**Diagonal-SPRITE and Its Applications for In Vivo Imaging at High Field**

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**Abstract:** Many solid or semi-solid tissues have ultra-short or short T2* components that make them virtually “invisible” to standard MRI techniques. For this reason, methods such as ultra-short echo time (UTE) techniques were developed to investigate and detect some of these short and ultra-short time components. Here we report on the development and implementation of a novel version of the Single Point Ramped Imaging with T1 Enhancement (SPRITE) technique in order to image mouse models. The method, called diagonal-SPRITE, was optimized within hardware limitations such as duty cycle and gradient strength at high field. Preliminary in vivo MR T2* value of organs of normal mouse models and related T2* maps were generated. In addition, the sequence was applied to highlight stomach short T2* after the application of a long T2* suppression pulse and to achieve a positive contrast in a SPIO-labelled cells experiment.

**INTRODUCTION**

Magnetic Resonance Imaging (MRI) is a powerful imaging modality for in vivo imaging of soft tissue, providing anatomical, functional, and biochemical information as well as unique high spatial resolution. However, standard MRI techniques do not provide good images of tissues with short and ultra-short T2* such as tendon or harder tissues such a bone. To investigate biological tissues with a T2* shorter than 1ms, ultra-short echo time (UTE) techniques have been developed [1]. Several disease states are likely to benefit significantly from UTE imaging techniques, including monitoring the development of fibrotic tissue in the liver, muscle and lungs, but also in the localization of superparamagnetic iron particles (SPIO) in cell tracking experiments.

Three types of UTE techniques are currently being developed. Recently a method known as sweep imaging with Fourier transformation (SWIFT) [2,3] has been prepared as an alternative to more robust radial projection reconstruction (PR) method [4,5]. The radial technique, initially developed for lung parenchyma imaging [6,7], has also been used to investigate atherosclerosis [8], boron-11 imaging [9] and many biological tissues such as bones [10]. A further method is based on Single Point Imaging (SPI), or Single Point Ramped Imaging with T1 Enhancement (SPRITE) techniques [11,12]. The SPRITE technique developed out of the solids NMR field, where research is being done into imaging solid materials such as Teflon tubing and concrete samples of different composition [13,14]. More recently it was applied in the clinic for sodium imaging of the brain [15]. In preclinical studies, an in vivo SPI work was carried out on lungs at 4.7T [16].

In this paper we focus on the SPRITE method. SPRITE has the advantage of being a single point imaging method with no frequency encoding window; it is therefore free from image distortions due to chemical shift and susceptibility at ultra-short acquisition times [17]. Because SPRITE is a pure phase encode technique, it is intrinsically a volume technique, requiring methods such as sensitivity of the RF coil or saturation bands to limit the field-of-view (FOV). It should also be noted that scan times are generally long in order to achieve high resolution. As the FOV and data acquisition time are directly related to the performance of the scanning system, gradient coil specifications (maximum strength and duty cycle) can also be a limiting factor. This represents a limit when SPRITE is used in the clinical setting as reported in the conical-SPRITE case [15] where ultra-short T2* images were obtained in 30 min at low resolution. No clear anatomical details could be observed. On the contrary, in pre-clinical high field system, due to the use of higher gradients strength, SPRITE can achieve good spatial resolution and anatomical details are distinguishable.

As preclinical work moves to higher field strengths to achieve better SNR, new challenges are emerging. These include: distortions due to Bo field inhomogeneities, susceptibility, chemical shift and eddy current artefacts. Some of the artefacts may be partly overcome by using single point imaging techniques such as SPRITE. However, the need to scan relatively small objects (mouse organs) and the need for good resolution together with existing hardware limitations, have limited the first proposed SPRITE method of Balcom et al. [12]. In order to work well within hardware limitations, a variant of the conventional SPRITE was created. Such technique makes maximum use of the gradient coil strength and duty cycle of a given system. The technique named diagonal-SPRITE utilizes a diagonal k-space trajectory.

To demonstrate the value of this novel SPRITE approach we have used the method in three ways. First to determine T2* maps of mouse tissues in vivo. Second, it was used to achieve positive contrast from SPIO-labelled cells injected into a hind leg of in vivo mice in order to differentiate the SPIO site from MRI signal voids and blood. Third, diagonal-SPRITE was applied with a T2* long suppression pulse in...
front in order to highlight short T2* components in stomach and liver.

**THEORY**

**Pulse Sequence Development Section**

The original SPRITE trajectory suggested by Balcom et al. [12] traverses k-space on a Cartesian grid. This is achieved by stepping one of the gradients through k-space in discrete steps while the other two gradients are held constant. The three phase encoding gradients position a point in a 3D time domain matrix. SPRITE has limited spatial resolution because it quickly runs up against gradient duty cycle limitations. On our gradient set, which has a strength of 20 G/cm and a 9% duty cycle, it was found that this is primarily due to exceeding the duty cycle limit from one of the constant gradients. In order to overcome this problem a novel hardware efficient k-space trajectory was developed.

A diagonal k-space trajectory was chosen and hence the name: diagonal-SPRITE (Fig. 1A). As for the original SPRITE method, diagonal-SPRITE reconstruction adopts no k-space regridding. The diagonal trajectory samples along a Cartesian grid. In fact, if diagonal-SPRITE is to cover a matrix of 128x128 pixels, the number of applied RF pulses and consequently the number of acquired points will be 128x128=16384. *Diagonal*-SPRITE is acquiring the entire k-space maintaining the characteristics of a single point imaging technique without an undersampling approach. Thus, image reconstruction can be generated by simply Fourier transforming k-space.

The diagonal trajectory steps two gradients dynamically at all times, while the gradient applied in the third direction is stepped, as for the original SPRITE, in a standard raster method (Fig. 1B).

*Diagonal*-SPRITE makes better use of gradient duty cycle when compared to SPRITE because none of the gradients are switched on for a long time at maximum strength unless an image volume of hundreds of slices is required. Therefore, there is a reduced demand on the hardware and as a result the resting time can be shorten. A resting time is left between detection of lines of k-space, in order to not exceed gradient duty cycle limits.

For large number (hundreds) of slices, the third gradient, which is usually run at low strength, may become large enough for its duty cycle to become the limiting factor, a situation which can be mitigated using longer resting time.

**Signal Equation and T1 Effects**

The signal equation for a steady-state technique such as SPRITE is generally defined as:

\[
S(t = T_p) = M_0 e^{-t/T_{2*}} \frac{1 - e^{-TR/T_1}}{1 - \cos \alpha e^{-TR/T_1}} \sin \alpha
\]  

where \( T_p \) is the phase encode time, TR is the time period between successive RF excitation pulses, \( \alpha \) is the flip angle and \( M_0 \) is the equilibrium longitudinal magnetization. This signal equation is valid for a coherent steady-state technique. The equation can remove the T1 dependency term in two cases: when the flip angle \( \alpha \) is small or when TR is long (of the same order of T1).

In the first case (small flip angles) where the flip angle is set to 2° or lower with a pulse length of 4 \( \mu \)s, Eq. (1) turns into a T2* dependent signal equation as follows:

\[
S(t = T_p) = M_0 e^{-t/T_{2*}} \sin \alpha
\]  

In the second case (long TR), if TR is long enough to accomplish the full magnetization recovery after excitation, every single k-space point can be related to the longitudinal magnetization at the equilibrium value \( M_0 \) and therefore to Eq. (2). Long TR, used by SPI techniques, cannot be used by SPRITE techniques especially when biological samples are under study, because of long scan time issues.

Small flip angles have the added advantage of avoiding T1 blurring effects. This effect was mentioned by Mastikhin et al. [18]. From such study the maximum number of points \( N_{\text{max}} \) sampled during acquisition of SPRITE in one direction was therefore defined by:

\[
N_{\text{max}} < \frac{\pi}{(TR/T_1 - \ln(\cos \alpha))}
\]  

Eq. (3) estimates the limit where a dynamic approach to steady state begins to decrease resolution. When *diagonal*-SPRITE is run at high magnetic field on biological tissues...
with an average T1 of 1500 ms, a flip angle of 2° and a TR of 1 msec. Nmax approaches 2000. Any matrix dimension less than Nmax ensures that no T1-effects are introduced into the image.

Multi-Tp Acquisition

In order to increase the volume efficiency of diagonal-SPRITE, we developed and implemented a multi-Tp acquisition. This is not the traditional multiple point acquisition where consecutive points are collected on the same FID at longer time phase. In the technique here described, diagonal-SPRITE applies as many 2° 4 μs RF pulses as needed to collect an entire line of k-space at a given Tp (for instance 0.3 ms) and at a longer Tp. Such scheme acquires the same k-space line at a different Tp with no dead time between the two acquired lines (Fig. 2). The process can be applied at more than two different Tps.

The resting time is dictated by various factors: the maximum gradient strength used, the gradients duration, the gradient duty cycle. This method gives a reduction in temporal minimum gradient strength used, the gradients duration, the gradient strength.

In multi-Tp acquisitions, images at different phase encode time are achieved without changing spatial resolution. The gradient strength is adjusted for each Tp as dictated by the following equation [11]

\[
\text{Gradient step}_{x,y,z} = \frac{1}{\gamma \cdot Tp \cdot FOV_{x,y,z}}
\]

where \(\gamma\) is the gyromagnetic ratio. At a fixed FOV and a changeable Tp, the modified parameter is the gradient strength G on each of the three axes.

Multi-Tp method is time efficient when compared to single diagonal-SPRITE acquisitions. For instance, if acquisitions, one at ultra-short Tp (0.3 ms) and the other at longer Tp (2 ms), are run separately, the total acquisition time can rise by 30% when compare with a multi-Tp scheme. This is due to the reduction of the resting time between a multi-Tp scheme and single acquisitions scheme. A single ultra-short acquisition at a Tp of 0.3 ms requires a resting time of 600 ms between detected lines of k-space when the maximum gradient strength on our scanner is used. Further, a 2 ms Tp acquisition requires a resting time of 100 ms. On the contrary, the multi-Tp scheme requires only a small resting time, usually 100 ms, due to the optimized use of gradient strength.

![Figure 2](image.png)

**T2* Suppression Pulse**

T2* suppression pulses are based on the observation that it is less demanding to excite long T2* species than to excite short T2* species [19]. Short T2* species are not affected by low-amplitude, long-duration pulses because their transverse magnetization decays faster than it is excited. As T2* increases, the excitation rate dominates the transverse relaxation. Thus, long-T2* species are fully excited and short-T2* species are relatively unaffected.

In terms of spectral linewidths, T2* is inversely proportional to the linewidth, meaning that short T2* species have broad linewidths and long T2* species have narrow linewidths. The narrow spectrum of a long T2* species is more easily covered by the RF spectrum and thus is easily excited. The broad short T2* species spectrum requires a wide bandwidth RF pulse to be excited. A narrow bandwidth RF pulse will fully excite the longer T2* species, but only partially excite the shorter T2* species, making it an effective long-T2* suppression pulse.

The pulse used in this study is a selective 90° pulse acting as a band selective pulse. The pulse was designed as a sinc function \((\sin(x)/x)\) and presented the same characteristics as a \(\pi/2\) pulse. It excited a narrow linewidth of the entire spectrum which corresponded to long T2* components when the pulse was applied to proton resonance. The pulse was applied between 5 and 8 ms, T2* species longer than such duration appeared well suppressed. Dephasers were placed before and after the T2* suppression pulse. While long T2* species were excited and dephased, the short T2* had time to decay and thus to restore the magnetization along Bo.

Off-resonance frequency signals, such as that of fat, were not suppressed.

**METHOD**

**In Vivo Scanning**

Scans were carried out on a 9.4T horizontal bore Varian Inova scanner (Palo Alto, CA) using a 25mm ID birdcage RF coil (Magnetic Laboratories, Oxford UK), gradient strength 200 mT/m, 9% duty cycle. All in vivo procedures were conducted in accordance with the UK Animals (Scientific Procedures) Act 1986. Anesthesia was maintained at 1.5% isoflurane/oxygen mix and body temperature was maintained using warm air (SA Instruments Stony Brook NY). None of the in vivo experiments were respiratory or ECG gated.

In the first experiment, a dead mouse was used to compare diagonal-SPRITE and SPRITE in terms of resolution and acquisition time. The acquisition time (15min 16sec) was maintained the same for the two experiments. The field of view varied such that for SPRITE it was 70x70x30 m and...
for diagonal-SPRITE was 30x30x30 m. The following parameters were used for both scans: TR was 0.9 ms (0.6 ms + Tp (phased encode time)); Tp was 0.3 ms and the resting time set to 250 ms. The matrix size was 120x120x21 and flip angle of 2° 4 μs. Duty cycle limit was not exceeded either in SPRITE or diagonal-SPRITE with such parameters. However, duty cycle values for the two techniques were estimated to approach the 9% of maximum duty cycle of the 9.4T scanner.

Diagonal-SPRITE was then applied to measure T2* of pulmonary vessels, stomach, spleen and gut in vivo experiments for normal mice. The FOV was maintained at 30x30x30 m, matrix size 120x120x21. TR was 0.6 ms + Tp and the resting time was set to 100 ms. 0.6 ms was the time required for gradient stabilization in order to avoid eddy current and gradient switching transient effects which might cause image distortion. In order to increase time efficiency a multi-Tp acquisition technique was used. For measuring the T2* of stomach and spleen, Tps were: 0.3, 0.7, 1, 2.5, 3.1 ms, acquisition time of 63 min. For gut, Tps were set to: 0.3, 0.8, 1.5, 2.5, 3.8 ms, acquisition time of 70 min. For lungs, Tps were: 0.3, 0.5, 1, 1.5 ms, acquisition time of 38 min. Every organ was imaged on 3 different normal mice.

The long T2* suppression pulse experiment was carried out on stomach and liver, where diagonal-SPRITE was run at an ultra-short Tp of 0.3 ms with and without pulse preparation. The pulse was applied for 8 ms at a power of 40.9 dB and flip angle of 150°.

The parameters for the two experiments were: Tp 0.3 ms, resting time 200 ms, matrix 120x120x21, slice thickness 1.4 mm, flip angle 1.6° 4 μs. The acquisition time was 13 min and 8 min for diagonal-SPRITE with T2* suppression pulse and diagonal-SPRITE without suppression pulse, respectively.

In the in vivo SPIO-labelled cells experiment, 4 mice (Balb/c, female, Harlan U. K.) were injected with biotinylated cells labelled with anti-biotin SPIO (2x10^6, 20 μl PBS) into the hind leg and control biotinylated cells (non-SPIO labelled cells) into the contralateral hind leg. Diagonal-SPRITE used such scanning parameters: FOV 30x30x30 m, 1 average, flip angle of 2°, matrix was 120x120x21, slice thickness 1.5 mm, resting time 100 ms and Tps 0.25, 0.7, 1.4, 2.8, 3.5 ms. Total scan time of 80 minutes. A subtraction method was used to highlight the short T2* of SPIOlabelled cells.

RESULTS

Axial images of the dead mouse with different resolution (0.58 mm in Fig. (3A), 0.25 mm Fig. (3B)) were obtained using the original SPRITE technique and diagonal-SPRITE. This difference in resolution is due to the efficiency of diagonal k-space trajectory compared with standard raster trajectory in scanning the same sample for the same length of time. The lung cavity, the heart and muscle can be observed in both images, although tissue details are better identified in Fig. (3B).

An example of transverse in vivo images through the abdomen obtained with diagonal-SPRITE is shown in Fig. (4). Gut (indicated by the full arrow), part of intestine (dotted arrow), a section of spinal cord (dashed arrow) and muscle are visible. The series of images, of the same slice with 5 different Tp acquisitions, emphasizes the shortening in signal intensity of gut. The T2* of this tissue corresponded to 1.16 ± 0.24 ms. The same behavior was found in pulmonary vessels, stomach and spleen (images not shown). The exponential decay of pulmonary vessels reported a T2* of 0.35 ± 0.03 ms. The T2* decay of spleen was found to be as short as 2.07 ± 0.15 ms. The T2* related to stomach was 2.7 ± 0.3 ms.

The T2* decay of stomach and gut appears to be dependent on the type of food used to feed the mice. In this case standard rodent chow was used. In addition, it has to be noted that the background noise observed in Fig. (4) increases due to the increasing motion experienced at Tp longer than 3 ms compared to that at ultra-short Tp. Susceptibility effects start to show up in images acquired with Tps longer than 1ms as can be seen around the gut. In addition, spin dephasing effects are more pronounced at long Tps.

Fig. (3). A) Transverse image of heart of a dead mouse by the use of standard-SPRITE and B) diagonal-SPRITE. The two methods used an acquisition of 15 min 16 sec. The same scanning parameter (TR, Tp, flip angle and slice thickness of 1 mm) were used for both standard and diagonal-SPRITE. However, due to duty cycle limitations, the minimum FOV used for standard-SPRITE acquisition was 70x70x30 m while in diagonal-SPRITE case it was reduced to 30x30x30 m with visible resolution changes.

Fig. (4). Gut (full arrow) signal decay for the 5 Tp acquired by diagonal-SPRITE. From a high signal intensity at ultra-short Tp, the signal decays to the level of background noise for Tp of 3.8 ms. Spinal cord (dashed arrow) and intestine (dotted arrow) are also shown. The same slice is reported for the 5 Tp acquisitions.

Fig. (5) shows a typical T2* map through the mouse abdomen. The bright areas identify structure with ultra-short
and short T2* components such as gut or bones, while dark areas identify the long T2*.

**Fig. (5).** T2* map of gut. The bright signal correspond to short T2* tissue values. Cortical bones and part of the intestine also present short T2* not evaluated in this work. The intensity scale reports T2* value from 0 to 30 ms.

Fig. (6A) shows stomach and subcutaneous fat under the skin. The signal from the liver is completely suppressed by the application of the long T2* suppression pulse. Also muscle reported zero signal. No signal is left from spinal cord because the Tp (0.3 ms) used was not short enough to detect the ultra-short bone components.

Fig. (6B), which represents the diagonal-SPRITE reference image without the use of the long T2* suppression pulse, reports signal from all the tissues. No contrast is seen between liver and stomach. Fig. (6A) is noisier than B due to the reduced SNR. This was due to both a reduction in stomach signal intensity after applying the pulse and to the total liver signal suppression.

**Fig. (6).** A) Liver and stomach image after long T2* suppression pulse was applied to the diagonal-SPRITE acquisition. Only stomach is seen as the T2* components from liver was completely suppressed. B) Diagonal-SPRITE reference image of liver and stomach without the use of long T2* suppression pulse.

Fig. (7) reports images from the SPIO-labelled cells in vivo experiment. At the shortest UTE acquisition (T_p= 0.25 ms) the SPIO-labelled cells showed high signal intensity similar to the surrounding tissue and control cells. However, with increasing T_p the signal from labelled cells drops significantly leading to negative enhancement. The subtraction between the image acquired at ultra-short T_p (0.25 ms) and the one acquired with a longer T_p (3.5 ms) gives rise to an image with positive contrast in the area where the SPIO-labelled cells were present (Fig. 7F).

**Fig. (7).** A-E Images acquired at different T_p, from an ultra-short T_p (0.25 ms) to a short (3.5 ms): A) area of interest in the hind leg where SPIO-labelled cells were injected; in E) the area drops in signal due to the shortening in T2*; F) subtraction between image A and E produces positive contrast that localizes SPIO-labelled cells.

**DISCUSSION**

Single point imaging (SPI) developed into SPRITE in order to speed up the acquisition. In this work the raster k-space trajectory has been modified in order to optimize the method for biological models within hardware limitations. A diagonal trajectory was chosen. The diagonal trajectory offers higher spatial resolution images compared to conventional SPRITE technique for the same acquisition time. This is due to the reduced constraint of diagonal-SPRITE on duty cycle and gradient strength.

The SPIO-labelled cells were confirmed to be in the leg muscle by histological Prussian blue staining (Fig. 8). The mean T2* value of SPIO-labelled cells for the 4 mice was 2.18 ± 1.07 ms. The control cells, which were injected in the contralateral hind leg, did not show a signal drop at longer T_p acquisitions and therefore no positive contrast was generated in the subtracted image.

**Fig. (8).** Photomicrograph showing Prussian blue staining (× 400 magnification) of the hind leg muscles of a C57BL/6 mouse following intramuscular injection of SPIO-labelled cells.
FOV between different $T_p$ acquisitions. This method is by a variation in gradient amplitude maintaining the same ages acquired with a single or a multi-$T_p$ method. The small flip angle (around 2º) helps to minimize such artefacts. The small flip angle also forces the SPRITE signal to be essentially $T_2^*$ dependent. Hence, it is appropriate to investigate the $T_2^*$ blurring which might influence the result image. Because on our 9.4T the read-out is as short as 4 μs for each single point, the $T_2^*$ decay is negligible during the acquisition window. Quantitative $T_2^*$ acquisition can accurately be accomplished.

Improving image quality and spatial resolution while maintaining a short scan time are desirable features for any imaging technique. The scan time for multiple $T_p$ acquisitions, was minimized by combining diagonal-SPRITE with the multi-$T_p$ method. When the $T_p$ changes, it is compensated by a variation in gradient amplitude maintaining the same FOV between different $T_p$ acquisitions. This method is slower than the multiple $T_p$ approach introduced by Halse et al. [20] but avoids extra blurring in the images.

The multi-$T_p$ used in this work, if compared to single acquisitions, significantly reduced the total detection time of the entire experiment especially in scanner systems, such as ours, where the duty cycle is low. Despite these reductions in acquisition time, the technique remains slow due to the acquisition of the entire FOV. In SPRITE the FOV is a function of the RF coil dimensions, which means that thin slices (1 mm) cannot be acquired in a short acquisition time (a few minutes) using a Cartesian trajectory. An option in further reducing scan time using diagonal-SPRITE might be to reduce the number of acquisition points, for example not detecting the corners of k-space. Pulse preparation might also reduce the acquisition time. For instance a saturation band pulse or a long $T_2^*$ suppression pulse, positioned in front of each acquisition line, might be useful. A third way to decrease acquisition time is the use of a smaller RF coil in order to reduce the FOV.

One of the potential criticisms of the multi-$T_p$ diagonal-SPRITE is the change in TR when the same k-space line is acquired at different $T_p$s. Once again no T1 effects are introduced in any of the images because the use of a small flip angle makes such artefacts negligible. This gives to the images acquired with a single or a multi-$T_p$ method the same quality, in terms of S/N and contrast. In addition, the duty cycle limit is not exceeded because in the acquisition the gradients strength is reduced from the ultra-short $T_p$ to the longer $T_p$.

Applications

Short $T_2^*$ values of organs of live normal mice, can be useful to detect baseline information when compared to $T_2^*$ values of disease models. Similarly baseline data can be compared to oxygen enhancement in pulmonary vessels, or when fibrosis and macromolecular deposition modify areas of the brain and liver. Inflammation of tissues such as tendons can be studied with UTE without the use of contrast agents.

The images of normal mouse organs demonstrate $T_2^*$ reproducibility, good sensitivity and good resolution; however when the $T_p$ time approaches 3 ms the images start to be noisier. This is due to motion artefacts important at long $T_p$ acquisitions. To achieve the lowest total scan time and therefore reduce the risk of physiological motion, non gated acquisitions are normally executed. However, respiratory gating or ECG gating needs to be applied when the motion blurs the image to the point that fine details are not distinguishable.

Another approach in highlighting short $T_2^*$ is given by the pulse suppression experiment. As reported in the result section, liver is completely suppressed leaving visible stomach and fat. Having reported that the short $T_2^*$ of stomach is of the order of $2.7 \pm 0.3$ ms at 9.4T, it was reasonable to detect signal from such an organ after the application of a long $T_2^*$ suppression pulse. However the total absence of liver $T_2^*$ components was unexpected. There are several possible explanations. First, mouse liver might only have long $T_2^*$ components and no ultra-short or short $T_2^*$, thus validating the suppression. Second, a partial volume effect can cover up small ultra-short $T_2^*$ tissues. Hardware performance limited spatial resolution, thus the technique could not resolve tissues smaller than the pixel size. Third, the long $T_2^*$ suppression pulse might not work as efficiently as expected, with some undesirable suppression of short $T_2^*$ components. Fourth, a magnetization transfer (MT) effect between long and ultra-short $T_2^*$ components might have reduced the signal intensity of the latter below coil sensitivity.

For the SPIO-labelled cells experiment, no gating was used to limit the scanning time, but details of mouse hind legs are seen. Diagonal-SPRITE collected short $T_2^*$ which would otherwise have decayed so fast so as to show extremely low signal by standard MRI techniques. In our experiment, it was found that the signal intensity from normal tissue, SPIO-labelled cells and control cells was of similar intensities at ultra-short $T_p$ values (0.25 ms). However, at longer $T_p$ (3.5 ms) the signal of the labelled and control cells differs, in fact the latter do not drop as dramatically as the signal from SPIO-labelled cells. We took advantage of subtraction (ie subtracting an image with long $T_2^*$ components from an image with short $T_2^*$ components) in order to obtain an image with only short $T_2^*$ components. The subtraction image and the subsequent positive contrast derived from SPIO-labelled cells, helps not only to localize but also to distinguish labelled cells from signal voids and blood. The differentiation is due to the lack of signal detected from voids area at short $T_p$ as well as at long $T_p$ acquisitions. On the contrary blood would show high signal intensity in a long $T_2^*$ acquisition.

CONCLUSION

The preliminary results, reported in this work, show that diagonal-SPRITE provides valuable preclinical in vivo imaging information with improvements in spatial resolution and a reduction in acquisition time when compared to the original SPRITE method. It achieved $T_2^*$ values of organs, dem-
onstrated in T2* maps, which are not easily achievable by standard MRI techniques. It shows two ways of highlighting short T2* using subtraction between images at different TPs or adopting a T2* suppression pulse in front of the sequence.

The positive contrast which helps to identify SPIO-labelled cells might be used in the future for all experiments which need to differentiate labelled cells from voids and blood. A positive contrast might also be achieved by the use of a long T2* suppression pulse which was shown to work well on our 9.4T scanner. It is worth noting however that although the use of a long T2* suppression pulse reduces the scanning time compared to the subtraction method, it is a more complicated method to implement. Despite this difficulty, RF pulse signal suppression should be considered the method of choice to separate short and long T2* components within the same tissue.

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