Water exchange rates of water-soluble manganese(III) porphyrins of therapeutical potential†

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The activation parameters and the rate constants of the water-exchange reactions of MnIII-TE-2-PyP⁵⁺ (meso-tetrakis(N-ethylpyridinium-2-yl)porphyrin) as cationic, MnIII-TnHex-2-PyP⁵⁺ (meso-tetrakis(N-n-hexylypyridinium-2-yl)porphyrin) as sterically shielded cationic, and MnIII-TSPP⁻ (meso-tetrakis(4-sulfonatophenyl)porphyrin) as anionic manganese(III) porphyrins were determined from the temperature dependence of ¹H NMR relaxation rates. The rate constants at 298 K were obtained as 4.12 ¥ 10⁷ s⁻¹, 5.73 ¥ 10⁸ s⁻¹, and 2.74 ¥ 10⁹ s⁻¹, respectively. On the basis of the determined entropies of activation, an interchange-dissociative mechanism (I_d) was proposed for the cationic complexes (ΔS_d = ~0 J mol⁻¹ K⁻¹) whereas a limiting dissociative mechanism (D) was proposed for MnIIITnHex-2-PyP⁵⁺ complex (ΔS_d = ±79 J mol⁻¹ K⁻¹). The obtained water exchange rate of MnIIITnHex-2-PyP⁵⁺ corresponded well to the previously assumed value used by Koenig et al. (S. H. Koenig, R. D. Brown and M. Spiller, Magn. Reson. Med., 1987, 4, 52–260) to simulate the ¹H NMRD curves, therefore the measured value supports the theory developed for explaining the anomalous relaxivity of MnIIITnHex-2-PyP⁵⁺ complex. A magnitude of the obtained water-exchange rate constants further confirms the suggested inner sphere electron transfer mechanism for the reactions of the two positively charged Mn(III) porphyrins with the various biologically important oxygen and nitrogen reactive species. Due to the high biological and clinical relevance of the reactions that occur at the metal site of the studied Mn(III) porphyrins, the determination of water exchange rates advanced our insight into their efficacy and mechanism of action, and in turn should impact their further development for both diagnostic (imaging) and therapeutic purposes.

Introduction

Reactive oxygen and nitrogen species have been implicated in numerous pathological conditions including inflammation, ischemia/reperfusion, hemorrhagic shock, autoimmune diseases, neurological disorders, radiation injuries, carcinogenesis and senescence. With Mn as an innocent metal center (lacking Fenton chemistry-related toxicity), and the porphyrin cyclic structure that provides both a high metal/ligand complex stability and the rich biologically relevant redox chemistry at the Mn site, Mn(III) porphyrins and their derivatives have been successfully developed as SOD mimics. The most efficacious ortho-Mn(III) N-alkylpyridylporphyrins possess electron-withdrawing positively charged quaternized pyridyl substituents close to the metal site which, alike SOD enzymes themselves, allows: (1) the MnIII/MnII reduction potential to be in the desirable range, (around +300 mV vs. NHE), and (2) electrostatic facilitation for the approach of negatively charged superoxide, O₂⁻. In addition to the antioxidant capacity, bioavailability of the Mn porphyrins is essential for their high in vivo ability to reduce oxidative stress injuries. Thus lipophilic analogues were later synthesized that more readily cross the lipid membranes and/or the blood–brain barrier. The most potent antioxidants in vitro⁶ and in vivo,¹⁻¹¹ the ortho isomers of Mn(III) N-alkyl-, N-methoxyalkyl- and N-tri(ethylene glycol)porphyrins, are also reactive towards biologically reactive oxygen or nitrogen species other than the superoxide anion, in particular towards peroxynitrite.¹⁴,¹⁵ The high reactivity of Mn(III) porphyrins towards small molecules, coupled with their ability to undergo fast axial-ligand substitution reactions while redox cycling, presents an essential aspect of both their in vivo and in vitro solution-phase chemistry. Though of insignificant SOD-like activity,⁶,¹⁷ the anionic MnTSPP⁻ possesses considerable ability to reduce peroxynitrite (log k_red(ONOO⁻) = 5.53),¹⁸ which justifies its use in studying oxidative stress injuries, particularly from mechanistic point of view (electrostatics, anion effect with respect to its level and site of accumulation and effects observed).

The axial interaction of water with the central metal ion is of fundamental importance in both the ligand-substitution and redox processes of metalloporphyrins in aqueous solution. In the case of iron–porphyrin complexes, the rates and the mechanism of water exchange were satisfactorily elucidated.¹⁹

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Fe^{III}TSP^3− as anionic, Fe^{III}TM-4-PyP^5+ (Fe(III) meso-tetraakis(N-methylpyridinium-4-y)porphyrin) as cationic and Fe^{III}TMPS^− (Fe(III) meso-tetraakis(sulfonatomesityl)porphyrin) as sterically shielded anionic complexes were studied. For all three complexes a dissociative mode of activation was suggested on the basis of positive activation entropy and activation volume. For the third complex the limiting dissociative (D) mechanism was proposed because of the presence of bulky mesityl groups, while for the first two complexes a dissociative interchange mechanism, I_D, was proposed. The water exchange of negatively charged Fe^{III}TSP^3− is five times faster than that of Fe^{III}TM-4-PyP^5+ indicating a labilizing effect of the increased electron density on the metal ion center.

Due to high biological and clinical relevance of the reactions of Mn(III) porphyrins with reactive species that occur at the metal site, we found it important to determine the water exchange rates for Mn(III) porphyrins that would further increase our insight into their efficacy and mechanism of action, and in turn impact their further development for both diagnostic (imaging, vide infra) and therapeutic purposes. Three complexes shown in the drawing (Fig. 1) are chosen in this study. Among them Mn^{III}TE-2-PyP^5+ and Mn^{III}TnHex-2-PyP^5+ have been the most widely studied and remarkably effective in decreasing oxidative stress injuries.\(^{11-15,17}\)

![Fig. 1 The structure of the Mn(III) porphyrins studied with axially coordinated water molecules indicated.](Image)

Since as early as 1924 it has been well accepted that both porphyrins and their metal complexes have affinity for malignant tissues.\(^{20}\) This feature has led to their extensive application in nuclear medicine, X-ray and MRI. With many papers describing the possible application of Mn(III) porphyrins as tumor imaging agent,\(^{21,22}\) A. G. Schering was awarded a patent for the use of Mn(III) porphyrins as tumor enhancing MRI contrast agents. The majority of the published papers dealt with Mn^{III}TSP^3−. In 1990 Nelson and his co-workers\(^{23}\) proposed the safe use of this porphyrin as an effective liver specific MRI contrast agent. Although other manganese metalloporphyrins were also studied and tested, to date only a non-porphyrin MnDPDP, manganese(ii) dipyridoxyldiphosphate, (Mangafodipir Trisodium™, initially synthesized by Rocklage et al. in 1989) reached phase III clinical trials for the detection of liver tumors. The Mn^{III}TSP^3− has a high relaxivity, which was found to be even higher when bound to a bovine serum albumin due to a preference of Mn monomeric high spin state.\(^{24}\) The application of Mn^{III}TSP^3− in human medicine is precluded by its relatively high toxicity. The effective dose for sufficient contrast (at 2 Tesla) is reported to be only five times lower than \(LD_{50}\).\(^{25}\) Therefore, Mn^{III}TSP^3− is considered useful in radioactive (\(^{51}\)Mn) nuclear medicine rather than in MRI.\(^{26}\)

It was also recently shown that Mn^{III}TE-2-PyP^5+ and Mn^{III}TnHex-2-PyP^5+ exhibit powerful potentials as molecular MR imaging probes for the prostate cancer detection and may be particularly effective at selectively detecting multifocal disease \textit{in situ}.\(^{27}\) Their unusually high \(T_1\) relaxivities in comparison with commercially available gadolinium chelates lowers the detection limits within the micromolar range at 7 Tesla. \textit{In vivo}, Mn^{III}TE-2-PyP^5+ and Mn^{III}TnHex-2-PyP^5+ selectively accumulate in prostate tumors with 10–11-fold greater relaxation changes than in normal prostate gland.\(^{27}\)

The relaxivity of Mn^{III}TSP^3− for water protons is unexpectedly high, as reported in many papers and summarized by Schaefe and Sharp,\(^{28}\) mainly because of the high inner sphere relaxation term. The NMR dispersion curves of \(T_1\) were described as unique for both shape and values.\(^{29}\) The results could not be interpreted by Solomon–Blombergen–Morgan theory.\(^{30,31}\) The relaxation behavior of Mn^{III}TSP^3− of \(S = 2\) spin state is particularly interesting from the relaxation theory point of view. Schaefe and Sharp\(^{28}\) showed in their study of NMRD profile in liquid and gel phases that the profiles could be satisfactorily described by accounting for the effects of Brownian motion on the splitting of the \(m = \pm 2\) and on the corresponding correlation time. The most extensive study of relaxivity of Mn^{III}TSP^3− in comparison to Fe^{III}TPP^− was published by Koenig et al.\(^{29}\) In calculations 10 ns, as the average residence time of the water in the coordination sphere for both iron and manganese porphyrin complexes, was applied. Since 1980 a value of \(1.4 \times 10^7 \text{s}^{-1}\) was cited for the water exchange rate constants of \(\text{Fe}^{III}\text{TPP}^−\).\(^{32}\) but it was later redetermined as \(2 \times 10^6 \text{s}^{-1}.\) From these values the average residence time is calculated as 100–500 ns for the iron complex; the higher limit seems to be more reliable. In spite of this remarkable progress in describing magnetic aspects of anomalous relaxivity of Mn^{III}TSP^3− another important factor, \textit{i.e.} its water exchange rate constant thus far has not been reported.

Herein, we describe the temperature dependent \(^{17}\)O NMR measurements performed in order to determine the residence time of axially coordinated water molecules in Mn(III) porphyrins and the values of the water-exchange activation parameters. Mechanistic conclusions regarding the water exchange of the studied Mn(III) porphyrins could be drawn from the experimental results. We also evaluated the reflections of the water-exchange kinetic results on the intimate mechanisms of the disproportionation/dismutation of superoxide and on the reduction of some other reactive oxygen and nitrogen reactive species.

**Experimental**

**Sample preparation**

All solutions were prepared by weight. The three investigated Mn(III) porphyrin complexes were synthesized according to reported procedures.\(^{14,17}\) They were dissolved in double ion-exchanged and ultrafiltered water (MILLIPORE from Milli-Q) and the pH, measured with a combined glass electrode calibrated with buffers (Metrohm), was adjusted by adding...
measured volumes of aqueous solutions of hydrochloric acid or sodium hydroxide of known concentrations. 17O-enriched water (10 at%, Amersham, UK) was added to the solutions for the 17O NMR measurements in order to improve sensitivity. The final concentrations of the solutions of Mn(III) porphyrins were 10.0 ± 0.5 mmol per dm³. Reference solutions were prepared from tetraethylammonium chloride (TEA) (Sigma-Aldrich). The temperature and the pH of the references were the same as of the porphyrin solutions.

17O NMR

Variable-temperature 17O NMR measurements were performed using Bruker DRX-400 (9.4 T, 54.2 MHz) or DRX-360 (8.5 T, 48.8 MHz) spectrometers. Homemade air-circulating and Eurotherm temperature control units were used to stabilize the temperature. The sample temperatures were corrected using the “mehanol thermometer” method. The samples were left for 15 min before each measurement to adjust to the given temperature. The change in temperature was always less or equal to 5 K. The samples were sealed in 5 mm NMR tubes. Longitudinal relaxation rates, 1/T₁, were obtained by the inversion recovery method and transverse relaxation rates, 1/T₂, directly from the line widths.

Results and discussion

The data were analysed according to Powel and Merbach’s previously published procedure. From the measured 17O NMR relaxation rates and angular frequencies of the Mn(III) porphyrin solutions, 1/T₁, 1/T₂, and ω, and of the TEA reference solutions, 1/T₁, 1/T₂, and ω, one can calculate the reduced relaxation rates and the chemical shift, 1/T₁, 1/T₂, and Δω, which may be written as in eqn (1)–(3) where 1/T₁m and 1/T₂m are the relaxation rates in the bound water, Δωm is the chemical shift difference between the bound water and bulk water (in the absence of a paramagnetic interaction with the bulk water), P_m is the mole fraction of bound water, and τ_m is the mean residence time (or inverse exchange rate, 1/τ_m = k_ex) of water molecules in the inner-coordination sphere. The total outer-sphere contributions to the reduced relaxation rates and chemical shift are represented by 1/T₁o, 1/T₂o, and Δω_o.

\[
\frac{1}{T_{1m}} = \frac{1}{P_m} \left[ \frac{1}{T_{1} - \Delta \omega} - \frac{1}{T_{1A}} \right] = \frac{1}{T_{1m} + \tau_m} + \frac{1}{T_{1o}} \tag{1}
\]

\[
\frac{1}{T_{2m}} = \frac{1}{P_m} \left[ \frac{1}{T_{2} - \Delta \omega} - \frac{1}{T_{2A}} \right] = \left( \frac{1}{T_{2m} + \tau_m T_{2m}^2} + \Delta \omega^2 \right) \frac{1}{\tau_m T_{2m}^2} + \Delta \omega_m \tag{2}
\]

\[
\Delta \omega_m = \frac{g_s \mu_B S (S + 1) B A}{3 k_B T} \tag{5}
\]

The relaxation rates of the bound water (1/T₁, 1/T₂) display an Arrhenius-like temperature dependence (eqn (4) with i = 1, 2) where A_m is a pre-exponential parameter (in s⁻¹) and the dimensions of E_m are the energies. The temperature dependence of the difference in chemical shift between the bound water and bulk water (Δω_o) is given by eqn (5) where g_s is the isotropic Landé g factor (g_s = 2.0 for Mn³⁺), S is the electron spin (S = 4/2 for Mn³⁺), A/h is the hyperfine or scalar coupling constant (in s⁻¹), and B is the magnetic field (in T). The binding time τ_m (or reciprocal exchange rate, k_ex) of water molecules in the inner sphere is assumed to obey the Eyring equation (eqn (6)) where ΔS^‡ and ΔH^‡ are the entropy and enthalpy of activation for the exchange process. The exchange rate at 298.15 K (k_ex) can be calculated by substituting this temperature into the Eyring equation.

\[
\frac{1}{T_{1m}} = \frac{A_m e^{-\Delta E_m^* / RT}}{1 + e^{-\Delta E_m^* / RT}} \tag{4}
\]

\[
\Delta \omega_m = \frac{g_s \mu_B S (S + 1) B A}{3 k_B T} \tag{5}
\]

\[
k_ex = \frac{k_B T}{h} e^{-\Delta S^‡ / R} e^{-\Delta H^‡ / RT} \tag{6}
\]

In order to validate our methodological approach, the exchange rate at 298.15 K and the activation parameters of the water-exchange reaction of Mn(II)aq in a slightly acidic solution were determined (Zn(II)aq was the reference) and were found to be in good agreement with previously reported data (see Table 1).

In highly basic solutions of Mn(III) porphyrin (pH = 12.0), the 17O relaxation rates and angular frequencies of bulk water were found to be within the experimental error range of the measured values of the reference solutions, indicating that the Mn(III) porphyrin complexes show no measurable relaxivity enhancement. That could be the case if no water molecule is coordinated to the inner-sphere of manganese(III) ion. Therefore, the proposed dominant porphyrin species at this pH for all three porphyrins must be the one in which only an oxo ligand is axially coordinated to the central Mn(III). Furthermore, the contributions of outer-sphere water molecules to the reduced relaxation rates and the chemical shift must be negligible as well.

In slightly acidic solutions (pH = 6.0), the temperature dependence of the reduced relaxation rates and the chemical shifts for the three Mn(III) porphyrin complexes are shown in Fig. 2. The maxima observed in the temperature dependence of 1/T₂, are characteristic of the changeover from the “fast exchange” regime at high temperatures, where T_{2m} is the dominant term in eqn (2), to the “slow exchange” regime at low temperatures, where τ_m is the dominant term. The changeover between the fast and the slow exchange limits is also manifested in Δω, the maxima in 1/T₂ corresponding to the points of inflection in Δω.

The exchange rates at 298.15 K and the activation parameters of the water-exchange reactions of three Mn(III) porphyrins were determined by nonlinear Levenberg–Marquard least-squares fit to the experimental curves and are given in Table 1. The temperature dependence of the reduced relaxation times and the chemical

| Table 1 The activation parameters and the rate constants at 298.15 K for the water exchange reactions of Mn(II)aq and three different Mn(III) porphyrin complexes |
|---------------------------------|-------------------|-----------------|-----------------|
| k_ex 10⁷/s⁻¹ | ΔH^‡ / kJ mol⁻¹ | ΔS^‡ / mol⁻¹ K⁻¹ | Ref. |
| MnTnHex-2-PyP5⁺ | 4.12 ± 10⁷ | 36 ± 3 | 4 ± 10 | this work |
| MnTnHex-2-PyP5⁺ | 5.73 ± 10⁷ | 34 ± 7 | -2 ± 23 | this work |
| MnTnHex-2-PyP5⁺ | 1.0 ± 10⁷ | 54 ± 5 | 79 ± 19 | this work |
| Mn(II)aq | 2.89 ± 10⁷ | 33.1 ± 0.6 | 9 ± 2 | this work |
| Mn(II)aq | 3.1 ± 10⁷ | 33.9 | 12.2 | 37 |
| Mn(II)aq | 2.1 ± 10⁷ | 32.9 | 5.7 | 37 |
outer-sphere terms to zero in the analysis, thus seven adjustable
parameters from Table 1.

The fastest exchange rate is observed for the negatively charged
complex: $k_{ex}^\text{-}\text{mMn}^\text{III}=$ \(2.74 \times 10^4\) s\(^{-1}\).
The coordinated water molecules of the positively charged \([\text{H}_2\text{O}]_2\text{Mn}^{\text{IIIT}}\text{TE-2-PyP}\)^+ and \([\text{H}_2\text{O}]_2\text{Mn}^{\text{IIIT}}\text{TnHex-2-PyP}\)^+ complexes are definitely
less labile with $k_{ex}^\text{-}\text{mMn}^\text{III}=$ \(4.12 \times 10^4\) s\(^{-1}\) and $k_{ex}^\text{-}\text{mMn}^\text{III}=$ \(5.73 \times 10^4\) s\(^{-1}\), respectively. Although the obtained first order rate
constants for the water exchange on \([\text{H}_2\text{O}]_2\text{Mn}^{\text{IIIT}}\text{TE-2-PyP}\)^+ and \([\text{H}_2\text{O}]_2\text{Mn}^{\text{IIIT}}\text{TnHex-2-PyP}\)^+ complex are an order of
magnitude lower than the second order rate constants of both
the one-electron oxidations of superoxide in neutral solution
(log $k = 7.76$ and 7.43, respectively) and the reductions of
ONOO$^-$ and CO$_2^-$ with \([\text{H}_2\text{O}]_2\text{Mn}^{\text{IIIT}}\text{TE-2-PyP}\)^+, (log $k = 7.53$
and 8.5, respectively), under in vivo and in vitro accessible
concentrations the inner sphere electron transfer mechanisms are not
precluded by the numerical values of the rates of water-
exchange reactions. Indeed, our study on the dependence of the
$k_{ex}^\text{-}\text{Mn}^\text{III} = \text{X}^\text{-}$ of different isomers of Mn(III) N-alkylpyridylporphyrins
upon the ionic strength indicates partial inner-sphere mechanism
for $\text{O}_2^-$ dismutation.\(^6,39\)

An additional confirmation of this statement can be invoked by
a detailed analysis of the reaction mechanism. The first step in the
inner-sphere electron transfer mechanism is assumed to be a fast
ion-pair formation equilibrium between the reactants, followed by
the rate determining step of the aqua ligand substitution on the
metal-ion center. According to the Fuoss equation, the ion-pair
formation constant for the $5^-$ and 1$^-$ charged ionic reactants
are the electron density of the metal centre and the steric
factors that determine the exchange rate constants are not entirely comparable because of the solvation
of the complex ions which statistically reduces the ligation rate constant for approximately an order of magnitude. That effect
alone is sufficient for bringing the rate constants to the same order of magnitude. Furthermore, it was shown that the aromatic groups
of the porphyrins provide a hydrophobic environment that may
also change the accessibility of central metal ions to the negatively
charged species involved in the above mentioned dismutation and
reduction processes,\(^4\) as well that the charges of the complexes are
either above or below the plane, which channels the negatively-
charged reactants toward the axial positions of Mn porphyrin
more efficiently.\(^6\) Both of the latter two effects are entirely ignored in the application of the Fuoss equation.

The activation enthalpy of the water-exchange reactions
decreases from 54 kJ mol\(^{-1}\) ([\([\text{H}_2\text{O}]_2\text{Mn}^{\text{III}}\text{TSPPP}\)^+] to
36 kJ mol\(^{-1}\) ([\([\text{H}_2\text{O}]_2\text{Mn}^{\text{IIIT}}\text{TE-2-PyP}\)^+] and to 34 kJ mol\(^{-1}\)
([\([\text{H}_2\text{O}]_2\text{Mn}^{\text{IIIT}}\text{TnHex-2-PyP}\)^+], but it is the effect of the activation
entropy that causes the opposite tendency in the exchange rates.
Namely, the smaller rate constants observed for the positively
charged complexes are due to a significant decrease in
entropy that causes the opposite tendency in the exchange rates.

According to van Eldik and his co-workers\(^5\) for iron(III)
oporphyrins, the crucial factors that determine the exchange rate
are the electron density of the metal centre and the steric
decompression of the complex. The dissociative activated water
exchange is faster when the electron density on the metal center is increased, weakening the metal–axial ligand (a water molecule) bonds, and when the peripheral substituents of the porphyrin are bulky sterically hindering the complex and extruding water molecules from the inner coordination sphere.

The peripheral 4-sulfonatophenyl substituents of the MnIII(TSPP) porphyrin are strong electron-donors that increase electron density at the central Mn(III) ion, which in turn stabilizes axial water ligands and promotes the limiting dissociative water-exchange mechanism similarly to the analogous Fe(III) complex. In the case of [(H₂O)₂MnIII-TE-2-PyP]⁺ and [(H₂O)₂MnIII-TnHex-2-PyP]⁺, the substituents on the porphyrin rings are among the strongest electron withdrawing groups due to the positively charged aromatic nitrogens in ortho positions. The electron withdrawing strengthens the metal–axial water bonds which dissociation must be assisted by an incoming water molecule. The dissociation occurs simultaneously with the approach of an outer-sphere water molecule, i.e., the bond cleavage parallels the new metal–water bond formation. Thus the operative mechanism for the assisted water-exchange in [(H₂O)₂MnIII-TnHex-2-PyP]⁺ is an interchange process resulting in a significantly lower reaction rate than observed for the [(H₂O)₂MnIII(TSPP)]⁺ complex. The lower activation enthalpy for the reactions of two positively charged complexes are therefore expected, since the metal–water bond cleavage in the transition state of the interchange mechanism is a concerted process with the new bond formation. Accordingly, the overall activation enthalpy must be significantly lower for the reaction following the interchange mechanism characterized by the dissociative mechanism that is likely to be the water-exchange of [(H₂O)₂MnIII(TSPP)]⁺–. However, it should be noted that based on our results, a clear distinction between the Iₓ and Iᵧ reaction mechanisms cannot be done; but is rather based on the similarity with the water exchange kinetics of the analogous iron(III) porphyrin complexes.

A rather small difference between the exchange rates of [(H₂O)₂MnIII-TE-2-PyP]⁺ and [(H₂O)₂MnIII-TnHex-2-PyP]⁺ complexes can be attributed to the different steric decomposition of two porphyrins. The hexyl side chains hinder the metal centre somewhat more than the ethyls do; therefore, if the associative mode of activation were operating, one would expect a decrease of the exchange rate due to a somewhat more difficult approach of the entering water molecule to the central metal ion. On the other hand, if the dissociative character of the interchange mechanism is proposed, the axial water molecules are expected to be slightly easier expelled from the [(H₂O)₂MnIII-TnHex-2-PyP]⁺ complex which is in agreement with the measured data. Still, the difference between the exchange rates is too small to be taken as a definite criterion for distinction between the Iₓ and Iᵧ mechanisms.

The introduction of a porphyrin chelate around the Fe(III) center stabilizes the coordinated water molecules in the axial position by inducing electron density on the metal center. It was reported that the water exchange rate constants on the Fe(III) porphyrins are ca. 10⁴ times larger than on the hexa-aqua system. Based on the similarity between the size of the high spin Fe(III) and Mn(III) ions, and the close resemblance of the intimate water exchange mechanisms observed for these two ions, the rate constant of the water exchange on aqua manganese(III) ion can be estimated to be in the range of ~10⁻¹⁰¹ s⁻¹, depending whether our data on [(H₂O)₂MnIIITSP]⁻ are compared to the rate constant of the water exchange on [(H₂O)₂FeIIITSP]⁻ reported in ref. 32 or in ref. 19, respectively.

In addition, we calculated the residence time of the water coordinated in [(H₂O)₂MnIIITSP]⁻ at room temperature to be 36 ns. Thus the applied 10 ns in the analysis of NMRD curves of relaxivity of water protons have not been far from the reality. Therefore, the interpretation of the anomalous relaxivity of [(H₂O)₂MnIII(TSPP)]⁻, which was recently confirmed by the density functional theory calculations of anomalous spin-polarization mechanism for high-spin manganese(III) porphyrin complexes, is probably correct, or at least it is not the result of the misused water residence time.

Conclusions

Via temperature-dependent ¹⁷O NMR measurements and adequate data treatment we found that the crucial factors that determine the rates and the mechanisms of the water-exchange reactions of manganese(III) porphyrins are identical to those for iron(III) porphyrins: the charge density of the metal centre and the steric decomposition of the complex. It was also observed that the rates of the water exchange for Mn(III) porphyrins are almost a magnitude higher than for Fe(III) porphyrins of similar structure and charge. Our results provide evidence that the outer-sphere contribution to the reduced relaxations rates and the chemical shift are negligible for the studied Mn(III) porphyrin solutions.

The suggested inner sphere electron transfer mechanism for the reactions of the two positively charged Mn(III) porphyrins with various biologically important free radicals is further confirmed by the magnitude of water-exchange rate constants obtained in the present study.

It is also worth mentioning that for [MnIII(TSPP)]⁻ the measured residence time of water molecules is in good agreement with the theoretically assumed from ¹H NMRD profiles.

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Notes and references
