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Human Whole Blood $^1$H$^2$O Longitudinal Relaxation with Normal and High-Relaxivity Contrast Reagents: Influence of Trans-Cell-Membrane Water Exchange

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Abstract

Purpose—Accurate characterization of contrast reagent (CR) longitudinal relaxivity in whole blood is required to predict arterial signal intensity in contrast-enhanced MR angiography (CE-MRA). This study measured the longitudinal relaxation rate constants ($R_1$) over a range of non-protein-binding and protein-binding CR concentrations in ex vivo whole blood and plasma at 1.5 and 3.0T under physiologic arterial conditions.

Methods—Relaxivities of gadoteridol, gadobutrol, gadobenate, and gadofosveset were measured for $[CR]$ from 0 to 18 mM [mmol(CR)/L(blood)]; the latter being the upper limit of what may be expected in CE-MRA.

Results—In plasma, the $^1$H$^2$O $R_1$ $[CR]$-dependence was non-linear for gadobenate and gadofosveset secondary to CR interactions with the serum macromolecule albumin, and was well described by an analytical expression for effective 1:1 binding stoichiometry. In whole blood, the $^1$H$^2$O $R_1$ $[CR]$-dependence was markedly non-linear for all CRs, and was well-predicted by an expression for equilibrium exchange of water molecules between plasma and intracellular spaces using $a$ $priori$ parameter values only.

Conclusion—In whole blood, $^1$H$^2$O $R_1$ exhibits a non-linear relationship with $[CR]$ over 0 to 18 mM CR. The non-linearity is well described by exchange of water between erythrocyte and plasma compartments, and is particularly evident for high relaxivity CRs.

Keywords

MRI; MR Angiography; Relaxation Rate; Relaxation Time; Contrast Agents

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Introduction

Background

Increased $T_1$-weighted water proton ($^1$H$_2$O) MR signal intensity during a gadolinium based contrast reagent (GBCR) bolus first-pass results from reduction of the longitudinal relaxation time constant ($T_1$). Reduced intravascular $^1$H$_2$O $T_1$, relative to that of the surrounding tissue, makes blood appear bright in spoiled gradient echo (SPGR) images. As a first approximation, it is typically assumed that blood $^1$H$_2$O signal intensity increases with increasing GBCR concentration, and therefore with increasing GBCR dose or injection rate. However, recent studies suggest this is not always the case.

Reduced image quality with increasing contrast reagent (CR) dose has been reported for reagents with relatively high relaxivity. For example, Schneider et al. (1) demonstrated better diagnostic performance in carotid, renal, and aorto-iliac contrast-enhanced MR angiography (CE-MRA) for single dose gadobenate dimeglumine compared with double dose. Possible explanations include lower than expected longitudinal ($R_1 = 1/T_1$) or dominant transverse ($R_2^* = 1/T_2^*$) relaxation rate constants at high CR concentration, [CR]. Each of these would reduce the relative blood pool signal. Preliminary reports of high concentration longitudinal and transverse relaxation rate constants in whole blood and plasma have been reported elsewhere (2,3). This study focuses on the mechanism of longitudinal relaxation.

In some environments, the $^1$H$_2$O $R_1$ value increases linearly with [CR] as given by Equation [1]:

$$R_1 = R_{10} + r_1[CR] \quad [1]$$

where $R_{10}$ is the rate constant in the absence of CR, and $r_1$ is the CR relaxivity. Whereas Eq. [1] is valid for homogeneous saline CR solutions and for non-protein binding reagent molecules in plasma, the relation in physiologic environments is more complex and non-linear. Most reported CR relaxivities have been measured in plasma rather than whole blood, or were measured over a relatively low [CR] range. For example, Rohrer et al. (4) measured several GBCR relaxivities in bovine plasma (0.47 – 4.7 T) and canine whole blood (1.5 T) between only 0.25 and 0.5 mM. The authors correctly caution that outside this range relaxivities may vary. In another often cited study, Pintaske et al. (5) measured gadopentetate dimeglumine, gadobutrol and gadobenate dimeglumine relaxivities from 0.01 – 16 mM in plasma. While plasma $r_1$ and transverse relaxivity ($r_2$) values are desirable constructs to advance our understanding of CR effects, they do not fully describe relaxation data from whole blood.

Theory

Water molecules in whole blood exchange very rapidly in equilibrium between the extracellular plasma space and the erythrocyte intracellular compartments (Figure 1). The kinetics of this anomalously fast transcytosome process can be characterized by the very small mean intracellular residence time ($\tau_i$), approximately 10 ms (6–9). Conventional
GBCRs are confined to the plasma space, thus creating two \(^1\)H\(_2\)O environments with potentially vastly different \(^1\)H\(_2\)O relaxation rate constants.

The \(^1\)H\(_2\)O MR behavior of a two-site exchange (2SX) system is described by the Bloch-McConnell equations (10). When [CR] is sufficiently small that observed longitudinal relaxation is mono-exponential \((R_1\) is single-valued: the MR system is in the fast-exchange-regime (FXR) condition), the \(R_1\) measured is given by Equation [2]:

\[
R_1 = \frac{1}{2} \left[ R_{1i} + R_{1o} + \frac{1}{\tau_i} + \frac{1}{\tau_o} \right] - \frac{1}{2} \left[ \left( \frac{1}{\tau_i} - \frac{1}{\tau_o} \right)^2 + \frac{4}{\tau_i \tau_o} \right]^{\frac{1}{2}}
\]  

\[2a\]

\[
p_i = 1 - p_o \tag{2b}
\]

\[
\frac{\tau_i}{\tau_o} = \frac{p_i}{p_o} \tag{2c}
\]

where \(R_{1i}, \tau_i, p_i\) and \(R_{1o}, \tau_o, p_o\) are the intrinsic rate constants, characteristic water residence times, and mole fractions (“populations”) in the intracellular (i) and extracellular (o, “outside”) spaces, respectively (11,12). When the longitudinal shutter-speed \(T_1^{-1} \equiv |R_{1o} - R_{1i}| \ll 1/\tau_i + 1/\tau_o\), the system is in the fast-exchange-limit (FXL), and Eq. [2] simplifies to Equation [3]:

\[
R_1 \xrightarrow{FXL} p_i R_{1i} + p_o R_{1o} \tag{3}
\]

(Appendix 1). Thus, in the limit of effectively infinitely fast exchange (e.g. \(\tau_i \rightarrow 0\)), the observed rate constant is the population-weighted average of the \(^1\)H\(_2\)O rate constants for the two compartments. In the case of extracellular GBCR, the FXL is equivalent to considering the CR as uniformly distributed throughout the entire blood volume (erythrocyte and plasma spaces) – tantamount to assuming the cell membranes are not present – fast water exchange “short-circuits” CR compartmentalization (13). In Eq. [3], the \(p_o\) term is the dilution factor for the effective distribution of CR throughout the entire volume. If the relaxation rate constant in the plasma space \(R_{1o}\) linearly increases with \([CR_o]\) (i.e. \(R_{1o} = r_{1o}[CR_o] + R_{1o0}\) in Eq. [3]), then the dependence of blood \(R_1\) on [CR] remains linear – the characteristic of the FXL.

As the shutter-speed increases, Eq. [2] describes a non-linear [CR]-dependence. The CR concentration is implicit in Eq. [2] as \(R_{1o} = r_{1o}[CR_o] + R_{1o0}\): the FXL condition is assumed for the extracellular \(^1\)H\(_2\)O system. Thus, the shutter speed \(T_1^{-1}\) is greater when \([CR_o]\) and/or \(r_{1o}\) are greater, because \(R_{1i}\) remains unchanged. For many years, it has been almost universally assumed that monomeric Gd\(^{3+}\) chelate CRs have such small \(r_{1o}\) magnitudes that no practical \([CR_o]\) value achieved \textit{in vivo} is large enough to move the blood \(^1\)H\(_2\)O system out of the FXL condition (13–15).

However, some newer GBCR’s interact with proteins. While a CR molecule is “bound” to a macromolecule, its tumbling speed is significantly reduced and the dipolar interaction

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between Gd\(^{3+}\) unpaired electrons and water hydrogen nuclear proton spins is much more effective. This causes an increased relaxivity, \(r_{10}\) (4,16–19). In general, there can be many different CR binding sites on each protein molecule, each with its own relaxivity (17). If one makes the simplifying assumption of an effective 1:1 CR:macromolecule stoichiometry, the relaxation rate constant for the CR-containing compartmental \(^1\)H\(_2\)O is given by Equation [4] (16):

\[
R_1 = R_{10} + r_{1f}[CR] + r_{1b}[CRM] \quad [4]
\]

where \(r_{1f}\) and \(r_{1b}\) are the unbound and bound CR relaxivities, respectively, \([CR]\) is the free CR concentration, and \([CRM]\) that of macromolecule-bound CR. This is the FXL equation for the system of CR exchanging on and off the macromolecule, M. Thus, \(r_{1b}\) and \([CRM]\) are the weighted average values for the bound relaxivities and CR concentrations in the different binding sites. For the 1:1 binding simplification, the equilibrium \([CR]\) and \([CRM]\) values of Eq. [4] can be written in terms of the total CR and macromolecule concentrations, \([CR_T]\) and \([M_T]\), respectively, and the mean equilibrium constant for binding, \(K_b\) (16) (Appendix 2). The interaction of such CRs with serum albumin in the blood plasma will increase their averaged relaxivity values. Thus, there is a chance that the shutter-speed, \(T_1^{-1}\), will be increased toward the \((1/\tau_i + 1/\tau_o)\) value [which is fixed near 160 s\(^{-1}\) by the physiological temperature, metabolic state, and hematocrit (Eqs. [2b, 2c])], and the blood \(^1\)H\(_2\)O signal will depart the FXL for the FXR condition. If this departure is significant, Eqs. [1] and [3] will not be valid. Equation [2], however, spans both the FXL and FXR conditions.

To accurately characterize GBCR relaxivity in physiologic whole blood, this study measured \(R_1\) associated with two “high relaxivity” reagents that bind/interact with serum albumin: gadobenate dimeglumine (MultiHance (MH); Bracco Imaging SpA, Milan, Italy) and gadofosveset trisodium (Ablavar (AB); Lantheus Medical, Billerica, USA); and two reagents that do not bind/interact with serum albumin: gadobutrol (Gadavist (GV); Bayer Healthcare, Leverkusen, Germany) and gadoteridol (ProHance (PH); Bracco Imaging SpA, Milan, Italy). These GBCRs were mixed in \textit{ex vivo} whole human blood and plasma, and studied at both at 1.5 and 3.0 T. The measurements were performed under physiologic conditions and over a \([CR]\) range spanning that of first-pass CE-MRA. The measured relaxation rates in whole blood were compared to those predicted by water exchange (Eq. [2]). The plasma relaxation rate constants were parameterized using the 1:1 stoichiometry binding model and used as empirical input to the water exchange prediction.

**Methods**

**Sample Preparation**

All experiments were performed with local institutional review board approval. Whole human blood from a single donor was obtained from the local blood bank, and the albumin concentration was measured in the laboratory. Blood was oxygenated with 95% O\(_2\) and 5% CO\(_2\) mixture \textit{via} a bubble diffuser and placed in 6-ml HDPE tubes (13 mm I.D.). These were embedded in 2% agar gel (Cat. no. S70210A, Fisher Scientific, Pittsburgh, USA) in two separate 35-tube phantoms. This allowed simultaneous measurement of the samples in
each tray. The four contrast reagents (PH, GV, MH, and AB) were added in 10 different concentrations (1, 2, 3, 4, 5, 6, 8, 10, 14, and 18 mM, measured per volume of whole blood) each, and the tubes sealed. Contrast agents were added in their respective formulary concentrations (AB: 0.25M, PH and MH: 0.5M, and GV: 1.0M), with whole blood then added to total volume of 6 mL. The associated, relatively small (<12%) increases in extracellular volume fractions due to the varying volumes of contrast solution added to each sample were accounted for in the analysis. The (30) remaining tubes were filled with saline or blood control samples. The phantoms were warmed to 37°C for the experiments and periodically mixed by inverting the samples between acquisitions until the whole blood measurements were completed. Following these measurements, the samples were set aside without movement for six hours to allow sedimentation of the red blood cells from the plasma, and the relaxation measurements were repeated for the plasma supernatant. Blood lab analysis (including oximetry) was performed before and after the whole blood relaxation measurements.

MR acquisitions

Relaxation measurements were performed using 1.5 and 3.0 T whole body scanners (Achieva, Philips Healthcare, Best, The Netherlands) and an 8-channel SENSE head RF coil for signal reception. $R_1$ was measured using an inversion-recovery (IR) Look-Locker pulse sequence with TR = 1 s, ΔTI = 4.8 – 5.0 ms, TE = 1.9 ms, flip angle = 8°, and number of averages = 3. Additional scan parameters were: matrix, 100 × 152; voxel size 2.0 mm × 1.0 mm × 3.0 mm; field of view, 200 mm × 152 mm; bandwidth, 1230 Hz/pixel. Average signal intensities from a region-of-interest (ROI) placed within each tube were fitted with mono-exponential recovery curves using non-linear least squares (Matlab, Mathworks, Natick, MA, USA) to yield the effective $R_1$. The standard Look-Locker correction was applied to obtain the ROI $R_1$ value in the absence of RF read excitation (20).

Fitting plasma relaxivity and predicting blood $R_1$ values

Plasma $r_1$ for PH and GV were determined by linear least squares fittings of their $^1$H$_2$O $R_1$ [CR]-dependences (Eq. [1]) with fixed $R_{10}$ values of 0.8 and 0.6 s$^{-1}$ for 1.5 and 3.0 T, respectively. For MH and AB, the data were fitted with the 1:1 albumin binding model (Eqs. [4] and [A2–5]). This utilized the analytical [albumin$_T$] value, 0.497 mM, $K_b$ values of 1.5 mM$^{-1}$ for MH (19) and 12.5 mM$^{-1}$ for AB, approximately accounting for all the binding interactions reported by Caravan et al. (17), and fixed $R_{10}$ values of 0.8 and 0.6 s$^{-1}$ for 1.5 and 3.0 T, respectively.

The [CR]-dependence of whole blood $^1$H$_2$O $R_1$ was predicted using Eq. [2]: with the fitted plasma $R_{1p}$ [CR]-dependence as the [CR$_o$]-dependence of $R_{1o}$, $\tau_i = 10$ ms, the analytical hematocrit fraction (0.36; adjusted for the added extracellular volume introduced by the contrast reagent solution) as $p_i$, and $R_{1i}$ fixed at 0.8 and 0.6 s$^{-1}$ for 1.5 and 3.0 T, respectively.
Results

Laboratory Analyses of Whole Blood Samples

After oxygenation of the blood and prior to the relaxation measurements, blood lab analyses measured: pH, 7.0; albumin, 3.3 g/dL (= 0.497 mM); hematocrit, 36%; pO$_2$, 169 mmHg; and sO$_2$, 97.3%. After whole blood relaxation measurements, oximetry yielded: pO$_2$, 197 mmHg; and sO$_2$, 97.9%.

Plasma $^1$H$_2$O $R_1$ [CR]-Dependence: the plasma $^1$H$_2$O Look-Locker IR ROI data are distinctly mono-exponential (see below). The dependences of plasma $^1$H$_2$O $R_{1p}$ on PH and GV concentrations were well fitted as linear (Eq. [1]) with $r_1$ values of 4.0 and 4.4 s$^{-1}$mM$^{-1}$, respectively at 1.5 T, and 3.7 and 3.9 s$^{-1}$mM$^{-1}$, respectively at 3.0 T. The results are shown in Figure 2a – 2d: [CR] is mmol(CR)/L(plasma) in these cases. These $r_1$ data values are in good agreement with literature values for PH and GV at physiologic temperature and these field strengths (4,5).

The plasma $^1$H$_2$O $R_1$ MH and AB concentration-dependences are non-linear; especially at very small [CR] values (16–19). These were fitted using the 1:1 binding model (Eqs. [4] and [A2–5]): with [M$_T$] = 0.497 mM, $K_b = 1.5$ and 12.5 mM$^{-1}$ for MH and AB, respectively, and fixed $R_{10}$ values of 0.8 and 0.6 s$^{-1}$ for 1.5 and 3.0 T, respectively. The fittings are shown in Figs. 2e – 2h. They match the data (circles) very well, and returned $r_1/M_{T}$ values of 5.1/18.1 and 9.2/48.3 s$^{-1}$mM$^{-1}$, respectively, for MH and AB at 1.5 T, and 4.6/12.1 and 7.0/15.6 s$^{-1}$mM$^{-1}$ at 3.0 T. As averaged binding site relaxivities, these are in good agreement with literature values considering field strength differences (16–19). The dashed ordinate asymptotic straight lines are shown in Figs. 2e – 2h. These illustrate clearly the model non-linearities below approximately 1 mM CR. We did not make measurements in this small [CR] region because we wanted to cover the greater range employed for CE-MRA. However, others have reported the non-linear plasma $^1$H$_2$O behavior with small [MH] (19) and [AB] (17) values. The curvature occurs at low [CR] because [M$_T$] is only 0.5 mM: the primary albumin CR binding sites become occupied at quite small [CR] values. The instantaneous slope of the $R_1$ vs. [CR] plot represents the relaxivity ($r_1$), and it is clear that $r_1$ is [CR]-dependent in the MH and AB cases. If one studied only small [CR] values, and used the asymptotic slopes, one would obtain very large $r_1$ values indeed (dominated by $r_{1b}$) for these “high-relaxivity” reagents. The slopes at larger [CR] values are not very different from those of PH and GV, because they are dominated by $r_{1f}$. As stated above, one could analyze the plasma $^1$H$_2$O $R_1$ [CR]-dependence with more complicated binding stoichiometry models (17): our $K_b = 12.5$ mM$^{-1}$ is actually the sum of AB step-wise $K_b$ values, since we go to such large [CR] values. With these, one could extract $r_1$ values for various different M-CR binding sites. However, for our purpose, the excellent descriptions of the experimental plasma $^1$H$_2$O $R_1$ data by either the linear, non-binding model (Eq. [1]) in the cases of Figs. 2a – 2d, or the non-linear, albumin-binding model (Eq. [4]) in the cases of Figs. 2e – 2h are perfectly sufficient. Although they are quite analytical fittings, they can be considered empirical fittings for our next step.
Whole Blood $^{1}$H$_2$O $R_1$ [CR]-Dependence

Examples of whole blood $^{1}$H$_2$O Look-Locker IR ROI data are shown in Figure 3. These are results for [AB$_T$] values (mmol(AB)/L(blood)) ranging from 1 to 18 mM. Semi-log plots of the $[M_0 - M(t)/M_1]$ inversion time ($t_1$) dependence are presented. The data (symbols) are fitted with a single-exponential IR expression having $M_0$, $M_1$, and $R_{1b}$ as variables. It is important to note the linearity of the data down to the noise threshold (~0.01). This demonstrates that the recoveries are mono-exponential: the $R_{1b}$ quantities are single-valued. The fitting-returned values for $R_1$ are displayed in Fig. 2 (plasma) and Figure 4 (blood).

In blood, there are two compartments, in which the CR and water molecules are not homogeneously distributed. Thus, the Fig. 3 mono-exponential recoveries mean that the exchange MR system is in the FXL or the FXR condition. It does not reach the slow-exchange-regime (SXR) condition, even at 18 mM [AB$_T$]. If it did, one would observe curvature beginning near log[$M_0 - M(t)/M_1$] ~ $p_i$ (in this study, $p_i = 0.36$) (21); this is the definition of the SXR condition (12). Thus, one important question is whether the whole blood $^{1}$H$_2$O signal falls into the FXL or the FXR condition? As justified above, since the 2SX Eq. [2] spans FXL and FXR, one must use it to describe the [CR$_T$]-dependence of blood $^{1}$H$_2$O $R_{1b}$. The measured blood hematocrit fraction (0.36, with correction for added extracellular volume due to each contrast reagent solution) was set equal to $p_i$, $\tau_i$ was taken as 10 ms, $R_{1i}$ was fixed at 0.8 and 0.6 s$^{-1}$ for 1.5 and 3.0 T, respectively, and the fitted plasma $^{1}$H$_2$O $R_1$ vs. [CR$_T$] curves for each of the four contrast agents at each field strength (Fig. 2) were used for the [CR$_0$]-dependences of $R_{1o}$. The blood $^{1}$H$_2$O $R_{1b}$ [CR$_T$]-dependence curves predicted by Eq. [2] are shown (solid) in Fig. 4: here, [CR$_T$] is mmol(CR)/L(blood). They agree with the data (circles) very well. It is important to note that these curves are not fittings: no parameters were varied. They are predictions based on the literature parameter values and our empirical plasma data: making the results all the more compelling. With actual fittings, the agreement would be even better. The dot-dashed lines display the predicted $R_1 = f([CR_T])$ if the FXL Eq. [3] is assumed. A similar result would be predicted by allowing $\tau_i$ to become vanishingly small in Eq. [2]. These do not agree with the data in any case, and especially not for MH and AB.

Discussion

As expected, plasma $^{1}$H$_2$O relaxation rate constants increased linearly with PH and GV concentrations (Figs. 2a – 2d). The observed relaxivities (slopes) (4.0 and 4.4 s$^{-1}$mM$^{-1}$ at 1.5 T, and 3.7 and 3.9 s$^{-1}$mM$^{-1}$ at 3.0 T, respectively) are slightly lower than values (4.1 and 5.2 s$^{-1}$mM$^{-1}$ at 1.5 T, and 3.7 and 5.0 s$^{-1}$mM$^{-1}$ at 3.0 T, respectively) reported elsewhere over a smaller [CR] range (0.25 to 0.5 mM) (4). This discrepancy may be related to very weak protein interactions that are more apparent over the smaller [CR] range (22). For MH and AB, the [CR]-dependence is distinctly non-linear (Figs. 2e – 2h). This is due to stronger binding of these reagents to plasma albumin. The effective 1:1 stoichiometric binding model (Eq. [4]) provides a parameterization of the non-linear behavior that can be used for predicting whole blood $^{1}$H$_2$O $R_1$ [CR]-dependency. Determination of true binding affinities and site relaxivities requires a multiple binding stoichiometry model. This has been performed elsewhere for AB (17). The application of the effective 1:1 binding model here
yields a more abrupt “break” in the theoretical curve at \([CR_T] = [MT_T] = 0.50\) mM for AB (Figs. 2g and 2h) than for MH (Figs. 2e and 2f), as the value of \(K_b\) is much greater for AB than for MH. More accurate characterization of the binding effects would require additional data points at lower concentrations but this was not the purpose of our study.

We believe the effective 1:1 binding model provides a sufficient description over the concentration ranges evaluated.

Contrary to the general expectation (13–15), constraint to the FXL condition (Fig. 4 dot-dashed lines) does not accurately describe the \([CR]-dependence of whole blood \(\text{H}_2\text{O}\) \(R_1\). This is particularly pronounced for MH and AB. While the PH and GV FXL lines match the data up to ~5 mM, the departure occurs at even lower concentrations for MH and AB. Generally speaking, departure from the FXL condition is quite evident whenever \(R_{1p}\) surpasses ~25 s\(^{-1}\). However, the 2SX Eq. [2] with \(\tau_i = 10\) ms, \(p_i = \text{Hct}/100\), and \(R_{1o} = R_{1p}\) describes well the data for all CRs at both field strengths (Fig. 4 solid curves), even though it is not fitted to them. Since this formulation spans both the FXR and FXL conditions, it is only prudent to use Eq. [2].

Deviation from the FXL condition is more pronounced for contrast reagents with high plasma relaxivity (Fig. 4) because such agents produce larger extracellular relaxation rate constants at a given concentration. In this situation, blood \(\text{H}_2\text{O}\) \(R_1\) values become influenced by the mean intracellular water molecule pre-exchange lifetime, \(\tau_i\). It has recently been shown that \(\tau_i\) is dominated by active trans-membrane water cycling that is driven by active trans-membrane osmolyte cycling (21). This opens the exciting possibility that \(\tau_i\) is a potential cellular metabolic activity imaging biomarker (23). In the current context, high relaxivity contrast agents may provide means for further study of erythrocyte metabolism.

The question remains open as to why the red blood cell \(\tau_i\) is anomalously small (7,24). To investigate further, equation [2] could be fitted to blood \(\text{H}_2\text{O}\) \(R_1\) CR titration data (such as in Fig. 4) obtained under different metabolic conditions and/or enzyme inhibitor levels.

The non-linear \([CR]-dependence of blood \(\text{H}_2\text{O}\) \(R_1\) described here has important implications for the interpretation of image data obtained during GBCR first-pass. In particular, optimization of both CE-MRA and dynamic contrast enhanced (DCE) MRI data analyses for tissue perfusion quantification rely on accurate knowledge of the \([\text{GBCR}]\) time-course during the first-pass. In CE-MRA, deliberate CR dose or injection rate modification to achieve the blood \(\text{H}_2\text{O}\) \(R_1\) desired for optimal image quality requires knowledge of the \([CR]-dependent \(r_1\) value. Quantitative DCE-MRI requires an accurate arterial input function (\(\text{AIF} \equiv [\text{GBCR}_p(t)]\) (12).

Any non-linear \([\text{GBCR}]\)-dependence of blood \(\text{H}_2\text{O}\) \(R_1\) due to erythrocyte water exchange effects must obviously be taken into account.

To summarize, the longitudinal relaxation rate constant of the water proton MR signal was measured in human plasma and oxygenated (arterial) whole blood under physiologic conditions titrated with various concentrations (1 – 18 mM per whole blood volume) of four different GBCR’s. Because the GBCR molecules are confined to the extracellular space, the whole blood \(\text{H}_2\text{O}\) \(R_1\) was influenced by the kinetics of water exchange between the plasma...
and intracellular spaces. At high GBCR concentrations, the $R_1$ values were smaller than predicted by the fast-exchange-limit (FXL) despite the extremely rapid erythrocyte transcytolemmal water exchange. The non-linearity of the $R_1$ [GBCR]-dependence at first-pass concentrations has important implications for CE-MRA and quantitative perfusion imaging using DCE. In addition, high relaxivity GBCR’s may serve as probes for erythrocyte transcytolemmal water exchange and cellular metabolic activity.

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List of Abbreviations

- **CR**: contrast reagent
- **CE-MRA**: contrast-enhanced MR angiography
- **$R_1$**: longitudinal relaxation rate constant
- **GBCR**: gadolinium based contrast reagent
- **SPGR**: spoiled gradient echo
- **$R_2^*$**: transverse relaxation rate constant
- **$R_{10}$**: relaxation rate constant in the absence of CR
- **$r_1$**: longitudinal relaxivity
- **$r_2$**: transverse relaxivity
- **$\tau_i$**: mean intracellular residence time
- **2SX**: two-site exchange
- **FXR, FXL, SXR**: fast-exchange-regime, -limit, slow-exchange-regime
- **$R_{1i}, R_{1o}$**: intracellular (i), extracellular (o, “outside”) rate constant
- **$\tau_i, \tau_o$**: intracellular (i), extracellular (o) characteristic water residence time
- **$p_i p_o$**: intracellular (i), extracellular (o) mole fraction (“population”)
- **$T_1^{-1}$**: shutter-speed
- **$r_{1o}$**: extracellular (o) relaxivity
- **$[CR_o]$**: extracellular (o) CR concentration
- **$R_{1o0}$**: extracellular (o) rate constant in absence of CR
- **$r_{1f}$**: unbound-CR relaxivity
- **$r_{1b}$**: bound-CR relaxivity
- **$[CRM]$**: equilibrium concentration of macromolecule-bound CR
[M] equilibrium concentration of unbound macromolecules
[CR_T] total CR concentration
[M_T] total macromolecule concentration
K_b mean equilibrium constant for binding
MH gadobenate dimeglumine (MultiHance)
AB gadofosveset trisodium (Ablavar)
GV gadobutrol (Gadavist)
PH gadoteridol (ProHance)
ROI region-of-interest
R_1p relaxation rate constant in plasma
pO_2 partial pressure of O_2
sO_2 O_2 saturation
IR inversion-recovery
[AB_T] total concentration of AB
t_I inversion time
R_1b rate constant in blood
Hct hematocrit
DCE dynamic contrast enhanced
AIF arterial input function

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Appendix 1. Longitudinal relaxation in the fast-exchange-limit

When water exchange in a two-compartment system is sufficiently rapid, that the system is well-characterized by mono-exponential longitudinal relaxation, the observed longitudinal relaxation rate constant $R_1$ is given by Equation [A1-1]:

$$R_1 = \frac{1}{2} \left[ R_{1i} + R_{1o} + \frac{1}{\tau_i} + \frac{1}{\tau_o} \right] = \frac{1}{2} \left\{ (R_{1i} - R_{1o})^2 + 2(R_{1i} - R_{1o}) \left( \frac{1}{\tau_i} - \frac{1}{\tau_o} \right) + \left( \frac{1}{\tau_i} + \frac{1}{\tau_o} \right)^2 \right\}^{\frac{1}{2}} \text{[A1-1]}$$

where $R_{1i}$ and $R_{1o}$ are the intrinsic $^1\text{H}_2\text{O}$ relaxation rate constants and $\tau_i$ and $\tau_o$ are the characteristic residence times of water molecules inside and outside the compartment, respectively (10,11). This is Eq. [2] in the text, and can be rewritten as Equations [A1-2] and [A1-3].

$$R_1 = \frac{1}{2} \left[ R_{1i} + R_{1o} + \frac{1}{\tau_i} + \frac{1}{\tau_o} \right] = \frac{1}{2} \left\{ (R_{1i} - R_{1o})^2 + 2(R_{1i} - R_{1o}) \left( \frac{1}{\tau_i} - \frac{1}{\tau_o} \right) \left( \frac{1}{\tau_i} + \frac{1}{\tau_o} \right)^2 + \left( \frac{1}{\tau_i} + \frac{1}{\tau_o} \right)^2 \right\}^{\frac{1}{2}} \text{[A1-2]}$$

$$R_1 = \frac{1}{2} \left[ R_{1i} + R_{1o} + \frac{1}{\tau_i} + \frac{1}{\tau_o} \right] = \frac{1}{2} \left\{ (R_{1i} - R_{1o})^2 + 2(R_{1i} - R_{1o}) \left( \frac{1}{\tau_i} - \frac{1}{\tau_o} \right) \left( \frac{1}{\tau_i} + \frac{1}{\tau_o} \right)^2 + \left( \frac{1}{\tau_i} + \frac{1}{\tau_o} \right)^2 \right\}^{\frac{1}{2}} + \left( \frac{1}{\tau_i} + \frac{1}{\tau_o} \right) \text{[A1-3]}$$

Setting $(R_{1i} - R_{1o})^2/(1/\tau_i + 1/\tau_o)^2 = 0$, the FXL condition (that is, the difference in relaxation rate constants is much less than the water exchange rate), yields Equation [A1-4].

$$R_1 \rightarrow \frac{1}{2} \left[ R_{1i} + R_{1o} + \frac{1}{\tau_i} + \frac{1}{\tau_o} \right] = \frac{1}{2} \left\{ 2(R_{1i} - R_{1o}) \left( \frac{1}{\tau_i} - \frac{1}{\tau_o} \right) \left( \frac{1}{\tau_i} + \frac{1}{\tau_o} \right)^2 + 1 \right\} \left( \frac{1}{\tau_i} + \frac{1}{\tau_o} \right) \text{[A1-4]}$$

Then, using the relationship $(1+x)^{1/2} = 1 + \frac{1}{2} x - ..., \text{ for } |x| < 1$, also in the FXL, yields Equation [A1-5].

$$R_1 \rightarrow \frac{1}{2} \left[ R_{1i} + R_{1o} + \frac{1}{\tau_i} + \frac{1}{\tau_o} \right] = \frac{1}{2} \left\{ 1 + \frac{(R_{1i} - R_{1o}) \left( \frac{1}{\tau_i} - \frac{1}{\tau_o} \right)}{\left( \frac{1}{\tau_i} + \frac{1}{\tau_o} \right)^2} \right\} \left( \frac{1}{\tau_i} + \frac{1}{\tau_o} \right) \text{[A1-5]}$$

This can be re-written as Equations [A1-6] then [A1-7].

$$R_1 \rightarrow \frac{1}{2} \left[ R_{1i} + R_{1o} + \frac{1}{\tau_i} + \frac{1}{\tau_o} \right] = \frac{1}{2} \left\{ \frac{1}{\tau_i} + \frac{1}{\tau_o} + \frac{(R_{1i} - R_{1o}) \left( \frac{1}{\tau_i} - \frac{1}{\tau_o} \right)}{\left( \frac{1}{\tau_i} + \frac{1}{\tau_o} \right)^2} \right\} \text{[A1-6]}$$
And finally, using the equilibrium mass action relationships, \( \frac{p_i}{p_o} = \frac{\tau_i}{\tau_o} \) and \( p_i + p_o = 1 \), yields Equations [A1-8] then [A1-9].

\[
R_1 \rightarrow \frac{1}{2} \frac{R_{1i} + R_{1o}}{R_{1i} - R_{1o}} \frac{(\frac{1}{\tau_i} - \frac{1}{\tau_o})}{(\frac{1}{\tau_i} + \frac{1}{\tau_o})} \quad [A1-7]
\]

For a two-compartment system in the fast-exchange-limit, Eq. [A1-9] describes the longitudinal relaxation rate constant as the population fraction-weighted average of the rate constants for the \( ^1H_2O \) signals from the two compartments. This result is used to describe the FXL condition in the text (Eq. [3]) and the dashed-dot lines in Fig. 4.

Note that the key assumption here is: \( (R_{1i} - R_{1o})^2/(1/\tau_i + 1/\tau_o)^2 \to 0 \) [i.e., \( T_1^{-1} \ll (\tau_i^{-1} + \tau_o^{-1}) \)]. This can happen because \( |R_{1o} - R_{1i}| \to 0 \) or because \( \tau_i \) (or \( \tau_o \)) \( \to 0 \). The biological intracellular \( \tau_i \) values are fixed by the physiologic temperature, the one-dimensional cell size, and the cell membrane water permeability coefficient, \( P_w \) (21), the latter being dominated by active trans-membrane water cycling (21). It is important to realize, however, that \( \tau_i \) (and \( \tau_o \)) is always finite: the erythrocyte \( \tau_i \) (10 ms) is anomalously small. Thus, the magnetic resonance FXL condition is realized in biology only where the shutter-speed \( T_1^{-1} \) \( \to 0 \), or is very small.

**Appendix 2. The plasma 1H2O longitudinal relaxation rate constant dependence on the concentration of protein-binding contrast reagent assuming effective 1:1 binding stoichiometry**

The binding equilibrium expressions are given in Equations [A2-1] – [A2-3]:

\[
K_b = \frac{[CRM]}{[CR][M]} \quad [A2-1]
\]

\[
[M_T] = [CRM] + [M] \quad [A2-2]
\]

\[
[CR_T] = [CRM] + [CR] \quad [A2-3]
\]

where \( K_b \) is the mean binding constant, \( [CR_T] \) is the total contrast reagent concentration, \( [CRM] \) is the equilibrium concentration of macromolecule-bound contrast reagent, \( [CR] \) is the equilibrium concentration of unbound contrast reagent, \( [M_T] \) is the total macromolecule concentration, and \( [M] \) is the equilibrium concentration of unbound macromolecules.

\[ K_b [CR]^2 + (K_b ([M_T] - [CR_T]) + 1) [CR] - [CR_T] = 0 \]  \[ \text{[A2-4]} \]

\[ [CR] = \frac{K_b [CR_T] - 1 - K_b [M_T] + \sqrt{(1 - K_b [CR_T] + K_b [M_T])^2 + 4K_b [CR_T]}}{2K_b} \]  \[ \text{[A2-5]} \]

Note that only one root of [A2-4] is valid. Combining Eqs. [A2-5], [A2-3], and [4] from the text, the plasma $^1$H$_2$O relaxation rate constant can be written as a function of $K_b$, $[CR_T]$, $[M_T]$, $r_{1f}$, and $r_{1b}$. In this work, $[CR_T]$ and $[M_T]$ are known by composition. This result has been presented previously (16).
Figure 1.
Illustration of blood trans-membrane water exchange between plasma and intracellular volumes. The exchange is characterized by $p_i$, $\tau_i$, $R_{1i}$ (mole fraction, residence time, and relaxation rate constant) for water inside the erythrocyte, and $p_o$, $\tau_o$, $R_{1o}$ for water in the plasma space. In the latter environment, the $^1$H$O$ longitudinal relaxation is catalyzed by dipole interaction with the paramagnetic extracellular contrast reagent (CR) and, in the case of a protein-binding reagent, CR bound to a macromolecule (M). Both the contrast reagent binding to macromolecules and water binding to CR or CRM are reversible processes.
Figure 2.
Contrast reagent concentration [CR]-dependence of plasma $^1$H$_2$O $R_1$ ($R_{1p}$) at 37°C and two field strengths. Circles give the measured $R_1$ values, and error bars the confidence intervals for the mono-exponential fittings of inversion-recovery data. Solid curves represent linear non-binding fittings (Eq. [1]) for gadoteridol (PH) and gadobutrol (GV), and non-linear protein-binding fittings (Eq. [4]) for gadobenate (MH) and gadofosveset (AB) (parameter values given in the text). The dashed ordinate asymptotes illustrate the MH and AB non-linearity at small [CR$_T$] values, and the relaxivity ($r_1$) values (slopes) that would result from linear fitting of only low [CR$_T$] data.
Figure 3.
Examples of whole blood $^1$H$_2$O$_b$ Look-Locker ROI inversion-recovery signal intensity, $M(t_1)$, data (symbols), plotted as $(M_0 - M(t_1))/M_1$ ($M_0$ and $M_1$ represent the magnetization at equilibrium and deviation from equilibrium immediately after inversion, respectively). Results are shown for 1, 2, 4, 8, and 18 mM AB, $[AB_T]$, at 3.0 T. Note the linearity of the data down to the noise threshold (∼0.01), indicating mono-exponential recovery. The slopes of the fits correspond to $R_1$, and are displayed in Fig. 2 (plasma) and Fig. 4 (blood).
Figure 4.

[CR]-Dependence of whole human blood $^1$H$_2$O $R_1$ ($R_{1b}$) (at 37°C, sO$_2$ 98%, pH 7.0, and two different field strengths) for gadoteridol (PH), gadobutrol (GV), gadobenate (MH), and gadofosveset (AB). Circles give the measured $R_1$ values and error bars the confidence intervals for mono-exponential fittings of inversion-recovery data. The solid curves are a priori predictions using the two-site-exchange Eq. [2] (parameter values are given in the text). Not being data fittings, the curves provide remarkably accurate predictions of the measurements. The dot-dashed lines are predicted by the fast-exchange-limit (FXL) Eq. 3. There is clear, systematic, deviation from the FXL prediction, most pronounced for MH and AB.

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