Surprising Complexity of the \([\text{Gd(AAZTA)}(\text{H}_2\text{O})_2]^-\) Chelate Revealed by NMR in the Frequency and Time Domains

Daniela Lalli,* Fabio Carniato, Lorenzo Tei, Carlos Platas-Iglesias,* and Mauro Botta

ABSTRACT: Typically, \(\text{Ln(III)}\) complexes are isostructural along the series, which enables studying one particular metal chelate to derive the structural features of the others. This is not the case for \([\text{Ln(AAZTA)}(\text{H}_2\text{O})_x]^- (x = 1, 2)\) systems, where structural variations along the series cause changes in the hydration number of the different metal complexes, and in particular the loss of one of the two metal-coordinated water molecules between Ho and Er. Herein, we present a \(^1\text{H}\) field-cycling relaxometry and \(^{17}\text{O}\) NMR study that enables accessing the different exchange dynamics processes involving the two water molecules bound to the metal center in the \([\text{Gd(AAZTA)}(\text{H}_2\text{O})_x]^-\) complex. The resulting picture shows one Gd-bound water molecule with an exchange rate ~6 times faster than that of the other, due to a longer metal–water distance, in accordance with density functional theory (DFT) calculations. The substitution of the more labile water molecule with a fluoride anion in a diamagnetic-isostuctural analogue of the Gd-complex, \([\text{Y(AAZTA)}(\text{H}_2\text{O})_2]^−\), allows us to follow the chemical exchange process by high-resolution NMR and to describe its thermodynamic behavior. Taken together, the variety of tools offered by NMR (including high-resolution \(^1\text{H}, ^{19}\text{F}\) NMR as a function of temperature, \(^1\text{H}\) longitudinal relaxation rates vs \(R_{10}\) and \(^{17}\text{O}\) transverse relaxation rates vs \(T\) ) provides a complete description of the structure and exchange dynamics of these Ln-complexes along the series.

INTRODUCTION

Magnetic resonance imaging (MRI) has rapidly emerged as one of the most important and widespread tools in diagnostic clinical medicine and biomedical preclinical research. This is due to several favorable properties that characterize the technique, including the great spatial resolution, typically of the order of millimeters in the clinical setting, the ability to produce cross-sectional images of the body with excellent soft-tissue contrast, and the absence of ionizing radiation such as those used in X-ray and computed tomography (CT) scan. Commonly, the various imaging techniques are associated with the use of suitable contrast media, which have the purpose of improving the signal-to-noise ratio and optimizing the visualization of the morphology and physiology. Despite the remarkable level of intrinsic contrast of MRI images, even this diagnostic modality largely uses contrast agents that, as it is well known, are based on \(\text{Gd(III)}\) complexes.\(^1\)–\(^3\) These exogenous probes possess high efficacy in accelerating the longitudinal relaxation rate \((R_L)\) of the water protons of the tissues in which they are distributed; therefore, they allow shortening of the measurement time and improvement of the contrast-to-noise ratio of MR images, thus facilitating the overall diagnosis. The clinically approved gadolinium(III)-based contrast agents (GBCAs) are complexes with octadentate polyanimo–polycarboxylate ligands in which the metal ion completes its coordination number \((\text{CN} = 9)\) by binding to one water molecule \((q = 1)\).\(^4\)–\(^6\) The high thermodynamic stability, kinetic inertness, and fast clearance of such complexes, which are essential prerequisites for their use in clinics, represent additional important benefits. Despite the many advantages associated with the use of GBCAs, their relaxivity values in clinical fields are only a fraction of those theoretically achievable.\(^7\) To compensate for their low efficacy, contrast agents are typically administered in relatively high doses to gain high-quality MRI images.\(^3\) On the other hand, the development of new GBCAs with enhanced efficacy, high thermodynamic stability, and kinetic inertness would allow reducing the injected doses. This would have a beneficial impact in terms of costs and could contribute to minimizing the possibility of long-term Gd deposition. The efficacy of a GBCA is associated with the relaxivity parameter \((r_1)\), which measures the \(R_I\) increase of the water protons per millimolar unit concentration of the paramagnetic ion and, for clinically used CAs, is about 5 mM\(^{-1}\) s\(^{-1}\) (at 1.5 T and 298 K).\(^1\) For this reason, the research toward the optimization of the values of \(r_1\) has been very active in the last 30 years and has led to an in-depth knowledge of the relationship between \(r_1\) and the...
molecular parameters that describe the structural aspects and dynamic processes in which the complexes are involved.

At the magnetic field values of clinical and preclinical relevance (approx. 1−7 T), \( r_t \) is essentially determined by the molecular tumbling rate \( (\tau_b) \), by the hydration state \( (q) \), by the average lifetime \( (\tau_M) \) of the coordinated water molecules, and by the electronic relaxation of the Gd\(^{3+}\) ion (described by the parameters \( \Delta^2 \) and \( \tau_V \)).\(^1,2\)

One of the possible ways to increase the efficiency of CAs is to increase the number of metal-bound water molecules, which can be achieved by the use of heptadentate ligands. However, the loss of one donor atom from the ligand comes at the cost of decreased thermodynamic stability and/or kinetic inertness. In addition, shortly after the initial studies, it was found that most of the \( q = 1 \) complexes show a pronounced tendency to form ternary complexes with oxyanions of biological relevance (e.g., carbonate, lactate, malonate, or oxalate) with the displacement of one or both water molecules.\(^8,9\) As a primary consequence, this is accompanied by a marked decrease in relaxivity, an effect opposite to the one sought. This process likely represents one of the steps involved in the metallation reaction in which Gd\(^{3+}\) is replaced by other endogenous metal ions (mostly Zn\(^{2+}\) and Cu\(^{2+}\)) under physiological conditions. Over the years, only a small number of complexes with \( q = 2 \) that do not strictly follow this behavior and feature improved properties have been reported.\(^10,11\) The family of hydroxypyridinone (HOPO)-based complexes (Scheme 1), developed by Raymond et al. since the mid-1990s, is a notable example. The ligands are heptadentate, leaving two open sites for water coordination in the eight-coordinate Gd-complexes.\(^12,13\) The complexes have considerable thermodynamic stability, high \( r_t \) values, and rapid exchange of the bound water. Furthermore, they do not readily form ternary complexes with coordinating anions.

Similar properties are found in the Gd-complex with an open-chain ligand containing three acetate arms and one picolinate group, reported by Caravan et al. (Scheme 1).\(^14\) Finally, a third class of \( q = 2 \) compounds is represented by the GdAAZTA-like complexes (Scheme 1).\(^15\) AAZTA and derivatives are mesocyclic heptadentate ligands that form complexes with the Gd\(^{3+}\) ions that are not only thermodynamically stable but also kinetically inert.\(^16,17\) [Gd(AAZTA)-(H\(_2\)O)_2]^- represents an excellent candidate for in vivo applications in preclinical studies, due to several favorable properties: easy and cheap to synthesize, it shows high thermodynamic stability, high relaxivity, and two inner-sphere water molecules in sufficiently fast water exchange with the bulk that are not displaced by bidentate anions in physiological media. Moreover, near physiological conditions, [Gd-(AAZTA)-(H\(_2\)O)_2]^- is significantly more kinetically inert than GdDTPA.\(^15,16\)

Despite AAZTA being reported in 2004,\(^15\) only recently it has been shown that when bound to different members of the Ln series, the resulting complexes display unexpected and unique chemical properties. A decrease of the hydration state \( (q) \) changes from 2 to 1) is observed along the Ln series, which is surprisingly accompanied by a remarkable decrease of the water exchange rate, ca. 2−3 orders of magnitude.\(^18\) Near the end of the series, the coordinated water molecule becomes so tightly bound to the metal, that its signal can be directly detected in high-resolution \(^1\)H NMR spectra of the Yb(III)-complex.\(^19\) The residence lifetime of the bound water molecule, measured by chemical exchange saturation transfer (CEST) experiments (150 \( \mu \)s), is 4 orders of magnitude higher than that of [Gd(AAZTA)-(H\(_2\)O)_2]^- .

Recently, a systematic study on the Ln(III) complexes with AAZTA highlighted that the structural change occurs along the Ln series between Ho and Er, due to the increased steric compression of the ligand on the bound water molecules upon decreasing the metal ion size.\(^20\) Most importantly, two coordinated water molecules in the [Ho(AAZTA)-(H\(_2\)O)_2]^- complex were detected by CEST experiments near 0 °C, which resonate at different chemical shift values \( (−47 \text{ and } −255 \text{ ppm}) \) and are characterized by significantly different water exchange rates \( (k_w=1/\tau_M=5.8 \times 10^3 \text{ s}^{-1}) \). This interesting observation has prompted us to reconsider the case of [Gd(AAZTA)-(H\(_2\)O)_2]^- and, using new NMR relaxometric data obtained with a much more intense magnetic field (11.7 instead of 2.1 T), to verify if there is any experimental evidence that the two water molecules can be characterized by different water exchange rates.

The topic that we address in this work is of interest both for the area of MRI biomedical imaging and, above all, for basic coordination chemistry as it considers fundamental aspects of the properties of lanthanide complexes.

### RESULTS AND DISCUSSION

**Relaxometric Characterization of [Gd(AAZTA)-(H\(_2\)O)_2]^-**

NMR spectroscopy is able to provide an accurate description of the water exchange process that occurs between the paramagnetic center and the bulk solvent. Exchange rates of the metal-bound water molecules \( (k_w) \) can be obtained via high-resolution NMR, by measuring the paramagnetic shift \( (\Delta \omega) \) and the linewidths of the \(^{17}\)O signal of the isotopically enriched bulk water \( (\Delta \omega/\Delta \nu/\Delta \nu_{1/2}(^{17}\text{O})) \), and hence its transverse relaxation rate \( (R_2=\pi^2/\Delta \nu_{1/2}(^{17}\text{O})) \), as a function of temperature. The so-obtained data can be analyzed with the Swift–Connick equations that describe the water exchange process between the two sites.\(^21,22\) As a first approximation, the

---

**Scheme 1. Chemical Structure of the Complexes Described in This Study**

![Chemical Structure of the Complexes Described in This Study](image-url)

---

https://doi.org/10.1021/acs.inorgchem.1c03194

Inorg. Chem. 2022, 61, 496−506
transverse relaxation rate can be considered inversely proportional to the sum of the residence lifetime of the water molecule coordinated to the metal center ($\tau_M$) and its transverse relaxation time ($R_2 \propto 1/(T_{2M} + \tau_M)$). On this basis, low-molecular-weight Gd(III)-chelates, characterized by $\tau_M$ values similar to $T_{2M}$ at physiological temperatures, are predicted to follow an "intermediate exchange regime", as reported for the [Gd(DTPA)(H$_2$O)]$^{3-}$ and [Gd(DOTA)·(H$_2$O)]$^{3-}$ complexes with a $\tau_M$ of about 300 ns at 298 K (Figure 1). Such chelates are expected to undergo a slow-to-

Figure 1. Temperature dependence of the $^17$O transverse relaxation rate calculated at 11.74 T for a monohydrated Gd(III)-complex 25 mM with a $\tau_M = 30$ ns, $\Delta H^0 = 40$ kJ mol$^{-1}$ (red profile) and a $\tau_M = 300$ ns, $\Delta H^0 = 48$ kJ mol$^{-1}$ (blue profile).

fast exchange transition in the temperature range between 280 and 350 K. Indeed, at low temperatures ($T = 280$ K), a decrease of the water exchange rate and a shortening of $T_{2M}$ occurs, which renders $\tau_M$ longer than $T_{2M}$, thus the dominant term in the denominator of the $R_2$ equation. In such exchange dynamics, described as the "slow-exchange regime", the $R_2$ values follow the same temperature dependence trend as $\tau_M$. $R_2$ is predicted to increase at rising temperatures ($280$ K $< T < 310$ K) to reach a maximum at $310$ K, coinciding with the slow-to-fast exchange transition point. A further temperature increase ($T > 310$ K) induces an increase of the water exchange rate and an increase of $T_{2M} (\tau_M \ll T_{2M})$, which is described as the "fast-exchange regime". Now $T_{2M}$ becomes predominant and dictates the behavior of the $R_2$ temperature dependence, which decreases for increasing temperatures (Figure 1).

While the temperature dependence of the $^17$O transverse relaxation rates previously measured on [Gd(AAZTA)·(H$_2$O)$_2$]$^{3-}$ at a low magnetic field strength ($B_0 = 2.1$ T) follows the expected behavior, the same measurements performed at higher magnetic fields (11.74 T) display an unusual trend in the lower temperature region (Figure 2). The increase in transverse relaxation rates at higher magnetic field strengths results in broader $^17$O NMR lines, measurable with better accuracy than at 2.1 T (Figure S1). This unraveled the presence of an unexpected trend in the relaxation profile at lower temperatures. Notably, an increase of $R_2$ becomes clearly observable for temperature values lower than 285 K (Figure 2). Such a trend resembles that previously observed for [Gd(HPDO3A)·(H$_2$O)], characterized by the presence of multiple isomers in slow exchange on the NMR timescale, each featuring different water exchange rates. Instead, in the case of [Ln(AAZTA)·(H$_2$O)$_x$]$^{3-}$ complexes, the existence of slow-exchange isomers is discarded by experimental evidence obtained from variable-temperature high-resolution NMR, where only one set of resonances is observable in the temperature range between 273 and 310 K. More likely, the behavior here reported could be owed to the presence of two Gd-coordinated water molecules with significantly different water exchange rates ($k_x = 1/\tau_M$). Such a hypothesis is in agreement with recent studies, in which the presence of two metal-bound water molecules with significantly different residence times were suggested by means of density functional theory (DFT) calculations and wave function analyses. In this case, the two Gd(III)-coordinated water molecules are expected to contribute to a different extent to the observed $^17$O-$R_2$ profile: the intermediate-exchanging species mainly contributes at intermediate temperatures, where the contribution of the faster-exchanging water molecule is almost absent. Conversely, the latter becomes predominant for $T$ lower than $\sim 280$ K, where the contribution of the slower-exchanging water is negligible. Previously, it was not possible to detect the presence of the faster exchange species at lower field strengths ($B_0 = 2.1$ T). On this basis, the $^17$O-$R_2$ data were reanalyzed by considering the presence in solution of two water molecules subject to different dynamics of exchange. To obtain a comprehensive description of such processes, a complete set of high-field $^17$O NMR measurements was collected, comprising water $^17$O-$R_2$ and chemical shift variations as a...
function of temperature (Figure 2), along with $^1$H nuclear magnetic relaxation dispersion (NMRD) profiles at three different temperatures (10, 25, and 37 °C) over the $^1$H Larmor frequency range 0.01 to 120 MHz (Figure 3). The $^1$H and $^{17}$O NMR data were simultaneously fitted to obtain more reliable results, the best-fit parameters are reported in Table 1. The data were well reproduced by considering that the two Gd(III)-bound water molecules are characterized by significantly different residence lifetimes ($\tau_M^A = 29$ ns and $\tau_M^B = 169$ ns) and enthalpy barriers associated with the exchange process ($\Delta H_M^A = 20$ kJ mol$^{-1}$ and $\Delta H_M^B = 29.5$ kJ mol$^{-1}$). It is worth noting that the average of these two residence lifetimes is very similar to that obtained previously under the assumption that the two coordinated water molecules are characterized by identical water exchange dynamics ($\tau_M = 90$ ns). The water molecule residing for shorter times on the metal center (A) exchanges ~6 times faster than the other, with a reaction enthalpy ~1.5 times lower. This suggests that the exchange reaction is mainly controlled by the energy cost needed to break one Gd–Ow bond, necessary to reach an eight-coordination transition state. It is reasonable to assume that breaking the Gd–Ow bond of the water molecule that is less strongly bound to the metal center, hence at a longer distance from it, requires the lowest energy. This is in line with previous predictions of Platas-Iglesias and co-workers, who calculated significantly different Gd–Ow distances for the two water molecules ($r_A = 2.505$ Å and $r_B = 2.484$ Å) differently positioned in the Gd(III)-coordination environment. The considerably more labile water molecule occupies a capping position, and the more tightly bound to the metal center resides in one of the vertices of the coordination polyhedron. This is in agreement with the labile capping bond effect introduced previously, which states that water ligands occupying sterically demanding capping positions are intrinsically labile. The calculated $^{17}$O hyperfine coupling constants show a slightly higher value for the more strongly bound and more slowly exchanging water molecule ($A_{Ow}/h = -3.8 \times 10^6$ rad s$^{-1}$) and $A_{Ow}/h = -3.9 \times 10^6$ rad s$^{-1}$), consistent with the DFT predictions. The rotational dynamics ($\tau_R = 74.0$ ps) and electronic parameters ($\Delta^2 = 2.6 \times 10^9$ s$^{-2}$; $\gamma = 30$ ps) characterizing the system are in agreement with previous studies reported in the literature.

The influence of the simultaneous presence of the two species is not evident in the dependence of the relaxivity ($r_1$) on the magnetic field strengths (Figure 3), as proton relaxivity is limited by the fast rotation of the complex in solution rather than by water exchange.

### Affinity Constant of the Fluoride Anion for the [Gd(AAZTA)(H$_2$O)$_2$]$^-$ Complex

Previous studies demonstrated that fluoride can directly compete with water molecules to coordinate lanthanide complexes and remain tightly bound. The halide affinity for the metal center can be sufficiently high to replace one or more inner-sphere water molecules. As a result, the paramagnetic metal ion loses efficiency in relaxing the bulk water protons, which leads to a significant decrease in relaxivity, conveniently detectable by relaxometry.

We investigated the binding of the fluoride anion to [Gd(AAZTA)(H$_2$O)$_2$]$^-$ by monitoring the change in the longitudinal relaxation rates of water protons ($R_1$), which

![Figure 3. $^1$H NMRD profiles of a 3.1 mM aqueous solution of [Gd(AAZTA)(H$_2$O)$_2$]$^-$, at pH 7.0, recorded at different temperatures (283 K (solid blue diamonds), 298 K (solid black diamonds), and 310 K (solid red diamonds)). The solid lines correspond to the fits of the data, as described in the text.](https://doi.org/10.1021/acs.inorgchem.1c03194)
occurs after adding increasing amounts of the halide to the aqueous solution of the complex. For this purpose, a 1 mM [Gd(AAZTA)(H₂O)₂]⁻ solution was titrated with NaF until variations of R₁ become negligible, which corresponds to an 800-fold excess of the halide. Changes of R₁ were measured at 32 MHz, 298, and 310 K (Figures 4 and S2). The so-obtained titration curves were analyzed following the proton relaxation enhancement (PRE) method, which provides access to the titration curves were analyzed following the proton relaxation method, which provides access to the changes of the apparent a enhancement (PRE) method, which provides access to the complex. At the end of the titration, a mixture of the binary adduct contribution to the relaxation was subtracted for each concentration of the binary adduct is estimated to be 12% of the total (Scheme 2). As a consequence, the relaxometric properties of the so-prepared solution arise from two different contributions: (1) one related to the preponderant ternary species having only one inner-sphere water molecule, which contributes to a minor extent to the observed relativity, and (2) the dominant contribution of the minor binary species with two coordinated water molecules. To characterize the relaxometric properties of the pure [Gd(AAZTA)(H₂O)F]²⁻, the complete formation of the ternary complex should be obtained, which can be only achieved using extremely high concentrations of the halide (≥1 M). However, it is known that high ionic strengths (>0.6 M) can substantially alter the relaxometric properties of the chelates in aqueous solutions, due to the strong salt-water interactions capable of modifying the structure and microviscosity of water and therefore, of changing the molecular reorientation rates of the complexes. Therefore, to investigate the relaxometric behavior of the [Gd(AAZTA)(H₂O)F]²⁻ complex, 0.6 M NaF was added to the solution of [Gd(AAZTA)(H₂O)₂]⁻, and the relaxometric contribution of the binary species, calculated on the basis of its mol fraction present in solution for each working temperature, was subtracted from the total, as explained below. Relaxometric Characterization of the [Gd(AAZTA)-(H₂O)F]²⁻ Ternary Complex. Relaxometric characterization of the ternary adduct, in which one water molecule is replaced with a fluoride anion, was performed with the aim of evaluating whether any changes in water exchange dynamics were detectable. To characterize the relaxometric properties of the ternary complex alone, a complete set of ¹H NMRD profiles and variable-temperature ¹⁷O NMR data was acquired on a [Gd(AAZTA)(H₂O)F]²⁻ solution, from which the binary adduct contribution to the relaxation was subtracted for each of the temperatures measured (Figures 5 and 6).

As already mentioned, the ternary complex was prepared by adding 0.6 M of NaF to a 1 mM [Gd(AAZTA)(H₂O)₂]⁻ solution. The concentration of the binary adduct present in such a solution was calculated for all of the measuring temperatures from the enthalpy of the fluoride–Gd-binding reaction, which was derived from the Kᵢ values reported in Table 2, assuming an Arrhenius behavior. Then, a solution of the binary complex having the same ionic strength as the ternary species was prepared by adding 0.6 M NaCl to a 1 mM [Gd(AAZTA)(H₂O)₂]⁻, and the relaxometric properties of the so-prepared solution arise from two di...
complex from that of the ternary species, to obtain the relaxation profiles of the monohydrated complex alone, free from possible influences of the relatively high ionic strength. The simultaneous analysis of the magnetic field dependence of $R_1$ of the water protons, and of the temperature dependence of $R_2$ and $\Delta \omega$ of $^{17}$O (Figures 5 and 6), was performed using the same approach described above, to obtain quantitative information on the exchange dynamics of the monohydrated complex. During the fitting procedure, the same set of parameters described above was fixed to the values used for the binary $[\text{Gd(AAZTA)(H}_2\text{O)}]^{2-}$ complex (Table 1), except for the hydration number that was set to 1.

By reasonably assuming that $F^-$ more easily substitutes the more labile water molecule, characterized by a shorter residence lifetime ($\tau_{M \text{A}} = 29$ ns), the longer bound distance from the metal center ($r = 2.505$ Å), and lower energy cost associated with breaking of the Gd–Ow bond ($\Delta H_{M \text{A}} = 20$ kJ mol$^{-1}$) and that its binding reduces the charge density at the metal center and labilizes the coordinated water molecule, we expect an acceleration of the water exchange rate. This is in agreement with the experimental data showing a remarkable ($\sim 50$-fold) decrease in the residence lifetime ($\tau_M = 3.4$ ns), followed by a slight reduction of the enthalpy associated with the exchange process ($\Delta H_M = 23.4$ kJ mol$^{-1}$) with respect to the more strongly bound water molecule of the binary complex. This behavior may be favored by the increase in the negative charge of the complex and by the steric interaction in the water coordination site that could destabilize the nine-coordinate ground state, thus diminishing the activation energy for the exchange process. However, the value of the hyperfine coupling constants ($A_\text{O}/\hbar = -3.8 \times 10^6$ rad s$^{-1}$) remains virtually unaffected.

DFT calculations were performed to rationalize the results of the relaxometric study (Tables S3–S5). The model systems investigated include a number of explicit second-sphere water molecules, while bulk solvent effects were considered using a polarized continuum model. Different computational studies evidenced that this mixed cluster/continuum approach is required to attain a proper description of the Ln–Ow and Ln–F bonds.$^{37,38}$

Calculations performed on the $[\text{Gd(AAZTA)(H}_2\text{O)}F]^{2-}$–$5\text{H}_2\text{O}$ system suggest that fluoride coordination may replace preferentially the water molecule that provides the weakest interaction with the metal ion (Figure 7), as the free energy

![Figure 5. $^1$H NMRD profiles of a 1.0 mM solution of $[\text{Gd(AAZTA)-(H}_2\text{O)}F]^{2-}$, in the presence of NaF 600 mM, recorded at different temperatures 283 K (solid blue diamonds), 298 K (solid black diamonds), and 310 K (solid red diamonds). The contribution to the relaxivity of the residual $[\text{Gd(AAZTA)(H}_2\text{O)}]^{3-}$ species was subtracted from the NMRD profiles of $[\text{Gd(AAZTA)(H}_2\text{O)}F]^{2-}$ for each temperature, as described in the text. The solid lines correspond to the fits of the data.](image)

![Figure 6. $^{17}$O transverse relaxation rates (top) and $^{17}$O chemical shift variation (bottom) as a function of the temperature of $[\text{Gd(AAZTA)-(H}_2\text{O)}F]^{2-}$–9.4 mM, in the presence of NaF 600 mM, measured at 11.74 T. The contribution to the relaxivity of the residual $[\text{Gd(AAZTA)(H}_2\text{O)}]^{2-}$ species was subtracted from the $^{17}$O profiles of $[\text{Gd(AAZTA)(H}_2\text{O)}F]^{2-}$ for each temperature, as described in the text. The solid lines correspond to the fits of the data.](image)

![Figure 7. Structure of the $[\text{Gd(AAZTA)(H}_2\text{O)}F]^{2-}$–system obtained using DFT calculations. Selected bond distances (Å): Gd(1)–F(1), 2.271; Gd(1)–N(1), 2.718; Gd(1)–N(2), 2.713; Gd(1)–N(3), 2.628; Gd(1)–O(1w), 2.495; Gd(1)–O(2), 2.414; Gd(1)–O(3), 2.445; Gd(1)–O(4), 2.456; and Gd(1)–O(5), 2.431.](image)
difference between the two ternary adducts is large (∼4.6 kcal mol⁻¹) (Tables S3 and S4). Fluoride coordination induces a significant lengthening of the Gd-Ow distance from 2.480 Å in [Gd(AAZTA)(H₂O)₂]⁻ to 2.495 Å in [Gd(AAZTA)-(H₂O)F]²⁻·SH₂O, using the same computational approach. This is in agreement with the variation of the water exchange observed experimentally. Thus, fluoride coordination appears to weaken the interaction between the remaining coordinated water molecule and the metal ion, thus facilitating water exchange following a dissociative mechanism. This is confirmed by the electron density calculated at the (3,−1) critical point characterizing the Gd−Ow bond (ρ_{BCP}), which decreases from 3.93 × 10⁻⁸ to 3.74 × 10⁻⁸ au upon fluoride coordination. Different computational studies on lanthanide complexes showed that lower ρ_{BCP} values characterize weaker Ln−Ow bonds.⁴⁰,⁴¹ The calculated Laplacian bond orders (LBOs) also point to weaker coordination of the water molecule in [Gd(AAZTA)(H₂O)F]²⁻·SH₂O (LBO = 0.100) than in [Gd(AAZTA)(H₂O)₂]⁻·4H₂O (LBO = 0.125).

Kinetics and Thermodynamics of the Fluoride Binding with [Y(AAZTA)(H₂O)₂]⁺. The presence of an NMR active ¹⁹F species transiently coordinated to the paramagnetic center in the ternary complex prompted us to study its binding interaction by high-resolution NMR lineshape analysis. However, the long electronic relaxation time that characterizes Gd(III) (τₑ ∼ 10⁻⁸ s) gives the paramagnetic metal ion an excellent relaxing capability, which causes severe broadening of the NMR lines of the neighboring nuclei. To access the ¹⁹F resonances necessary to evaluate the exchange rate between the metal-bound and free fluoride, we substituted Gd(III) with the diamagnetic analogue Y(III). Despite the structure of the Ln(III)AAZTA complexes varying as a function of the lanthanide contraction, the rather similar ionic radii of Gd(III) and Y(III) are expected to preserve the coordination geometry and the water dynamics properties. However, we cannot exclude the presence in solution of two species with different hydration numbers (q = 1, 2) in fast exchange. By means of variable-temperature high-resolution ¹⁹F NMR measurements, it was possible to access the kinetic information on the exchange rate of the chemical reaction, and therefore the thermodynamic parameters associated with fluoride exchange.

A quantitative evaluation of the chemical exchange rate was obtained on a sample containing a mixture of the fluoride-free and fluoride-bound species that are characterized by distinct ¹⁹F resonances, by monitoring variations of the line shapes induced upon temperature changes. The sample was prepared to ensure a ~1 to 1 molar ratio of fluoride in the free (F⁻_{free}) and bound (F⁻_{bound} = [Y(AAZTA)-(H₂O)F]²⁻) form at the lowest working temperature, to obtain a system with a symmetrical two-site exchange reaction. This facilitates measurements of the linewidths and therefore determination of the fluoride exchange rate (see Experimental Section). High-resolution one-dimensional (1D) ¹⁹F NMR spectra were acquired by varying the temperature between 275 and 350 K. The simultaneous presence of two distinct sets of signals associated with the F⁻_{bound} (−72 ppm) and F⁻_{free} (−120 ppm) species indicates that the exchange between the free and bound species is slow on the NMR timescale (Figure 8). The third signal resonating at −75 ppm was assigned to the trifluoroacetate (TFA) present in the solution of the complex after deprotection.

It is worth noting that DFT calculations performed on the [Y(AAZTA)(H₂O)F]²⁻ system provide a calculated ¹⁹F NMR shift of −82.7 ppm, in reasonable agreement with the experimental value of −72 ppm. Calculations on the F⁻·18H₂O system give a calculated ¹⁹F chemical shift of −125.3 ppm, in excellent agreement with the experiment. These results clearly indicate that DFT provides a reasonable model for the structure of the fluoride adduct in solution.

The integration of the two peaks allows calculating the relative concentration of the F⁻_{bound} and F⁻_{free} species for all of the measured temperatures. Significant variations of the linewidths of the fluoride-free and bound forms were detected only above 325 K (Figure S4), suggesting that the fluoride exchange rate increases at increasing temperatures, and tends to reach the coalescence point for temperatures higher than 350 K (Figure 8). The fluoride exchange rate was determined by fitting the ¹⁹F integrals and linewidths at various temperatures, with respect to the TFA reference signal, through the dynamic NMR (DNMR) LineShape Analysis tool (version 1.1.2) implemented in Bruker’s Topspin 3.2 (Figure S4). The fluoride exchange rate is 2280 s⁻¹ at 325 K (Table S2), 1 order of magnitude faster than that found for a similar system, a tripositive Y(III)DOTA-tetraamide derivative at 298 K.⁴⁴ This difference in rates is attributable to the electrostatic repulsion occurring between the fluoride anion and the metal center in the negatively charged [Y(AAZTA)-(H₂O)₂]⁻ complex, which significantly accelerates the dissociation reaction with respect to the tripositive complex.

The plot of the logarithm of the kinetic constant as a function of temperature provides a linear correlation that allows estimating the thermodynamic parameters of the exchange reaction. The enthalpy (ΔH°) and entropy (ΔS°) variations of the reaction are extracted from the slope and intercept of the Eyring plot (Figure 9) and enable determining the variation of the Gibbs free energy (ΔG°) associated with the exchange process (Table S2).

The enthalpy difference extracted from the plot (ΔH° = 32.26 kJ mol⁻¹) is lower than that reported for other tripositive Y(III)DOTA-tetraamide derivatives (ΔH° = 45–47 kJ mol⁻¹), suggesting that the energy cost for the bond breaking between
the fluoride anion and the metal center in a negatively charged complex is lower than that of a tripositive chelate.
Conversely, the entropy variation ($\Delta S = -82.2 \text{ J mol}^{-1} \text{ K}^{-1}$) is significantly different from that reported for Y(III)-DOTA-tetraamide derivatives ($\Delta S = -41$ to 58 $\text{J mol}^{-1} \text{K}^{-1}$) indicating that the fluoride–metal ion interaction is appreciably affected by the nature of the ligand substituents, as well as the structure and the charge of the complex.

**CONCLUSIONS**
The complex $[\text{Gd(AAZTA)}(\text{H}_2\text{O})_2]^{-}$ represents one of the few bis-aquated Gd$^{3+}$ derivatives that displays favorable properties for *in vivo* applications as an MRI contrast agent. Bis-hydrated Gd(III)-complexes have been developed with the aim of improving the efficacy of GBCAs, which in principle may allow the injection of lower Gd$^{3+}$ doses in clinical practice and biochemical research. Only recently, unpredictable and unique coordination properties have been proven to characterize Ln(III)-complexes of AAZTA, which have not yet been fully characterized and understood. These concern the decrease of the hydration state along the lanthanide series due to the lanthanide contraction, which is unexpectedly associated with a significant decrease of the exchange rate of the remaining inner-sphere water molecule. This behavior is similar to that shown by the eight-coordinate $[\text{Ln(PDTA)}-(\text{H}_2\text{O})_2]^{-}$ derivatives, but in contrast to that observed for the nine-coordinate $[\text{Ln(DTPA–BMA)}(\text{H}_2\text{O})]$ and $[\text{Ln(DO3A)}-(\text{H}_2\text{O})_2]$ complexes. The detailed NMR studies reported here, allowed us to reveal subtle details of the coordination chemistry of $[\text{Gd(AAZTA)}(\text{H}_2\text{O})_2]^{-}$ and to address one of the questions that remained unanswered, that is whether the two water molecules coordinated to the metal ion are characterized by different residence times in the metal coordination sphere. The new $^1\text{H}$ NMRD and particularly $^1\text{O}$ NMR data, collected at higher magnetic field strengths, provided clear evidence that the two identical water ligands have substantially different exchange rates, one being considerably more labile than the other.
Such finding is supported by DFT calculations predicting that the faster-exchanging water molecule is more distant from the metal center and occupies the sterically hindered capping position of the coordination polyhedron, while the slower-exchanging water, located at one of the vertices of the coordination polyhedron, is closer to the metal center.
In addition, it has been shown that the more labile water molecule can be replaced by fluoride to form a $[\text{Gd(AAZTA)}-(\text{H}_2\text{O})F]^{2-}$ adduct, which is characterized by a fast water exchange rate ($\tau_M = 3.4$ ns at 298 K). Fluoride exchange is conversely very slow and could be analyzed by high-resolution NMR using the diamagnetic Y(III) derivative ($\tau_M = 1.4$ ms at 298 K).
The resulting picture clarifies the mechanism by which the increased steric compression induced by the contraction of the ionic radius along the Ln series favors the loss of the more labile water molecule. As a result, the coordinated water molecule with a lower steric hindrance interacts more tightly and resides for longer times at the metal center.
These results, in addition to their relevance to the coordination chemistry of f-elements, also indicate an approach to improve the effectiveness of this complex as an MRI probe for clinical and preclinical applications. One of the possible avenues to take could be inducing structural changes of the coordination geometry to drive the selective acceleration of the slow-exchanging water molecule while keeping the other one in an optimal exchange regime. This would induce significant relaxivity gains if the rotational dynamics is slowed down, for instance by interaction with human serum albumin (HSA). It is worth noting that lipophilic GdAAZTA derivatives retain the two coordinated water molecules when bound to HSA. In fact, theoretical predictions estimate an increase in $r_1$ for macro-molecular GdAAZTA derivatives ($\tau_M = 5$ ns) at the clinically relevant magnetic fields of 1.5 and 3 T (Figure 10). At 1.5 T, these simulations predict a relaxivity gain of $\sim$14% (from $r_1 = 69.5$ to 79.2 $\text{mM}^{-1} \text{s}^{-1}$) by reducing the residence time of the water molecule with $\tau_M = 169$ ns to 29 ns. Acceleration of water exchange results in an $r_1$ increase of $\sim$33% when taking as a reference a system in which both water molecules are characterized by $\tau_M = 169$ ns ($r_1 = 59.7$ $\text{mM}^{-1} \text{s}^{-1}$). The effect is less pronounced at 3 T assuming $r_6 = 5$ ns but more important for an intermediate $r_6$ of 0.5 ns (Figure S5). Therefore, in light of these new findings, we envisage that the relaxation properties of GdAAZTA derivatives can be

![Figure 9](https://doi.org/10.1021/acs.inorgchem.1c03194)

**Figure 9.** Eyring plots for the exchange rates of the fluoride-free and $[\text{Y(AAZTA)}(\text{H}_2\text{O})_2]^{-}$-bound form. Dashed lines represent the fitted data.

![Figure 10](https://doi.org/10.1021/acs.inorgchem.1c03194)

**Figure 10.** Relaxivity values calculated for a bis-hydrated Gd(III) complex with a $r_6 = 5$ ns and different water residence times. All other parameters are those listed for $[\text{Gd(AAZTA)}(\text{H}_2\text{O})_2]^{-}$ in Table 1. Vertical dashed lines indicate magnetic fields of 1.5 and 3 T.
optimized by modifying the ligand structure, for example, by replacing one or two carboxylic moieties with more sterically demanding donor groups (as propionate or phosphonate arms).44

Taken together, the results reported in this paper represent a step forward toward an understanding of the structural and dynamic properties of lanthanide complexes with polyamino— polycarboxylate ligands, an important family of complexes because of their biomedical applications.

**EXPERIMENTAL SECTION**

**NMR Experiments.** ¹H, ¹³C, and NMR spectra of the ligands and their precursors were recorded at 298 K using a Bruker AVANCE III 500 spectrometer equipped with a 5 mm double resonance TXI probe.

**Preparation of the [Ln(AAZTA)(H₂O)₂]⁻ Complexes.** The AAZTA ligand was kindly provided by Bracco Imaging S.p.A. (purity >99%).

[Ln(AAZTA)(H₂O)₂]⁻ complexes were prepared by adding 1.1 equiv of LnCl₃ salts to an aqueous solution of the AAZTA ligand at pH = 6. After the addition, the pH was adjusted to 6.0 with dilute NaOH and the solution was stirred at room temperature (r.t.) for 12 h. Then, the pH was increased to 10 by means of 0.1 M NaOH, and the solution was stirred for 3 h, to promote the precipitation of the uncomplexed Ln(III) as insoluble hydroxides. The solution was centrifuged (10 000 rpm, 5 min, r.t.), the supernatant was filtered through 0.2 μm filters and neutralized with dilute HCl. The concentration of Ln(III) complexes was evaluated by ¹H NMR measurements, using Evans’ method. The 1D ¹H NMR spectrum of the diamagnetic [Y(AAZTA)(H₂O)₆]⁻ complex demonstrates the high purity of the complex (>95%) (Figure S6).

**Fluoride Affinities for the [Gd(AAZTA)(H₂O)₆]⁻ Complex.** Association constants of [Gd(AAZTA)(H₂O)F]⁻ were determined by relaxometric titrations. The titrations were performed at 32 MHz, at two different temperatures (298 and 310 K), by adding increasing amounts of the NaF salt to 1 mM [Gd(AAZTA)(H₂O)₆]⁻ solution. Relaxivity values were measured after each addition for monitoring the formation of the ternary complex. A total of 13 points were collected for each titration. The final concentration of NaF was 0.8 M, corresponding to an 800-fold molar excess of the anion.

The same titration was performed in identical conditions, at 298 K, by adding increasing amounts of NaCl to a 1 mM [Gd(AAZTA)(H₂O)₆]⁻ solution. NMRD profiles were acquired on the ternary [Gd(AAZTA)-(H₂O)₆]⁻ complex, prepared by adding 0.6 mM NaF salt to a 1 mM [Gd(AAZTA)(H₂O)₆]⁻ solution, and on the binary [Gd(AAZTA)-(H₂O)₆]⁻ complex, prepared by adding 0.6 mM NaCl salt to a 1 mM [Gd(AAZTA)(H₂O)₆]⁻ solution (Figure S3a).

**Fluoride Affinity for the [Y(AAZTA)(H₂O)₆]⁻ Complex.** The rate of fluoride exchange with [Y(AAZTA)(H₂O)₆]⁻ was determined by ¹⁹F high-resolution NMR measurements. The ternary complex was prepared by titrating a 160 mM [Y(AAZTA)(H₂O)₆]⁻ solution with increasing amounts of NaF at 275 K. After each addition, high-resolution 1D ¹⁹F spectra were acquired and the integrals of the ¹⁹F NMR signals of the fluoride-free and bound forms were measured to monitor the formation of the [Y(AAZTA)(H₂O)F]³⁻ complex. At the end of the titration, an equal molar ratio of the fluoride-free and bound forms was reached, corresponding to a NaF concentration of 280 mM. 1D ¹⁹F spectra were acquired by varying the temperature from 275 to 350 K. Delay times (D1) were set to 10 s to make integration quantitative. Spectra were phased and baseline before analysis. The rate of exchange of fluoride was calculated fitting the ¹⁹F line shapes of the F_free and F_bound species referred to the TFA peak, at various temperatures through the dynamic NMR (DNMR) LineShape Analysis module (version 1.1.2) implemented using Bruker’s Topspin 3.2, as detailed in Figure S4.

**Relaxometric Measurements.** ¹/T₁, ¹H nuclear magnetic relaxation dispersion (NMRD) profiles were acquired using two different instruments, one operating at lower field strengths (0.01 to 10 MHz) and the other at higher field strengths (20–120 MHz). Low field data were measured using a fast-field cycling (FFC) Stelar SMARTTracer relaxometer (Stelar s.r.l., Mede, PV, Italy) equipped with a silver magnet. High-field measurements were collected with a high-field relaxometer (Stelar) equipped with an HTS-110 3T Metrology cryogen-free superconducting magnet. The measurements were performed using the standard inversion recovery sequence (2 experiments and 2 scans) with a typical 90° pulse width of 3.5 μs and the reproducibility of the data was within ±0.5%. The temperature was controlled with a Stelar VTC-91 heater airflow equipped with a copper—constantan thermocouple (uncertainty of ±0.1 K).

**¹⁷O Measurements.** Variable-temperature ¹⁷O NMR measurements were recorded on a Bruker AVIII 500 spectrometer equipped with a 5 mm probe and standard temperature control unit. Solutions containing 2.0% of the ¹⁷O isotope (Cambridge Isotope) and 10% D₂O for the external lock were used. The observed transverse relaxation rates were calculated from the signal full width at half-maximum (Δν₁₂). The bulk magnetic susceptibility contribution was subtracted from the ¹⁷O NMR shift data using the ¹H NMR shifts of the ¹⁷O-free and bound forms referred to the TFA peak, at various temperatures through the dynamic NMR (DNMR) LineShape Analysis module. The integration grid was set with the integral = ultra fine tting the ¹⁷F line shapes of the complexes. The resolution 1D ¹⁷O NMR spectra were phased and baselined before analysis. The rate of exchange of fluoride was calculated fitting the ¹⁷O line shapes of the F_free and F_bound species referred to the TFA peak, at various temperatures through the dynamic NMR (DNMR) LineShape Analysis module (version 1.1.2) implemented using Bruker’s Topspin 3.2, as detailed in Figure S4.

**ASSOCIATED CONTENT**

**Supporting Information**

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.inorgchem.1c03194.

NMRD and ¹⁷O data; relaxometric titration; ¹H NMR spectrum of the ligand; and Cartesian coordinates obtained with DFT calculations (PDF).

**AUTHOR INFORMATION**

Corresponding Authors

Dania Lalli — Dipartimento di Scienze e Innovazione Tecnologica, Università del Piemonte Orientale 'A.'
Apply the correction factors for inorganic chemistry. 

ACKNOWLEDGMENTS

This research was supported by the Università del Piemonte Orientale (Ricerca locale FAR2019). F.C., L.T., and M.B. acknowledge the financial support from the Ministero dell’Università e della Ricerca (PRIN 2017A2KEPL project “Rationally designed nanogens embedding paramagnetic ions as MRI probes”). This work was carried out within the framework of the COST CA15209 Action “European Network on NMR Relaxometry”.

REFERENCES

(1) Merbach, A. S.; Helm, L.; Toth, E. The Chemistry of Contrast Agents in Medical Magnetic Resonance Imaging; John Wiley & Sons, Inc., 2013.
(2) Helm, L.; Morrow, J. R.; Bond, C. J.; Carniato, F.; Botta, M.; Braun, M.; Baranay, Z.; Pujales-Paradelo, R.; Regueiro-Figueroa, M.; Esteban-Gómez, D.; Platas-Iglesias, C.; Scholl, T. J. Contrast Agents for MRI: Experimental Methods; Royal Society of Chemistry, 2018; pp 121–242.
(3) Wahser, J.; Gale, E. M.; Rodríguez-Rodriguez, A.; Caravan, P. Chemistry of MRI contrast agents: current challenges and new frontiers. Chem. Rev. 2019, 119, 957–1057.
(4) Aime, S.; Botta, M.; Terreno, E. Gd (III)-Based Contrast Agents for MRI. Advances in Inorganic Chemistry; Elsevier B.V., 2005; Vol. 57, pp 173–237.
(5) Bellin, M.-F. MR contrast agents, the old and the new. Eur. J. Radiol. 2006, 60, 314–323.
(6) Helm, L. Optimization of gadolinium-based MRI contrast agents for high magnetic-field applications. Future Med. Chem. 2010, 2, 385–396.
(7) Caravan, P.; Farrar, C. T.; Frullano, L.; Uppal, R. Influence of molecular parameters and increasing magnetic field strength on relaxation of gadolinium-and manganese-based T1 contrast agents. Contrast Media Mol. Imaging 2009, 4, 89–100.
(8) Aime, S.; Botta, M.; Crich, S. G.; Giovenzana, G.; Pagliarini, R.; Sisti, M.; Terreno, E. NMR relaxometric studies of Gd (III) complexes with heptadentate macrocyclic ligands. Magn. Reson. Chem. 1998, 36, S200–S208.
(9) Botta, M.; Aime, S.; Barge, A.; Bobba, G.; Dickens, R. S.; Parker, D.; Terreno, E. Ternary complexes between cationic Gd(III) chelates and anionic metabolites in aqueous solution: an NMR relaxometric study. Chem.—Eur. J. 2003, 9, 2102–2109.
(10) Messeri, D.; Lowe, M. P.; Parker, D.; Botta, M. A stable, high relaxation, diaqua gadolinium complex that suppresses anion and protein binding. Chem. Commun. 2001, 24, 2742–2743.
(11) Pellegrati, L.; Zhang, J.; Drahos, B.; Villette, S.; Suzenet, F.; Guillaumat, G.; Petoud, S.; Tóth, E. Pyridine-based lanthanide complexes: towards bimodal agents operating as near infrared luminescent and MRI reporters. Chem. Commun. 2008, 6591–6593.
(12) Raymond, K. N.; Pierre, V. C. Next generation, high relaxation gadolinium MRI agents. Bioconjugate Chem. 2005, 16, 3–8.
(13) Datta, A.; Raymond, K. N. Gd–hydroxypropyridonate (HOPO)-based high-relaxivity magnetic resonance imaging (MRI) contrast agents. Acc. Chem. Res. 2009, 42, 938–947.
(14) Gale, E. M.; Kenton, N.; Caravan, P. [Gd(CyPic3A)(H2O)2]−: a stable, bis (aquated) and high-relaxity Gd (III) complex. Chem. Commun. 2013, 49, 8060–8062.
(15) Aime, S.; Calabi, L.; Cavallotti, C.; Gianolio, E.; Giovenzana, G. B.; Losi, P.; Maiocchi, A.; Palmisano, G.; Sisti, M. [Gd-AAZTA]−: a new structural entry for an improved generation of MRI contrast agents. Inorg. Chem. 2004, 43, 7588–7590.
(16) Baranay, Z.; Uggeri, F.; Giovenzana, G. B.; Bényei, A.; Brücher, E.; Aime, S. Equilibrium and kinetic properties of the lanthanoids (III) and various divalent metal complexes of the heptadentate ligand AAZTA. Chem.—Eur. J. 2009, 15, 1696–1705.
(17) Vagner, A.; Gianolio, E.; Aime, S.; Maiocchi, A.; Tóth, L.; Baranay, Z.; Tei, L. High kinetic inertness of a bis-hydrated Gd-complex with a constrained AAZTA-like ligand. Chem. Commun. 2016, 52, 11235–11238.
(18) Karimi, S.; Tei, L.; Botta, M.; Helm, L. Evaluation of water exchange kinetics on [La(AAZTA)-Ph(NO3)(H2O)]3− complexes using proton Nuclear Magnetic Resonance. Inorg. Chem. 2016, 55, 6300–6307.
(19) Castelli, D. D.; Tei, L.; Carniato, F.; Aime, S.; Botta, M. [Yb(AAZTA)(H2O)]3−: an unconventional ParamEST MRI probe. Chem. Commun. 2018, 54, 2004–2007.
(20) Baranay, Z.; Castelli, D. D.; Platas-Iglesias, C.; Esteban-Gómez, D.; Bényei, A.; Tei, L.; Botta, M. Combined NMR, DFT and X-ray studies highlight structural and hydration changes of [La(AAZTA)]3− complexes across the series. Inorg. Chem. Front. 2020, 7, 795–803.
(21) Swift, T. J.; Connick, R. E. NMR-Relaxation Mechanisms of O2− in Aqueous Solutions of Paramagnetic Cations and the Lifetime of Water Molecules in the First Coordination Sphere. J. Chem. Phys. 1962, 37, 307–320.
(22) Leigh, J., Jr. Relaxation times in systems with chemical exchange: Some exact solutions. J. Magn. Reson. (1967) 1971, 4, 308–314.
(23) Powell, D. H.; Dhubhghaill, O. M. N.; Pubanz, D.; Helm, L.; Lebedev, Y. S.; Schlaepfer, W.; Merbach, A. E. Structural and dynamic study of 17O NMR, EPR, and NMRE studies of monomeric and dimeric Gd6 Complexes of interest in Magnetic Resonance Imaging: an integrated and theoretically self-consistent approach. J. Am. Chem. Soc. 1996, 118, 9333–9346.
(24) Delli Castelli, D.; Caligara, M. C.; Botta, M.; Terreno, E.; Aime, S. Combined high resolution NMR and 1H and 17O relaxometric study sheds light on the solution structure and dynamics of the lanthanide (III) complexes of HPDO3A. Inorg. Chem. 2013, 52, 7130–7138.
(25) Regueiro-Figueroa, M.; Platas-Iglesias, C. Toward the prediction of water exchange rates in magnetic resonance imaging...
contrast agents: a density functional theory study. J. Phys. Chem. A 2015, 119, 6436–6445.

(26) Peters, J. A. The reliability of parameters obtained by fitting of 1H NMRD profiles and 17O NMR data of potential GD1-based MRI contrast agents. Contrast Media Mol. Imaging 2016, 11, 160–168.

(27) Solomon, I. Relaxation Processes in a System of Two Spins. Phys. Rev. 1955, 99, 559–565.

(28) Solomon, I.; Bloemendal, N. Nuclear Magnetic Interactions in the HF Molecule. J. Chem. Phys. 1956, 25, 261–266.

(29) Bloemendal, N. Proton relaxation times in paramagnetic solutions. J. Chem. Phys. 1957, 27, 572–573.

(30) Bloemendal, N.; Morgan, L. O. Proton Relaxation Times in Paramagnetic Solutions. Effects of Electron Spin Relaxation. J. Chem. Phys. 1961, 34, 842–850.

(31) Freed, J. H. Dynamic effects of pair correlation functions on spin relaxation by translational diffusion in liquids. II. Finite jumps and independent T1 processes. J. Chem. Phys. 1978, 68, 4034–4037.

(32) Rodriguez-Rodriguez, A.; Regueiro-Figueroa, M.; Esteban-Gomez, D.; Rodriguez-Blas, T.; Patinec, V.; Tripier, R.; Tirsoc, G.; Carniato, F.; Botta, M.; Platas-Iglesias, C. Definition of the labile capping bond effect in lanthanide complexes. Chem.—Eur. J. 2017, 23, 1110–1117.

(33) Blackburn, O. A.; Kenwright, A. M.; Beer, P. D.; Faulkner, S. Axial fluoride binding by lanthanide DTMAs complexes alters the local crystal field, resulting in dramatic spectroscopic changes. Dalton Trans. 2015, 44, 19509–19517.

(34) Blackburn, O. A.; Routledge, J. D.; Jennings, L. B.; Rees, N. H.; Kenwright, A. M.; Beer, P. D.; Faulkner, S. Substituent effects on fluoride binding by lanthanide complexes of DOTA-tetraamides. Inorg. Chem. 2016, 55, 6231–6239.

(35) Payne, K. M.; Wilds, J. M.; Carniato, F.; Botta, M.; Woods, M. On Water and its Effect on the Performance of T1-Shortening gadolinium hydroxypropyridonide MRI contrast agents. Inorg. Chem. 2000, 39, 5747–5756.

(46) Frisch, M.; Trucks, G.; Schlegel, H.; Scuseria, G.; Robb, M.; Cheeseman, J.; Scalmani, G.; Barone, V.; Mennucci, B.; Petersson, G. Gaussian 09, revision D.01; Gaussian, Inc.: Wallingford, CT, 2009.

(47) Frisch, M. J.; Trucks, G. W.; Schlegel, H. B.; Scuseria, G. E.; Robb, M. A.; Cheeseman, J. R.; Scalmani, G.; Barone, V.; Petersson, G. A.; Nakatsuji, H.; Li, X.; Caricato, M.; Marenich, A. V.; Bloino, J.; Janesko, B. G.; Gomperts, R.; Mennucci, B.; Hratchian, H. P.; Ortiz, J. V.; Izmaylov, A. F.; Sonnenberg, J. L.; Williams, D.; Ding, F.; Lipparini, F.; Egidi, F.; Goings, J. F.; Peng, B.; Petrone, A.; Henderson, T.; Rasanenge, D.; Zakrzewski, V. G.; Gao, J.; Rega, N.; Zheng, G.; Liang, W.; Hada, M.; Ehara, M.; Toyota, K.; Fukuda, R.; Hasegawa, J.; Ishida, M.; Nakajima, T.; Honda, Y.; Kitao, O.; Nakai, H.; Vreven, T.; Throssell, K.; Montgomery, J. A.; Jr.; Peralta, J. E.; Ogliaro, F.; Bearpark, M. J.; Heyd, J. J.; Brothers, E. N.; Kudin, K. N.; Staroverov, V. N.; Krish, T. A.; Kobayashi, R.; Normand, J.; Raghavachari, K.; Rendell, A. P.; Burant, J. C.; Iyengar, S. S.; Tomasi, J.; Cossi, M.; Millam, J. M.; Klene, M.; Adamo, C.; Cammi, R.; Ochterski, J. W.; Martin, R. L.; Morokuma, K.; Farkas, O.; Foresman, J. B.; Fox, D. J. Gaussian 16, revision C01; Gaussian, Inc.: Wallingford, CT, 2016.

(48) Dolg, M.; Stoll, H.; Savin, A.; Preuss, H. Energy-adjusted pseudopotentials for the rare earth elements. Theor. Chim. Acta 1989, 75, 173–194.

(49) Andrae, D.; Haeusermann, U.; Dolg, M.; Stoll, H.; Preuss, H. Energy-adjusted ab initio pseudopotentials for the second and third row transition elements. Theor. Chim. Acta 1990, 77, 123–141.

(50) Rega, N.; Cossi, M.; Barone, V. Development and validation of reliable quantum mechanical approaches for the study of free radicals in solution. J. Chem. Phys. 1996, 105, 11060–11067.

(51) Lu, T.; Chen, F. Multiftn: a multifunctional wavefunction analyzer. J. Comput. Chem. 2012, 33, 580–592.

(52) Neece, F. Software update: the ORCA program system, version 4.0. Wiley Interdiscip. Rev.: Comput. Mol. Sci. 2018, 8, No. e1327.

(53) Bazyrsa, M.; Sadlej, A. J. Two-component methods of relativistic quantum chemistry: from the Douglas–Kroll approximation to the exact two-component formalism. J. Mol. Struct.: THEOCHEM 2001, 573, 181–200.

(54) Reizer, M. Douglas–Kroll–Hess Theory: a relativistic electrons-only theory for chemistry. Theor. Chem. Acc. 2006, 116, 241–252.

(55) Ahlrichs, R.; May, K. Contracted all-electron Gaussian basis sets for atoms Rb to Xe. Phys. Chem. Chem. Phys. 2000, 2, 943–945.

(56) Tao, J.; Perdew, J. P.; Staroverov, V. N.; Scuseria, G. E. Climbing the density functional ladder: Nonempirical meta–generalized gradient approximation designed for molecules and solids. Phys. Rev. Lett. 2003, 91, No. 146401.

(57) Xing, Y.; Jindal, A. K.; Regueiro-Figueroa, M.; Le Fur, M.; Kervarec, N.; Zhao, P.; Kovacs, Z.; Valencia, L.; Perez-Lourido, P.; Tripier, R. On the relationship between NMR chemical shifts of thermally polarized and hyperpolarized 89Y complexes and their solution structures. Chemistry 2016, 22, 16657.

(58) Ditchfield, R. Molecular orbital theory of magnetic shielding and magnetic susceptibility. J. Chem. Phys. 1972, 56, 5688–5691.

(59) Helgaker, T.; Jaszunski, M.; Ruud, K. Ab initio methods for the calculation of NMR shielding and indirect spin-spin coupling constants. Chem. Rev. 1999, 99, 293–352.

(60) Neese, F.; Wennmohs, F.; Hansen, A.; Becker, U. Efficient, approximate and parallel Hartree–Fock and hybrid DFT calculations. A ‘chain-of-spheres’ algorithm for the Hartree–Fock exchange. Chem. Phys. 2009, 356, 98–109.

(61) Stonyach, G. L.; Auer, A. A.; Izsak, R.; Neese, F. Self-consistent field calculation of nuclear magnetic resonance chemical shielding constants using gauge-including atomic orbitals and approximate two-electron integrals. J. Chem. Theory Comput. 2018, 14, 619–637.