## Absolute Oxygen R<sub>1e</sub> Imaging In Vivo with Pulse Electron Paramagnetic Resonance

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**Purpose:** Tissue oxygen  $(O_2)$  levels are among the most important and most quantifiable stimuli to which cells and tissues respond through inducible signaling pathways. Tumor  $O_2$  levels are major determinants of the response to cancer therapy. Developing more accurate measurements and images of tissue  $O_2$  partial pressure (p $O_2$ ), assumes enormous practical, biological, and medical importance.

**Methods:** We present a fundamentally new technique to image  $pO_2$  in tumors and tissues with pulse electron paramagnetic resonance (EPR) imaging enabled by an injected, nontoxic, triaryl methyl (trityl) spin probe whose unpaired electron's slow relaxation rates report the tissue  $pO_2$ . Heretofore, virtually all in vivo EPR  $O_2$  imaging measures  $pO_2$  with the transverse electron spin relaxation rate,  $R_{2e}$ , which is susceptible to the self-relaxation confounding  $O_2$  sensitivity.

**Results:** We found that the trityl electron longitudinal relaxation rate, R<sub>1e</sub>, is an order of magnitude less sensitive to confounding self-relaxation. R<sub>1e</sub> imaging has greater accuracy and brings EPR O<sub>2</sub> images to an absolute pO<sub>2</sub> image, within uncertainties.

**Conclusion:** R<sub>1e</sub> imaging more accurately determines oxygenation of cancer and normal tissue in animal models than has been available. It will enable enhanced, rapid, noninvasive O<sub>2</sub> images for understanding oxygen biology and the relationship of oxygenation patterns to therapy outcome in living animal systems. **Magn Reson Med 72:362–368, 2014.** © **2013 Wiley Periodicals, Inc.** 

**Key words:** oxygen; imaging; EPR; pulse; R<sub>1</sub>; spin lattice relaxation; in vivo; tumor

#### INTRODUCTION

Importance of Electron Paramagnetic Resonance  $\mathsf{O}_2$  Images

Molecular oxygen,  $O_2$ , is a crucial molecular determinant of states of human health and disease. One third of human deaths are due to diseases of  $O_2$  deprivation such as ischemic heart disease and cerebrovascular disease (1). Regions of low  $O_2$ , hypoxia, thought to be a universal characteristic of solid tumors (2), reduce the efficacy of radiation therapy in their treatment (3). The importance of organismal response to hypoxia is reflected in the hundreds of genes regulated by hypoxia inducible factor-1 (HIF-1) (4,5). As will be seen in animal model tumor images presented in this study, large  $pO_2$  gradients separate regions of high and low  $pO_2$  throughout a tumor. In vivo quantification of tissue  $pO_2$  in disease states, thus, requires imaging to fully characterize the oxygenation state of the entire tumor. Electron paramagnetic resonance (EPR)  $O_2$  images may define the mechanism by which hypoxia creates resistance to radiation, which remains elusive (2). pO2 EPR images registered with either stereotactic biopsies or gene induction images can define quantitative gene induction response to local pO<sub>2</sub> - in vivo - in states of health and disease (6).

# $\mathsf{R}_1$ Images Solve Confounding Self-relaxation of $\mathsf{O}_2$ EPR Images

Heretofore, in in vivo EPR O2 images, pO2 has been measured by means of the broadening of the EPR spectral lines of an injected spin probe predominantly through Heisenberg spin exchange (HSE) with local O<sub>2</sub> (7). The spin probe is a paramagnetic trityl molecule (8) with a single narrow EPR line whose relaxation times are long enough to enable pulse imaging. Spin packet spectral line width (LW) broadening in a continuous wave (CW) experiment is physically equivalent to increasing the transverse magnetization relaxation rate  $(R_{2e})$  in a pulse experiment.  $LW = (\gamma_e T_{2e})^{-1} = R_{2e}/\gamma_e$ ;  $\gamma_{\rm e} \equiv$  gyromagnetic ratio of the electron (9). Both R<sub>2e</sub> and  $R_{1e}$  depend linearly on pO<sub>2</sub> (8,10,11). The spin probe collision rate with extremely rapidly relaxing  $O_2$ , increases R<sub>2e</sub>, destroying the magnetization phase coherence. EPR images of  $R_{2e}$  yielded quantitative  $O_2$  images in vivo (12-15) providing approximately 1-mm resolution, and a method less susceptible to confounding biologic variation than other techniques (2).

HSE between the unpaired electrons of  $O_2$  and spin probe similarly increases the spin probe electron spinlattice or longitudinal relaxation rate  $R_{1e}$ , which measures the loss of the trityl spin magnetization energy to the lattice, predominantly to the  $O_2$  (7). However, HSE self-relaxation processes, HSE interactions between spin probes, affect  $R_{1e}$  and  $R_{2e}$  differently. HSE between two trityl molecules with differing EPR frequencies produces additional precession phase shifts that increase the  $R_{2e}$ (7). HSE does not alter the total energy of an interacting spin probe pair, and, therefore, does not affect the electron spin system energy—the longitudinal magnetization. Use of image pulse sequences dependent on  $R_{1e}$  reduces probe self-relaxation by nearly an order of magnitude. At

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FIG. 1. IRESE pulse sequence for R<sub>1e</sub> imaging

this level, essentially the only physiologic  $R_{\rm 1e}$  relaxation rate increase is due to  $O_2.$ 

#### Imaging of Trityl Concentration Is Difficult

Trityl concentration, referred to as [trityl], is related to the amplitude of the trityl signal in each voxel and might be useful to correct the effect of [trityl] self-relaxation on  $R_{2e}$ . However the accuracy in determining [trityl], using local signal amplitude is complicated by the extracellular fluid compartment distribution of trityl (16) and the variability of the extracellular volume fraction in the chaotic tumor environment. Trityl distribution volume varies from voxel to voxel making [trityl] correction difficult, blurring  $R_{2e}$ -based pO<sub>2</sub> accuracy. We find, here,  $R_{1e}$  pO<sub>2</sub> images are far less [trityl] dependent and more accurate than  $R_{2e}$ -based images.

### Imaging Longitudinal Relaxation Rate $R_{1e}$ Is Straightforward and Accurate with Pulse EPR

In saline solutions and very low [trityl], R<sub>1e</sub> is close to R<sub>2e</sub> at physiologic temperatures. Although quantitative oxygen  $R_{1e}$  EPR measurements (11,17) and images (18) have been reported these were continuous wave studies. Pulse imaging is more efficient, shows better precision and, as we will show in this study, accuracy (19). This work is the first to demonstrate direct, quantitative, pulse in vivo  $R_{\rm 1e}$  images. The long electron relaxation times of the trityl OX063 (8) with  $T_{\rm 1e}$  and  $T_{\rm 2e}$  of  $\sim$ 7  $\mu$ s at hypoxic conditions, and consequent slow relaxation rates enable pulse  $R_{\rm 1e}$  images with current RF technology. We read out R<sub>1e</sub> from each voxel using inversion recovery (IR) (20), imaging the recovering longitudinal magnetization as a function of time after the inversion pulse using electron spin echo (ESE) imaging (14,21)—a tomographic frequency-encoding method. We thus refer to inversion recovery with ESE readout as IRESE.

The inversion recovery sequence IRESE (Fig. 1), inverts the spin polarization with a  $\pi$ -pulse preceding the ESE  $\pi/2$ - $\pi$  detection sequence. The recovery is measured as a function of the delay T (Fig. 2) between the inversion pulse and the ESE detection sequence. The coefficient A in the equation in Figure 2 is less than or equal to, but near, two. Values less than 2 account for incomplete EPR line inversion. A single echo time  $\tau$  is fixed at an optimum slightly longer than the imager dead time.

This study compares  $R_{1e}$  with  $R_{2e}$  imaging using a 250 MHz pulse imager (14). We demonstrate that in vivo  $R_{1e}$  imaging shows much higher absolute  $pO_2$  accuracy, overcoming the limitations imposed by [trityl] relaxation on  $R_{2e}$  imaging.

#### METHODS

#### Imaging Methods

A pulse 250 MHz imager (14) was enhanced with a passive transmit-receive switch (22) and  $\pi/2$ - and  $\pi$ - pulses of equal duration/bandwidth (23). The imager was controlled with SpecMan4EPR v. 1.1.6 (24).

Image acquisition time and spatial resolution were fixed at, respectively, 10 minutes and 1.5 mm. Spatial resolution of the ESE and IRESE images were defined by the Rayleigh resolution criterion (14,25). We used the same pulse sequences for phantoms and for animal imaging. The standard deviation of the relaxation times/rates in homogeneous phantoms, excluding two outer layers to avoid partial volume artifacts, was used as an estimation of relaxation time/rate errors. Tables 1 and 2 present parameters of the  $R_{1e}$  and  $R_{2e}$  sequences. The repetition time, T<sub>R</sub>, for ESE sequences was adjusted to keep constant the delay between the last pulse in a sequence and the first pulse of the next sequence. For accurate  $R_{1e}$  in IRESE images the image recorded at infinite recovery time T was equated to an image recorded without an inversion pulse. For voxel intensity fitting to an exponential recovery function, a T equal to  $36 \ \mu s$  was assigned to this image.

For images, (14,26) 208 equal solid angle projections (27) were acquired; gradient  $|\vec{G}| = 15$  mT/m; field of view = 4.24 cm. A 53 baseline projections obtained with 1.5 mT lower main field were acquired with every fourth projection and subtracted from the previous four projections. Projection numbers were expanded with four-fold cubic B-spline interpolation (27) and were filtered with a three-dimensional Ram-Lak filter cutoff at one half the Nyquist frequency. Voxels with amplitude less than 15% maximum at the shortest delay were eliminated (thresholded) (14).

The system frequency band-pass function for each acquisition technique was measured using zero gradient sample signal amplitude at 50 spanning  $B_0$  fields (14). Projections were normalized using this function. Image



FIG. 2. Simulation of a single voxel's signal amplitude dependence on T in IRESE sequence.  $T_{1e}=5~\mu s$  was used.

Table 1Pulse Sequences and Imaging Protocols

Protocol	Description				
Nonimaging two-pulse ESE	$\pi/2$ -τ-π-τ-echo; 35 ns $\pi/2$ and $\pi$ RF pulses; 80 τ's logarithmically spaced between 630 ns and 14 μs; 16-step phase cycling; 70 μs repetition time; echo is integrated; τs are measured in random order.				
Nonimaging IRESE	π-T-π/2-τ-π-τ-echo; 35 ns π/2 and π RF pulses; τ = 630 ns; 16-step phase cycling for detection sequence; 80 Ts are spaced logarithmically between 0.5 μs and 32 μs; 80 μs repetition time; echo is integrated; Ts are measured in random order.				
Two-pulse ESE imaging	$\pi/2$ -τ-π-τ-echo; 35 ns $\pi/2$ and $\pi$ RF pulses; time trace 1500 points with 4 ns dwell time; 16-step phase cycling, 37472 acquisitions per τ, including phase cycling; 5 τs are spaced logarithmically between 0.63 μs and 2.4 μs; $T^{LF}_{R} = 10.37$ μs; $ G  = 15$ mT/m; imaging time 10 minutes.				
IRESE imaging	$\pi$ -T- $\pi/2$ - $\tau$ - $\pi$ -τ-echo; 35 ns $\pi/2$ and $\pi$ RF pulses; time trace 1500 points with 4 ns dwell time; $\tau = 630$ ns; 16-step phase cycling applied only for detection sequence, 9600 acquisitions per T, including phase cycling; 8 Ts are spaced logarithmically between 0.41 µs and 14 µs; T <sup>LF</sup> <sub>R</sub> = 25 µs;  G  = 15 mT/m; imaging time 10 minutes.				

analysis was performed with in-house software written using MATLAB (Mathworks, Inc., Natick, MA).

The spatial resolution of an image can be quantified by the response of an image to an abrupt step function change in sample density fitted with the Gauss error function (erf(x/ $\sqrt{2\sigma}$ )). The width of this error function  $\sigma = 1.5$  mm is an estimate of the ESE image spatial resolution. IRESE image has the same spatial resolution.

#### Spin Probe

Spin probe was the trityl OX063 radical methyl-tris[8-carboxy-2,2,6,6-tetrakis[2-hydroxyethyl]benzo[1,2-d:4,5-d']bis [1,3]dithiol-4-yl]-trisodium salt, GE Healthcare (Little Chalfont, Buckinghamshire, UK). Phantoms with 1 mM spin probe in saline were glass cylinders 9.5 mm i.d., 45 mm long. The 0%  $O_2$  sample was degassed by means of freeze-pump-thaw; the 9.3%  $O_2$  sample was bubbled with a 9.3%:90.7% oxygen-nitrogen mixture and epoxy sealed. For measuring oxygen concentration dependence, the phantoms were bubbled in situ with humidified gas mixture. Thirty minutes or more were given for oxygen equilibration during these measurements.

#### Nonimaging versus Imaging Conditions

Acquisition of spatial information requires considerable time, which, for in vivo imaging, is limited by the animal physiology. Therefore, imaging protocols have to balance the precision of the measurements with the experiment's duration. As a result, the relaxation times in imaging protocols are estimated from only five points ( $R_{2e}$ ) or eight points ( $R_{1e}$ ) on the decay curve. Such restrictions do not apply for nonimaging measurements on phantoms, which have larger numbers of delays (T) from the inversion pulse (80 versus 5  $R_{2e}$  – 8  $R_{1e}$  for imaging) and wider  $T_R$  interval (at least 5 hypoxic  $T_{1e}$  for nonimaging measurements were repeated ~ 50 times, 20 s for each measurement; data were fitted independently, and the average was presented.

#### Animal Imaging

FSa fibrosarcomas were grown on the legs of 6- to 8week-old C3H mice (HSD, Indianapolis, IN) immobilized with a partial circumference vinyl polysiloxane cast (GC Dental Products, Kasugai, Japan) (28). For Figure 4, OX063 was injected IV 0.56 mmol/kg followed by infusion at 0.63 mmol/kg/h. In Figure 5, after each  $R_{1e}/R_{2e}$  image pair additional 0.21 mmol/kg OX063 was injected, and infusion was increased by 0.35 mmol/kg/h, to a maximum of 3.85 mmol/kg/h. This was performed on three animals with consistent results. Tumor was defined by  $T_2$  enhancement in RARE MRI registered with EPR images (29).

Animal experiments followed USPHS policy, and were approved by the Institutional Animal Care and Use Committee.

#### RESULTS

# In Vitro Studies Show Reduction of $\rm R_{1e}$ Sensitivity to Confounding [trityl] Relaxation by Nearly an Order of Magnitude Relative to That of $\rm R_{2e}$

We performed studies in vitro by using the OX063 spin probe, as described in Methods (15,30,31). Sodium chloride concentration affects the relaxation times (8). Thus, normal saline (145 mM NaCl) resembling animal condition, was used as solvent (1). Temperature, was kept at 37°C, as in rodents and humans (32). These conditions affected the relaxation times and made them substantially different from the measurements on the same spin probe dissolved in water and performed at room temperature; see for example (33). In saline sodium ions allow the tri-acid OX063, whose charge is -3 at physiologic pH, to approach more closely to each other than in water, increasing [trityl] dependent transverse selfrelaxation. The dependence of OX063 relaxation rates on  $pO_2$  is given in Figure 3a. The experimental data were corrected for the effect of spin probe concentration, referred to as [trityl] by applying linear relation between

Table 2Parameters of Pulse Sequences

		RF		Transmitted
		power	Bandwidth	average
	Pulse length	[VV]	[MHz]	power [W]
2pESE	35 ns,π/2	39.6 (π/2),	8.7	0.57
(T <sub>2e</sub> )	and $\pi$	158.5 (π)		
IRESE	35 ns,π/2	39.6 (π/2),	8.7	0.42
(T <sub>1e</sub> )	and $\pi$	158.5 (π)		



FIG. 3. Relaxation rates of OX063 dissolved in saline at 37°C. **a:** [trityl] corrected dependence (concentration independent) of relaxation rates on pO<sub>2</sub>. The data were obtained using 0.46 mM sample. The relaxation rates were then extrapolated to zero [trityl] by subtracting 0.46 mM·0.165·10<sup>6</sup> s<sup>-1</sup>/mM = 76·10<sup>3</sup> s<sup>-1</sup> for R<sub>2e</sub>, and 0.46 mM·36.3·10<sup>3</sup> s<sup>-1</sup>/mM = 17·10<sup>3</sup> s<sup>-1</sup> for R<sub>1e</sub>. Best fit: R<sub>1e</sub> =  $8.9\cdot10^3 \text{ s}^{-1}$ /torr \*pO<sub>2</sub> +  $1.6\cdot10^5 \text{ s}^{-1}$ ; R<sub>2e</sub> =  $8.9\cdot10^3 \text{ s}^{-1}$ /torr \*pO<sub>2</sub> +  $1.6\cdot10^5 \text{ s}^{-1}$ ; R<sub>2e</sub> =  $8.9\cdot10^3 \text{ s}^{-1}$ /torr \*pO<sub>2</sub> +  $1.6\cdot10^5 \text{ s}^{-1}$ ; R<sub>2e</sub> =  $8.9\cdot10^3 \text{ s}^{-1}$ /torr \*pO<sub>2</sub> +  $1.6\cdot10^5 \text{ s}^{-1}$ ; R<sub>2e</sub> =  $0.165\cdot10^6 \text{ s}^{-1}$ /mM [trityl] +0.16·10<sup>5</sup> s<sup>-1</sup>; R<sub>2e</sub> =  $0.165\cdot10^6 \text{ s}^{-1}$ /mM [trityl] +0.16·10<sup>5</sup> s<sup>-1</sup>; R<sub>2e</sub> =  $0.165\cdot10^6 \text{ s}^{-1}$ /mM [trityl] +0.16·10<sup>5</sup> s<sup>-1</sup>; R<sub>2e</sub> =  $0.165\cdot10^6 \text{ s}^{-1}$ /mM [trityl] +0.16·10<sup>5</sup> s<sup>-1</sup>; R<sub>2e</sub> =  $0.165\cdot10^6 \text{ s}^{-1}$ /mM [trityl] +0.16·10<sup>5</sup> s<sup>-1</sup>; R<sub>2e</sub> =  $0.165\cdot10^6 \text{ s}^{-1}$ /mM [trityl] +0.16·10<sup>5</sup> s<sup>-1</sup>; R<sub>2e</sub> =  $0.165\cdot10^6 \text{ s}^{-1}$ /mM [trityl] +0.16·10<sup>5</sup> s<sup>-1</sup>; R<sub>2e</sub> =  $0.165\cdot10^6 \text{ s}^{-1}$ /mM [trityl] +0.16·10<sup>5</sup> s<sup>-1</sup>; R<sub>2e</sub> =  $0.165\cdot10^6 \text{ s}^{-1}$ /mM [trityl] +0.16·10<sup>5</sup> s<sup>-1</sup>; R<sub>2e</sub> =  $0.165\cdot10^6 \text{ s}^{-1}$ /mM [trityl] +0.16·10<sup>5</sup> s<sup>-1</sup>; R<sub>2e</sub> =  $0.165\cdot10^6 \text{ s}^{-1}$ /mM [trityl] +0.16·10<sup>5</sup> s<sup>-1</sup>; R<sub>2e</sub> =  $0.165\cdot10^6 \text{ s}^{-1}$ /mM [trityl] +0.16·10<sup>5</sup> s<sup>-1</sup>. The 95% confidence intervals for fit parameters are: ±  $1\cdot10^4 \text{ s}^{-1}$  for offsets, ±  $1.2\cdot10^2 \text{ s}^{-1}$ /torr for O<sub>2</sub> and ±  $7\cdot10^3 \text{ s}^{-1}$ /mM for concentration proportionality coefficients.

[trityl] and relaxation rate (Fig. 3b). It should be noted that relaxation rates of spin probe in water show weaker and nonlinear dependence on [trityl] (data not shown). The reasons for that are under investigation. There was no significant difference between the slopes of the linear dependences of  $R_{1e}$  and  $R_{2e}$  on  $pO_2$ . However, there was a dramatic difference in the slopes of the dependences of  $R_{1e}$  and  $R_{2e}$  on spin probe concentration, referred to as [trityl] (Fig. 3b).  $R_{2e}$  versus [trityl] had nearly five times greater slope than did  $R_{1e}$ .

The quantitative dependence of  $R_{2e}$  on [trityl] is much weaker than on  $pO_2$  or  $O_2$  concentration, referred to as  $[O_2]$ . At 37°C,  $[O_2]$  in saline solution is 219  $\mu$ M at 159 torr or ~1.4  $\mu$ M/torr. Thus, the sensitivity of trityl  $R_{2e}$  to  $[O_2]$ is ~6.3·10<sup>6</sup> s<sup>-1</sup>/mM, but only 0.165·10<sup>6</sup> s<sup>-1</sup>/mM for [trityl].

The results of  $R_{1e}$  imaging and relaxation measurements of phantoms with 0% and 9.3%  $O_2$  (71 torr  $pO_2$  at 37°C and atmospheric pressure) are summarized in Table 3. These  $[O_2]$  bracket relevant values from  $pO_2$  studies of hypoxia in animals (15,34). The relaxation times determined under nonimaging conditions are given in the footnotes of the table. No significant difference between average relaxation times determined under imaging and nonimaging conditions was found.

Residual dependence of  $R_{1e}$  on [trityl] is likely due to spectral diffusion, the interaction of excited spins with the other spins whose EPR frequencies are beyond the excitation bandwidth (20). Such spins may be from trityl molecules containing <sup>13</sup>C (1.109% natural abundance) producing hyperfine lines split up to 12 G from the central line (35), considerably beyond the excitation bandwidth of the pulse sequence. The results of  $R_{2e}$  measurements using ESE are given in the table for comparison. To estimate errors, the standard deviation of the relaxation times was estimated for homogeneous phantom voxels, excluding regions with edge artifacts. The  $R_{2e}$  image had a standard deviation smaller than that of the  $R_{1e}$  imaging method. However, for 0% O<sub>2</sub> the standard deviation of the  $R_{1e}$ image approached that of the  $R_{2e}$  image.

#### R<sub>1e</sub> in the Tumor of a Live Animal Shows Far More Freedom from [trityl] Relaxation

For a demonstration of the robustness of  $R_{1e}$  imaging on a live animal tumor (Fig. 4b), we compared the IRESE image slice with the same slice from the  $R_{2e}$  image (Fig. 4a), obtained from a two-pulse ESE image on the same animal 10 min later. The brighter, richly colored areas in both of the images are well oxygenated, whereas the darker, more intensely blue areas are hypoxic. The tumor

Table 3

Precision of Images Estimated as a Standard Deviation of Relaxation Times in an Image of Homogeneous Phantom<sup>a</sup>

	0% pO <sub>2</sub>			9.3% pO <sub>2</sub>		
Pulse sequence	Average T <sub>2e</sub> or T <sub>1e</sub> [µs]	Standard deviation of $T_{2e}$ or $T_{1e}$ [µs]	Standard deviation of R <sub>2e</sub> or R <sub>1e</sub> [x 10 <sup>3</sup> s <sup>-1</sup> ]	Average T <sub>2e</sub> or T <sub>1e</sub> [μs]	Standard deviation of $T_{2e}$ or $T_{1e}$ [µs]	Standard deviation of $R_{2e}$ or $R_{1e}$ [x $10^3$ s <sup>-1</sup> ]
TR measurer	ments (T <sub>2e</sub> and R <sub>2e</sub> )					
2pESE	3.1	0.2	7.9 <sup>b</sup>	1.25	0.07	46 <sup>b</sup>
SLR measurements (T <sub>1e</sub> and R <sub>1e</sub> )						
IRESE	5.0	0.3	7.9 <sup>b</sup>	1.31	0.15	90 <sup>b</sup>

<sup>a</sup>Images are obtained in 10 min on phantoms containing 1 mM OX063 dissolved in normal saline at 37°C. No [trityl] correction is applied. Nonimaging relaxation times: for 0%  $O_2$  sample  $T_{2e} = 2.99$  mgr;s (two-pulse ESE),  $T_{1e} = 4.82$  µs (IRESE); for 9.3%  $O_2$  sample  $T_{2e} = 1.24$  mgr;s (two-pulse ESE),  $T_{1e} = 1.33$  mgr;s (IRESE).

 $^{b}7.9 \cdot 10^{3} \text{ s}^{-1}$  corresponds to 0.9 torr; 46  $\cdot 10^{3} \text{ s}^{-1}$  corresponds to 5 torr; 90  $\cdot 10^{3} \text{ s}^{-1}$  corresponds to 10 torr.



FIG. 4. a-c: Sagittal slice (0.7 mm) of a mouse leg bearing a tumor. **a:**  $R_{2e}$  image from ESE. **b:**  $R_{1e}$  image from IRESE. **c:** [trityl] image from ESE. The tumor contour is obtained from a registered MRI. **d-f:** Stacked histograms of  $R_{2e}$  (d),  $R_{1e}$  (e), and the difference between  $R_{1e}$  and  $R_{2e}$  (f).

outline from a registered  $T_2$ -weighted MRI is shown in both the images as red contours. The outlines and general oxygenation patterns are very similar. However, absolute voxel  $pO_2$  values differ considerably. Figure 4c presents the [trityl] image obtained with two-pulse ESE. Figures 4d and 4e are histograms of the  $R_{2e}$  and  $R_{1e}$ relaxation rates, respectively. The red-colored histograms are from the voxels enclosed by the red tumor contours in Figures 4a–c. The blue histogram bars are from the leg tissue outside the tumor contours. They are added to the tumor histogram bars so that the ultimate height of the blue plus red histogram bars represents the total number of image voxels with the indicated relaxation rate.

The mode of the overall  $R_{1e}$  distribution from the tumor shown in Figure 4e is nearly  $0.35 \cdot 10^6 \text{ s}^{-1}$  smaller than that from the  $R_{2e}$  image. In the distributions of the  $R_{1e}$  and  $R_{2e}$ , there are two components with different modes and widths, the tumor and the residual leg area. The sharper component with the lower mode, localized primarily in the tumor area, is more clearly distinguished in the  $R_{1e}$  image. That component is associated with the narrower distribution of relaxation rates expected in the hypoxic tumor area and shows the improved performance of  $R_{1e}$  imaging. The slowest  $R_{1e}$ observed in the animal experiments (Fig. 4e) are very close to the rates found in deoxygenated phantoms. This is expected because tumors are frequently hypoxic. R<sub>2e</sub> rates, however, are  $\sim~0.35{\cdot}10^{6}~s^{-1}$  higher than  $R_{1e}$  rates (Fig. 4f). Because the relaxation rates due to different mechanisms are additive, the shift of the distribution toward higher values indicates that R<sub>2e</sub> images are more susceptible to  $O_2$ -independent relaxation than is  $R_{1e}$ . At room temperature and at a negligible [trityl] the R<sub>2e</sub> and R<sub>1e</sub> are similar.

Ultimately, the advantage of  $R_{1e}$ -based  $pO_2$  imaging over  $R_{2e}$ -based  $pO_2$  imaging lies in its reduced susceptibility to confounding variation from [trityl]-induced relaxation. We demonstrated this by artificially increasing [trityl] by increasing the rate of trityl infusion to an animal and assessing the changes in the relaxation rates in selected regions. Although the local tissue oxygenation in subvolumes of tumors is known to vary (36,37), we assumed the average tissue oxygenation during the experiment to be constant. We justify this assumption below based on the results. In the absence of changes in oxygenation all changes in the relaxation rates should be from [trityl] relaxation effect. As a surrogate for the local [trityl], we used the voxel signal intensity obtained from  $R_{2e}$  images normalized to the intensity of voxels in a 1 mM phantom. The voxel signal intensity is R<sub>2e</sub>-independent because the spin echo amplitude was extrapolated to  $\tau = 0$ . In Figure 5, the change in relaxation rates was imaged by alternating ESE and IRESE images. The [trityl], estimated from the ESE images, in animal tissues was stabilized before imaging by comparison of consecutive image intensities. Two regions of the image, one directly in the tumor volume and the other just outside the tumor volume in a well-perfused region, were compared for each image type. The regions consisted of a cube of 3  $\times$  3  $\times$  3= 27 voxels, as seen in the leftmost image which shows a sagittal slice including the regions. The  $R_{2e}$  and  $R_{1e}$  are plotted to the right of the image slice from each region. The [trityl] dependence coefficients of  $R_{2e}$  and  $R_{1e}$  are indicated along with their uncertainties determined from the scatter of values about the regression lines. The  $R_{2e}$  values, for each location, depend strongly on the apparent [trityl] while the R<sub>1e</sub> values are independent of [trityl] to within the value uncertainties shown. The effect of [trityl] on R<sub>2e</sub> is considerably stronger than the effect of cycling hypoxia (36,37) and, more importantly is observed not only in tumor but in all anatomic areas.

#### DISCUSSION

This work demonstrates that  $R_{1e}$  images have nearly an order of magnitude reduced sensitivity to self-relaxation in vivo and, thus, higher accuracy. The precision of  $R_{1e}$  and  $R_{2e}$  images is very similar, especially at low  $[O_2]$ . Figure 5 shows that  $R_{2e}$  does indeed depend on [trityl] in vivo. The trend is clearly visible because the range of [trityl] in Figure 5b is far larger than typically found in in vivo images. The slopes of  $R_{2e}$  [trityl] dependences in Figure 5 ranging from  $0.2 \cdot 10^6$  to  $0.9 \cdot 10^6$  s<sup>-1</sup>/mM are



FIG. 5.  $R_{1e}$  ( $^{\circ}$ ) and  $R_{2e}$  ( $^{\bullet}$ ) in a mouse versus [trityl] as spin probe is infused at different rates. [trityl] is obtained by normalization of animal signal intensity in each voxel on the voxel intensity of phantom with 1mM concentration. A sagittal slice of  $R_{1e}$  image of the tumorbearing leg is shown on the left. Leg profile and tumor contour are obtained from registered  $T_2$ -weighted MRI.  $R_{1e}$  and  $R_{2e}$  [trityl] dependences for two areas [1] and [2] obtained by averaging voxels in  $\sim 8 \text{ mm}^3$  cube (27 voxels) are shown. Error bars are the  $R_{1e}$  and  $R_{2e}$  standard deviations. Area 1 is located in the tumor, and area 2 is in the muscle. The slopes of [trityl] dependence and 50% confidence intervals are given in the plots. [Color figure can be viewed in the online issue, which is available at wileyonlinelibrary.com.]

considerably stronger than that found in the phantoms  $(0.165 \cdot 10^6 \text{ s}^{-1}/\text{mM})$ . In part, this may be due to our underestimation of the real spin probe concentration. The tri-acid spin probe cannot penetrate the cell membrane at physiologic pH (16). Its extracellular distribution volume can be as small as 30% of the tissue volume (38) increasing the actual concentration by a factor of three relative to that based on voxel intensity. The [trityl] dependence of R<sub>1e</sub> is five times smaller in phantoms and is undetectable in vivo, Figure 5, areas [1] and [2]. The pO<sub>2</sub>, inferred from the R<sub>1e</sub> images in each voxel are, to within the uncertainty, accurate and are the absolute measurements of tissue or tumor oxygenation.

As noted above, virtually all in vivo EPR  $O_2$  images have exploited sensitivity of  $R_{2e}$  to local pO<sub>2</sub>, except for one qualitative study (18). The present study demonstrates, for the first time, pulse EPR O<sub>2</sub> imaging in live animals that is sensitive to the spin probe electron  $R_{1e}$ . The long relaxation times of trityls and the O<sub>2</sub>-induced changes in  $R_{1e}$  are much more favorable than with the nitroxides used in the past as an infusible spin probe for  $O_{\rm 2}$  imaging in vivo (39,40).  $R_{\rm 1e}\text{-based}$   $O_{\rm 2}$  imaging is superior to  $R_{2e}$  or  $R^*_{2e}$  imaging modalities because of the reduced sensitivity to [trityl]. The [trityl] in vivo is difficult to control and, given the unknown volume in each voxel occupied by the extracellular trityl spin probe, is difficult to determine accurately. The accuracy of  $R_{1e} O_2$ images is thus improved by virtually eliminating sensitivity to trityl concentration.

#### CONCLUSIONS

 $R_{\rm 1e}$  EPR  $O_2$  imaging with trityl spin probes is feasible and has the same precision as  $R_{\rm 2e}$  imaging. The absolute accuracy of this method is superior to  $R_{\rm 2e}$  imaging because of the smaller confounding effect of environmental parameters other than  $O_2$  on  $R_{\rm 1e}$ . Therefore,  $R_{\rm 1e}$  EPR imaging should be considered as a primary method for oxymetry. Ten-minute images of a 1 mM hypoxic sample provide 1 torr  $pO_2$  resolution with 1–1.5 mm spatial resolution (14,19).

The improved accuracy of the  $O_2$  images, presented here, should further enhance animal studies of tissue and tumor oxygenation. The low radiofrequency at which these studies have been undertaken and the large animal tumors already studied with  $R_{2e} pO_2$  images (29) argue for the distinct possibility for measurements in human tumors.  $pO_2$  images can direct therapy of cancers with radiation. The enhanced accuracy of  $R_{1e} pO_2$  imaging shown here argues strongly for this technique to be incorporated in future animal and perhaps human images.

#### REFERENCES

- Longo DL, Fauci AS, Kasper DL, Hauser SL, Jameson JL, Loscalzo J, editors. Harrison's principles of internal medicine, 18th edition, Volume1. New York, NY: McGraw Hill; 2011. 3610 p.
- Tatum JL, Kelloff GJ, Gillies RJ, et al. Hypoxia: importance in tumor biology, noninvasive measurement by imaging, and value of its measurement in the management of cancer therapy. Int J Radiat Biol 2006;82:699–757.
- Brizel DM, Sibley GS, Prosnitz LR, Scher RL, Dewhirst MW. Tumor hypoxia adversely affects the prognosis of carcinoma of the head and neck. Int J Radiat Oncol Biol Phys 1997;38:285–289.
- Semenza GL. Hypoxia-inducible factor 1: master regulator of O2 homeostasis. Curr Opin Genet Dev 1998;8:588–594.
- 5. Semenza GL. HIF-1, O(2), and the 3 PHDs: how animal cells signal hypoxia to the nucleus. Cell 2001;107:1-3.
- 6. Elas M, Hleihel D, Barth ED, Haney CR, Ahn KH, Pelizzari CA, Epel B, Weichselbaum RR, Halpern HJ. Where it's at really matters: in situ in vivo vascular endothelial growth factor spatially correlates with electron paramagnetic resonance pO2 images in tumors of living mice. Mol Imaging Biol 2011;13:1107–1113.
- Molin YN, Salikhov KM, Zamaraev KI. Spin exchange: principles and applications in chemistry and biology. Goldanskii VI, Gomer R, Schafer FP, Toennies JP, editors. Berlin: Springer-Verlag; 1980. 242 p.
- Ardenkjaer-Larsen JH, Laursen I, Leunbach I, Ehnholm G, Wistrand LG, Petersson JS, Golman K. EPR and DNP properties of certain novel single electron contrast agents intended for oximetric imaging. J Magn Reson 1998;133:1–12.
- Abragam A. Principles of nuclear magnetism. Oxford: Oxford University; 1961.
- Popp CA, Hyde JS. Effects of oxygen on electron-paramagneticresonance of nitroxide spin-label probes of model membranes. J Magn Reson 1981;43:249–258.
- Subczynski WK, Hyde JS. The diffusion-concentration product of oxygen in lipid bilayers using the spin-label T1 method. Biochim Biophys Acta 1981;643:283-291.
- Halpern HJ, Yu C, Peric M, Barth E, Grdina DJ, Teicher BA. Oxymetry deep in tissues with low-frequency electron-paramagnetic-resonance. Proc Natl Acad Sci U S A 1994;91:13047–13051.
- Subramanian S, Devasahayam N, Murugesan R, Yamada K, Cook J, Taube A, Mitchell JB, Lohman JA, Krishna MC. Single-point (constant-time) imaging in radiofrequency Fourier transform electron paramagnetic resonance. Magn Reson Med 2002;48:370–379.
- Epel B, Sundramoorthy SV, Mailer C, Halpern HJ. A versatile high speed 250-MHz pulse imager for biomedical applications. Concepts Magn Reson Part B Magn Reson Eng 2008;33B:163–176.

- 15. Elas M, Williams BB, Parasca A, Mailer C, Pelizzari CA, Lewis MA, River JN, Karczmar GS, Barth ED, Halpern HJ. Quantitative tumor oxymetric images from 4D electron paramagnetic resonance imaging (EPRI): methodology and comparison with blood oxygen leveldependent (BOLD) MRI. Magn Reson Med 2003;49:682–691.
- 16. Williams BB, al Hallaq H, Chandramouli GV, Barth ED, Rivers JN, Lewis M, Galtsev VE, Karczmar GS, Halpern HJ. Imaging spin probe distribution in the tumor of a living mouse with 250 MHz EPR: correlation with BOLD MRI. Magn Reson Med 2002;47:634–638.
- Kusumi A, Subczynski WK, Hyde JS. Oxygen-transport parameter in membranes as deduced by saturation recovery measurements of spinlattice relaxation-times of spin labels. P Natl Acad Sci U S A 1982; 79:1854–1858.
- Hama Y, Matsumoto KI, Murugesan R, Subramanian S, Devasahayam N, Koscielniak JW, Hyodo F, Cook JA, Mitchell JB, Krishna MC. Continuous wave EPR oximetric imaging at 300 MHz using Radiofrequency power saturation effects. Antioxid Redox Signal 2007;9:1709–1716.
- Epel B, Sundramoorthy SV, Barth ED, Mailer C, Halpern HJ. Comparison of 250 MHz electron spin echo and continuous wave oxygen EPR imaging methods for in vivo applications. Med Phys 2011;38: 2045-2052.
- 20. Schweiger A, Jeschke G. Principles of pulse electron paramagnetic resonance. Oxford: Oxford University Press; 2001.
- Mailer C, Sundramoorthy SV, Pelizzari CA, Halpern HJ. Spin echo spectroscopic electron paramagnetic resonance imaging. Magn Reson Med 2006;55:904–912.
- 22. Sundramoorthy SV, Epel B, Mailer C, Halpern HJ. A passive dualcirculator based transmit/receive switch for use with reflection resonators in pulse electron paramagnetic resonance. Concepts Magn Reson B 2009;35B:133–138.
- 23. Quine RW, Tseytlin M, Eaton SS, Eaton GR. A very fast switched attenuator circuit for microwave and R.F. applications. Concepts Magn Reson Part B Magn Reson Eng 2010;37B:39–44.
- 24. Epel B, Gromov I, Stoll S, Schweiger A, Goldfarb D. Spectrometer manager: a versatile control software for pulse EPR spectrometers. Concepts Magn Reson B 2005;26B:36–45.
- Ahn KH, Halpern HJ. Simulation of 4D spectral-spatial EPR images. J Magn Reson 2007;187:1–9.
- Ahn KH, Halpern HJ. Spatially uniform sampling in 4-D EPR spectral-spatial imaging. J Magn Reson 2007;185:152–158.
- Ahn KH, Halpern HJ. Comparison of local and global angular interpolation applied to spectral-spatial EPR image reconstruction. Med Phys 2007;34:1047–1052.

- 28. Haney CR, Fan X, Parasca AD, Karczmar GS, Halpern HJ, Pelizzari CA. Immobilization using dental material casts facilitates accurate serial and multimodality small animal imaging. Concepts Magn Reson B 2008;33B:138–144.
- 29. Epel B, Haney CR, Hleihel D, Wardrip C, Barth ED, Halpern HJ. Electron paramagnetic resonance oxygen imaging of a rabbit tumor using localized spin probe delivery. Med Phys 2010;37:2553–2559.
- 30. Matsumoto K, Subramanian S, Devasahayam N, Aravalluvan T, Murugesan R, Cook JA, Mitchell JB, Krishna MC. Electron paramagnetic resonance imaging of tumor hypoxia: enhanced spatial and temporal resolution for in vivo pO2 determination. Magn Reson Med 2006;55:1157–1163.
- 31. Elas M, Ahn KH, Parasca A, Barth ED, Lee D, Haney C, Halpern HJ. Electron paramagnetic resonance oxygen images correlate spatially and quantitatively with oxylite oxygen measurements. Clin Cancer Res 2006;12:4209–4217.
- 32. Krogh A. The comparative physiology of respiratory mechanisms. Philadelphia: University of Philadelphia Press; 1941.
- Owenius R, Eaton GR, Eaton SS. Frequency (250 MHz to 9.2 GHz) and viscosity dependence of electron spin relaxation of triarylmethyl radicals at room temperature. J Magn Reson 2005;172:168–175.
- 34. Elas M, Bell R, Hleihel D, et al. Electron paramagnetic resonance oxygen image hypoxic fraction plus radiation dose strongly correlates with tumor cure in FSA fibrosarcomas. Int J Radiat Oncol Biol Phys 2008;71:542–549.
- Bowman MK, Mailer C, Halpern HJ. The solution conformation of triarylmethyl radicals. J Magn Reson 2005;172:254–267.
- 36. Yasui H, Matsumoto S, Devasahayam N, Munasinghe JP, Choudhuri R, Saito K, Subramanian S, Mitchell JB, Krishna MC. Low-field magnetic resonance imaging to visualize chronic and cycling hypoxia in tumor-bearing mice. Cancer Res 2010;70:6427–6436.
- Matsumoto S, Yasui H, Mitchell JB, Krishna MC. Imaging cycling tumor hypoxia. Cancer Res 2010;70:10019–10023.
- Ganong WF. Review of medical physiology. San Mateo, CA: Appleton and Lange; 1987.
- Halpern HJ, Yu C, Peric M, Barth E, Grdina DJ, Teicher BA. Oxymetry deep in tissues with low-frequency electron paramagnetic resonance. Proc Natl Acad Sci U S A 1994;91:13047–13051.
- Velan SS, Spencer RG, Zweier JL, Kuppusamy P. Electron paramagnetic resonance oxygen mapping (EPROM): direct visualization of oxygen concentration in tissue. Magn Reson Med 2000;43:804–809.