INTRODUCTION

Dynamic-contrast-enhanced (DCE) MRI is widely used in order to assess vascular function, and has received much recent attention in clinical oncology trials of vascular targeted agents, amongst other applications.\(^1,2\) DCE-MRI is used to measure the uptake of an injected contrast agent over several minutes in a tumour, and appropriate modelling of these data can yield quantitative measures of clinically informative vascular properties.\(^3,4\)

In order to compute such measures of vascular function an arterial input function (AIF) is required. The AIF describes the concentration of contrast agent in the vascular space as a function of time. Ideally this would be measured for each visit in each patient to account for the natural variations in the AIF which have been observed in adults,\(^5\) and are particularly pronounced in paediatric subjects.\(^6\) An accurate measurement method for the concentration of contrast agent is, therefore, desirable but has proven challenging in practice. The use of a population-averaged AIF has been suggested\(^7\) but variations in patient AIF between visits will be erroneously propagated to changes in the tissue vascular parameters.\(^7\) Despite this, comparisons of the repeatability and sensitivity to treatment effects obtained using population and individually measured AIFs have shown that in practice using a population-averaged AIF gives better repeatability and comparable treatment sensitivity to individually measured AIFs.\(^9\) This result was obtained using data acquired with a spoiled gradient echo sequence, and it implies that AIF measurement errors obtained with this widely used sequence are on average larger than natural variations in the AIF. More recently, a functional form of a population AIF which retains the characteristics of an individualised AIF has been described but the measurement of a patient-specific AIF remains sought after.\(^10\)

An obvious way to obtain a patient-specific AIF is to monitor a blood vessel in the imaging volume.\(^11-15\)
However, in order to ensure observable uptake in the tissues of interest, the concentration of contrast agent in the feeding blood vessels is so high that it cannot be measured accurately, i.e. the relationship between high concentrations of contrast agent and signal is non-linear. To avoid this, a pre-bolus experiment\textsuperscript{16,17} has been proposed and pursued by many in the field whereby a 1/10th dose of contrast agent is delivered prior to the main examination. The observed contrast agent concentration is then scaled to approximate the AIF during the main examination.

However, imaging flowing blood remains challenging, and this is especially true for the spoiled gradient echo sequences that are widely used for DCE-MRI acquisitions. Where quantitative contrast agent concentration measures are required, \( T_1 \) quantification of the spoiled gradient echo signal must be performed, which means that the MR signal must be in a steady state.\textsuperscript{18,19} In tissues, the contrast agent dynamics are such that the steady state assumption applies with good accuracy, but this may not be true for the flow velocities observed in blood vessels typically used for measuring AIFs. This is because the blood entering the field of view during the volume excitation delivers spins that have been subject to insufficient excitations to reach a steady state. In addition, blood flow in major arteries is highly pulsatile, which further complicates the measurement.

Techniques that use the phase of the imaging signal to measure the AIF\textsuperscript{20–23} avoid many of the difficulties associated with inflow effects. However, the phase of the signal has lower precision due to reduced signal-to-noise. Foltz \textit{et al.}\textsuperscript{24} compares phase and magnitude methods and additionally notes that phase information is not readily available on most imaging units meaning that phase methods are not easily accessible to many users. Hence, this paper focuses on the magnitude methods which are more widely available.

Existing work using the magnitude signal has largely been limited to static phantoms with analytical models\textsuperscript{25} or with constant flow rates such as Roberts \textit{et al.} 2011.\textsuperscript{26} Experiments were conducted by van Schie \textit{et al.} 2018,\textsuperscript{27} using a low (in the context of that seen in major arteries), constant flow velocity and a method to potentially correct these errors was proposed. Garpebring \textit{et al.}, 2011,\textsuperscript{28} used a phantom with a Gaussian-like flow velocity waveform to examine the effects of inflow and non-ideal RF spoiling and made several recommendations, including orientating the images in parallel with the main \( B_0 \)-field, but confirmed the that flow effects remain an obstacle. Rather than correct the imaging signal, Han \textit{et al.}, 2011,\textsuperscript{29} proposed deliberately exciting spins before entering the imaging volume to reduce the signal error; work which was validated using a low velocity pulsatile phantom. A phantom that used relatively low velocity but pulsatile flow is described in the aforementioned work by Foltz \textit{et al} 2019,\textsuperscript{24} and was used to compare measurements of the passage of a bolus of contrast agent using both phase and magnitude-based DCE-MRI and DCE-CT. Ning \textit{et al.}, 2018,\textsuperscript{30} has applied and expanded much of this work in animal models to demonstrate improvements in the repeatability of pharmacokinetic modelling.

What is missing from the literature, which would help guide future directions in this area, is a empirical characterisation of the measurement error introduced by flow across the entire range of flow velocities seen in major human arteries. Furthermore, the impact of physiologically realistic pulsatile flow seen over the course of the cardiac cycle needs to be understood. The work presented here (1) examines the impact of flow on \( T_1 \) quantification across the range of physiologically realistic flow rates, (2) directly assesses the effect of pulsatile flow, similar to that seen in the major arteries that are commonly used to measure arterial input functions, and (3) uses a validated virtual phantom to examine the likely errors that would be seen \textit{in vivo} with pulsatile flow during the passage of a contrast agent causing rapidly changing \( T_1 \).

**METHODS AND MATERIALS**

The aim of this work is to not only quantify the measurement error introduced by flow on quantitative imaging of \( T_1 \), but to assess the impact of these errors within the context of measurement of an AIF for DCE-MRI. Specifically, the suggestion of a prebolus experiment whereby the \( T_1 \) of blood in a major artery is dynamically measured is explored. Assessing the implications of these errors is further complicated as the \( T_1 \) of the blood is rapidly changing. This study uses a physical phantom in order to accurately reproduce a range of both constant and physiologically realistic pulsatile waveforms in a tube of similar diameter to a large artery such as the carotid or femoral. These arteries usually run in parallel with the main \( B_0 \)-field and physical phantom design and subsequent image orientations were chosen to mimic this (and follow the recommendations in Garpebring \textit{et al}, 2011\textsuperscript{28}). Although the imaging signal, and subsequent \( T_1 \) measurement is expected to vary across the image space, i.e. be more accurate at more efferent locations, to maximise signal and reduce noise, we focus on a section of the image space at the centre of the physical phantom, which is in the middle of the coil. This approach enables us to validate the virtual phantom, which can be used to describe errors in different locations and with different imaging parameters. However, \textit{in vivo} the \( T_1 \) of the pulsatile blood flow is also changing as the bolus of contrast agent passes. This is very difficult to repeatedly simulate with a physical phantom. Hence, the results from the physical phantom with a constant underlying \( T_1 \) were used to validate a virtual phantom whose underlying \( T_1 \) could be easily changed to simulate those expected in a pre-bolus experiment.

This section first describes the construction and set up of the physical phantom, the flow waveforms produced by the phantom, and the methodology used to establish a baseline \( T_1 \) measurement of the blood-mimicking fluid (BMF). Then, the imaging sequences and image processing methodologies are defined, these being designed to replicate clinically feasible practices whilst isolating the errors introduced by flow.

Finally, the design of the virtual phantom is discussed and its use described.

**Physical phantom**

The physical phantom was designed to reproduce the scenario of blood flowing through a major artery in a patient. A tube of similar
diameter to that of a major artery was chosen and attached to a pump capable of delivering both continuous flow velocities and pulsatile waveforms similar to those seen in vivo. The practicalities of building such a phantom in a clinical imaging centre placed a number of constraints on the design and these will be discussed below. A diagram of the experimental setup is shown in Figure 1. An MR compatible, programmable syringe pump was used to generate either constant flow or a repeated pulsatile waveform in a closed loop of tubing containing the manufacturer supplied and required BMF consisting of a 4:5 mix of glycerol and water which has a similar viscosity to blood. The pump was controlled by a PC, and both were sited outside the magnet room with the connecting tubes passing through a waveguide. A 6m long length of flexible plastic tubing (6.35 mm internal diameter, similar to the femoral artery) formed a closed loop between the input and output ports of the programmable pump. A section of the tubing was held straight inside a short length of rigid Perspex tubing which passed between three loading bottles containing CuSO₄ solution (1 litre each, consisting of 770 mg CuSO₄, 1 ml arquad (1%), 0.15 ml H₂SO₄, in 23 cm tall, 9.5 cm diameter plastic bottles) placed inside the head coil at the isocentre of the imaging unit (Figure 2).

In vivo, the blood flowing into the imaging volume in a major artery has recently passed through the heart twice and traversed the lungs. Therefore, it is assumed that it has been in the main B₀-field of the magnet for a long length of time relative to its T₁ and enters the imaging volume close to fully polarised. Its spin history outside the imaging volume is difficult to know but it is expected that it will have received some excitations from the B₁ field, but not enough to have reached saturation. In contrast, the fluid entering the imaging volume in the physical phantom will have flowed from the pump outside the imaging room. To more closely replicate the B₀ and B₁ fields that blood in a patient experiences, an early iteration of the physical phantom had a coil of pipe at the centre of the magnet through which the BMF flowed before entering the imaging volume. However, such a long length of pipe of this diameter (6.35 mm) generated such resistance to flow that the pump was unusable. Therefore, the long coil of pipe was replaced with a short length of wider diameter pipe (19 mm), allowing the pump to run up to 100 cm.s⁻¹ in the 6.35mm pipe where measurements are made, and reducing the flow velocity in the 19mm diameter section of pipe. In terms of the magnetic field history of the BMF, the 19 mm pipe can be thought of as simulating the passage of the blood through the lungs and heart before entering the imaging volume, i.e. experiencing the B₀ field but outside the imaging volume. It is calculated that the BMF experienced greater than 4 x T₁ periods before being imaged.

The continuous flow velocities tested were chosen to span the physiological range of instantaneous flow velocities seen in major arteries: 0, 3, 6, 9, 13, 16, 32, 47, 63, 79, 95 cm. s⁻¹ (unfortunately the image data for the four higher velocities were lost for the 2D sequence). Figure 3 shows the pre-programmed flow waveforms used in this study which are representative of those seen in humans in the femoral (red) and carotid (blue) arteries. The mean flow velocity of these waveforms was 31% and 12% of the peak velocities over the cycle - it is the mean flow velocity which is reported in the results from pulsatile flow. The mean flow velocities of these pulsatile waveforms were chosen to be the same as those used above for continuous flow velocities.

Establishing the T₁ of static BMF Images were obtained using a Siemens Magnetom Aera 1.5 T. Baseline T₁ measurements of the BMF were obtained using an inversion recovery sequence at 1.5 T with the following parameters: TR: 10000 ms, TE: 1.2 ms, pixel size: 2.3 mm, FA: 6°, slice thickness: 8 mm, acquisition matrix: 128 x 128, bandwidth: 490 Hz/pixel, TI: 0, 131, 150, then 200–1300 in increments of 100, then 1500–4000 in increments of 500 ms.
Implications for quantifying MRI T1 relaxation in in flowing blood

The spoiled gradient echo imaging parameters for the 3D/2D protocol were TR = 3.0/5.5 ms, TE = 0.95/1.21 ms, 2.3 × 2.3 mm pixels, 108 × 128/128 × 128 matrix size, 248 × 294/294 × 294 mm field of view, slice thickness 5 mm, bandwidth 650/1395 Hz/pixel, and GRAPPA factor of 2. The flip angles were 3.5° and 7° for the 3D sequence and 5° and 10° for the 2D sequence, and these were chosen to bracket the Ernst angle (Wang et al 1987, Imran et al 1999 for the T1 of the BMF (found in section 2.2) and TRs used for each sequence - the Ernst angles were 4.9° for the 3D and 6.7° for the 2D sequences. The 3D sequence consisted of 14 coronal slices per volume acquired in 2.9 s, and 10 low flip angle pre-contrast followed by 50 high flip angle volumes were acquired dynamically for each flow scenario. The 2D sequence used a single coronal slice (the central slice in Figure 4) acquired in 0.7 s and 50 low flip angle pre contrast images followed by 50 high flip angle dynamic images were obtained.

Figure 3. Plot of physiological waveforms provided by the Compuflow 1000MR. Negative flow rate indicates reverse flow in the tubing.

Figure 4. Coronal images were converted into axial slices. The coronal images on the left depict signal from the pipe running along the z axis. The central axial slice intersects the centre of the pipe, hence the widest area of signal (grey). The slice immediately below and above the central slice intersect a smaller cross-section of the pipe and so have narrower columns of signal. The top and bottom image do not intersect with the pipe at all. Reformating the slices this way enabled a ROI to be drawn to isolate the signal from the pipe.

Image processing

All image analysis and simulations were conducted using MATLAB (The Mathworks, Natick, MA). A cylindrical (3D) or rectangular (2D) volume of interest (VOI) was constructed for both the 2D and 3D images to select voxels that are inside the straight section of tubing and within ±10 mm of the centre of the (coronal) image plane along the axis of the tubing. This was trivial for the 2D sequence images. For the 3D sequence, the image volume was axially reformatted and circular regions of interest (ROI) were drawn inside the cross-section of the tubing on the reformatted slices (Figure 4). This was done on each reformatted slice that corresponded to the ±10 mm extent defined above.

For each experimental condition a single signal value for the low flip-angle images was obtained by averaging over the VOIs, then averaging these over the repeated acquisitions (as required by the Fram method, initial images not in a steady-state were excluded – either one or three images for the 3D and 2D sequences respectively). The average signals from the VOI were then obtained for each of the dynamic images. Dynamic T1 estimates for each experimental condition were produced using the Fram equation:

\[ T_{1-\text{True}} = \frac{1}{\text{TR}} \left( \log \left( \frac{S_2}{\sin(\theta_2)} - \frac{S_1}{\sin(\theta_1)} \right) \right)^{-1} \]  \hspace{1cm} (1)

(1) where \( \theta_{1/2} \) are the low and high flip-angles (LFA, HFA) and \( S_{1/2} \) are the corresponding measured signals.

Flip angle correction

The flip angle a spin experiences is a function of the B1 field. As the B1 field is not homogeneous in practice, the flip angle a spin experiences varies from the user defined value.\(^{35}\) This error will be propagated through the Fram equation leading to an inaccurate measurement of T1. As the aim of this work was to investigate errors introduced by flow, the true flip angle experienced by every spin was estimated, as follows, to allow for more accurate T1 estimation using the Fram method. Firstly an inversion recovery sequence using 19 inversion times between 0 and 4000 ms was used to establish the T1 of static BMF (see section 2.2 - a method which is not susceptible to B1 field inhomogeneities. Adapting equation 1 to include an inhomogeneity factor \( f \) we have:

\[ T_{1-\text{True}} = \frac{1}{\text{TR}} \left( \log \left( \frac{S_2}{\sin(\theta_2)} - \frac{S_1}{\sin(\theta_1)} \right) \right)^{-1} \]  \hspace{1cm} (2)
Using equation 1, estimates of $T_1$ were generated from the simulated signal for each spin in the virtual imaging volume. The number of excitations the spin would have experienced at each point in the imaging volume is governed by: 

$$n = \frac{\text{distance into imaging volume of voxel}}{(\text{TR} \times \text{velocity})}.$$ 

Assuming a constant magnetic field whilst in the image volume, the above equation was used to calculate the simulated signal for each spin as used in the physical phantom. As with the physical phantom, simulations were repeated using the mean Parker population AIFs ± 1 standard deviation and at flip angles of 3.5°/7° and 3°/16°. The aim of this analysis was to provide an estimate for the range of likely error in vivo.

**RESULTS**

The $T_1$ of BMF

The results of the IR experiment indicated that the BMF had a $T_1$ of 812 ms which was used as the true value of the BMF in further analysis.

**Flip angle correction**

The calculated correction factor for each voxel along the length of the tube is shown in Figure 5.

**$T_1$ quantification using spoiled gradient echo sequences**

The results of measuring the $T_1$ of static BMF with the physical phantom are shown in Table 1, for constant flowing BMF in Figure 6 and Table 2, and for pulsatile flow in Figure 7.

**Virtual spoiled gradient echo signal and flow**

The signal produced from BMF entering the virtual phantom as it flows into the imaging volume is shown in Figure 8 for both low and high flip angle simulated sequences. The corresponding $T_1$ values estimated from these signals using equation 1 are also shown.

**$T_1$ measurement flow response using a spoiled gradient echo sequence with the physical and virtual phantoms**

Figure 6 shows that the $T_1$ estimated using the Fram method has a non-linear relationship with the flow velocity in the physical phantom when measured with 2D and 3D sequences (solid lines). This relationship is also replicated with the virtual phantom (dashed) although the peak of the overestimate is shifted to a higher flow velocity. The error introduced by flow became statistically significant ($p = 0.05$) at 3.7 cm.s$^{-1}$ for 3D and 17.7 cm.s$^{-1}$ for 2D.

When the flow is pulsatile, the results from the physical phantom show that the measured $T_1$ varies with the mean flow.
velocity (Figure 7). Pulsatile flow increases the standard deviation of the measures with the more pulsatile femoral waveform causing greater measurement variance than the carotid. The flow velocities stated are the mean velocities produced by the waveforms, i.e. 31% and 12% of peak flow velocity for the carotid and femoral waveforms respectively. The increased variance is particularly prominent with the 2D sequence. Despite the increased variance, the mean measures from the pulsatile flows show a similar pattern to the constant flow velocities.

The virtual phantom was used to simulate the measured $T_1$ of a range of underlying $T_1$s. The contours in Figure 9 join lines of equal measured $T_1$. For example, the results show that a measured $T_1$ of 1500 ms (Figure 9, yellow) could represent an underlying $T_1$ of between 1100 ms and 1600 ms for flow velocities between 0 cm s$^{-1}$ and 60 cm s$^{-1}$.

Simulated pre-bolus measurement
To demonstrate the effect of flow related bias on the AIF curve, a Parker population AIF scaled by 1/10$^{th}$ was used to simulate a pre-bolus experiment$^{16,17}$, see Figure 10. Error bars indicate the range of values obtained using the virtual phantom at four key time points for constant flow velocities between 0 cm s$^{-1}$ and 75 cm s$^{-1}$, which are similar to those found in the aorta.

DISCUSSION
It has been shown that accurate measures of contrast agent concentration are desirable for the fitting of models to uptake curves.$^{3,39}$ Various workers measure the passage of a bolus of MRI contrast agent using spoiled gradient echo sequences in order to obtain an AIF for model fitting. This work assesses the error due to flow that exists in these measures. By quantifying and describing the sources of error, we inform the discussion around the strategies being employed as the field strives towards more accurate quantitative measures of vascular function.

It is accepted within the community that current techniques are sensitive to treatment in the context of whole-trial analyses. However, uncertainty in the measurement of the AIF undermines the interpretation of models which purport to be...
Figure 7. Plots showing the mean and standard deviation of $T_1$ over 50 measurements as a function of the mean flow velocity (over cardiac cycle) for pulsatile waveforms, 3D (upper) and 2D (lower). Mean flow velocity seen in an adult male is around 30 cm s$^{-1}$ in the carotid artery and around 10 cm s$^{-1}$ in the femoral artery. True $T_1$ is 812 ms.

Figure 8. Plot of simulated low (red) and high (blue) flip angle signals obtained from flowing virtual BMF. LFA/HFA = 3.5°/7°, TR = 3 ms and flow velocity = 5 cm s$^{-1}$. The signals are used to create the $T_1$ curve via the Fram equation are shown in green. The BMF produces high signal as it arrives in the imaging volume (left) and reduces to a steady state as it receives more excitations and passes through the volume. The HFA signal for the BMF is higher than the LFA signal when it first enters the imaging volume but reaches a steady-state below the signal from the LFA sequence. The differing behaviours of these signals as they approach a steady state results in the non-linear measured $T_1$ shown. “True” $T_1$ is 1200 ms.

Figure 9. Contour plots of measured $T_1$ for a range of $T_1$ and flow velocities simulated using flip angles of 3.5° and 7° (top), and 3° and 16° (bottom), with a TR of 3 ms. The contours join points of the same measured $T_1$.

quantifying sought-after measures of vascular function on an examination-by-examination or even pixel-by-pixel basis.

Following the order of the results, this section discusses the impact of using flip angle correction to isolate the error introduced by flow, the quantification of $T_1$ using the physical phantom and how these measures were used to validate the virtual phantom. Finally, a summary of the implications for those working in the field of DCE-MRI is discussed.

Flip angle correction
The type of flip angle correction profiles seen in Figure 5 for the coronal plane are entirely consistent with those shown in previous work. The correction factor profiles assume that all of the error in $T_1$ quantification is because of an inhomogeneous $B_1$ field. The error on the absolute measurement observed here should be noted for future studies using the double-flip angle.
Figure 10. Graph showing a simulated pre-bolus AIF (blue), i.e. a typical Parker population AIF at 1/10th dose - the blue error bars indicate the population standard deviation. (courtesy of G. Parker). The virtual phantom is used to find the maximum and minimum gadolinium concentration that would be measured as the flow rate is varied from 0 to 75 cm s$^{-1}$ at key points as the bolus passes. The simulated measures are offset for clarity. The simulated measures are also obtained for a Parker population functions ±1 standard deviation. It is clear that the bolus with a larger peak (+1 standard deviation) is less susceptible to flow induced measurement errors. It is also clear how sensitive the measures are to choice of flip angles.

$T_1$ quantification using spoiled gradient echo sequences

There is increased standard deviation in the measurement of $T_1$ with the 2D sequence compared to the 3D (Figures 6 and 7). This is likely due to the reduced signal to noise due to its decreased measurement volume and image acquisition time.

Virtual spoiled gradient echo signal and flow

The non-linear relationship between flow velocity and measured $T_1$ can be explained using Figure 8, which describes the relationship between the measured imaging signals and distance into the imaging volume for a constant flow velocity using the virtual phantom; i.e. how the signals change as they experience more excitations. As the spins travel further into the imaging volume (from left to right), they are subject to more excitations and each curve monotonically reaches its steady-state at a rate that depends on the sequence flip-angle. If the flow velocity is increased, the number of excitations experienced when the spins have reached isocentre will be reduced, i.e. the curves in Figure 8 will be stretched along the x-axis away from the origin. The Fram method (equation 1) produces a measurement of $T_1$ from a function of the low and high flip angle image signals and as shown in Figure 8, the high flip angle signal initially has a higher signal magnitude than the low flip angle signal, but reaches a lower magnitude at steady state. This results in the $T_1$ measurement rising and then overshooting the correct measurement, before falling to the correct value as the distance into the slice increases.

$T_1$ measurement of non-pulsatile flow using a spoiled gradient echo sequence with the physical and virtual phantoms

The non-linear relationship between flow velocity and $T_1$ measurement in the physical phantom is qualitatively replicated by the virtual phantom, but the curves in Figure 6 for the virtual phantom are shifted to the right. It is hypothesised that this discrepancy is due to the virtual phantom simulating a perfectly homogeneous RF excitation profile across the imaging volume. The actual excitation profile experienced by the physical phantom is influenced by a variety of factors such as $B_0$ and slice profile variations. Importantly, excitations of unknown magnitude affect the B1 before it enters the imaging volume. Therefore, the spin history of the magnetisation at isocentre is different to that simulated by the virtual phantom. None of these factors are easily mitigated, especially within the clinical context. A subject in the scanner will have nearly all their blood within the main $B_0$-field and close to the imaging volume, thus it will be equally difficult to characterise the spin-history of blood before it enters the imaging volume.

AIF measurement

There are large uncertainties in the measurement of $T_1$ due to flow. These errors are non-linearly related to flow across the range of mean velocities seen in large vessels. These difficulties are even further exacerbated by pulsatile flow and in the case of fast 2D imaging due to the temporal variation in mean velocity during a signal acquisition. Flow pulsatility would be difficult to model in practice and subsequent changes in measured $T_1$ are largely treated, incorrectly, as part of the “noise”. However, the observed increased variance introduced by pulsatile versus constant flow was not large when using the 3D sequence. It may be possible to partially correct the measures from a slower imaging sequence if the mean flow velocity is known. However, for faster imaging sequences, the mean flow velocity over each individual image acquisition would be required.

In practice a bolus injection of contrast agent is used. While the rate of change of contrast during the equilibrium Phase is negligible compared to the effects of pulsatile flow, the same is not true during the initial passage of the bolus in the major vessels (around 10–15 sec). The random interaction between the change in $T_1$ due to changes in contrast agent concentration and the $T_1$ measurement error introduced by pulsatile flow further complicates the accurate measurement of $T_1$.

Figure 10 shows the errors introduced by flow and the sensitivity of the errors to flip angle. This should be considered when designing sequences. The sequence with the 3° and 16° flip angles is less susceptible to flow induced measurement errors than the 3.5° and 7° flip angle measures. Additionally, the variation in experienced flip angles across the imaging volume (Figure 5) should also be considered - these variations may impact attempts at correcting flow affected signals.

Pre-bolus measurement

For a fixed $T_1$, the $T_1$ that will be measured varies as a function of flow velocity. Thus, pulsatile flow will yield a range of measured
would vary non-linearly from 900 ms to ~766 ms as the flow and pulsatilities which are similar to those observed in vivo. However, the underlying T₁ of blood were lower to begin with, the plots on Figure 9 show that less measurement variability would be introduced by flow. For example, the range of T₁ that would be measured if the underlying T₁ were 900 ms would vary non-linearly from 900 ms to ~766 ms as the flow velocity is increased from 0 cm.s⁻¹ to ~50 cm.s⁻¹.

When using a separate acquisition to measure the AIF (such as the already discussed pre-bolus experiment), less error would be introduced by flow if it were performed after the main DCE-MRI acquisition since the contrast agent-administered for the main DCE-MRI examination will be well mixed in the blood, thereby reducing its baseline T₁. This may be particularly useful in avoiding bias in the establishment of the baseline T₁. However, in this scenario, the decrease in the T₁ of blood prior to the AIF measurement will mean a smaller contrast between this signal and the subsequent peak signal, leading to a greater uncertainty in the T₁ measures. Further investigation is required to assess whether measuring the AIF in this way would be worthwhile.

Summary and implications for those working in DCE-MRI
This paper assesses the expected inaccuracy in T₁ measurement due to physiologically realistic pulsatile flow using a physical phantom which is used to validate a virtual phantom. Results from both phantoms inform the discussion and should be of interest to those working in DCE-MRI and particularly those looking to use model fitting to obtain quantitative measures. We expand on previous work, particularly by Roberts et al. and more recently by van Schie et al., by examining flow velocities and pulsatilities which are similar to those observed in vivo.

The effect of flow on T₁ measurement using the double flip angle method
The effects of flow on T₁ measurement are large compared to the changes which would be observed during the passage of a bolus of contrast agent and are non-linear with flow velocity. ROIs used for measuring AIFs should be placed as far away as possible from the entry points of flow to the imaging volume to reduce this effect.

Model fitting
The ability to obtain absolute quantitative measures of vascular function by fitting models to DCE-MRI data is questionable given that no strategy has yet been shown to accurately measure the AIF. Reporting of the absolute model parameter values which purport to be related to function, rather than changes from a baseline, should be discouraged.

Correcting the signal from flowing spins
If the mean flow velocity were known, e.g. through phase difference techniques, it seems feasible to partially correct the T₁ measurements. Peeters et al. and Van Schie, 2018 proposed methods to do this by obtaining the number of excitations a flowing fluid has received at a point and then correcting for it. This work further illuminates difficulties in applying this technique in vivo given the large variations in T₁ measurement which pulsatile flow is shown to introduce. Such strategies would need to account for the range of excitations observed over the cardiac cycle. Arteries with lower pulsatility may present easier targets for potential correction of the inflow-effect. It may be that the use of complementary quantitative flow techniques may inform the choice of artery and could provide data, which would assist in the correction of the in-flow effect over the cardiac-cycle on a per-patient/examination basis. Given the large variation in pulsatilities and flow velocities observed across subjects, along with the variance in the prior number of excitations that may be experienced by incoming blood as it passes through potentially tortuous feeding arteries near to the imaging volume, such corrections would need to be bespoke for a particular examination.

Sequence and examination design
Higher flip angles reach saturation more quickly, as the signal is a function of \( \cos(\alpha) \) (see equation 3), thus reducing the in-flow-effect. This should be a consideration during sequence design along with other factors discussed in De Neayer et al. Measurement error due to the inflow effect may be reduced if the AIF is measured after the main DCE-MRI investigation (discussed above).

Alternative approaches to AIF measurement in a vessel
Even if measurement error could be reduced to zero, the measurement of AIF in a feeding vessel is still a proxy for the plasma curve in the tissue. Therefore, solving the inflow effect issue would not yield a perfect model fit. In the context of clinical trials, a population AIF has been shown to provide results where the repeatability is low enough that clinically useful changes can be measured. Therefore, it is suggested that rather than pursuing further improvement to these techniques, i.e. towards more quantitative measures, more effort should be spent developing methods for tissue-level AIF estimation, similar to that of Schabel et al. Reference tissue approaches are attractive in principle but their effectiveness in practice is contingent on the contrast dynamics of the two regions being complimentary, which cannot always be guaranteed.

Proposed radial acquisitions have significant advantages over Cartesian imaging in terms of spatial and temporal resolution, and this work indicates that similar evaluations to assess the robustness of radial acquisitions to pulsatile flow should be undertaken.

CONCLUSION
Quantitative T₁ measurement using spoiled gradient echo sequences in blood is severely compromised by its flow. The relationship between measurement error and flow is highly non-linear which seems likely to complicate any attempt to correct this, especially given the pulsatility seen in arteries. Of particular note for DCE-MRI, these findings indicate that any AIF obtained for the fitting of pharmacokinetic models from monitoring the
$T_1$ of blood in an artery is likely to be inaccurate. Thus, the value of such strategies is undermined.

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