Implementation and validation of ASL perfusion measurements for population imaging

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Purpose: Pseudocontinuous arterial spin labeling (pCASL) allows for noninvasive measurement of regional cerebral blood flow (CBF), which has the potential to serve as biomarker for neurodegenerative and cardiovascular diseases. This work aimed to implement and validate pCASL on the dedicated MRI system within the population-based Rotterdam Study, which was installed in 2005 and for which software and hardware configurations have remained fixed.

Methods: Imaging was performed on two 1.5T MRI systems (General Electric); (I) the Rotterdam Study system, and (II) a hospital-based system with a product pCASL sequence. An in-house implementation of pCASL was created on scanner I. A flow phantom and three healthy volunteers (<27 years) were scanned on both systems for validation purposes. The data of the first 30 participants (86 ± 4 years) of the Rotterdam Study undergoing pCASL scans on scanner I only were analyzed with and without partial volume correction for gray matter.

Results: The validation study showed a difference in blood flow velocity, sensitivity, and spatial coefficient of variation of the perfusion-weighted signal between the two scanners, which was accounted for during post-processing. Gray matter CBF for the Rotterdam Study participants was 52.4 ± 8.2 ml/100 g/min, uncorrected for partial volume effects of gray matter. In this elderly cohort, partial volume correction for gray matter had a variable effect on measured CBF in a range of cortical and subcortical regions of interest.

Conclusion: Regional CBF measurements are now included to investigate novel biomarkers in the Rotterdam Study. This work highlights that when it is not feasible to purchase a novel ASL sequence, an in-house implementation is valuable.

Keywords
arterial spin labeling, cerebral blood flow, flow phantom, population imaging
1  INTRODUCTION

Measurement of regional cerebral blood flow (CBF) with pseudocontinuous arterial spin labeling (pCASL) is a cost-effective and safe option for repeated assessment of regional cerebral perfusion without the need to administer contrast media. This technique allows for quantification of CBF in ml/100 g/min and is, therefore, especially attractive for large and longitudinal population-based studies in which healthy participants undergo MRI scans at multiple time points along the course of the study period.

The Rotterdam Study is a prospective population-based cohort study ongoing since 1990 in the city of Rotterdam in The Netherlands, which targets cardiovascular, endocrine, hepatic, neurological, ophthalmic, psychiatric, dermatological, otolaryngological, locomotor, and respiratory diseases. The Rotterdam Scan Study is a substudy and focuses specifically on (repeat) brain MRI examinations to study neurodegenerative and cerebrovascular disease and was officially embedded in the core protocol of the Rotterdam Study in 2005, when a dedicated MRI system was installed in the Rotterdam Study Research Centre. To maintain consistency in image acquisition, the MRI hardware and software have not changed since.

Evidence for regional CBF as a biomarker in healthy aging and cardiovascular disease, as well as neurodegeneration is steadily building in the literature. However, the original MRI protocol of the Rotterdam Scan Study only contained a phase contrast assessment of blood flow through major brain feeding arteries, assessing total CBF. The reason for this is that, despite ASL first being introduced in the early 1990s, the adoption of ASL as one of the standard sequences available on MRI systems is only a recent development due to the 2015 consensus paper.

Here we present the process of implementing and validating pCASL on the MRI system within the Rotterdam Study Research Centre. The validation of this pCASL sequence was done by CBF measurements of a novel perfusion phantom and healthy volunteers and comparing these to CBF measurements with a novel product pCASL sequence on one of the 1.5T MRI systems at the main site of the Erasmus Medical Centre in Rotterdam. In addition, the first 30 ASL datasets were assessed for their ability to measure regional CBF, corrected for partial volume estimates of gray matter, in the eldest cohort of participants of the Rotterdam Study.

We present this work to illustrate how local, multidisciplinary collaborations between MR physicists, engineers, and radiologists can result in successful implementation of a novel imaging technique. The need for ASL in the Rotterdam Study was given by a radiologist and epidemiologist and the process of programming and compiling the ASL sequence was done by physicists. During the optimization of the image analysis pipeline, the role of the engineer was to show that the sequence was working as intended (introduction of the phantom experiment) and communicate between the physics and radiologists teams in optimizing the processing pipeline. The final implementation was done via close collaboration between the MR Physics and Population-Based Imaging groups at the department of Radiology and Nuclear Medicine of the Erasmus MC. This work shows that porting a novel MRI sequence to an older MRI system is valuable and, therefore, may be beneficial for sites that do not have the option to purchase novel ASL sequences for their MRI systems and software versions.

2  METHODS

All imaging was performed on 1.5T MRI systems (General Electric Healthcare, Milwaukee, WI, USA). Experiments were conducted under approval of the institutional ethics committee, in compliance with the declaration of Helsinki.

2.1 Implementation and validation of ASL in population based imaging

Imaging was performed on two different 1.5T MRI systems; (a) the population imaging scanner, a Signa EXCITE with software version 11 (released November 2003) and an 8-channel head coil; (b) an Optima MR450w 1.5T with software version DV26 (released June 2017) and a 24-channel head coil, which includes the General Electric product version of pCASL. The implementation of pCASL at scanner I was done in-house, converting the product pCASL sequence from software version DV26 to 11. This was done by mimicking the product sequence as much as possible hereby keeping up with the theoretical background of the implementation of the product sequence. This was a debugging process that required downgrading novel functions and input structures from DV26, ensuring that the sequence could be compiled for software version 11. This particular part of porting back the sequence is highly dependent on both software versions and required close collaboration with GE. The final step in coding the in-house pulse sequence was to fine-tune the unbalancing of the radiofrequency (RF) gradients used to generate the label and control images. This process ensures that the effects of phase shifts between RF pulses in the pseudocontinuous labeling caused by magnetic field inhomogeneities at the labeling plane are minimalized, ensuring that the labeling efficiency of the pCASL scan was as high as possible. Please note that we cannot publish any code of the pulse sequence, since this is GE proprietary information. However, via the GE research collaboration portal, we are open to collaborations in which this experience is shared.
Acquisition parameters were the same for both pCASL implementations; 3D spiral gradient echo read-out with background suppression, 512 points per arm, eight arms, reconstruction diameter = 200 mm, reconstruction matrix 128 x 128 x 30, slice thickness 4 mm, post label delay (PLD) 1.525 s, label duration 1.45 s, number of excitations = 3, echo time = 0.011 s, repetition time = 4.6 s. Each pCASL acquisition included a proton density weighted image for normalization (the same acquisition parameters, but no labeling applied).

2.2  |  Phantom experiment

A perfusion phantom (beta test version of QASPER, Gold Standard Phantoms, London, UK), which allows for setting variable continuous flow rates, was used. Measurements were done on both MRIs systems at two different days, 2 weeks apart. Per session, 12 pCASL scans were made in random order for different flow rates (200-475 ml/min, steps of 25 ml/min). Voxelwise perfusion-weighted images (ΔM/M0) were calculated for each pCASL scan.\(^9\) Median values were calculated over a circular region of interest (ROI), which was placed over the perfusion region of the phantom via visual inspection. A scaling factor between the two sequences was calculated by dividing the ROI median ΔM/M0,0.26 by the ROI median ΔM/M0,11.

2.3  |  Healthy, young volunteer experiment

Three young healthy volunteers (all female, <27 years) underwent pCASL scanning on both MR systems on the same day. Perfusion-weighted pCASL images before quantification were compared by calculating the signal-to-noise ratio (SNR) of ΔM as the mean perfusion-weighted signal in the GM divided by the standard deviation of a circular ROI (radius of 10 voxels) well outside the brain\(^12\) and by calculating the spatial coefficient of variation (CoV) within the whole brain GM ROI.\(^13\) The whole brain GM mask was obtained via segmentation of a T1-weighted structural scan (fast in FSL), which was linearly registered to the pCASL image (flirt in FSL) and only included voxels with partial volume estimates for GM (PVEGM) > 70%. CBF maps were quantified with oxford_asl (FSL, version 6.0.1, Oxford, UK),\(^7\) taking into account the scaling factor between the two implementations resulting from the phantom experiment. In addition, 16 cortical and sub-cortical ROIs from the Harvard-Oxford atlas (thresholded at partial volume estimates > 25%) were non-linearly transformed from MNI standard space to the participant’s pCASL space and used to calculate the absolute differences in ROI CBF calculated between scanner I and scanner II.

2.4  |  Application of ASL imaging in the Rotterdam scan study

Thirty elderly participants (20 females, mean age 86 ± 4 years) of the RS-I cohort in the Rotterdam Study (study entry in 1989-1993) underwent scanning on scanner I only (Sep-Nov 2018). SNR, CoV, and ROI average CBF were calculated in the same manner as for the young volunteers. To show the potential and advantage of being able to investigate regional CBF, a comparison was made of CBF calculated with and without PVE correction to assess the effect of local GM volume (BASIL within FSL\(^14\)).

3  |  RESULTS

3.1  |  Implementation and validation of pCASL

The flow phantom results illustrate a difference in flow sensitivity of ΔM/M0 between the two pCASL implementations, with the pCASL on scanner I being more sensitive to increasing flow velocities (Figure 1). The scaling factor between the two pCASL implementations has an exponential relationship, with an asymptote at 30.9 (Figure 1B).

For the healthy, young volunteers the SNR of the perfusion-weighted signal directly resulting from image reconstruction was smaller and CoV was larger for scanner I than for scanner II (Table 1 and Figure 2). When taking the scaling factor and additional smoothing into account, group averaged CBF\(_{GM}\) (uncorrected for PVE of GM) for scanner I and II in the healthy volunteers were similar: 63.3 ± 1.8 ml/100 g/min and 62.2 ± 2.0 ml/100 g/min, respectively. Figure 3 shows the differences in CBF between the two scanners for the whole brain GM and 16 cortical and sub-cortical ROIs calculated for all three volunteers. Note that, except for the brainstem, ROI differences in CBF for the two scanners are on the order of ±6 ml/100 g/min.

3.2  |  Application of ASL imaging in the Rotterdam scan study

The dataset of one elderly participant was excluded due to severe head motion during the pCASL acquisition. Group averaged SNR of the perfusion-weighted signal was lower for the elderly participants than for the younger, while spatial CoV of the perfusion-weighted signal for the whole brain GM was higher than for the young volunteers. Both values are reported in Table 1. Group averaged whole brain GM CBF for the 29 remaining elderly participants was increased when correcting for PVE\(_{GM}\): 52.4 ± 8.2 ml/100 g/min vs 63.0 ± 8.9 ml/100 g/min, uncorrected and corrected for PVE\(_{GM}\).
respectively (paired t test, *P* < .001). Regional investigation showed that the increase in GM CBF with partial volume correction varies across the brain, ranging from no significant change in CBF in brainstem (paired t test, *P* = 0.353) and cerebellum (paired t test, *P* = 0.345) to a little over 50% in the caudate nucleus (Figure 4).

### DISCUSSION

This work shows the in-house implementation and validation of pCASL on a population imaging MRI system, with an outdated software platform. The first 30 pCASL datasets acquired in the Rotterdam Study highlight the potential of using pCASL for monitoring regional cerebral perfusion in population imaging, even on older software platforms.

The flow phantom scans led to the inclusion of a scaling factor in the quantification of CBF in the pCASL scans of the population-based scanner. The results of the phantom experiment indicate that labeling efficiency decreases at lower blood flow velocities (<30 cm/s) in our in-house implementation of pCASL. This is likely due to the difference in gradient coils between the two scanners. The asymptote of the scaling factor between the two different implementations (Figure 1B) suggests that, in the healthy population, a constant scaling factor is sufficient to correct for this difference in labeling efficiency. However, care should be taken in participants of the Rotterdam Study with blood flow velocities through the major cerebral arteries (<30 cm/s) in the labeling plane of the pCASL sequence. Within the Rotterdam Scan Study protocol phase contrast measurements of the flow velocities through the internal carotid and basilar arteries are measured. Therefore, the non-constant scaling factor as measured with the flow phantom is being incorporated in the CBF analysis of the complete cohort.

The use of a flow phantom is highly recommended when comparing pCASL implementations, because it allows for an objective manner to test the reproducibility of the CBF measurements between two different scanners. However, should a flow phantom not be available, it is important to note that, in healthy volunteers, cerebral perfusion is variable due to several physiological parameters, including blood caffeine levels, blood pressure, hormone levels, etc. In absence of a flow phantom, it is important to compare sequences/scanners in light of previously reported reproducibility of ASL-based measurements of CBF.

There are different reconstruction engines implemented in the software versions of scanner I and scanner II, which results in different inherent scaling of the perfusion-weighted signal compared with the M0 image as well as different smoothing of images directly resulting from the reconstruction. The reconstruction engine as implemented in scanner II (DV26), does not allow for the reconstruction of the raw ΔM images and therefore it was not feasible to investigate ΔM images without smoothing for scanner II. The differences in smoothing between the scanners leads to the higher spatial CoV in the perfusion-weighted images of scanner I, clearly seen in the healthy volunteers (Figure 2), which is the reason for including additional spatial smoothing in the post processing of these images.
FIGURE 2  Example of perfusion weighted (ΔM) images of one young volunteer. Note the difference in smoothing occurring during the reconstruction of the images. On scanner I with software version 11 (top row), no smoothing is implemented during reconstruction. On scanner II with software version DV26 (bottom row), smoothing occurs during reconstruction of the images.

FIGURE 3  Absolute differences in regional CBF estimates between scanner I and scanner II in healthy volunteers. Note that these differences are calculated in ml/100 g/min. The dotted line is there as a reference at 0, that is, represents no difference in ROI average CBF between scanner I and scanner II. GM = gray matter, BS = brainstem, FL = frontal lobe, OL = occipital lobe, TL = temporal lobe, PL = parietal lobe, Ce = cerebellum, Ins = insular cortex, ACC = anterior cingulate cortex, PCC = posterior cingulate cortex, Tha = thalamus, Cau = caudate nucleus, Put = putamen, Pal = pallidum, Hip = hippocampus, Am = amygdala, Ac = Accumbens.

FIGURE 4  Top: Group averaged (N = 29) regional CBF values for elderly participants of the Rotterdam Scan Study, uncorrected and corrected for partial volume estimates of gray matter (GM). The error bars indicate the standard deviation across the group. The dotted line indicates the group averaged, uncorrected value for whole brain grey matter CBF. Bottom: The percentage in CBF per ROI after partial volume correction. GM = gray matter, BS = brainstem, FL = frontal lobe, OL = occipital lobe, TL = temporal lobe, PL = parietal lobe, Ce = cerebellum, Ins = insular cortex, ACC = anterior cingulate cortex, PCC = posterior cingulate cortex, Tha = thalamus, Cau = caudate nucleus, Put = putamen, Pal = pallidum, Hip = hippocampus, Am = amygdala, Ac = Accumbens. *Significantly larger than 0, P < .001. (Bonferroni-corrected P-value = .0029)
images. The higher SNR for the perfusion-weighted signal on scanner II is likely caused by both the scaling and smoothing differences during reconstruction. However, note that the SNR for $\Delta M$ for young volunteers at scanner I is already in line with the results for 3D acquisitions as reported by Vidorreta et al.\textsuperscript{12} (and using the latest reconstruction engine clearly improves this value). Rather than changing these aspects of the reconstruction engines, which can lead to revoking CE/FDA marking of the system, we opted for adapting the post-processing pipeline of the pCASL data for scanner I.

Limited differences were found between regional CBF measured with pCASL on scanner I and scanner II when taking the smoothing and scaling differences between the two software versions into account in the processing pipeline of perfusion-weighted images from scanner I. The variability of CBF measured with ASL at different scanners is known to fluctuate in the order of >17%, as reported previously.\textsuperscript{15} We therefore deemed the ±6 ml/100 g/min (<10% of the whole brain GM average) an acceptable range of fluctuation for the validation of this exam. The exception to this is the region of the brainstem, with a lower average CBF in scanner I. From previous work it is known that ASL requires optimisation for accurate brainstem CBF measurement.\textsuperscript{16} Our validation study highlights this aspect and indicates that it may not be feasible to make inferences about CBF in the brainstem within the Rotterdam Scan study. Note that extending the number of young healthy volunteers could have aided a more thorough comparison in regional CBF measurements between scanner I and scanner II. However, the combination of the phantom and young volunteer results gave us confidence that the pCASL implementation on scanner I was functioning appropriately and application in the Rotterdam Scan Study followed.

The added value of measuring regional CBF within the Rotterdam Scan Study is illustrated by the first datasets of the elderly cohort. The regional PVE\textsubscript{GM} correction for CBF (Figure 4) illustrates the capability of assessing the effect of local GM volume on CBF measurement and highlights that the underestimation of underlying GM CBF due to partial volume effects is dependent on the ROI. The latter can be explained by the aspect that for subcortical gray matter structures (such as the caudate nucleus) the voxel size within the ASL sequence allows for limited voxels solely consisting of GM. The possibility of taking these regional variations in GM volume into account for the CBF measurements will allow for disentangling the effect of changes in measured CBF due to a true change in GM perfusion or to GM atrophy.\textsuperscript{14} Note that this is valuable information in the context of the longitudinal Rotterdam Study and the incorporation of the current ASL processing pipeline within the analysis of the complete dataset of this cohort is now in progress. Future work with this data set, therefore, includes investigating the effect of aging and GM atrophy on regional CBF measurements.

A limitation of this work is the PLD of 1525 ms, which we chose based on the $T_1$ of blood at 1.5T (1350 ms), the knowledge that the majority of the participants being scanned in the Rotterdam Scan Study are healthy, and to keep the imaging protocol consistent across all cohorts. Although this PLD led to SNR and CoV values comparable with the literature for the young volunteers, the lower SNR and higher CoV for the elderly participants may be improved with a longer PLD, for example, 2000 ms as recommended by the ASL White Paper.\textsuperscript{7} This is based on the assumptions that the longer PLD will lead to less macro-vascular artifacts within the perfusion-weighted images, because it would accommodate the expected longer arterial arrival times occurring in healthy aging\textsuperscript{7} and that this affects the SNR and CoV in a stronger positive manner than the decrease in signal of the labeled blood because of $T_1$ relaxation during the prolonged PLD. In summary, this work illustrates the successful application of an in-house implementation of pCASL in the Rotterdam Scan Study, including validation of this sequence with a novel flow phantom. The regional CBF data that are now acquired within the study population will be used in the future to investigate novel biomarkers of neurodegenerative and cardiovascular disease in the Rotterdam Study.

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CONFLICT OF INTEREST

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