingly, such selectivity is partially retained upon non-covalent (i.e. intercalation) mixture formed by native DNA duplex and a nucleoside in the monomeric form. Exploiting the electron spin properties of such different and rich binding mode scenarios, 1D/2D pulsed EPR experiments have been used and tailored to differentiate among the different modes. An unusual correlation of dispersion of hyperfine couplings and strength of the binding mode(s) is described.

Metal ions play important roles in many ribozyme-mediated examples of scission and ligation reactions but their precise role in those reactions is not yet fully understood. The presence of an imidazole moiety in biological molecules has focused attention towards imidazole-modified nucleosides and the corresponding complex formed with copper (II) ions. Selected histamidine/histidine modified 2'-deoxyribadenosines (histam'dA or hi'dA) have been found to be strong chelators for Cu(II) ions. Their efficiency as chelators has been proven for the DNA cleavage reaction, revealing the intercalative ability of the complex Cu(II)/ligand. Cu(II) ions have been also extensively used as cofactors for DNAzymes (deoxyribozymes) for cleavage or ligation reactions, either on RNA or DNA strands. The highly densely modified DNAzyme sequence has been also tested and metal-independence has been demonstrated. Covalently modified nucleoside has been used as paramagnetic probes (i.e. copper ions) for the determination of interspin distances; intercalation of copper(II) ligand has been studied mainly by CW EPR for elucidating the structure of DNA-fiber. To make this scenario more challenging, ligation and cleavage reactions often make use of "metal soup" as cofactor or metal-lanthanide binary cofactor, and the role of different cofactors in the catalytic process still remains unknown. In order to localize the binding region(s) over DNA or RNA sequences, selective chemical modifications have been tested; such an approach has revealed itself to be tedious and does not solve the major issue concerning the identification of binding sites of metal and lanthanide cofactors with respect to the nucleotide structure, with a particular focus on single or multiple sites. Taking into account that such systems very often contain paramagnetic species, NMR spectroscopy is not the most suitable method for the analysis of the structural features of these complexes. Absorption and fluorescence spectral studies are more conventional methods to analyze these complexes, but they only provide a global view of the system under investigation and very often do not provide information on the nature of the nucleus/nuclei forming the coordination sphere of the metal center. Electron paramagnetic resonance (EPR), known also as electron spin resonance (ESR), offers a plethora of experiments for detection of short- and long-range interactions. Hyperfine spectroscopy is leading the structural analysis of the catalytic cores of several proteins, especially those containing iron-sulfur clusters as initiators of radical reactivity.
Among these, the more conventional pulsed EPR technique, H*yperfine Sublevel Cor*relation spectroscopy (HYSCORE) and the emerging technique ELDOR-Detected NMR (EDNMR) are methods of choice for evaluating interactions of the paramagnetic species and its surrounding nuclei. In particular, E*lectron-electron DOuble Resonance (ELDOR)-detected NMR (EDNMR) is gaining popularity due to the latest developments in wave-generator and multi-frequency approaches.20,21. This sequence was first developed in Schweiger’s labora-

tory22, and uses selective microwave pulses which simultaneously pump EPR and NMR transitions of the spin manifold, so called spin forbidden transitions, where both the electron and nuclear spins change their projection direction. As compared to conventional Electron Nuclear DOuble Resonance (ENDOR) techniques it has a much higher sensitivity and does not exhibit nucleus- or pulse-dependent spectral artifacts23,24. Historically, the wide-

spread adoption of EDNMR has been hampered by what is termed the “central blind spot”. The limited resolution is mitigated by performing the EDNMR experiment at higher magnetic fields (>9 GHz) as demonstrated by a number of recent studies24. Furthermore, the multi-frequency approach can combine the advantages of low field optimized experiments (HYSCORE) with those at higher field (EDNMR)25,26. EPR/ESR hyperfine spectroscopic methodologies have been successfully combined with the analysis of metalloenzymes that exhibit high selectivity and specificity for catalytic processes involving radical species27. The restricted alphabet of nucleic acids com-

pared to amino acids for proteins is often associated with lower selectivity and specificity, and even for ribozymes, the main role for the catalytic process is extensively recognized with the protein moiety associated with metal cofactors. However, a minimalistic approach for testing binding modes is related to the analysis of emerging and promising structures such as for DNAzymes (deoxyribozymes) where the role of the metal cofactor remains to be understood, as well as that of lanthanide cofactor28. The discovery of new structures, very often associated with a serendipitous approach29, is severely lacking with respect to an accurate recognition and description of (i) binding mode(s) and (ii) binding site(s) from the perspective of the DNA structure30, which may provide a lower differentiation than that available in the catalytic core of proteins.

In this work two main aspects have been taken into account concerning the incorporation of a copper(II) chelator into DNA sequences. First, a short overview of two different nucleosides containing an imidazole moiety is reported; second, the binding modes of the modified nucleoside have been analyzed by two different scenarios: (i) covalent attachment to the DNA sequence, and (ii) non-covalent (intercalation) mixture of nucleoside and Cu2+. The two structures differ in the connection of the imidazole ring to the nucleoside: in one case an ethyl-bridge has been used and in the second structure a urea-ethyl-bridge has been used, connecting the nucleoside moiety to the imidazole moiety. A crowded scenario of binding modes (and binding sites) involves: (a) copper(II) free in solution, (b) the monomeric form of the nucleoside combined with Cu2+, (c) native DNA duplex combined with Cu2+, (d) covalently modified DNA sequence with one singly modified nucleotide combined with Cu2+, (e) native (unmodified) DNA sequence with non-covalently attached (intercalation-type) monomeric nucleoside, combined with Cu2+. Those five different binding modes have been correlated to hyperfine spectroscopy (interactions electron-nuclei) to implement the core of such a fast and straightforward protocol, in order to confirm (or discard) the formation of a complex and to provide preliminary information on the nature of the nuclei that comprise the coordination sphere.

Results

Binding efficiency retained. The analysis of different binding modes focused on two ligands, shown in Fig. 1a,b. The overall study involved the insertion of the monomeric form of the imidazole moiety into DNA duplexes (indicated in Fig. 1c), both as covalently attached and in a non-covalent mixture, in order to estimate the different binding modes for Cu2+. In Fig. 2a,b details of ligands L1 and L2 are depicted. An ethyl bridge, introduced into the structure L3, has replaced the ethyl-urea bridge between the nucleoside and the imidazole fragment that characterizes the modified nucleoside L1. Figure 2c,d illustrate the DNA sequences containing one modified nucleotide and the native sequence, respectively. The L1 monomer was analyzed by CW and pulsed EPR spectroscopy. As for the L1 ligand, the binding mode of copper(II) was tuned by different pH values and different ligand/copper ions ratios (Supplementary Information S1 shows how the EPR/ESR spectra are affected by pH changes in the mixture 1:2, Cu2+/L). In this study we aim to go beyond the impact of the pH values on the coordination of copper ion(s); testing different Cu2+/ligand ratios, we can guarantee a full coordination at a ratio 5:1 (ligand/Cu2+), as shown in Supplementary Information S2 and S3. Furthermore, operating at this ratio, a very similar trend has been observed for the two different ligands, as reported in Fig. 2e; the subsequent analysis has been then focused on the solution at pH 7.45 and with a ratio 5:1 (ligand/Cu2+). Such reduced influence of pH within the range 4.00–10.3 increases the amount of species with complete copper ion coordination sphere, ensuring a negligible effect on species distributions4,6. Those distributions involve the equilibrium between bound and free states of the copper ions, as well as monomeric and dimeric forms of the formed adducts. The analysis of monomeric L1 and L2 with Cu2+ has been combined with extensive DFT computational studies. The hyperfine couplings obtained by those calculations have been used as starting values for the simulation of spectral features. Indeed, in Fig. 3b,c the LUMO and SOMO models, respectively, derived by DFT calculations are depicted. The LUMO model would be the best molecular representation of the distribution of the hyperfine coupling constants, while the SOMO model confirms the occupied orbital for the unpaired electron of this paramagnetic complex. Furthermore, in Fig. 3d the comparison between experimental and simulated CW EPR spectrum is reported for the complex Cu2+/L1, at pH 7.45. In Fig. 3e the isotropic hyperfine couplings for this complex are reported, both from DFT computations and EasySpin fitting procedures (in brackets)31.

Slight discrepancies exist between experimental and simulated spectra, caused by the presence of minor spec-

ies in the equilibrium of bound and free states. The comparison of the six complexes (L1 and L2 ligands at six different pH values) is provided in the Supplementary Information S4. The full comparison of the monomeric forms of the two ligands demonstrates the retained efficiency of the structure L1, where the binding properties are analogous to those of L1, even in the absence of the urea bridge. For the simulation of the monomeric form
Figure 1. (a,b) Schematic view of attached modes for the imidazole moiety to the nucleoside. (c) Simplified schematic view of duplex and paramagnetic center (Cu$^{2+}$) involved in this study. (d) Free copper(II) in solution added to the unmodified duplex, L$_1$ and L$_2$ covalently attached to the duplex and L$_1$ and L$_2$ used as intercalating agents.

Figure 2. (a) Detailed structure of the old ligand L$_1$, connecting the imidazole moiety and the nucleoside via an ‘ethyl-urea-bridge’. (b) Detailed structure of the ligand L$_2$, connecting the imidazole moiety and the nucleoside via an ‘ethyl-bridge’. (c) Covalently modified DNA sequence. (d) Native DNA sequence formed by 20 nucleotides. (e) CW EPR spectra for the monomeric form of ligands L$_1$ and L$_2$, recorded at three different pH values (4.00, 7.45 and 10.3).
(i.e. at pH 7.45, for L$_2$) six nuclei were used: $^{63}$Cu, two $^1$H, two strongly coupled $^{14}$N (hyperfine coupling around 20 MHz for both atoms) and one weakly coupled $^{14}$N, placed in the so-called remote position of the imidazole moiety. The agreement between the experimental and simulated spectrum prompted us to extend the comparison of the DFT computed hyperfine couplings with the experimental spectra of L$_1$ and L$_2$ ligands. Tables composing the ensemble (Supporting Information S4) support the slight differences detected for the monomeric form of the ligands. A very similar trend is observed for L$_1$ and L$_2$: the ethyl-bridge does not perturb the formation of the complexes. Supporting Information S5 summarizes the six structures derived by DFT computational analysis, in the same manner as that single example depicted in Fig. 3b,c. Details about DFT calculations and optimizations are provided in the Methods section.

Among the hydrogen atoms coupled to the copper paramagnetic center, strongly coupled exchangeable protons have not been detected (Supporting Information S6). This supports the scenario where several heteronuclear couplings (i.e., $^{14}$N) are strongly correlated to the copper paramagnetic center.

**Echo field sweep as a first diagnostic differentiation.** The analysis of different echo field sweep spectra recorded at low field (9 GHz) provides a first overview on the strength of the copper binding. The five spectra in Fig. 4 show a typically axial g-tensor for the copper center; slight deviation from axial to rhombic are shown in Supporting Information S4 (Tables summarizing the parameters used for EPR spectra simulations). The broadening of the perpendicular component of the tensor (g$_\perp$) is observed especially in Fig. 4b,e (red and orange lines, respectively). Compared to free copper in solution (Fig. 4a, black line), the complex formed by monomer L$_1$ and copper ions is expected to provide such broadening. In case of the orange line (Fig. 4e), it concerns the mixture formed by copper ions, native DNA duplex and monomeric L$_2$, interacting in a non-covalent manner with respect to the DNA sequence (intercalation). Such a three-component complex generates a small broadening compared to the monomeric L$_2$/Cu$^{2+}$ complex, but higher than that generated by duplexes, both native or singly-modified (Fig. 4c,d, respectively). Surprisingly, the covalently attached ligand L$_2$ does not guarantee a strong coordination, even if the modified DNA sequence is used with a molar excess (5:1).

**HYSCORE experiments.** Five solutions at pH 7.45 were then analyzed by 2D HYSCORE experiments for the detection of weak hyperfine couplings (<10 MHz). Two main regions can be identified for the hyperfine couplings, involving $^{14}$N and $^1$H nuclei. The 1:2 ratios (Cu$^{2+}$/L$_1$) also indicated the relative amounts of those nuclei with respect to three different pH values (Supporting Information S7, S8 and S9). Focusing on the solution at pH 7.45, five different solutions were studied. In Fig. 5a the HYSCORE sequence is depicted. Figure 5b shows the free
Cu$^{2+}$ in solution, without any monomeric ligand or DNA sequence. The main hyperfine coupling detected for
this mixture is related to $^1$H, with a Larmor frequency centered at 14 MHz and a hyperfine coupling of 5 MHz.
A completely different scenario is offered by the monomeric L$_2$ ligand and Cu$^{2+}$ at pH 7.45 (Fig. 5c). The main
hyperfine coupling observed is that of the remote $^{14}$N (with a coupling of 2.7 MHz); the dynamics of the complex
and the distribution of radical species in this sample cover the entire region of single and double transitions
(besides the combination frequencies), rendering this crowded region quite difficult to disentangle, but confirming
the binding proposed by CW experiments. The spectra depicted in Fig. 5d,e show $^1$H to be the predominant
nucleus within the hyperfine pattern; 5d is the native duplex added to Cu$^{2+}$ solution at pH 7.45 while the 5e is the
modified duplex containing one L$_2$ ligand (covalently attached) added to Cu$^{2+}$ solution at pH 7.45. No significant
differences were detected between these two samples. The presence of one singly modified nucleotide with a
20-mer DNA sequence does not guarantee the same binding exhibited by the monomeric L$_2$ ligand. Such a loss of
binding strength is unexpected, if compared with the monomeric form of the ligand. An attempt to recover the
coupling with $^{14}$N centers was made by preparing a non-covalent mixture (intercalation of L$_2$). The ligand L$_2$ was
indeed added (still with a ratio 5:1 with respect to the Cu$^{2+}$ ion) to the native DNA sequence. The weak coupling
is then detected in the order of 3.0 MHz.

**ELDOR-Detected NMR (EDNMR) experiments.** The observed B values for EDNMR experiments have been
selected on the echo-field sweep spectra recorded at 34 GHz (Supporting Information S10). The differences
among different samples are here less pronounced, due to the additional broadening generated at higher field/
higher frequency. For the EDNMR recorded at 34 GHz, a crucial role is played by the blind spot generated by the
low frequency couplings. It was then necessary to optimize the width of the central blind spot; making it nar-
row enough so as not to obscure signals from any low-$\gamma$ nuclei present while at the same time using an ELDOR
pulse of sufficient power to excite the formally forbidden electron and nuclear transitions. Consequently, Q-band
EDNMR is always a compromise between the signal-to-noise ratio and the width of the central blind spot. For
our Q-band studies, the most convenient setup consisted of a 9000–400–800 ns pulse scheme (pulse sequence on
the top-left of the figure). Thus, Fig. 6 shows a series of Q-band EDNMR spectra of different Cu$^{2+}$ complexes;
namely the hexa-aqua complex ([Cu(H$_2$O)$_6$]$^{2+}$) (Fig. 6a), the mixture of Cu$^{2+}$ with L$_2$ ligand (Fig. 6b). In addi-
tion to the signal corresponding to the $^1$H nuclei (49.7 MHz), two broad signals corresponding to strongly cou-
ppled $^{14}$N were observed at 17.2 and 24.7 MHz, respectively. Signals within the blind spot regions (Supplementary
Information S11) and related to the setup ambiguity have not completely assigned. The signals at 2.52 and
4.55 MHz have been identified as single-quanta (sq) EDNMR transitions of the $^{14}$N nucleus ($\nu_{14N} = 3.7 MHz$,
$I = 1$), as they are centered around the $\nu_{14N}$ Larmor frequency and split by 1.5 MHz, in agreement with HYSOCORE
experiments (Supplementary Information S11b). Even if for the single-quanta transitions there is a severe overlap
with the blind spot region, the EDNMR spectra contain a significant set of signals also for frequencies higher than 10 MHz, due to the $^{14}$N nuclei coupled with different hyperfine coupling constants. This allows us to focus on the spectral regions higher than 10 MHz and to observe informative differences. As for the HYSCORE experiments, Fig. 6c,d do not exhibit remarkable differences; the presence of one covalently attached ligand (L$_2$) within the DNA sequence does not show the same coordination sphere of the ligand in the monomeric form. By analyzing the three component mixture (orange line, Fig. 6e), a clear splitting of signal is observed, allowing for differentiation between $^{14}$N deriving from the native DNA duplex and $^{14}$N deriving from the monomeric ligand L$_2$. The echo-detected field-swept spectra (EDFS) are shown as insets on the top-right of Fig. 6. The HTA pulse was applied to the $B$ value indicated on the spectrum (blue arrow). In order to verify the orientation effect of the g-tensor with respect to the EDNMR experiments, a second $B$ value was chosen, as shown in Fig. 7. For the g-component centered at 1080 mT the splitting mentioned above is even more pronounced. The signals corresponding to $^{14}$N nuclei of monomeric ligands (16.2 and 23.7 MHz) are joined by the signal of the $^{14}$N (duplex, at 30.6 MHz) and by those of the monomeric ligand in the non-covalent mixture (19.2 MHz). In contrast to strongly coupled nuclei, weakly coupled $^1$H nuclei, observed as signals at $\pm 51.4$ MHz, are not subject to asymmetries depending on the magnetic field positions. This is because the hyperfine coupling of weakly coupled $^1$H nuclei is smaller than the inhomogeneous EPR linewidth. This effect allows us also to differentiate among remote and local hydrogen atoms, as shown for Cu$^{2+}$ in solution at two different observed B-values (Supplementary Information S12). To complete and support this discussion, especially as far as the conformational properties of imidazole ring in the major groove of the DNA duplex with one modified nucleoside are concerned, Molecular Dynamics (MD) simulations have been performed (Supporting Information S14 and S15). Thus, for the label attached onto deoxyadenosine residue at ninth position, as the structure displayed in Fig. 2c, the dihedral angle defined by the four atoms marked in the inset of Figs S14 and S15 was monitored. The distribution of the dihedral angle suggests four conformations with major populations at $-180^\circ$ and $+180^\circ$ and minor populations at $-60^\circ$ and $+60^\circ$. With respect to the standard $B$-DNA structure, the minor population conformations are buried deeper into the major groove. Only the major conformations at $-180^\circ$ and $+180^\circ$ are more exposed. The flexibility/rigidity of the linker might determine the accessibility and functional efficiency of the imidazole ring in its interaction with Cu$^{2+}$ ions.

Figure 5. (a) 2D HYSCORE pulse sequence; the experiments for the 5 samples indicated in b–f are recorded at 20 K. (b) CuCl$_2$·2H$_2$O in a cacodylated buffer solution (pH 7.45). (c) Monomeric L$_2$ and Cu$^{2+}$. (d) Native DNA duplex and Cu$^{2+}$. (e) Covalently modified DNA containing one L$_2$ ligand and Cu$^{2+}$. (f) Three components mixture containing native DNA sequence, monomeric L$_2$ (non-covalently added) and Cu$^{2+}$.
Figure 6. ELDOR-detected NMR spectra recorded at Q-band frequencies ($\nu_0 = 34.025$ GHz) and 20 K. The magnetic field was selected to be at the position of the maximum of the echo-FS spectra. The EDNMR are recorded on the following samples: (a) CuCl$_2\cdot$2H$_2$O in H$_2$O. (b) Monomeric L$_2$ ligand ('ethyl-bridge') with CuCl$_2\cdot$2H$_2$O in a cacodylated buffer solution (pH 7.45). (c) Native (unmodified) DNA sequence with CuCl$_2\cdot$2H$_2$O in a cacodylated buffer solution (pH 7.45), using an excess 5:1. (d) Singly modified DNA sequence (L$_2$ ligand covalently attached) with CuCl$_2\cdot$2H$_2$O in a cacodylated buffer solution (pH 7.45), using an excess 5:1. (e) Monomeric L$_2$ ligand mixed with native DNA sequence (NON-covalently attached) with CuCl$_2\cdot$2H$_2$O in a cacodylated buffer solution (pH 7.45), using an excess 5:1. $^1$H signals are centered around $|\pm \nu_{1H}| = 51.4$ MHz, while the $^{14}$N signals are to the different values of $|\pm \nu_{1H}| = 51.4$ MHz, while the $^{14}$N signals implies the contribution of the native duplex and the L$_2$ ligand, both in the covalent and non-covalent binding mode. In all cases the strength of the detection pulses was selected to yield a maximum echo intensity.

Figure 7. ELDOR-detected NMR spectra recorded at Q-band frequencies ($\nu_0 = 34.025$ GHz) and 20 K. The magnetic field was selected to be at the position indicated on the echo-FS spectra (top-left). Other experimental details as for Fig. 6.
**Discussion**

Looking at the monomeric structures of the two ligands, the absence of the urea bridge on the L2 structure does not affect the binding efficiency. Indeed, the ethyl (alkyl) bridge among the imidazole unit and the nucleoside can still guarantee a strong coordination to the paramagnetic center as that of L1, as demonstrated by different EPR parameters. Furthermore, degradation of the urea bridge within more complex oligomers is then avoided by using ligand L1, containing exclusively the ethyl bridge attached to the nucleoside. The binding modes identified in this work suggest a preferred affinity to nitrogen nuclei, even if the role of the phosphate backbone remains elusive, with respect to the major/minor grooves of DNA. However, it paves the way for several different approaches, including selective inhibition of binding sites by amplifying or reducing the intercalative efficiency of those ligands (L1 or L2 in the monomeric form). It seems that the imidazole moiety within the duplex has lower coordination propensity compared to the free monomeric ligand. Furthermore, the EPR/ESR studies strongly suggest a reduced accessibility of the imidazole moiety that can be overcome by using it as intercalating agent.

The overall study proposed in this work suggests that the non-covalent mixtures can guarantee the suitable binding affinity towards copper ions. It would be extremely interesting to monitor and follow cleavage and ligation upon different binding modes to identify those nuclei mainly responsible for the catalytic core of DNAzymes/deoxyribozymes, as well as ribozymes and other catalytic nucleic acids. Recently, the impact of tethered imidazole on duplex stabilization has been reported in an aptamer-type construction; such enhanced stabilization could be also key to the non-covalent approach illustrated in this work. The modularity in the binding approach has also been reported for terpyridine/aptamer conjugates; a suitable spatial configuration for a substrate within a flexible DNA region is even able to increase the affinity for Cu²⁺ and Fe³⁺ during the oxidation of dopamine to aminochrome. Even for those very recent examples a spectroscopic analysis that can achieve atomistic resolution is severely lacking.

This work paves the way for a series of promising and novel experiments. First of all, the entire set of structures here proposed exhibit natural isotopic abundance; this means that in future, selective coupling can be switched on/off by using ¹³C, ³¹H, ¹⁵N or ¹⁷O enriched isotope analogues. For example, the number of transitions for ¹⁴N/¹⁵N will greatly simplify the HYSORE and EDNMR spectra (Supplementary Information S13). Secondly, self-cleaving DNA sequences can be monitored during the cleavage reaction by applying two different approaches: a single modified nucleotide can be mixed to the native DNA sequence and the catalytic effect of Cu²⁺ can be checked in terms of hyperfine couplings with the nuclei in close proximity. For the EPR pulsed experiments, the EDNMR can be further improved by higher-powered ELDOR pulses. The combination frequencies involving the simultaneous excitation of two different nuclei can make the interpretation of spectra more difficult. However, by simulation of the EDNMR spectra, as well as comparison with more conventional ENDOR experiments and integrative combination of DFT data, a description at atomic scale can be obtained. It is possible to state that the loss of affinity for singly labelled oligonucleotides strongly supports a multi-sites binding scenario, as confirmed by signal splittings in the EDNMR spectra. Fascinating opportunities arise for transferring the approach here proposed from the structure to the reactivity of catalytic DNA fragments. Investigations in this direction are ongoing and will be reported in due course.

**Methods**

**Synthesis of DNA sequence.** Synthesis of L₁ and L₂ ligands was performed and adapted from ref.³. Solid-phase synthesis of DNA duplexes singly modified with deoxyadenosine-L₁ (or –L₂) was carried out according to procedures described elsewhere.

**Sample preparation for EPR experiments.** Duplexes have been formed by heating DNA strands at 90°C for 3 minutes and slowly cooled down to room temperature. For the different pH, three buffers were used: Sodium Acetate (pH 4.00), Cacodylate (pH 7.45), Glycine/NaOH (pH 10.3). 10% (v/v) of glycerol was added before freezing the sample in liquid Nitrogen. With respect to Cu²⁺ solution (CuCl₂), a molar excess of monomeric ligand or double helix was used, up to an excess of 1:10. 100 mL solutions were used in 4 mm tubes (X-band) and 3 mm tubes (Q-band), respectively.

**EPR experiments.** Continuous Wave (CW) X-Band measurements were carried out using an X-band Bruker E500 instrument (9.4 GHz, TE₀₁₂ resonator) equipped with a nitrogen flow cryostat. All CW experiments were recorded at 120 K and with a shot repetition rate of 100 Hz, unless stated otherwise. Pulsed EPR experiments at X-band were performed on a Bruker ELEXYS E-580X-band spectrometer with a SuperX-FT microwave bridge and a Bruker ER EN4118X-MD4 dielectric resonator. Cryogenic temperatures (20 K) were obtained by the use of an Oxford flow cryostat. The field-swept EPR spectra were recorded by electron spin echo (ESE) detection; electron−spin−echo (ESE)-detected EPR experiments were carried out with the pulse sequence: π/2−τ−π−τ−echo. For the X-band experiments the mw pulse lengths τ₁₂ = 16 ns and τ₁₂ = 32 ns and a τ value of 200 ns were used. A two-step phase-cycle was applied to remove all unwanted echoes.

The Hyperfine Sublevel Correlation (HYSCORE) experiments were carried out using the pulse sequence π/2−τ−π−π−τ−π−t₁−π−t₂−π−π−τ−τ−echo. The time traces of the HYSCORE spectra were baseline corrected using a third-order polynomial, apodized with a Hamming window and zero-filled. After two-dimensional Fourier transformation, the absolute value spectra were calculated. A four-step phase cycle (for X-band experiments) was used to remove unwanted echoes.

The ELDOR-detected NMR (EDNMR) measurements were performed at the Q-band frequency using the pulse sequence HTA−T−π/2−τ−τ−π−τ−π−τ−echo. A high turning angle (HTA) microwave pulse was applied at the frequency ντ₂. The detection Hahn echo pulse sequence π/2−τ−π−τ−π−τ−echo was matched with the cavity resonance microwave frequency νr, and the spectra were acquired via continuous sweeping of the HTA frequency ντ₂ at fixed B₀. The length of the HTA pulse was optimized to 9 μs. Between the HTA pulse and the detection
sequence a delay $\tau_d$ of 204 ns was included. In the ESE detection sequence the same pulse lengths and interpulse delay as in the field-swept EPR experiments were applied. To ensure sufficient sensitivity, the echo integration window $\tau_{int}$ was set to 500 ns. No phase-cycle was used in the EDNMR experiments. The simulations of the EPR spectra were performed with the Easyspin software package (version 5.2.2)\textsuperscript{31}.

**DFT calculations.** All DFT calculations for L\textsubscript{1} and L\textsubscript{2} ligands bound to Cu\textsuperscript{2+} ions were performed with the ADF (Amsterdam Density Functional) code developed by E. J. Baerends and co-workers\textsuperscript{44}. Triple-zeta basis sets have been used throughout. All geometry optimizations were performed in vacuo relying on the Generalized Gradient Approximation (GGA) VVPB exchange-correlation (XC) potential (VWN + BP: Vosko, Wilk & Nusair\textsuperscript{45}) + corrective terms by Becke\textsuperscript{46} for the exchange, and Perdew\textsuperscript{37} for the correlation) with ADF grid precision 6. Subsequently, the computation of hyperfine coupling tensors relied on the use of the B3LYP exchange-correlation potential\textsuperscript{38,39} (20% of Hartree-Fock–Fock exchange) turning on the (non-relativistic) ADF option ESR.

**MD simulation.** 3D structure of Imidazole labelled adenosine monophosphate was generated in Gadget\textsuperscript{40}. Structure was capped at O\textsubscript{3}, and phosphate oxygens using methyl groups before geometry optimization. Using the ORCA package\textsuperscript{41}, the structure was optimized at restricted Hartree-Fock level of theory using 6–31G* basis set\textsuperscript{42}. Using amber tools\textsuperscript{18}, AM1-BCC charges were calculated and generalized amber force field topology compatible with gromacs2018.2\textsuperscript{28} package was generated for labelled deoxyadenosine. Structure for duplex DNA was generated using make-na server (http://structure.usc.edu/make-na/server.html). Imidazole-derivative molecule was fitted onto deoxyadenosine residue, using the purine ring atoms (only carbon and nitrogen) for the least square fitting, and coordinates of imidazole ring were transferred to pdb file creating the labelled DNA molecule. DNA molecule was modelled using parmbsc1 forcefield\textsuperscript{44}. Labeled duplex was placed in a dodecahedron box with 1.0 nm distance between the box walls and DIMA. Simulation box was solvated with tip3p water molecules and salt concentration was set to 0.15 M using Na\textsuperscript{+} and Cl\textsuperscript{−} ions. Simulation system was energy minimized using steepest descent algorithm until the largest force was smaller than 1000 kJ/mol/nm, followed by temperature equilibration to 300 K in 100 ps using Berendsen thermostat with a tau-t of 0.1 ps. Pressure was equilibrated to 1 atm in 1 ns using Berendsen barostat and temperature was regulated using velocity-rescaling thermostat at 300 K.

Using the equilibrated structure, three 100 ns production run simulations were started in which temperature was regulated using velocity-rescaling thermostat and pressure with Parrinello-Rahman barostat at 300 K and 1 atm using tau-t of 1 ps and tau-p of 2 ps. Structures were saved every 10 ps. Further details of MD analysis are provided in Supporting Information S16.

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experiments, analyzed data and wrote the paper. All authors edited the manuscript.

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Author Contributions

J.-M.M. performed the DFT analysis on the monomeric ligands L1 and L2. H.A. prepared the samples for the C.W. EPR experiments. S.C.D. performed the MD analysis on the singly-labelled duplex. G.S. designed pulsed EPR experiments, analyzed data and wrote all. All authors edited the manuscript.

Additional Information

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