

Calibration of arterial spin labeling data—potential pitfalls in post-processing

Joana Pinto¹  | Michael A. Chappell^{2,3}  | Thomas W. Okell²  | Melvin Mezue² | Andrew R. Segerdahl² | Irene Tracey^{2,4} | Pedro Vilela⁵ | Patrícia Figueiredo¹ 

¹Institute for Systems and Robotics and Department of Bioengineering, Instituto Superior Técnico, Universidade de Lisboa, Lisbon, Portugal

²Wellcome Centre for Integrative Neuroimaging, FMRIB, Nuffield Department of Clinical Neurosciences, University of Oxford, Oxford, United Kingdom

³Institute of Biomedical Engineering, Department of Engineering Science, University of Oxford, Oxford, United Kingdom

⁴Nuffield Division of Anaesthetics, Nuffield Department of Clinical Neuroscience, University of Oxford, Oxford, United Kingdom

⁵Imaging Department, Hospital da Luz, Lisbon, Portugal

Correspondence

Joana Pinto, Institute for Systems and Robotics-Lisboa, Department of Bioengineering, Instituto Superior Técnico, Universidade de Lisboa, Av. Rovisco Pais 1, 1049-001 Lisbon, Portugal.
Email: joanasequeirapinto@tecnico.ulisboa.pt

Funding information

Portuguese Foundation for Science and Technology (FCT), Grant/Award Numbers: PTDC/BBB-IMG/2137/2012, PD/BD/135114/2017 and UID/EEA/50009/2019; Engineering and Physical Sciences Research Council UK, Grant/Award Numbers: EP/P012361/1; Wellcome Trust, Grant/Award Numbers: 203139/Z/16/Z.

Purpose: To assess the impact of the different post-processing options in the calibration of arterial spin labeling (ASL) data on perfusion quantification and its reproducibility.

Theory and Methods: Absolute quantification of perfusion measurements is one of the promises of ASL techniques. However, it is highly dependent on a calibration procedure that involves a complex processing pipeline for which no standardized procedure has been fully established. In this work, we systematically compare the main ASL calibration methods as well as various post-processing calibration options, using 2 data sets acquired with the most common sequences, pulsed ASL and pseudo-continuous ASL.

Results: Significant and sometimes large discrepancies in ASL perfusion quantification were obtained when using different post-processing calibration options. Nevertheless, when using a set of theoretically based and carefully chosen options, only small differences were observed for both reference tissue and voxelwise methods. The voxelwise and white matter reference tissue methods were less sensitive to post-processing options than the cerebrospinal fluid reference tissue method. However, white matter reference tissue calibration also produced poorer reproducibility results. Moreover, it may also not be an appropriate reference in case of white matter pathology.

Conclusion: Poor post-processing calibration options can lead to large errors in perfusion quantification, and a complete description of the calibration procedure should therefore be reported in ASL studies. Overall, our results further support the voxelwise calibration method proposed by the ASL white paper, particularly given the advantage of being relatively simple to implement and intrinsically correcting for the coil sensitivity profile.

KEYWORDS

ASL, calibration, cerebral blood flow, kinetic modeling, MRI

This is an open access article under the terms of the Creative Commons Attribution License, which permits use, distribution and reproduction in any medium, provided the original work is properly cited.

© 2019 The Authors. *Magnetic Resonance in Medicine* published by Wiley Periodicals, Inc. on behalf of International Society for Magnetic Resonance in Medicine

1 | INTRODUCTION

Arterial spin labeling (ASL) is a non-invasive MRI technique that provides quantitative images of tissue perfusion, by using magnetically labeled blood water protons as an endogenous blood flow tracer.¹⁻³ ASL is acknowledged to have great potential as a completely non-invasive quantitative perfusion imaging technique, but its implementation has been challenging because of the intrinsically low SNR. This has motivated the development of a multitude of signal acquisition and processing strategies that aim to overcome this limitation as well as the publication of a ASL implementation consensus paper.⁴ Although pseudo-continuous ASL (pCASL) is the recommended labeling strategy, pulsed ASL (PASL) is still a commonly used technique.⁵⁻⁹

In principle, cerebral blood flow (CBF) can be quantified based on a single time delay measurement—the post-labeling delay (PLD) in pCASL and the inversion time (TI) in PASL—provided that it is long enough relative to the transit time of the label bolus between the arteries and the capillaries, the so-called arterial transit time (ATT). However, if this condition cannot be assumed, which is the case in pathologies presenting delayed arterial transit times, multiple-PLD/TI measurements are necessary.¹⁰⁻¹² Furthermore, if an appropriate kinetic model is fitted to the data, the multiple-PLD/TI strategy allows the assessment of not only CBF but also ATT¹³ and potentially arterial blood volume (aBV),^{14,15} which may be parameters of interest in their own right. In any case, to obtain CBF measures in absolute units, it is necessary that the relative CBF (CBF_{rel}) images should be normalized by the equilibrium magnetization of arterial blood (M_{0a}), which is usually derived from the equilibrium magnetization measured in tissue (M_{0t}). Critically, it has been demonstrated through theoretical analysis that the 2 factors that CBF quantification using ASL is most sensitive to are M_{0a} estimation and labeling efficiency.¹⁶

The current recommendation for ASL calibration involves the acquisition of a separate proton-density weighted image followed by the extraction of M_{0t} by extrapolation as a function of the repetition time (TR). This is subsequently converted to an M_{0a} image by first applying spatial smoothing and then dividing the image by the brain average brain–blood water partition coefficient (λ).⁴ Nevertheless, several other strategies might be used depending on the acquisition scheme. If no background suppression is used (e.g., PICORE),¹⁷ it is also possible to obtain M_{0t} directly from the ASL data by averaging the control images at a particular PLD/TI. In the specific case of multiple-TI PASL, it is possible to obtain M_{0t} by fitting a saturation-recovery curve to the multiple-TI control images. Additionally to the selection of a calibration image, it is necessary to decide whether to compute a voxelwise M_{0a} value, or a single average value across a homogeneous reference region, usually gray matter (GM), white matter (WM), or cerebrospinal fluid

(CSF).^{18,19} For the practical implementation of the calibration pipeline, a number of subtler choices must also be made, however, these are rarely discussed or even reported in ASL studies. Although some calibration method comparisons have been reported,^{16,18-20} the impact and potential pitfalls of the corresponding processing options have yet to be investigated.

Here, we systematically compare the impact of using different calibration post-processing pipelines on the quantification of perfusion and its reproducibility, using both PASL and pCASL acquisitions.

2 | THEORY

Calibration is required to obtain CBF (mL/100 g/min) in absolute units from ASL measurements, through normalization by M_{0a} (and α , the labeling/inversion efficiency)

$$CBF = \frac{CBF_{rel}}{\alpha M_{0a}} \times 6000. \quad (1)$$

Calibration therefore entails estimation of M_{0a} and it is achieved in 2 main steps: (1) generation of M_{0t} map, and (2) derivation of M_{0a} from M_{0t} map.

2.1 | Generation of M_{0t} map

The method used for generating the M_{0t} map depends on the acquisition scheme and options chosen. There are 3 main strategies: long TR calibration scan (*LongTR*), ASL control averaging (*CtrlAvg*), and control saturation recovery (*SatRec*).

2.1.1 | Long TR calibration scan (*LongTR*)

The *LongTR* approach is based on a separately acquired long TR scan, which approximates the equilibrium magnetization of tissue in each voxel i , $M_{0t}(i)$. This map should be corrected for the amount of T_1 relaxation at the TR at each voxel, according to

$$M_{0t,corr}(i) = \frac{M_{0t}(i)}{1 - e^{-\frac{TR_{long}}{T_{1,i}}}}, \quad (2)$$

where $T_{1,i}$ corresponds to the tissue-specific T_1 in each voxel i and TR_{long} is the corresponding TR of the M_{0t} scan. This correction depends on a separate tissue segmentation procedure. Using a long TR minimizes the impact of the T_1 correction, which may become negligible for sufficiently long TR values.

2.1.2 | Averaging of control images (*CtrlAvg*)

If no background suppression is applied, it is also possible to estimate M_{0t} based on the control images. This can be carried out by averaging control images at a fixed TI ($M_{ctrl}(i)$), which should then be corrected for the amount of T_1 relaxation

during TI, at each voxel, to yield a corrected map of the tissue equilibrium magnetization

$$M_{0t,corr}(i) = \frac{M_{ctrl}(i)}{1 - Ae^{-\frac{TI}{T_{1,t}}}}. \quad (3)$$

Furthermore, in this approach, the inversion–saturation efficiency (A) is also a variable. A value of 0.90 was chosen as it was the approximate average value of A estimated in *SatRec*.

For the generation of the M_{0t} map, one can argue that this method is less appropriate, mainly because the value of TI is not chosen for this purpose and therefore it is not sufficiently long to minimize the dependency on the values of T_1 and A .

2.1.3 | Saturation recovery of control images (*SatRec*)

If no background suppression is applied, it is possible to estimate M_{0t} by fitting a saturation-recovery curve to the series of control images, M_{ctrl} (TI), if multiple TIs are sampled and the acquisition sequence includes presaturation (which is the case of Q2TIPS PASL).¹⁷ This strategy allows estimation of M_{0t} as well as T_{1t} and saturation efficiency (A) (“asl_calib”, <https://fsl.fmrib.ox.ac.uk/fsl/fslwiki/BASIL/>).

2.2 | Derivation of M_{0a} from M_{0t} map

Once an M_{0t} map is generated, the next step is the derive M_{0a} . Two types of methods are commonly used for that purpose: the reference tissue (RT) and the voxelwise (Voxel).

2.2.1 | Reference tissue methods (RT-CSF, RT-WM, RT-GM)

The reference tissue (RT) approach consists of computing a single M_{0a} value based on a homogeneous tissue region (GM, WM, or CSF). The first step is to correct the M_{0t} (as well as CBF) maps for the RF head coil sensitivity profile. In case the coil sensitivity profile is not available, an approximate correction may be carried out by normalizing the M_{0t} (and CBF) maps using a bias image estimated directly from the data using appropriate post-processing tools (e.g., FAST).²¹

The next step is to define an appropriate RT mask across which an average M_{0t} value is computed, $\langle M_{0t} \rangle_{rt}$. This usually involves tissue segmentation based on a structural image (T_1 -weighted image or T_{1t} map in case this has been estimated) followed by registration of the RT mask with the M_{0t} map. Finally, the M_{0a} value is obtained by normalizing $\langle M_{0t} \rangle_{rt}$ with the respective λ , while correcting for the difference in T_2^* relaxation between the reference tissue and arterial blood (associated with the M_{0t} measurement), yielding

$$M_{0a} = \frac{\langle M_{0t} \rangle_{rt} e^{TE \left(\frac{1}{T_{2,t}^*} - \frac{1}{T_{2,a}^*} \right)}}{\lambda_{rt}}, \quad (4)$$

where $T_{2,a}^*$ is the transverse relaxation time of arterial blood, $T_{2,rt}^*$ is the transverse relaxation time of the reference tissue, and λ_{rt} is blood–brain water partition coefficient of the corresponding reference tissue.

2.2.2 | Voxelwise method (Voxel)

In this method, an M_{0a} map is obtained through direct extrapolation from the M_{0t} value in each voxel, by normalizing with the respective tissue λ , while correcting for the difference in T_2^* relaxation between the respective tissue and arterial blood, according to

$$M_{0a}(i) = \frac{M_{0t}(i) e^{TE \left(\frac{1}{T_{2,t}^*} - \frac{1}{T_{2,a}^*} \right)}}{\lambda_t}, \quad (5)$$

where $T_{2,t}^*$ and λ_t are the transverse relaxation time and the blood–brain water partition coefficient, respectively, of the tissue in each voxel. Regarding λ_t , in principle, a different λ value exists in each voxel because λ differs as a function of tissue type and hence the voxel’s partial volume estimates (PVEs). For that reason, λ_t should be a PVE-weighted average of λ . PVEs are usually obtained through tissue segmentation using appropriate tools (e.g., FAST).²¹ The M_{0a} map is then smoothed, to minimize noise contributions.

2.3 | Simplifications

A number of assumptions and approximations to the theoretically ideal calibration just described are usually undertaken to overcome additional complexities in both data acquisition and processing. These are described here for each of the main calibration methods, and the respective post-processing options that are tested in this work are indicated, relative to the default options derived from the theoretical considerations above. All options are summarized in Table 1.

2.3.1 | Generation of M_{0t} map

Long TR calibration scan (LongTR)

The current recommendation indicates that the T_1 relaxation correction should only be carried out if $TR < 5$ s.⁴ However, the consensus paper also recommends the use of a voxelwise approach, in which case this may not have a significant impact (as we will show). We therefore tested the impact of not correcting the M_{0t} images for T_1 relaxation (vs. correcting, the default).

TABLE 1 Summary of calibration methods and post-processing options tested

Method	Options		
Generation of M_{0r} map			
PASL- <i>SatRec</i>	presaturation efficiency	A estimated (default) A = 100%	
PASL- <i>CtrlAvg</i>	TI value	long TI ~2400 ms (default) short TI ~800 ms	
	T_1 correction	yes (default) no	
	presaturation efficiency	A = 90% (default) A = 100%	
	T_1 correction	yes (default) no	
pCASL- <i>LongTR</i>	T_1 correction	yes (default) no	
Derivation of M_{0a} from M_{0r} map			
Reference tissue CSF	RT mask	restrictive (default)	restrictive threshold ($PVE_{CSF} \geq 0.9$) intersected with MNI atlas ventricles mask
		intermediate	less restrictive threshold ($PVE_{CSF} \geq 0.6$) intersected with MNI atlas ventricles mask
		extensive	less restrictive threshold ($PVE_{CSF} \geq 0.6$)
	bias correction	yes (default)	
		no	
Reference tissue WM	RT mask	restrictive (default)	restrictive threshold ($PVE_{WM} \geq 0.9$)
		extensive	less restrictive threshold ($PVE_{WM} \geq 0.6$)
	bias correction	yes (default)	
		no	
Reference tissue GM	RT mask	restrictive (default)	restrictive threshold ($PVE_{GM} \geq 0.9$)
		extensive	less restrictive threshold ($PVE_{GM} \geq 0.6$)
	bias correction	yes (default)	
		no	
Voxel	smoothing	FWHM = 10.5 mm (default)	
		FWHM = 17.5 mm	
		no	
	λ value	PVE-weighted average (λ_{pve}) (default)	$\lambda_{pve}(i) = \lambda_{GM} PVE_{GM}(i) + \lambda_{WM} PVE_{WM}(i) + \lambda_{CSF} PVE_{CSF}(i)$
		brain average ($\lambda_{avg} = 0.9$) tissue-specific (λ_{tspec})	$\lambda_{avg}(i) = \lambda_{GM} \times 0.5 + \lambda_{WM} \times 0.5$ $\lambda_{tspec}(i) = \begin{cases} \lambda_{GM} if \text{ GM} \\ \lambda_{WM} if \text{ WM} \\ \lambda_{CSF} if \text{ CSF} \end{cases}$

Control averaging (CtrlAvg)

One of the crucial points when using the *CtrlAvg* approach is the value of the TI because of its strong interaction with the T_1 correction. We therefore compared using a longer (default; 2400 ms) and a shorter (800 ms) value, while correcting or not correcting for T_1 .

When acquisition sequences include presaturation pulses (such as Q2TIPS PASL), quantification of the presaturation efficiency, A , might be an important factor. For this

approach, we compared A of 90% (default; the approximate average value of A estimated in *SatRec*, see below) with A of 100%.

Control saturation recovery (SatRec)

In the *SatRec* approach it is also possible to estimate A as part of model fitting, which should be the ideal option. Therefore, we compared estimation of A (default) with having a fixed A of 100%.

2.3.2 | Derivation of M_{0a} from M_{0t} map

Reference tissue methods (RT-CSF, RT-WM, RT-GM)

Whenever using the RT method, the choice of a tissue region of interest may range between being very restrictive to being more inclusive, resulting in considerably different PVEs and potentially very different calibrations. Here, we explored the impact of using a restrictive mask (default) with less restrictive masks. In particular, a restrictive tissue mask was obtained from the respective PVE maps in ASL space, through the application of stringent thresholds ($PVE_{\text{tissue}} \geq 0.9$). Additionally, the CSF mask was further intersected with a mask of the lateral ventricles derived from the MNI atlas.²² Furthermore, regarding coil sensitivity correction, we considered the bias field approach as default (see Discussion) and compared this with the option of no correction.

Voxelwise method (Voxel)

In the Voxel method, smoothing of the M_{0t} map by a median filter with FWHM = 10.5 mm (3×3) was considered default and was compared with using a median filter with FWHM = 17.5 mm (5×5) as well as with the option of no smoothing. Regarding λ , the ideal PVE-weighted λ (default) was compared with using a single brain value (0.9), the average of GM and WM λ values, and a tissue-specific λ (Table 1). For illustration, a summary of the Voxel method post-processing and options is displayed in Figure 1.

3 | METHODS

The impact of the different calibration methods and respective post-processing options was evaluated in terms of average GM CBF quantification as well as its test–retest reproducibility.

3.1 | Data acquisition

Test–retest multiple-PLD/TI pCASL and PASL data sets were previously acquired.^{15,23} The PASL study was approved by the Hospital da Luz Ethics Committee, and all subjects gave written informed consent in accordance with the Declaration of Helsinki. The pCASL study was carried out under an agreed technical development protocol approved by the Oxford University Clinical Trials and Research Governance office, in accordance with International Electrotechnical Commission and United Kingdom Health Protection Agency guidelines.

In the PASL study, 9 healthy volunteers (22.9 ± 5.6 y, 4 males) were studied on a 3T Siemens Verio whole-body MRI system (Erlangen, Germany) using a 12-channel-receive head RF coil, on 2 sessions separated by ~ 1 wk.¹⁵ In the pCASL study, 8 healthy volunteers (28.3 ± 2.5 y, 6 males) were studied on a 3T Siemens Verio whole-body MRI system (Erlangen) using a 32-channel receive head RF coil, on 3 occasions separated by 1 wk and 1 mo.²³ A reference image with no labeling or background suppression, TR = 6s, and all other parameters identical to the pCASL scan, was collected for calibration. In both data sets, T_1 -weighted structural images were acquired from each subject for tissue segmentation and registration purposes. Acquisition details for both data sets are summarized in Table 2.

3.2 | Data analysis

Image analysis was conducted using FSL (FSL5.0.1, <http://fsl.fmrib.ox.ac.uk/fsl/fslwiki/FSL>) and MATLAB (2013a, <http://mathworks.com>). The multiple-PLD/TI ASL images in each data set were aligned with each other by motion correction using MCFLIRT.²⁴ For each data set, at each PLD/TI, the control images and the pairwise differences between control and label images were averaged across repetitions, yielding

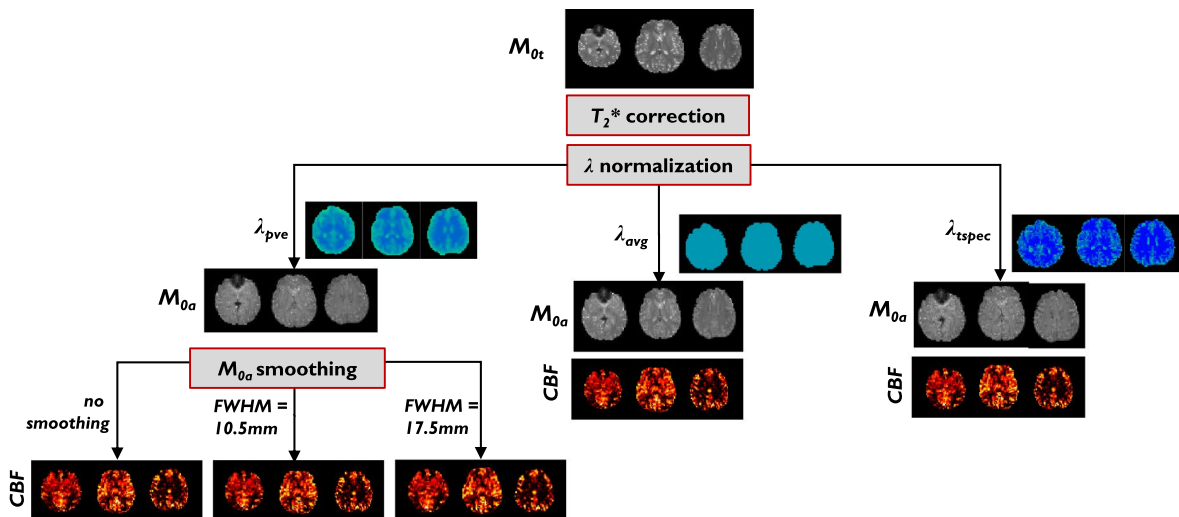


FIGURE 1 Illustrative individual example of the voxel M_{0a} derivation post-processing options

TABLE 2 Summary of the main acquisition parameters for the PASL and pCASL data sets

	PASL (PICORE-Q2TIPS ¹⁷)	pCASL
Readout	2D multi-slice GE-EPI	2D multi-slice GE-EPI
Background suppression	no	yes
TR	2500 ms	4000 ms
TE	19 ms	13 ms
Number of slices	9	24
Slice time	50.0 ms	45.2 ms
Voxel size	$3.5 \times 3.5 \times 5.0 \text{ mm}^3$	$3.5 \times 3.5 \times 5.0 \text{ mm}^3$
Labeling parameters	labeling slab thickness = 100 mm labeling bolus duration = 750 ms [*] labeling slab gap = 18.8 mm	labeling duration = 1400 ms
TI/PLD values	400–2400 ms, in steps of 200 ms (11 values)	250–1500 ms, in steps of 250 ms (6 values)
Control-label pairs	8 for each TI	8 for each PLD

^{*}The Q2TIPS module allowed limiting the labeling to a maximum of 750 ms by adjusting TI_1 and TI_{1s} for each TI: for $TI < 1000$ ms, $TI_1 = TI_{1s} = TI - 25$ ms, and for $TI > 1000$ ms, $TI_1 = 750$ ms and $TI_{1s} = 900$ ms.

time series of mean control images (M_{ctrl} (PLD/TI)) and mean magnetization difference images (ΔM_{diff} (PLD/TI)) as a function of PLD/TI, respectively. Additionally, for the PASL data set, off-resonance effects caused by imperfect inversion slice profile in 2D multi-slice imaging were corrected.¹¹

The structural images were segmented using FAST²¹ to estimate tissue masks and corresponding partial volume estimate (PVE) maps. Both types of image were co-registered to the ASL space of each subject and session using a linear transformation (FLIRT)²⁴ (see Registration and Tissue Segmentation in the Supporting Information). For the PASL data sets, the T_1 map derived from the saturation recovery approach was used as reference image, instead of the ASL data, because it provided more tissue contrast. An extended kinetic model including a tissue contribution and an intravascular arterial compartment was fitted to ΔM_{diff} (PLD / TI), yielding maps of relative CBF and aBV, and ATT^{14,25} (see Kinetic Modeling in the Supporting Information).

3.3 | Calibration comparisons

Calibration of the data in each subject and session was carried out using the combinations of options and parameters listed in Tables 1 and 3. For the purpose of comparing calibration options, GM-average CBF values were obtained using a GM mask derived for each subject and session using the GM mask obtained from segmentation, co-registered to ASL space. Statistically significant differences between methods and options were tested using repeated-measures ANOVA. When significant effects were found, post-hoc analysis using pairwise t tests were carried out ($P < 0.05$). The reproducibility of the GM average CBF measurements was also assessed, by computing the inter- and intra-subject coefficients of variation (CV_{inter} and CV_{intra}) for the median CBF values across GM (see Coefficients

TABLE 3 Parameters used in post-processing calibration of the ASL data sets

	Parameters		
	GM	WM	CSF
T_1 (s)	1.3	1.0	4.3
T_2^* (ms)	60	50	400
λ	0.98	0.82	1.15

of Variation in the Supporting Information).^{15,23,26,27} Significant differences in CV across M_{0a} estimation methods for each M_{0t} map generation method were assessed using jackknife resampling, followed by 1-way ANOVA and post-hoc pairwise t tests between methods ($P < 0.05$).

4 | RESULTS

First, we present the quantification and reproducibility results obtained for the main calibration methods using the default post-processing options. Second, the impact of variations in the post-processing options is assessed.

4.1 | Comparison between calibration methods

Figure 2 shows illustrative examples of individual CBF maps obtained using the *SatRec* and *CtrAvg* M_{0t} generation methods in PASL and the *LongTR* M_{0t} generation method in pCASL and each of the M_{0a} derivation methods (RT-CSF/RT-WM/RT-GM/Voxel) with the corresponding default post-processing options. Minor differences can be observed in the Voxel compared to the RT methods, particularly in the

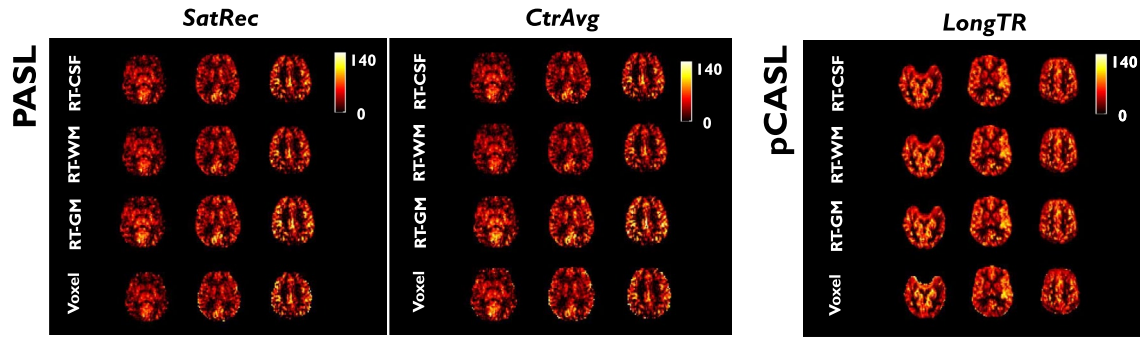
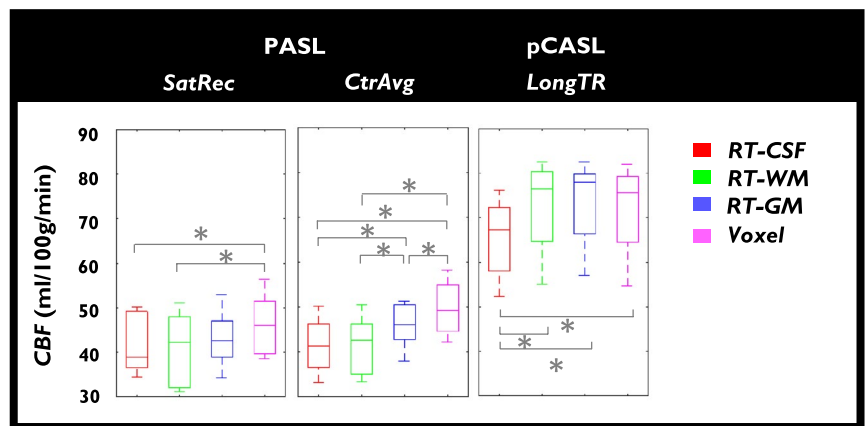


FIGURE 2 Illustrative examples of individual CBF (ml/100g/min) maps obtained using the 4 M_{0a} derivation methods (RT-CSF/RT-WM/RT-GM/Voxel), the *SatRec* and *CtrAvg* M_{0t} generation methods in PASL (left), and the *LongTR* M_{0t} generation method in pCASL (right), with the respective default post-processing options

FIGURE 3 Group results for the GM average CBF values obtained using the different calibration methods for session 1 of the PASL and pCASL data sets, with the respective default options. Bottom and top edges of box plots represent the 25th and 75th percentiles, respectively. *Significant differences as assessed by pairwise t tests ($P < 0.05$)



borders of CBF maps. This can be explained by the amplification of noise that occurs on the voxelwise division by the M_{0t} map carried out in the Voxel method.

The group results for the average GM CBF values, obtained using the different calibration methods with the respective default options, for the first session of each data set, are presented in Figure 3. Small CBF differences across the different M_{0a} derivation methods were observed although these were significantly different at times. In particular, for PASL *SatRec*, only RT-CSF and RT-WM methods were significantly different from the Voxel method. For PASL *CtrAvg*, CBF values were significantly different across methods except between RT-CSF and RT-WM methods. For pCASL *LongTR* approach, only CBF values from the RT-CSF method were significantly different from the other M_{0a} derivation methods.

Finally, the reproducibility metrics of the average GM CBF values for the main calibration methods, using the default post-processing options, are presented in Figure 4. All methods were found to exhibit good reproducibility, with pCASL being superior to PASL, and the RT-WM method generally yielding the worst values. A significant main effect of M_{0a} derivation method was found for each

M_{0t} map generation method. Subsequent post-hoc analysis yielded significant differences between several M_{0a} derivation methods. The Voxel and RT-CSF methods yielded the lowest CV_{intra} values, whereas the Voxel and RT-GM methods yielded the lowest CV_{inter} values, with significant differences in several cases. The RT-WM method systematically carried out worst for both CV_{intra} and CV_{inter} , often with significant differences.

4.2 | Impact of calibration post-processing options

The group results for the GM average CBF values obtained when varying the calibration post-processing options are presented in Figure 5, for each of the main calibration methods tested on session 1 of each data set. For each method, there was a significant main effect of the post-processing option; subsequent pairwise comparisons revealed significant differences between some options, as indicated in the plots.

When applying the *SatRec* method for M_{0t} generation, the RT-CSF approach yielded the highest CBF differences across methods, with the extensive mask producing significantly greater CBF values relative to the default option that

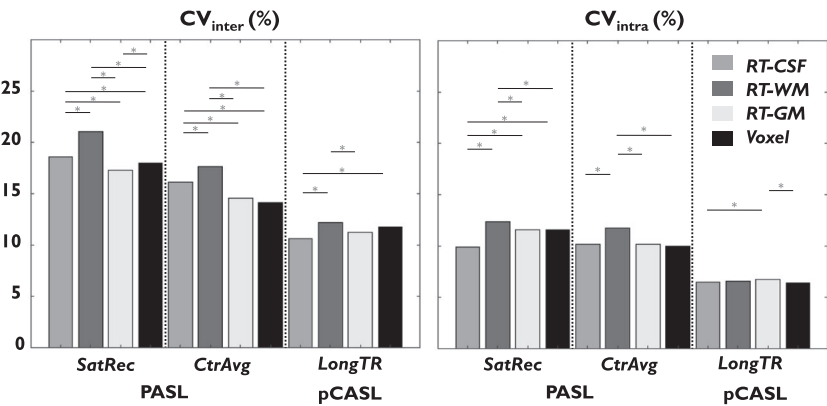


FIGURE 4 Reproducibility metrics (CV_{inter} , CV_{intra}) of GM average CBF values, obtained for each data set, and each of the main calibration methods, using their default post-processing options. *Significant differences as assessed by pairwise t tests ($P < 0.05$)

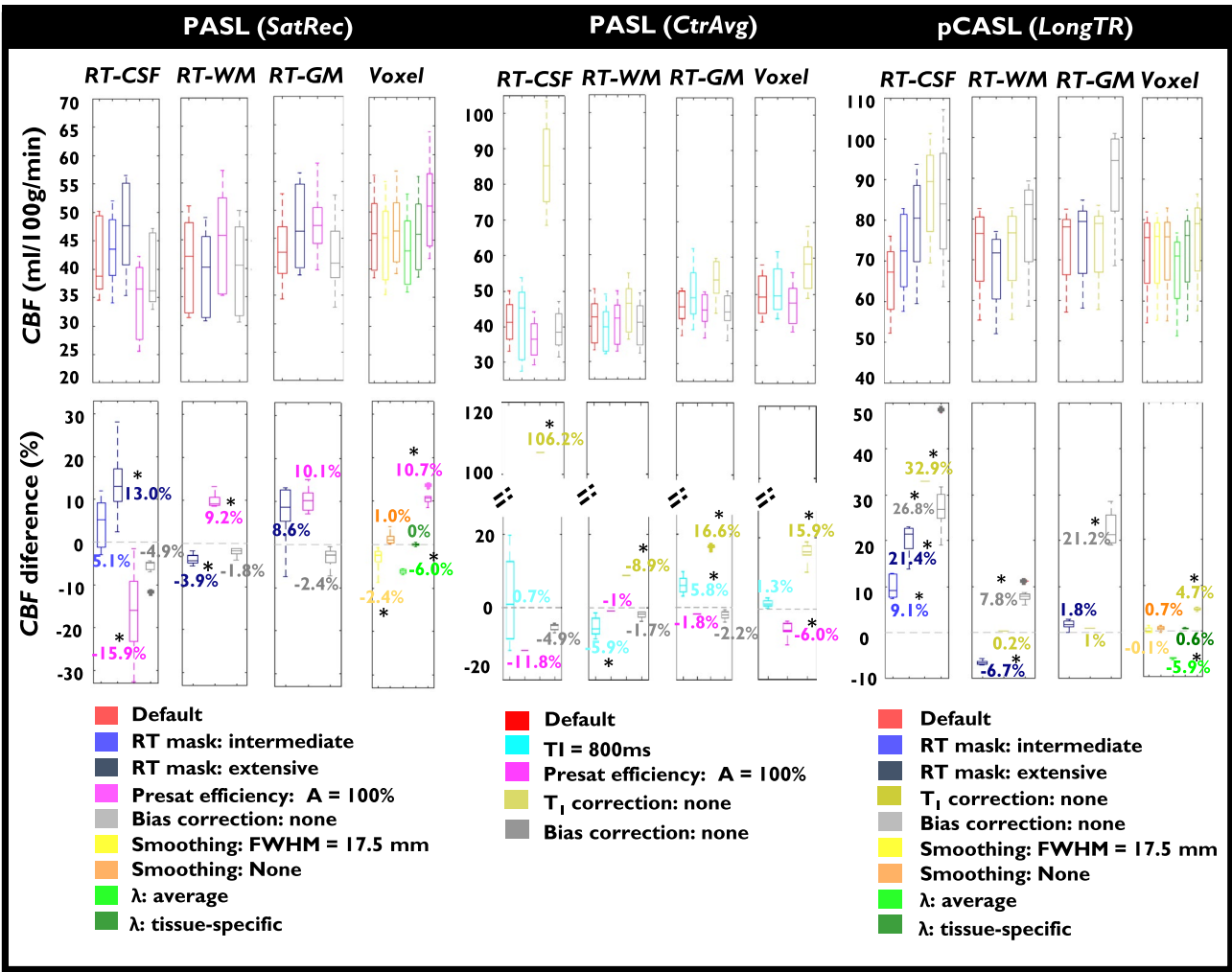


FIGURE 5 Group results for the GM average CBF values (top) and differences relative to default options (bottom), obtained when varying the calibration post-processing options, for each of the main calibration methods tested, in session 1 of PASL-SatRec (left), PASL-CtrAvg (middle), and pCASL-LongTR (right). Box plots represent the 25th and 75th percentiles, respectively. *Significant differences as assessed by pairwise t tests ($P < 0.05$)

uses a much more restrictive mask. WM masking options resulted in a significant CBF decrease, but only of ~5% relative to the default value. Fixing $A = 100\%$ instead of allowing its estimation ($A \sim 90\%$) yielded significant CBF differences relative to default option for all methods except for RT-GM.

Not correcting for field inhomogeneities using bias correction had small and non-significant impact on CBF quantification, although the impact was higher when using the RT-CSF approach compared with the other RT approaches. For the Voxel approach, an increase in kernel size for spatial

smoothing led to significant but small effects on CBF. Not applying any smoothing at all yielded non-significant differences. Finally, different options for the value of λ also resulted in significant but small CBF differences when using an average λ , whereas non-significant differences were observed when using a tissue-specific λ .

In contrast with the *SatRec* approach, the *CtrAvg* approach revealed great sensitivity to some post-processing options, particularly the T_1 correction, and more so when using the RT-CSF method. Significant increases in CBF as high as ~100% were found when using the RT-CSF method if no T_1 correction was carried out. When using $TI = 800$ ms, significant CBF differences were found for RT-WM and RT-GM methods compared to the default option.

When applying the pCASL *LongTR* method for M_{0r} generation, differences in CBF values were obtained with the different options, particularly when using the RT-CSF method. When not performing T_1 correction, the RT-CSF method yielded a significantly greater difference compared with the default options, with the Voxel method yielding a significant but much smaller difference. Consistently with the PASL *SatRec* results, less restrictive masks led to important and significant CBF differences, except for the RT-GM method, particularly in the case of the extensive CSF mask. Also, similarly with PASL *SatRec* but to a higher extent, bias correction had greater and always significant impact on all RT methods, particularly on RT-CSF and RT-GM. When using the Voxel method, the different spatial smoothing options yielded non-significant differences relative to the default option. Non-significant differences were also observed when using a tissue-specific λ but, in contrast, average λ yielded a statistically significant difference.

5 | DISCUSSION

We systematically compared the main ASL calibration methods as well as the associated post-processing options, in terms of CBF values and corresponding test–retest reproducibility.

5.1 | Comparison between calibration methods

When comparing the different calibration methods using the default post-processing options, minimal differences were observed in CBF within each data set (Figure 2). When focusing on the GM average CBF values (Figure 3), relatively small differences were found despite being significant in some cases.

Overall, the RT-CSF method yielded the lowest CBF values across M_{0r} estimation methods. Although, for the PASL *SatRec* approach, only RT-CSF and RT-WM methods were

significantly different from the Voxel method, for the pCASL *LongTR* approach, significantly lower CBF values were obtained when using the CSF as a reference tissue, compared with other strategies for deriving M_{0a} from M_{0r} . This finding may be explained by the fact that our pCASL data set, obtained using a 32-channel head coil, suffered from B_1 field inhomogeneities that were not fully accounted for by bias field correction. In fact, this interpretation is corroborated by the observation that, in the case of pCASL, when using less restrictive CSF masks (including the whole lateral ventricles rather than just a few central voxels where coil sensitivity is much lower), the CBF values increased considerably and became comparable to the ones obtained using the Voxel approach.

It should be noted that the bias field correction approach used here is not the ideal method for accounting for the coil sensitivity profile. Nevertheless, this strategy serves the purpose of evaluating the relative differences between calibration methods and post-processing options. In fact, it clearly emphasizes the impact of sensitivity correction on CBF, particularly when using the RT-CSF approach. Furthermore, the CBF values obtained with RT methods, when corrected for the bias field, become closer to the ones obtained with the Voxel method, which intrinsically corrects for this issue. Nevertheless, further work is necessary to fully estimate the impact of this strategy on calibration methods. Methods for the estimation of the coil sensitivity profile have been proposed,^{28,29} but these require the acquisition of additional data using imaging sequences that are not widely available. Alternatively, a proton density image acquired with a body coil can also be used to obtain an approximate sensitivity map.³⁰ However, this method is limited by the fact that it only corrects for non-uniformity in the receive field, not the transmit field, which is also not spatially uniform at field strengths of 3T or higher, and this correction may not account for other factors (e.g., prescan transmit settings).

It is possible to circumvent RF field inhomogeneity issues altogether by using the voxelwise method. In this case, M_{0a} is computed at each voxel based on the measured M_{0r} image and it is therefore affected by the same RF coil sensitivity as the ASL measurements themselves. Normalization of the estimated CBF map by the M_{0a} map therefore intrinsically corrects for coil sensitivity variations. This well-known effect has been explicitly reported,²⁰ with observed signal loss in anterior regions on both proton-density and perfusion-weighted ASL images being successfully restored in the final perfusion maps by calibration using a Voxel approach. Nevertheless, the voxelwise division intrinsic to this approach might lead to increased variability in CBF quantification because of noise amplification and also to edge and/or partial volume effects.³¹

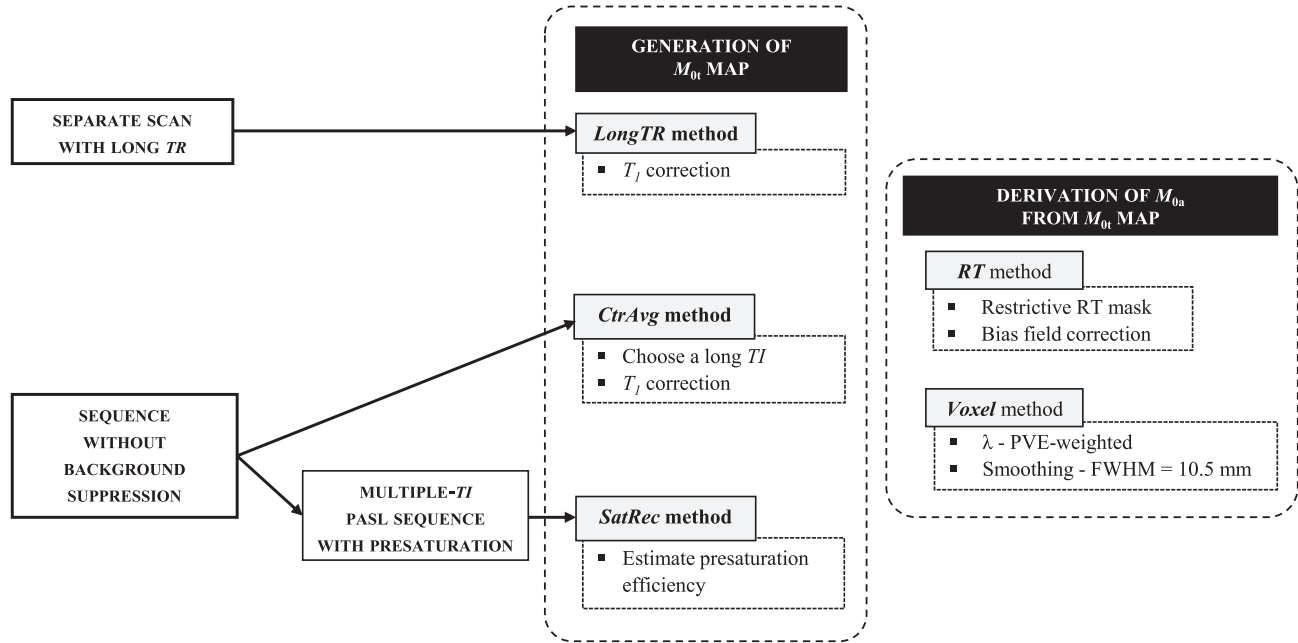


FIGURE 6 Summary of the main calibration methods, according to the data acquisition procedure (left) and with indication of the corresponding recommended options, for the generation of the M_{0r} map (middle) and the derivation of M_{0a} from the M_{0r} map (right)

5.2 | Impact of calibration post-processing options

We observed that some of the processing options had great impact on the final results whereas others were relatively unimportant. Interestingly, very few studies have investigated the impact of different calibration strategies, and critically, they are not in agreement. One study compared the 3 M_{0a} derivation schemes that we also tested here (RT-CSF, RT-WM, and Voxel) using data collected with 3 single-TI (1400 ms) PASL sequences (including PICORE-Q2TIPS) and found ~35% higher CBF values for the RT-CSF relative to the Voxel method.¹⁸ Another study also observed a discrepancy in CBF quantification when using different calibration methods based on single-PLD pCASL data, with RT-WM yielding the lowest CBF values and Voxel the highest (~20% difference).²⁰ A third study compared different calibration methods on CASL and pCASL data and found only minor differences. However, they reported slightly lower CBF values using RT-CSF compared with RT-WM possibly because of not correcting for the uneven sensitivity profile of the head coil.¹⁹

Our results highlight that one of the processing options with greatest impact was the correction of the M_{0r} map for incomplete T_1 relaxation in each tissue based on the TR of the respective image acquisition, particularly when using the RT-CSF method. In the current guidelines, this correction is only recommended when $TR < 5$ s.⁴ However, this recommendation is coupled with the use of the Voxel method for extrapolating M_{0a} , in which case we do not observe large effects of T_1 correction. In fact, the impact of T_1 correction is much greater if the RT-CSF derivation method is used, because CSF

has a much longer T_1 than GM or WM. In this case, a TR longer than 16 s would be required for an almost full recovery of the longitudinal magnetization. Therefore, we believe that this correction should be applied even if TR is longer than 5 s, when the RT-CSF calibration method is used. A related option with relatively high impact is the value of PLD/TI used to extract M_{0r} . In the case of the ContrAvg approach in PASL, using a lower TI leads to greater sensitivity in CBF quantification. For PASL, the presaturation efficiency value was also important, because this is sequence-specific and therefore should be chosen accordingly. The impact of the bias correction approach is highly dependent, not only on the M_{0r} derivation method, but also on the type of ASL data set and the RF coil sensitivity of the acquisition. In contrast with the 12-channel RF coil used for the PASL data set, the 32-channel RF coil used for the pCASL data set displayed a relatively heterogeneous sensitivity profile, leading to significant impact of bias correction, particularly when using RT-CSF and RT-GM methods.

Overall, our results indicate that when acquiring multiple-TI PASL data, the SatRec approach is preferable to the ContrAvg approach because the latter strongly depends on the choice of PLD/TI value and on T_1 correction. In terms of M_{0a} derivation, the RT-CSF approach was the most sensitive to variations in the processing options, making this approach more prone to CBF discrepancies. In addition, variability in the CBF differences across subjects was also greater than with other M_{0a} derivation methods. The RT-CSF method is highly dependent, not only on T_1 correction as discussed before, but also on the tissue mask and whether or not bias correction is carried out. The RT-WM and RT-GM methods

are also dependent on these parameters but to a lesser extent. Notwithstanding, the RT-WM approach might not be feasible whenever WM lesions are present, such as in multiple sclerosis or small vessel disease. On the other hand, the RT-GM approach is highly influenced by PVE, rendering the use of restrictive GM masks mandatory. Nevertheless, approaches for PVE correction most often target only the relative CBF images and not M_0 measures.^{31,32} Therefore, we believe that this correction should not affect our results concerning the comparison of calibration methods. Similarly to the RT-WM and RT-GM methods, the Voxel method yielded relatively small CBF differences across different processing options. In fact, some options had negligible impact on CBF quantification, namely the degree of smoothing. As expected, the use of an averaged, fixed λ underestimates GM CBF. A summary scheme of recommended calibration procedures, as well as the corresponding options that should be applied and/or reported, is depicted in Figure 6.

5.3 | CBF values, spatial distributions and reproducibility

The maps obtained in our study exhibit spatial distributions that are consistent with the expected variations across tissues and brain regions.^{5,15,33} All average GM CBF values found in our study were within the wide range of values reported in the literature. However, the PASL data set systematically produced lower CBF values than the pCASL data set. Surprisingly, in a study using 5 different commonly used ASL sequences, the opposite pattern was seen between single-PLD pCASL and multiple-TI PASL using the RT-CSF method (25/40 mL/100 g/min for pCASL/PASL).⁶ In another study, similar values were found for both PASL and pCASL single-PLD/TI sequences (62/60 mL/100 g/min for PASL/pCASL), using different calibrations strategies (extra calibration scan for PASL/control averaging for pCASL).³⁴ The differences between pCASL and PASL observed here are of unknown cause and are likely explained partially by specific implementation issues.

The inter- and intra-subject coefficients of variation obtained in our study for GM average CBF are all within intervals of good reproducibility. For PASL, $CV_{intra} \sim 10\text{--}12\%$ is in agreement with the literature. A multiple-TI QUASAR ASL study using a model-free approach reported within-week $CV_{intra} \sim 10\%$,³⁵ whereas a study from our group achieved within-week CV_{intra} values in the range of 10–21%.¹⁵ A single-TI PASL study, using a sequence similar to the 1 used in our study, found within-week $CV_{intra} \sim 6\%$.⁵ Regarding pCASL, our results of $\sim 6\text{--}7\%$ are also in accordance with the literature. A single-PLD pCASL study using 2 different PLD values yielded $CV_{intra} \sim 5\text{--}10\%$, which increased with PLD.³⁶ Consistently, a multiple-PLD pCASL study obtained $CV_{intra} \sim 5\%$ for average week-repeat GM CBF value.²³

5.4 | Limitations and future work

Although our approach aimed to systematically and comprehensively test all relevant processing options in the calibration methods used, options regarding relatively less important parameters could also be tested, in particular the tissue-specific T_2^* used to correct for T_2^* decay when deriving the M_{0r} maps. Another option that could be tested is the use of subject-specific or voxelwise T_1 values, which could be estimated directly when using the *SatRec* approach in PASL or acquired separately. Importantly, some of the calibration methods may not be appropriate to use in certain clinical populations. For example, in patients with WM abnormalities, such as in multiple sclerosis or small vessel disease, the RT-CSF or RT-GM methods may be preferable to the RT-WM.

Furthermore, we acknowledge that some options could not be tested because of the lack of data availability. In particular, the *LongTR* approach should be also compared using PASL, to determine the best method for M_{0r} generation. Additionally, a more precise approach for correction of the coil sensitivity profile should be carried out. Furthermore, we also acknowledge that the EPI images suffer from susceptibility-induced geometric distortions that might affect image co-registration. This can be circumvented by acquiring a B_0 field map and using this to correct such distortions. Finally, a prospective study assessing the impact of calibration strategies by acquiring both PASL and pCASL on the same subjects should be conducted to further validate our results.

6 | CONCLUSION

In conclusion, we found that considerable discrepancies in CBF values can be obtained when using poor choice simplifications in the post-processing calibration of pCASL and PASL data. Nevertheless, strategies based on reference tissue or voxelwise methods both perform well when carefully implemented in terms of CBF quantification and reproducibility. In general, the greatest sensitivity was found for correction for incomplete T_1 relaxation when using proton density reference images for calibration. Correction for RF field inhomogeneities also had great impact, as did the value of presaturation efficiency, whereas the values of brain–blood water partition coefficient had moderate impact. In contrast, the degree of spatial smoothing applied to the calibration images or the mask used for the reference tissue had minor effects. The voxelwise approach, as proposed in the ASL white paper, has the advantage of simplicity, while the reference tissue methods are more vulnerable to sensitivity and relaxation correction errors, in particular when using CSF. Nevertheless, our results emphasize the need for consistent post-processing options across studies as well as the need for a complete description of the various post-processing

calibration options so that absolute CBF quantification is effectively achieved. Failure to take the impact of these fine calibration options into account would seriously compromise the use and applicability of ASL perfusion imaging.

ACKNOWLEDGMENTS

This work was supported by the Portuguese Foundation for Science and Technology (FCT) grants PTDC/BBB-IMG/2137/2012, PD/BD/135114/2017 and UID/EEA/50009/2019; the Engineering and Physical Sciences Research Council UK (EP/P012361/1) and the Wellcome Trust (203139/Z/16/Z). The authors wish to express their gratitude to the European Cooperation in Science and Technology-Arterial Spin Labelling Initiative in Dementia (COST-AID Action BM1103). We are also grateful for the insightful comments and recommendations of the Referees.

DATA AVAILABILITY STATEMENT

The underlying data associated with figures in this study are available from the Oxford University Research Archive (ORA-Data) (DOI: 10.5287/bodleian:xQx8Vnv5e). For enquiries regarding the imaging data contact: ARM for the pCASL data (andrew.segerdahl@ndcn.ox.ac.uk) or PF for the PASL data (patricia.figueiredo@tecnico.ulisboa.pt).

ORCID

Joana Pinto  <https://orcid.org/0000-0002-6863-1883>

Michael A. Chappell  <https://orcid.org/0000-0003-1802-4214>

Thomas W. Okell  <https://orcid.org/0000-0001-8258-0659>

Patrícia Figueiredo  <https://orcid.org/0000-0002-0743-0869>

REFERENCES

- Detre JA, Leigh JS, Williams DS, Koretsky AP. Perfusion imaging. *Magn Reson Med*. 1992;23:37–45.
- Williams DS, Detre JA, Leigh JS, Koretsky AP. Magnetic resonance imaging of perfusion using spin inversion of arterial water. *Proc Natl Acad Sci U S A*. 1992;89:212–216.
- Alsop DC, Detre JA. Reduced transit-time sensitivity in noninvasive magnetic resonance imaging of human cerebral blood flow. *J Cereb Blood Flow Metab*. 1996;16:1236–1249.
- Alsop DC, Detre JA, Golay X, et al. Recommended implementation of arterial spin-labeled perfusion MRI for clinical applications: a consensus of the ISMRM perfusion study group and the European consortium for ASL in dementia. *Magn Reson Med*. 2015;73:102–116.
- Wang Y, Saykin AJ, Pfeuffer J, et al. Regional reproducibility of pulsed arterial spin labeling perfusion imaging at 3T. *NeuroImage*. 2011;54:1188–1195.
- Gevers S, van Osch MJ, Bokkers RPH, et al. Intra- and multicenter reproducibility of pulsed, continuous and pseudo-continuous arterial spin labeling methods for measuring cerebral perfusion. *J Cereb Blood Flow Metab*. 2011;31:1706–1715.
- Mutsaerts HJMM, Steketee RME, Heijtel DFR, et al. Reproducibility of pharmacological ASL using sequences from different vendors: implications for multicenter drug studies. *MAGMA*. 2015;28:427–436.
- Gil-Gouveia R, Pinto J, Figueiredo P, Vilela PF, Martins IP. An arterial spin labeling MRI perfusion study of migraine without aura attacks. *Front Neurol*. 2017;8:280.
- Pimentel M, Vilela P, Sousa I, Figueiredo P. Localization of the hand motor area by arterial spin labeling and blood oxygen level-dependent functional magnetic resonance imaging. *Hum Brain Mapp*. 2013;34:96–108.
- MacIntosh BJ, Lindsay AC, Kyliantiras I, et al. Multiple inflow pulsed arterial spin-labeling reveals delays in the arterial arrival time in minor stroke and transient ischemic attack. *Am J Neuroradiol*. 2010;31:1892–1894.
- Figueiredo P, Clare S, Jezzard P. Quantitative perfusion measurements using pulsed arterial spin labeling: effects of large region-of-interest analysis. *J Magn Reson Imaging*. 2005;21:676–682.
- Gonzalez-At JB, Alsop DC, Detre JA. Cerebral perfusion and arterial transit time changes during task activation determined with continuous arterial spin labeling. *Magn Reson Med*. 2000;43:739–746.
- Buxton RB, Frank LR, Wong EC, Siewert B, Warach S, Edelman RR. A general kinetic model for quantitative perfusion imaging with arterial spin labeling. *Magn Reson Med*. 1998;40:383–396.
- Chappell MA, MacIntosh BJ, Donahue MJ, Günther M, Jezzard P, Woolrich MW. Separation of macrovascular signal in multi-inversion time arterial spin labelling MRI. *Magn Reson Med*. 2010;63:1357–1365.
- Sousa I, Vilela P, Figueiredo P. Reproducibility of the quantification of arterial and tissue contributions in multiple postlabeling delay arterial spin labeling. *J Magn Reson Imaging*. 2014;40:1453–1462.
- Wu WCC, St Lawrence KS, Licht DJ, Wang DJ. Quantification issues in arterial spin labeling perfusion magnetic resonance imaging. *Top Magn Reson Imaging*. 2010;21:65–73.
- Luh WM, Wong EC, Bandettini PA, Hyde JS. QUIPSS II with thin-slice TI1 periodic saturation: a method for improving accuracy of quantitative perfusion imaging using pulsed arterial spin labeling. *Magn Reson Med*. 1999;41:1246–1254.
- Cavuşoğlu M, Pfeuffer J, Uğurbil K, Uludağ K. Comparison of pulsed arterial spin labeling encoding schemes and absolute perfusion quantification. *Magn Reson Imaging*. 2009;27:1039–1045.
- Chen Y, Wang Z, Detre JA. Impact of equilibrium magnetization of blood on ASL quantification. In *Proceeding of the 19th Annual Meeting of ISMRM*, Montreal, Canada, 2011. p. 300.
- Fazlollahi A, Bourgeat P, Liang X, et al. Reproducibility of multiphase pseudo-continuous arterial spin labeling and the effect of post-processing analysis methods. *NeuroImage*. 2015;117:191–201.
- Zhang Y, Brady M, Smith S. Segmentation of brain MR images through a hidden Markov random field model and the expectation-maximization algorithm. *IEEE Trans Med Imaging*. 2001;20:45–57.
- Evans AC, Collins DL, Mills SR, Brown ED, Kelly RL, Peters TM. 3D statistical neuroanatomical models from 305 MRI volumes. In *Proceedings of the IEEE Conference Record Nuclear Science Symposium and Medical Imaging Conference*, San Francisco, CA, 1993. p. 1813–1817.
- Mezue M, Segerdahl AR, Okell TW, Chappell MA, Kelly ME, Tracey I. Optimization and reliability of multiple postlabeling

- delay pseudo-continuous arterial spin labeling during rest and stimulus-induced functional task activation. *J Cereb Blood Flow Metab.* 2014;34:1919–1927.
24. Jenkinson M, Bannister P, Brady M, Smith S. Improved optimization for the robust and accurate linear registration and motion correction of brain images. *NeuroImage.* 2002;17:825–841.
 25. Chappell MA, Woolrich MW, Kazan S, Jezzard P, Payne SJ, MacIntosh BJ. Modeling dispersion in arterial spin labeling: validation using dynamic angiographic measurements. *Magn Reson Med.* 2013;69:563–570.
 26. Pinto J, Jorge J, Sousa I, Vilela P, Figueiredo P. Fourier modeling of the BOLD response to a breath-hold task: optimization and reproducibility. *NeuroImage.* 2016;135:223–231.
 27. Tjandra T, Brooks J, Figueiredo P, Wise RG, Matthews PM, Tracey I. Quantitative assessment of the reproducibility of functional activation measured with BOLD and MR perfusion imaging: implications for clinical trial design. *NeuroImage.* 2005;27:393–401.
 28. Dai W, Robson PM, Shankaranarayanan A, Alsop DC. Sensitivity calibration with a uniform magnetization image to improve arterial spin labeling perfusion quantification. *Magn Reson Med.* 2011;66:1590–1600.
 29. Ahlgren A, Wirestam R, Knutsson L, Petersen ET. Improved calculation of the equilibrium magnetization of arterial blood in arterial spin labeling. *Magn Reson Med.* 2018;80:2223–2231.
 30. Okell TW, Chappell MA, Kelly ME, Jezzard P. Cerebral blood flow quantification using vessel-encoded arterial spin labeling. *J Cereb Blood Flow Metab.* 2013;33:1716–1724.
 31. Chappell MA, Groves AR, MacIntosh BJ, Donahue MJ, Jezzard P, Woolrich MW. Partial volume correction of multiple inversion time arterial spin labeling MRI data. *Magn Reson Med.* 2011;65:1173–1183.
 32. Asllani I, Borogovac A, Brown TR. Regression algorithm correcting for partial volume effects in arterial spin labeling MRI. *Magn Reson Med.* 2008;60:1362–1371.
 33. Mutsaerts HJ, van Osch MJP, Zelaya FO, et al. Multi-vendor reliability of arterial spin labeling perfusion MRI using a near-identical sequence: implications for multi-center studies. *NeuroImage.* 2015;113:143–152.
 34. Chen Y, Wang D, Detre JA. Test-retest reliability of arterial spin labeling with common labeling strategies. *J Magn Reson Imaging.* 2011;33:940–949.
 35. Petersen ET, Mouridsen K, Golay X. The QUASAR reproducibility study, part II: results from a multi-center arterial spin labeling test-retest study. *NeuroImage.* 2010;49:104–113.
 36. Wu B, Lou X, Wu X, Ma L. Intra- and interscanner reliability and reproducibility of 3D whole-brain pseudo-continuous arterial spin-labeling MR perfusion at 3T. *J Magn Reson Imaging.* 2014;39:402–409.

SUPPORTING INFORMATION

Additional supporting information may be found online in the Supporting Information section at the end of the article.

FIGURE S1 Illustrative examples of MPRAGE images registered to ASL space (bottom) and respective CSF (blue) and GM (yellow) masks retrieved from MPRAGE image segmentation (top), for the PASL and pCASL datasets

How to cite this article: Pinto J, Chappell MA, Okell TW, et al. Calibration of arterial spin labeling data—potential pitfalls in post-processing. *Magn Reson Med.* 2020;83:1222–1234. <https://doi.org/10.1002/mrm.28000>