Arterial Spin Labeling Perfusion of the Brain: Emerging Clinical Applications

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Supported in part by COST Action BM1103 on ASL in dementia
Essentials
- Due to the close link between brain metabolism and perfusion, patterns of FDG PET closely resemble those of ASL (page 11)
- ASL complements structural information of standard MRI in neurocognitive decline and may allow early diagnosis of dementia (page 11)
- ASL may detect mismatch perfusion in acute stroke, and identify tissue at risk in chronic cerebrovascular disease (page 13)
- ASL may detect arterio-venous shunting in arterio-venous malformation and fistulas (page 16)
- ASL may be used to localize the epileptogenic focus in seizure disorders (page 17).
- In neoplasms, ASL may replace standard Gadolinium enhanced dynamic susceptibility contrast (DSC) notably in patients with allergies, renal insufficiency or children (page 18)
- ASL may detect subtle functional changes in psychiatric disorders such as post-traumatic stress disorder or mild traumatic brain injury, where standard structural MRI typically provides no detectable anomaly (page 21)
- Emerging new applications of ASL include super-selective ASL to map vascular territories and cerebrovascular reserve imaging (page 32)

Disclosure
No conflicts of interest
Abstract

Arterial spin labeling (ASL) is an MR imaging technique used to assess cerebral blood flow (CBF) non-invasively by magnetically labeling inflowing blood. In this article we review the main labeling techniques, notably pulsed and pseudo-continuous ASL as well as emerging clinical applications. In dementia, the pattern of hypoperfusion on ASL closely matches the established patterns of hypometabolism on FDG-PET due to the close coupling of perfusion and metabolism in the brain. This suggests that ASL might be considered as an alternative for FDG, reserving PET to be used for the molecular disease-specific amyloid and tau tracers. In stroke, ASL can be used to assess perfusion alterations both in the acute and the chronic phase. In arteriovenous malformations and dural arteriovenous fistulas, ASL is very sensitive to detect even small degrees of shunting. In epilepsy, ASL can be used to assess the epileptogenic focus, both in the peri- and inter-ictal period. In neoplasms, ASL is of particular interest in cases in which gadolinium-based perfusion is contraindicated (e.g., allergy, renal impairment) and holds promise to differentiate tumor progression from benign causes of enhancement. Finally, various neurologic and psychiatric diseases including mild traumatic brain injury or posttraumatic stress disorder display alterations on ASL images in the absence of visualized structural changes. In the final part, we review current limitations and future developments of ASL techniques to improve clinical applicability, such as multiple inversion time ASL sequences to assess alterations of transit time, reproducibility and quantification of CBF, and to measure cerebrovascular reserve.
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Introduction

Arterial spin labeling (ASL) is an MR imaging technique that enables the measurement of brain perfusion non-invasively at the tissue level. Benefiting from the contrast of inflowing magnetically-labeled blood, ASL obviates the need for an exogenous contrast agent. Although the principle of ASL was introduced in early 1990s (1-3) and is feasible on low-field MR systems, ASL greatly benefits from the improved signal-to-noise ratio (SNR) of modern high-field MRI systems (4). The increasing availability of 3T scanners as well as the development of improved pulse sequences and multi-channel receiver array coils has led to a rapidly growing interest in ASL within the past few years, paving the way for widespread application in neurological and psychiatric disorders. The current review will start by briefly summarizing the essential technical requirements, and then discuss in more detail emerging clinical applications of ASL (such as dementia, stroke, vascular malformations, epilepsy, tumors and psychiatry), and concludes with a critical review of the current limitations and future perspectives of brain ASL imaging.

Essential technical considerations

Basic Concepts of ASL

ASL is based on the principle of magnetically labeling inflowing arterial blood protons prior to their entry into the tissue of interest. As such, it can be viewed as a tracer technique (5, 6), with water acting as the natural endogenous tracer to estimate tissue perfusion. The label is created by applying radio-frequency (RF) pulses to invert the bulk magnetization of the blood water
protons. Images are acquired after the labeling and inflow period using rapid acquisition techniques such as echo-planar imaging (EPI), gradient and spin echo (GRASE) imaging, or 3-dimensional fast spin-echo (3D-FSE) imaging using a stack-of-spirals approach (7-9). A pair of images is always acquired: a labeled image, in which the blood water magnetization is inverted, and a control image, in which the blood water magnetization is not inverted. The signal difference between labeled and control images is proportional to the amount of magnetization inverted and delivered to the tissue. If all the labeled blood has arrived at the imaging voxel at the time of image acquisition, the signal difference will be proportional to cerebral blood flow (CBF). The current main implementations of ASL are pulsed and pseudo-continuous labeling.

**Pulsed ASL**

In PASL, the arterial blood water is labeled using a short adiabatic inversion pulse. The labeling pulses are on the order of ~10ms and designed to invert the blood water instantaneously in a particular region, typically located inferior to the brain (Figure 1A). After labeling, a post-label delay period is required (also known as the inflow time or TI for PASL), during which time the inverted blood moves from the labeling region into the brain, loosing gradually its label through longitudinal T1 relaxation. For this reason, PASL is intrinsically a lower SNR technique than is pseudo-continuous ASL (pCASL). The control acquisition for PASL consists of applying an RF pulse with equivalent power to the labeling pulse, but which has a net zero effect on the blood water magnetization in the labeling region.
Pseudo-continuous ASL

In pCASL, a long labeling period (1-2 sec) is made up of a train of very short (≈1ms) pulses (9). This train of short pulses is designed to invert the inflowing blood magnetization in an adiabatic or pseudo steady-state manner (Figure 1B). It is useful to think of the blood being continuously inverted as it flows through a ‘labeling plane’ in the inferior-superior direction. If the phase of every second pulse in the pCASL pulse train is shifted by 180°, the flowing blood water is minimally perturbed, and thus enables acquisition of non-labeled control images. pCASL has recently been adopted as the labeling method of choice for clinical imaging, due to its ease of implementation and high SNR (10).

Pro and cons: PASL vs pCASL

The main advantage of pCASL over PASL is its higher SNR - a substantial benefit in a technique that is intrinsically SNR-limited. The reason for this can be understood by considering the degree of labeling of the arterial blood as it arrives at the tissue, and how this changes during the inflow time and/or post-labeling delay (Figure 2). For PASL, a slab of arterial blood is inverted at TI=0, and subsequently undergoes T1 relaxation during the inflow time. For pCASL, the arterial blood is continuously inverted as it passes through the labeling plane, which means that all the blood has the same amount of T1 decay when it arrives at the tissue, irrespective of when it was labelled. However, the benefits of pCASL are not as large in practice as would be predicted theoretically, due to: (i) the inversion efficiency of the RF pulses used in PASL, higher than that achieved using the pCASL flow-induced adiabatic
inversion pulse train; (ii) the sensitivity of the pCASL labeling train to off-
resonance effects, which can cause apparent regional hypoperfusion if
different feeding arteries are affected differently; (iii) the longer arterial arrival
times for pCASL for which the labeling plane is required to intersect major
feeding arteries with flow in a known direction, generally placed distally to the
PASL labeling slab. Consequently, for any given study, it is always worth
considering the specific pros and cons of pCASL and PASL in order to choose
the most appropriate option.

The critical importance of the Post-label delay time

An important aspect of all ASL techniques is the introduction of the PLD (or
TI) between the end of the labeling pulse and the time of image acquisition
(11). If the PLD is longer than the longest transit time between the tagging
plane and the imaging volume, the ASL signal becomes insensitive to
variations in arterial arrival time, as long as the blood and tissue T1 values are
similar (true for grey matter but not white matter) (11, 12). This enables
cerebral blood flow (CBF) quantification and minimizes the appearance of
intravascular signal in the ASL images, yet at the expense of SNR, due to the
T1 relaxation of the bolus (T1\text{blood}=1600\text{ms at 3T}). In addition, the choice of
PLD depends on the subject’s age, with older subjects showing longer arterial
arrival times. The recommended PLD for pediatric and adult clinical
populations are 1500 ms and 2000 ms, respectively (10).

Alterations of transit time - multi TI ASL
Most clinically available ASL sequences use only a single delay between labeling and image acquisition, based on original values calculated from healthy young adults. In case of proximal vessel occlusion, there is a delayed arrival of blood in the parenchyma, which may falsely suggest a reduced relative CBF as estimated by ASL (Figure 3) (13) and increased ASL signal in the feeding arterial vessels, known as “arterial transit artifact” (14). Reduced cardiac output, as seen frequently in elderly populations, leads to similar effects, e.g. in the vascular borderzone regions (15). The use of multi-TI ASL sequences aim to overcome this methodological shortcoming, but due to the longer scan times required, they are currently not recommended in daily clinical practice (16, 17).

Other parameters affecting ASL can be found in the online supplement.

ASL in neurodegenerative diseases

General considerations - coupling of brain perfusion and brain metabolism

Alzheimer Dementia (AD) is the most common type of dementia. According to the model of neurodegeneration proposed by Jack et al. (18), brain metabolic alterations precede structural abnormalities during cognitive decline in AD. This explains the considerable interest in the assessment of brain glucose metabolism based on FDG-PET (18F-2-fluoro-2-deoxy-D-glucose - positron emission tomography) in the work-up of dementia. This technique has identified typical patterns of metabolic alterations in various types of dementia. For example, AD is typically associated with hypometabolism in bilateral parietal and temporal regions sparing the occipital lobe, while dementia with
Lewy bodies also has bilateral parietal and temporal and additionally occipital hypometabolism (19). One disadvantage of FDG-PET is that the alterations in blood glucose metabolism are not specific for a given disease. This has led to the development of disease-specific tracers of amyloid- and tau-PET, which were recently introduced for clinical use. As a consequence, there is a need for an alternative technique, which might be used as a surrogate marker to replace FDG-PET in the work-up of dementia. ASL is a very promising alternative technique for two reasons. First, perfusion and metabolism are typically tightly coupled in the brain (20, 21). This perfusion-metabolism coupling implies that the known patterns of hypometabolism in FDG-PET in dementia can be transferred to patterns of hypoperfusion on ASL MRI. Correspondingly, there is a good correlation between brain perfusion measured using ASL (22-26) and brain metabolism measured by FDG PET (27-30), for example in the domain of MCI and AD. Second, structural MRI is routinely performed in many patients during the workup of cognitive decline. Adding an ASL sequence to the existing MR protocol is minimally demanding for the patient, and does not require additional radiation or use of contrast agents, is less costly and allows for the assessment of brain structure and perfusion in one imaging session. Moreover, if ASL could replace FDG PET, this would enable more disease-specific PET imaging with the newly available amyloid or tau tracers.

**Normal aging**

In order to be able to assess reductions in perfusion measured by ASL in cognitive decline, it is first necessary to understand the evolution of ASL
imaging appearances during normal aging. A study of 44 healthy participants from age 4 to 78, found the highest grey matter perfusion in children (97 ± 5 ml/100 g/minute) which had already deceased in adolescents (79 ± 3 ml/100 g/minute). After the age of around 20, grey matter perfusion remained rather constant at 58 ± 4 ml/100 g/minute until the age of around 80 (31). Another study of 38 elderly healthy individuals (mean age 82.2 ± 3.7) found variable patterns of decreased and increased perfusion (32). Restom and colleagues compared 15 healthy young adults around 25 years of age to 12 healthy elderly controls around 75 years old, and found a significantly lower CBF at rest in the elderly (33). Rusinek et al. focused on the hippocampus during normal aging from 26 to 92 years due to the particular interest of this structure in dementia, and found no statistically significant age or gender effect (34).

More recently, ASL was used in 148 consecutive control subjects and 65 MCI cases around 76 years of age (35). About half of the control participants developed subtle cognitive decline during follow-up. ASL imaging, performed at baseline, revealed decreased perfusion in the posterior cingulate cortex in those individuals who later developed subtle cognitive decline, indicating the predictive value of ASL for the earliest form of cognitive decline. Interestingly, the ASL pattern of those deteriorating controls was similar to MCI cases, indicating that these individuals were initially able to maintain an intact cognitive status through their cognitive reserve despite already present reductions in CBF.

Alzheimer’s disease and mild cognitive impairment
Alzheimer’s disease (AD) is associated with gray matter volume loss in the mesial temporal and parietal lobes. In earlier stages of the disease, such as mild cognitive impairment (MCI) or preclinical AD, atrophy can be minimal though metabolic imaging (FDG-PET) may already be abnormal (18, 36).

ASL studies in established AD have consistently shown a reduction in CBF (22-25, 30, 37, 38) in a posterior parietal distribution, including the precuneus, posterior cingulate, angular gyrus, and superior parietal gyrus (25). The ASL pattern is remarkably similar (Figure 4) to the pattern of the hypometabolism seen with FDG-PET (39) and both modalities have similar diagnostic performance (38). The information from ASL persists after correction for local gray matter atrophy, illustrating its independent diagnostic value, although uncorrected ASL maps may be preferred in a clinical setting to increase the effect-size. Similar patterns of hypoperfusion have been observed in subjects with MCI, and preliminary evidence suggests that abnormal perfusion in the precuneus can predict conversion to AD (40), though larger studies are required for confirmation.

*Fronto-temporal lobar dementia, Lewy Body Dementia, and Vascular Dementia*

Less is known about the potential of ASL in dementias other than AD. In subjects with fronto-temporal lobar dementia, Due et al. (30) found right frontal hypoperfusion in 21 patients, similar to FDG-PET, while perfusion in parietal regions and posterior cingulate was preserved relative to AD. These findings were confirmed in a study comparing AD and fronto-temporal lobar
dementia (41), with both ASL and FDG showing a clearly differentiating pattern in fronto-temporal lobar dementia compared to AD (Figure 4).

In dementia with Lewy bodies, Taylor et al (42) found posterior hypoperfusion in the posterior cingulate and higher-order visual association areas in 15 dementia with Lewy bodies patients; no comparison with AD was made. In the related Parkinson’s Disease Dementia disorder, Le Heron et al. (43) found the pattern of posterior hypoperfusion to largely overlap with AD patients, hinting at a similar mechanism of neurodegeneration.

Vascular dementia by definition is caused by impaired cerebral perfusion and widespread decreases in CBF have been found especially in bilateral frontal and parietal areas in subjects with vascular dementia (44, 45) and subjects with confluent incidental white matter ischemic changes (46). The pattern overlaps even more with AD in post-stroke patients (47), though this may well be due to concomitant (unveiled) Alzheimer pathology. A related point to note here is the limited ability of ASL to measure white matter perfusion. Due to the low perfusion levels and long arterial arrival times in white matter, the SNR of the ASL signal is typically very low, and it is not possible to obtain reliable estimates of CBF unless the acquisition parameters are specifically optimized (48-50).

ASL in Cerebrovascular Disease
Early on, ASL techniques were recognized to be valuable for imaging brain ischemia. Indeed, following the National Institute of Neurological Disorders and Stroke (NINDS) trial (51) and the development of diffusion imaging, the potential of acquiring brain perfusion images without contrast was recognized. This allows perfusion imaging in patients with contraindications to IV contrast, such as renal failure, and simplifies the acquisition of quantitative CBF, which is challenging with contrast agent-based perfusion techniques, such as dynamic susceptibility contrast (DSC).

Initial reports of ASL imaging in stroke demonstrated proof-of-principle, but were limited by their inability to image the entire brain (52). Since the widespread use of whole-brain, high SNR ASL sequences, several studies have shown reasonable concordance of ASL with DSC measures of perfusion deficit, primarily time-based parameters such as the mean transit time and time-to-peak of the residue function (Tmax) (Figure 5). (53-56). Vessel-selective ASL has the additional advantage of being more sensitive to identify perfusion changes induced by collateral flow: (57) in stroke patients. Routine ASL can also show perfusion changes in Moyamoya patients (58). It has also been applied to cases where treatment was initiated on the basis of the presence of collateral flow (59).

The ability of ASL to measure severe hypoperfusion or determine perfusion in regions with prolonged arterial arrival times remains a critical issue for imaging patients with cerebrovascular disease. In these situations, there can be regions with a lack of ASL signal intensity in the ischemic lesion that
render reconstruction of summary maps difficult. However, for the detectability of stroke, ASL techniques have been shown to be at least equivalent to DSC techniques, and can be used for a mismatch approach with DWI images (54, 60). In our experience, it is particularly helpful when ASL shows normal perfusion or hyperperfusion (as is sometimes seen after vessel recanalization), since it eliminates the possibility of a diffusion-perfusion mismatch that would reflect tissue at-risk of infarction, thus directly affecting patient management.

In patients with known cerebrovascular disease, alternative ASL imaging strategies could be considered, including long-label long post-label delay single time-point ASL and multi-delay ASL. The former will mitigate the problem of slow flow not arriving at the tissue bed by the time of imaging, resulting in CBF underestimation. However, differences in the decay of the label when it is within the blood, with its longer T1 compared with tissue, can lead to misinterpretation, with adequate CBF that arrives later than expected appearing hyperintense. Multi-delay ASL, in which ASL images with several post-label delay times are acquired, offers the potential to measure both perfusion and arterial arrival time, and use the arrival time information to improve the accuracy of the CBF maps. This addresses some of the problems of long-label long-delay ASL, but at the expense of SNR per unit time. Practically, this means that multi-delay ASL scans will have reduced spatial resolution unless scan efficiency is improved or much longer imaging times are accepted. One promising approach to improve the time efficiency of multi-delay ASL is Hadamard encoding (61). Hadamard encoding is a method in
which blocks of labeling are mixed with blocks of non-labeling, with different permutations of the labeling and control blocks. This method enables reconstruction of images with different delay times in a more time-efficient manner, leading to higher SNR, particularly for transit time mapping. (61, 62).

ASL has also been applied to chronic cerebrovascular disease, including carotid stenosis and occlusion (14, 63, 64) and Moyamoya disease (58, 65-68). Such diseases are characterized by near normal CBF, but often striking abnormalities of arterial arrival time, since blood arrives at the parenchyma through alternative pathways. For this reason, many of these studies have availed themselves of multi-delay ASL approaches, to correct CBF values for arrival time, and to map transit time abnormalities directly (Figure 6). Given the relatively long delays, either the use of extremely long label and post-label delay times or multi-delay ASL is critical for a proper perfusion measurement in these diseases. This being said, even with state-of-the-art techniques, there are a few patients with very severe Moyamoya disease, in which ASL imaging of CBF is practically infeasible due to the extremely circuitious and delayed arrival time of the blood flow, due to loss of the magnetic label during the transit between the labeling region and the brain parenchyma.

ASL in AVM and fistula

One unexpected but useful clinical application of ASL imaging is the ability to identify lesions with arteriovenous shunting. Because >90% of water is extracted during the first pass through the capillaries, and because the mean
dwell time of water molecules once extracted from the vascular into the extravascular space is several minutes, most of the labeled water decays within the brain parenchyma, and does not reach the veins. This leads to the excellent depiction of parenchymal perfusion and the absence of vascular signal on most ASL scans. However, if the capillary bed is absent, as in the two most common shunting lesions of the CNS, the arteriovenous malformation (AVM) and dural arteriovenous fistula (AVF), ASL signal appears in venous structures (69). This can be quite striking, given the high blood volume of the veins (Figure 7).

While large AVM’s and dural AVF’s are not difficult to diagnose with routine anatomical MR imaging, small lesions or evolution of lesions following embolization therapies can be very challenging to identify. Le et al. demonstrated that identification of small (<2 cm) lesions was significantly improved if readers were allowed to view ASL in addition to conventional imaging (70). In fact, this may be more sensitive than conventional angiography, as venous signal on ASL has been seen in AVMs in the setting of intracranial hemorrhage where initial angiograms were negative, possibly due to mass effect. It has also been shown that identification of such lesions outside the brain, within the upper neck region, is also possible with ASL (71). Finally, shunting physiology has been identified in about 10% of developmental venous anomalies, suggesting that transitional lesions might be more common than is generally appreciated (72).
One challenge for ASL when using venous signal to identify shunt lesions is that conspicuity depends on the precise timing parameters (i.e., the labeling duration and the PLD. While the use of recommended parameters \(10\) appears fairly sensitive for detecting most lesions, it is possible that in either slow or fast shunting lesions, venous ASL signal could be suboptimal or missed. Therefore, there is interest in multi-delay ASL, in which images at different labeling duration and/or post label delay are acquired as part of a single acquisition. While these sequences were first developed for better CBF quantification, the ability to optimize the sensitivity for the detection of venous signal is also enhanced. However, spatial resolution is reduced compared with single-delay ASL for a given imaging duration, and thus any gains might be offset by this lower resolution. ASL-based multi-delay MR angiographic sequences have also been developed, focusing more on direct vessel visualization \(73\). As such, these latter sequences would compete more directly with conventional angiography, with comparable temporal resolution without the associated risks.

**ASL in Epilepsy**

The main interest of ASL in the context of epilepsy is to locate a potential epileptogenic focus. ASL might further be helpful in the differential diagnosis for example in assessing for a stroke in the acute stage. During the acute peri-ictal period, the CBF is typically increased due to pathologic neuronal activity \(74\), while in the chronic inter-ictal period, CBF is typically reduced as the epileptogenic region typically is less functional and active as compared to
the normal brain tissue (75-79).

Nuclear medicine techniques such as ictal and inter-ictal SPECT and PET are often performed to detect metabolic changes related to epileptic activity. Since standard structural MR imaging is routinely performed, ASL might be of interest as an additional functional marker of cerebral perfusion with minimal additional effort. Hypoperfusion on ASL corresponds to hypoperfusion on interictal PET and electrophysiological data (75, 76, 80), indicating the potential use of ASL for epileptic focus localization (Figure 8).

**ASL in neoplasms**

Hemodynamic changes are present in many CNS neoplasms, and in general, CBF and cerebral blood volume (CBV) increase with tumor grade. Historically, much of our MR-based knowledge about these hemodynamic changes has been gained from DSC and dynamic contrast enhancement (DCE) imaging (81-87). This has had two main consequences: (1) most literature has focused on relative CBV changes, because the DSC measurement of CBV is more straightforward compared with CBF, and (2) most of our data is of relatively poor spatial resolution and prone to artifacts, since the rapid passage of the intravascularly-confined tracer necessitates rapid, sometimes suboptimal, imaging. ASL cannot measure relative CBV; instead, CBF measurements are more straightforward. While CBV and CBF are related via the Grubb’s law ($CBV = CBF^{0.38}$) in normal tissue (88), it is likely that this relationship is altered in tumors, which may form abnormal blood vessels that alter the CBV/CBF
ratio. Also, since there is no requirement to accurately track a rapid intravascular bolus of contrast, the ASL experiment is more flexible, such that SNR, spatial resolution, amount of tolerable image distortion, and imaging time may be traded off against one another. Lastly, since water has high permeability in the normal and neoplastic tissue, there is no need with ASL (unlike DSC) to employ complicated leakage-correction algorithms to obtain quantitative results.

**Primary CNS Neoplasms**

GBM is the most common high-grade CNS tumor in adults, and is associated with high metabolism and CBF (Figure 9). Initial reports demonstrated that ASL and DSC show largely concordant results in terms of identifying this tumor, and distinguishing it from contrast-enhancing mimics (89-92). Higher CBF in GBM correlates with genetic markers (such as epidermal growth factor receptor) and is associated with shorter progression free survival time (93). Lower grade tumors typically demonstrate lower CBF. CBF changes may portent transformation to a more aggressive phenotype. In fact, some have suggested that CBF quantification provides a better estimate of event-free survival for a wide range of gliomas than does a histological grading scale (94). One exception to this rule of increasing CBF and tumor grade is oligodendroglioma, a grade 2 tumor that often demonstrates increased CBF.

**Secondary CNS lesions**

Most metastatic brain lesions show similar or lower glucose uptake and CBF compared with gray matter. However, several metastatic lesions demonstrate
increased CBF, and share the feature of high vascularity. These include renal cell carcinoma (95), angiosarcoma, hemangioblastoma (96), and melanoma, and in these cases, ASL may be helpful (Figure 10).

**Assessing treatment response**

The quantitative nature of ASL has enabled studies of CBF changes over time or with treatment, to assess its impact on prognosis and outcome. A particularly important and challenging distinction to make is between radiation necrosis and recurrent tumor in patients with new or increasing contrast enhancement. Since radiation necrosis is associated typically with reduced CBF and most recurrent tumors demonstrate increased CBF, perfusion imaging may be helpful to distinguish the two entities. DSC and MR spectroscopy have also been used for this distinction, with some success (97-99). ASL is likely to improve upon DSC perfusion measures, given its better imaging performance around resection cavities (where blood products can affect DSC results). It is also less sensitive to the presence of surrounding large vessels, which can make it challenging to quantify CBF in the actual lesion (Figure 11).

Using clinical and imaging endpoints, Oszunar et al. (100) suggested that ASL is superior to FDG PET and DSC for distinguishing tumor recurrence from radiation necrosis in gliomas, with a sensitivity of 94%. In the early time period following resection (< 4 wks), where any contrast enhancement is probably due to pseudo-progression, again ASL appeared more useful than DSC (101). Another technique that can evaluate both initial presentation and treatment
changes in tumors is DCE imaging, in which the rate of contrast agent leakage is estimated.

ASL in psychiatric and other neurologic disorders
An increasing number of investigations have assessed ASL in various psychiatric and neurologic diseases.

Depression
In a study of 25 medication-naive adolescents with major depressive disorder, Ho et al., found a complex pattern of hypoperfusion in frontal, limbic, paralimbic, and cingulate yet hyperperfusion within the subcallosal cingulate, putamen, and fusiform gyrus (102). The authors conclude that adolescents with major depressive disorder differ in their baseline perfusion in executive, affective, and motor networks. Several studies assessed adults with depression and found reduced CBF in the default mode network (103) (a brain network initially described based on CBF maps (104) that is consistently found in resting fMRI studies (105)) as well as bilateral subgenual anterior cingulate cortex, left prefrontal dorso-median cortex, and left subcortical areas (putamen, pallidum and amygdala) (106). Lui et al. further subdivided patients with refractory depressive disorder from patients with non-refractory depressive disorder (107). Non-refractory depressive disorder subjects versus controls had reduced CBF in the left prefrontal cortex and increased CBF mainly in the limbic-striatal areas. Conversely, refractory depressive disorder subjects had decreased CBF predominantly in the bilateral frontal and
bilateral thalamic regions. The direct comparison of the two depressive disorder groups further revealed higher CBF mainly in the limbic-striatal areas. Finally, late-life depression is associated with increased white matter CBF yet conserved grey matter CBF (108). In conclusion, these results indicate that different subtypes of depression are associated with specific patterns of CBF alterations, which furthermore change over the lifespan.

**Psychosis**

Patients with schizophrenia compared to controls had a pattern of increased CBF in left putamen/superior corona radiata and right middle temporal gyrus yet decreased CBF in bilateral precuneus and middle frontal gyrus (109). Moreover, a sub-division into patients according to symptomatology revealed an association of negative symptoms (including difficulties in thinking or coming up with ideas, decreased ability to start initiate tasks, lowered levels of motivation or drive, lack of interest in other people) with reduced CBF in bilateral superior temporal gyrus, cingulate gyrus, and left middle frontal gyrus, and an association of positive symptoms (including delusions, hallucinations, disorganized speech or behavior) with increased CBF in cingulate gyrus and superior frontal gyrus and decreased CBF in precentral gyrus/middle frontal gyrus. A study by Ota et al. found a different pattern of reduced relative CBF in the left prefrontal and bilateral occipital cortices compared to the healthy volunteers (110). Finally, Kindler et al. observed decreased mean CBF in the frontal and temporal regions and significantly increased default mode network connectivity in the precuneus in schizophrenia patients. Currently it is impossible to disentangle whether these
observed differences in ASL based alterations in CBF in schizophrenia patients are due to different stages and subtypes of the disease, methodological, or a combination of both.

Posttraumatic stress disorder
Posttraumatic stress disorder (PTSD) has attracted considerable interest in recent years. Schuff et al. assessed 17 male veterans with PTSD and observed increased relative CBF in primarily right parietal and superior temporal cortices (111). Li et al. implemented a more complex setup in veterans nearly 20 years after the 1991 Gulf War, contrasting baseline ASL with a physostigmine challenge (112). Physostigmine significantly decreased hippocampal relative CBF in controls and veterans with syndrome 1 (impaired cognition), yet significantly increased hippocampal relative CBF in veterans with syndrome 2 (confusion-ataxia) and 3 (central neuropathic pain), suggesting that ASL might differentiate subtypes of PTSD types.

Mild traumatic brain injury
Mild traumatic brain injury (MTBI) is another entity that has attracted considerable interest, notably related to the discussion of brain alterations in professional football and soccer players. Conventional MRI sequences are not sensitive enough to detect structural brain changes, and consequently an increasing number of investigations used advanced imaging techniques such as ASL or diffusion weighted imaging (113). In 2009, Ge et al. demonstrated reduced CBF in bilateral thalamus in patients with MTBI (114). Later on, the same group extended these findings by combining ASL to assess CBF and
diffusion tensor (DTI) and diffusion kurtosis (DKI) imaging to assess white matter, again detecting substantial alterations in both perfusion and white matter integrity in MTBI patients focusing on the thalamus (113). Interestingly these alterations persisted more than 9 months after injury. Doshi et al. assessed the acute stage of MTBI and found increased in regional CBF in the left striatum, and in frontal and occipital lobes (115). Finally, Wang et al. investigated pediatric MTBI and found reduced CBF in bilateral frontotemporal regions (116). In summary, these results indicate that ASL might be a sensitive marker to assess MTBI-related alterations in the brain, which are not detectable by conventional MRI techniques. The discrepancies between these studies in turn also implies the need for strict standardization of both data acquisition and data analysis.

**Pain and migraine**

Owen et al. investigated healthy volunteers during painful thermal stimulation and observed bilateral CBF changes including the insula, secondary somatosensory, and cingulate cortices, as well as the supplementary motor area, in an attempt to quantify pain-induced perfusion alterations in the brain (117). A similar approach by Maleki et al. found alterations in relative CBF in somatosensory cortex, anterior cingulate cortex, anterior insula, hippocampus, amygdala, thalamus, and precuneus. Taken together, these studies suggest that ASL might be used as an operator-independent marker in subjective pain syndromes. Accordingly, Liu et al. observed in patients with postherpetic neuralgia, a pattern of increased CBF in left striatum, right thalamus, left primary somatosensory cortex, left insula, left amygdala, left
primary somatomotor cortex, and left inferior parietal lobule yet decreases CBF in the frontal cortex (118).

Migraine is of particular interest with respect to ASL imaging. Although controversial, it is thought that migraine attacks might involve vascular dysregulation. Correspondingly, mapping the brain CBF by ASL might reveal insights into the pathophysiology of migraine. Kato et al. performed ASL imaging in a patient during a migraine attack, after treatment with a triptan and in the attack-free period (119). During the migraine attack, CBF was reduced in bilateral median thalamic areas including hypothalamus and increased in the frontal cortex as compared to baseline. 30 minutes after treatment initiation, CBF perfusion improved in thalamus and hypothalamus. A case series by Pollock et al. demonstrated regional cerebral hyperperfusion in 3/11 patients during a headache episode that corresponded to previous aura symptoms (120). While these pilot studies demonstrated interesting preliminary findings, larger scale studies are clearly needed to investigate whether ASL might contribute to a better understanding of the pathophysiology of pain, notably a potential vascular component of migraine.

Novel extensions of ASL

Super-selective ASL to map vascular territories (aka selective territory mapping)

Contrary to contrast-enhanced CT perfusion or DSC, ASL provides the opportunity to selectively label vascular territories, such as left carotid, right carotid and vertebrobasilar territories (121, 122) or even their more distal branches. This can be of interest from a scientific perspective, for example, to
assess the impact of vascular variants of the circle of Willis on perfusion of the brain tissue, such as caudate and lenticular nucleus and thalamus (123). More importantly, territorial ASL might have clinical value, e.g., to determine whether multiple acute vascular lesions are in the same vascular territory as argument for distant cardio-embolic events in several vascular territories versus local stenotic origin for a single vascular territory or for neurosurgical planning of e.g. aneurysm clipping or tumor resection.

*Cerebrovascular Reserve Imaging*

The ability of an organ to adapt to a physiological challenge, particularly one that stresses its limits, can be measured using reserve studies. Such imaging may unmask deficits that are not apparent on routine images. Myocardial stress-rest imaging following either exercise or pharmaceutical vasodilatory challenge has been a mainstay of cardiac imaging for decades. A similar cerebrovascular “stress test” can be performed, using either physiological (i.e., CO$_2$ inhalation or breath-hold) and pharmaceutical vasodilation. In particular, acetazolamide, a carbonic anhydrase inhibitor, causes 20-50% CBF increase in normal subjects. However, patients in whom the cerebrovasculature is already maximally dilated to maintain baseline CBF may not be able to augment their CBF further in response to an acetazolamide challenge. In fact, those with very severe disease may demonstrate reduced CBF in response to challenge, a phenomenon known as cerebrovascular steal.
Limited data using non-MRI techniques, including transcranial Doppler, SPECT, and PET suggest that demonstration of poor reserve is an important risk factor for subsequent cerebrovascular events (124). Given its non-invasive nature, ASL is an ideal method to image CBF changes in response to a challenge (14, 125). Since these patients may have large artery stenosis or occlusion, the use of multi-delay ASL to quantify both arrival time and CBF is recommended. An example of reserve imaging with multi-delay ASL in a patient with Moyamoya disease with cerebrovascular steal of the bilateral anterior circulation is shown in Figure 12.

Functional MRI and ASL

Functional MRI studies generally rely on the blood oxygen level-dependent (BOLD) effect (126) and the neurovascular coupling (127). Because fMRI does not directly measure neuronal activation but the indirect vascular response, alterations of the neurovascular coupling might influence the resulting fMRI BOLD signal, for example already in normal aging (33, 128). The combination of ASL and fMRI might disentangle vascular and neuronal contributions of the BOLD response (128). Moreover it is possible to use the time-series of the ASL raw data to calculate fMRI activation maps (129) or resting-state functional connectivity networks known from fMRI (105). Although the quality of these ASL based results is generally below that of classic EPI BOLD fMRI, this might be interesting from a clinical perspective as ASL can be used to calculate relative CBF maps and resting-state functional connectivity networks of a reasonable quality without
additional cost or scanning time which is particularly helpful in elderly patients (Figure 13).

Conclusions
Although ASL has been around for over two decades, it only recently began to make the transition from a research tool to clinical use due increasing awareness of radiologists and clinicians of its capability and technical improvements that have made this approach more reliable and available as product sequences on MRI platforms. For many diseases, including dementia, vascular diseases, neoplasms, and various psychiatric diseases, ASL provides additional and complimentary information to that available from structural MRI imaging. Current methodological developments aim to increase the robustness and decrease inter-scanner variability of CBF estimation.

Online supplement
*Quantification of CBF and local alterations in CBF estimation*
A number of investigations directly compared CBF estimation in ASL to 15O H2O PET as reference standard (16, 130, 131) or combined PET-MRI technology (132, 133). Overall, these studies showed good correlations between CBF estimation in ASL and PET. Furthermore, most studies showed a systematic bias of relative CBF estimation in ASL. In addition, the
placement of the imaging slab substantially changes the relative CBF estimation in ASL and the correlation with water PET (134) indicating the need for rigorous standardization of image acquisition. Moreover, the direct comparison between two different ASL sequences (135) demonstrated a good global correspondence between the relative CBF estimated by both techniques but also pointed to systematic local differences between the results, even if acquired with both sequences during the same session. Note that differences in the calibration method used explained most of the results in this case. It is likely that different ASL sequences on different MRI machines with different hardware will show even larger differences in the estimated relative CBF maps. This implies that individual ASL sequences might have variations in local sensitivity, which in turn highlights the need for strict standardization of image acquisition and calibration of each ASL sequence, e.g. using flow phantoms. This is of particular interest in neurodegenerative diseases, as disease-related alterations in relative CBF, notably at early stages of the disease, are well in the range of normal variability between different ASL sequences even when these sequences (135) are performed with recently recommended protocols (10).

Partial volume correction

Another potential issue with ASL is the relatively low spatial resolution typically used (on the order of a few millimeters in each direction), which makes it nearly impossible to obtain any relative CBF measure located uniquely within the cortical ribbon (136). Due to the large difference in perfusion values between gray and white matter (in a ratio of 3:1), even a
small amount of white matter within a voxel located primarily within gray matter will lead to an artificially low perfusion value. For this reason, a series of post-processing techniques have been developed, using the assumption that the relative CBF only varies slowly within the gray matter (and white matter) to separate the contributions from each tissue type to the perfusion maps (136). This method might be particularly important when focusing on statistical differences between patients presenting with gray matter atrophy, enabling reduced perfusion to be differentiated from an increased proportion of white matter relative CBF in a thinning cortical ribbon (though one could argue from a pragmatic viewpoint that it does not matter why perfusion is abnormally low).

Reproducibility, Inter-individual Variability and SNR

For many years, developments of ASL focused on methodological improvements and did not investigate the important issue of intra- and inter-subject reproducibility. One of the first large-scale reproducibility studies was published in 2010, based on the QUASAR sequence (137). The mean gray matter CBF was determined to be 47.4±7.5 ml/100 g/min in that study, with a standard deviation between the different sites SDs = 1.8 (CI: 1.0–3.3) ml/100 g/min, a between-subject standard variation of 5.5 (CI: 5.0–6.2) ml/100 g/min and a within-subject standard deviation of 4.7 (CI: 4.5–5.0) ml/100 g/min with a corresponding repeatability of 13.0 ml/100 g/min. Such results - based on more than 280 subjects - showed that using a single sequence in different research centers was possible, and demonstrated that the reproducibility
achieved by ASL was comparable to equivalent nuclear medicine methods (137).

Note that for this study all data were acquired on identical 3T scanners from a single manufacturer. Since then, numerous other studies have been published, comparing different sequences on a single manufacturer (138), near identical sequences on different manufacturers (139), and combinations in between. The general results confirm the same level of reproducibility, within and between subject standard deviations as originally found in the QUASAR reproducibility study (137). Potential site-specific variations in gray matter CBF values might actually be problematic, and current research is focusing on eliminating such differences to avoid bias in future multicenter trials (10).

1.5T versus 3T

Since ASL is a low SNR technique, it benefits substantially from any possible increase in SNR. SNR increases approximately linearly with field strength, and therefore 3T is expected to increase SNR by a factor of two relative to 1.5T (4). In addition, T1 also increases with field strength, resulting in a slower disappearance of the labeled blood at 3T (T1blood = 1.65s) than at 1.5T (T1blood=1.35s). Both effects will act together to increase SNR, and the ASL white paper therefore recommended the use of higher field strengths for ASL (10).

2D versus 3D acquisition schemes
The most widely used image acquisition scheme for ASL is EPI, which is a multi-slice 2D technique. The intrinsic SNR advantage of 3D acquisition schemes renders them an attractive alternative for the imaging module of ASL sequences. Various approaches have been implemented, including 3D GRASE (140), 3D spiral FSE (9) and 3D EPI (141). Although 3D image readout methods have some challenges regarding image acquisition speed and resolution, overall, they are advantageous for ASL, for several reasons. In addition to the intrinsic SNR increase gained from exciting the entire volume with each read-out, they have the additional advantage that it is possible to define a precise time at which the whole volume is imaged. This contrasts with 2D techniques such as multi-slice EPI, where each brain slice is acquired at a different time, and consequently has a different PLD. As well as simplifying CBF quantification and avoiding any potential saturation of inflowing blood by previous slice excitation pulses, a single TI/PLD enables optimal background suppression to be achieved over the whole imaging slab, therefore providing higher SNR and improved data quality for clinical use.

Variation of ASL CBF due to caffeine and other vasoactive substances

Vasoactive substances are frequently prescribed notably in the elderly, and may alter CBF. It may be less appreciated that even substances such as caffeine, which are widely distributed in beverages such as cola or coffee, significantly reduce the CBF estimated in ASL imaging, by about 20% in young adults (142) and even more in elderly controls (by approximately 23-25%) and mild cognitive impairment (MCI) (~30%) (143, 144). Disease-related alterations in relative CBF, for example in dementia, are considerably smaller
than caffeine-induced reduction of global CBF. Consequently, vasoactive substances must be considered as a potential confound for quantitative analyses of relative CBF, and compensation strategies such as normalizing with respect to whole brain average relative CBF might be worth considering, depending on the clinical context. These observations indicate that in particular in conditions in which relatively subtle alterations of relative CBF are expected, notably dementia or psychiatric diseases, patients should be asked to refrain from caffeine ingestion 4-6 h prior to MR imaging.

Decreased cardiac output / alterations in hematocrit

Decreased cardiac output might lead to delayed arrival of the labeled blood, which might be misinterpreted as reduced perfusion. Therefore, care needs to be taken when scanning patients with potential cardiac issues, as the entire circulation system might be affected. Other systemic alterations might directly affect brain perfusion, such as a change in hematocrit. Indeed, the normal physiological response to a reduced amount of oxygen carrying capacity will be to increase blood flow systemically throughout the body, and in particular to the most important organs, such as the brain. The problem for ASL is that a reduced hematocrit will also be accompanied by an increase in blood relaxation time $T_{1b}$ (145, 146), leading to a potential overestimation of perfusion measured by ASL, as the label will disappear at a slower rate than assumed. Note that these issues are particularly relevant in neonates (147) and in patients with erythrocyte-related diseases, such as Sickle Cell Disease (148).
ASL labeling schemes: (a) in PASL, an inversion slab is placed proximal to the imaging volume to label blood in the arterial feeding vessels supplying the brain. The pulse is short (~10ms) and all the blood is inverted simultaneously; (b) in pCASL, the inflowing arterial blood is continuously inverted as it flows through the labeling plane, via a process known as flow-induced adiabatic inversion. The pCASL labeling pulse train is typically applied for a period of ~1-2s.
Differences in labeling degree of ASL bolus for PASL (left column) and pCASL (right column). The top row shows the temporal profile of the bolus (1=fully inverted; 0=fully relaxed). Since the PASL inversion slab is inverted at a single point in time (t=0 on this graph), all the inflowing arterial blood undergoes the same amount of T1 recovery at all time points after this. In pCASL, blood is labeled as it flows through the inversion plane and recovers en route to the imaging volume. A-D shows the degree of labeling remaining at several time points after the (start of) labeling (t=0): A t=0; B t = arterial arrival time (ATT); C t = bolus duration (τ); D t = ATT + τ. Color scale represents the range from fully inverted (red) to fully relaxed (blue). It can be seen that the pCASL labeling process produces a bolus with a higher overall degree of inversion than PASL, resulting in a higher intrinsic SNR for pCASL.
Figure 3

Example of an underestimation of relCBF in ASL due to a proximal vessel stenosis: The estimated relative CBF based on DSC perfusion is within normal limits (A). In contrast, the relative CBF estimated using a standard single TI ASL sequence demonstrates a marked reduction in the left anterior and middle cerebral artery territories (B). The origin of this discrepancy is the increased perfusion delay, illustrated in the DSC-derived delay map (C), which exactly matches the altered perfusion in ASL, due to the presence of a high-grade stenosis of the left internal carotid artery (time of flight, D). The underlying principle is explained in E. The normal perfusion time-series of DSC imaging (solid black line) is shifted to the right due to the presence of a proximal vessel stenosis and slower collateral flow (dotted line). As DSC imaging acquires an entire time-series, this shift in the bolus arrival simply cases a shift of the estimated curve, and the relative CBF can be accurately estimated in DSC. In contrast, a single TI ASL sequence with a standard TI (indicated by the vertical line) will under-estimate the true perfusion simply because it is too early with respect to the peak of the perfusion curve.
FIGURE 4 Examples of dementia.

Transverse FDG and ASL images of a healthy individual (first row, male, age 57, mini mental state examination 30), patient with AD (second row, male, age 52, MMSE 19), and patient with fronto-temporal lobar dementia (third row, female, age 53, MMSE 26). Functional images show predominant prefrontal abnormalities in FTD and parietal abnormalities in AD. ‘Red color’ reflects normal metabolism and perfusion.
50 year-old woman presenting with stroke with 14 hrs of right hemiparesis and aphasia, national institute of health stroke scale (NIHSS) of 9 at the time of imaging. Patient later found to have extracranial left internal carotid artery dissection. (A) demonstrates irreversibly damaged tissue on DWI within the left caudate and putamen. Using a multidelay ASL sequence capable of acquiring both (B) CBF and (C) arterial transit time images, a larger region of perfusion abnormality is identified. (D) The region of perfusion abnormality (time to the maximum of the residue function, Tmax) on conventional dynamic susceptibility contrast images is concordant with the findings on the ASL images.
Example of multi-delay ASL imaging. A fixed labeling duration of 2000 ms is used, but on subsequent images, different post-label delays (PLDs) ranging from 700 to 3000 ms are employed, yielding ASL difference (control – label) images as shown in (a). From this data, and the use of general kinetic modeling, one can simultaneously measure (b) an arrival-time corrected CBF and (c) the arterial arrival time itself. In this patient, there is near symmetric CBF but clear arterial arrival delay in the right hemisphere, as shown by the higher values on the arrival time map.
51 year-old man with exertional headaches, imaged at 1.5T with an AV fistula. (A) T2 images demonstrate very subtle flow voids in the inferior frontal lobe (arrows). (B) ASL images demonstrate very bright linear signal in the region of the right inferior frontal lobe, which extends to the right cavernous sinus, indicative of an arteriovenous shunt lesion. (C) Collapsed and (D) source MR angiographic images confirm the presence of an ethmoid dural fistula, which was recognized only after the observation of the abnormal ASL signal within the venous structures draining the fistula. This case also demonstrates that while 3T is preferable to 1.5T, relevant clinical information can be obtained with ASL at 1.5T.
Comparison of ASL MRI and 18FDG PET in epilepsy. Each vertical column represents a single patient's imaging studies. A: preoperative axial T1 weighted MRI. B: raw 18FDG PET scan images. C: raw pASL CBF maps. D: image fusion of T1MRI and 18FDG PET. E: image fusion of T1MRI and pASL. Arrows indicate regions of focal cortical dysplasia that are epileptogenic.

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78 year old man with unresectable anaplastic astrocytoma (WHO Grade III), seen on (A) post-contrast T1-weighted and (B) FLAIR images. (C) ASL demonstrates increased CBF in the region of the tumor. (D) Increased CBV is also visible on bolus dynamic susceptibility contrast (DSC) imaging (black arrow), though it is more difficult to appreciate due to the extensive number of arterial and venous vessels that surround the tumor. The findings of high CBF are characteristic of high-grade (Grades III and IV) glial neoplasms.
FIGURE 10

T2 (A), post-contrast T1 (B), and ASL (C) images in a 40 yo man with von Hippel Lindau disease and multiple hemangioblastomas, status post multiple prior surgeries. Arrows point out the extremely high CBF measured with ASL even in very small lesions.
FIGURE 11

57 yo man with lung cancer brain metastasis, 15 months status post resection, radiation, and chemotherapy. (A) Axial T1Gd shows a new enhancing lesion with (B) axial FLAIR demonstrating extensive vasogenic edema in the region of the resection cavity. (C) ASL demonstrates no increase in CBF in this region, similar to findings on (D) FDG PET. These findings were considered to represent radiation necrosis and the patient is being followed serially rather than with re-resection.
29 year-old woman with bilateral Moyamoya disease. Both FLAIR (A) and diffusion-weighted images (B) show evidence of prior infarcts in the deep white matter on the left side. Multi-delay ASL images acquired at baseline demonstrates normal CBF in the bilateral anterior circulation (C). However, imaging 10 min following 1 g of IV acetazolamide (D) demonstrates the expected CBF increase in the posterior circulation, but marked reduction in CBF in both anterior circulations, compatible with cerebrovascular steal. This finding has been associated with a high risk of subsequent cerebrovascular events and prompted bilateral direct superficial temporal artery-middle cerebral artery bypass.
FIGURE 13

Functional connectivity analysis on the ASL raw-data in 154 healthy elderly controls and 66 mild cognitive impairment (MCI) cases. Note that for illustrative purposes, the raw ASL data were directly analyzed using tensorial independent component in FSL (www.fmrib.ox.ac.uk/fsl/). The spatial representation of one of the independent components is illustrated in A. The on-off labeling pattern of the raw ASL data is clearly visible in the corresponding temporal representation in B. Adding the known prior knowledge of the on-off time-course could further improve the resulting functional connectivity networks.
References


24. Dai W, Lopez OL, Carmichael OT et al. Mild cognitive impairment and alzheimer disease: patterns of altered cerebral blood flow at MR


40. Chao LL, Buckley ST, Kornak J et al. ASL perfusion MRI predicts
cognitive decline and conversion from MCI to dementia. Alzheimer Dis Assoc Disord 2010; 24(1):-19-27.


48. van Gelderen P, de Zwart JA, & Duyn JH. Pitfalls of MRI measurement


53. Viallon M, Altrichter S, Pereira VM et al. Combined use of pulsed arterial spin-labeling and susceptibility-weighted imaging in stroke at 3T. Eur Neurol 2010; 64(5):-286-296.


56. Hernandez DA, Bokkers RP, Mirasol RV et al. Pseudocontinuous arterial


109. Pinkham A, Loughead J, Ruparel K et al. Resting quantitative cerebral
blood flow in schizophrenia measured by pulsed arterial spin labeling perfusion MRI. Psychiatry Res 2011; 194(1):64-72.


125. Bokkers RP, van Osch MJ, van der Worp HB et al. Symptomatic carotid artery stenosis: impairment of cerebral autoregulation measured at the
brain tissue level with arterial spin-labeling MR imaging. Radiology 2010; 256(1):-201-208.


139. Mutsaerts HJ, van Osch MJ, Zelaya FO et al. Multi-vendor reliability of


**Figure 1** ASLarterial spin labeling labeling schemes. A, In PASLpulsed ASL, an inversion slab is placed proximal to the imaging volume to label blood in the arterial feeding vessels supplying the brain. The pulse is short (∼10 msec) and all the blood is inverted simultaneously. B, In PCASLpseudocontinuous ASL, the inflowing arterial blood is continuously inverted as it flows through the labeling plane by means of a process known as flow-induced adiabatic inversion. The PCASLpseudocontinuous ASL labeling pulse train is typically applied for a period of approximately 1–2 seconds.
Figure 2 Differences in labeling degree of ASL arterial spin labeling bolus for PASL pulsed ASL (left column) and PCASL pseudocontinuous ASL (right column). The top row shows the temporal profile of the bolus (1 = fully inverted; 0 = fully relaxed). Since the PASL pulsed ASL inversion slab is inverted at a single point in time (t = 0 on this graph), all the inflowing arterial blood undergoes the same amount of T1 recovery at all time points after this. In PCASL pseudocontinuous ASL, blood is labeled as it flows through the inversion plane and recovers en route to the imaging volume. A–D show the degree of labeling remaining at several time points after the start of labeling (t = 0): A, t = 0; B, t = arterial arrival time (ATT); C, t = bolus duration (τ); D, t = ATT + τ. Color scale represents the range from fully inverted (red) to fully relaxed (blue). It can be seen that the PCASL pseudocontinuous ASL labeling process produces a bolus with a higher overall degree of inversion than does the PASL pulsed ASL, resulting in a higher intrinsic SNR signal-to-noise ratio for PCASL pseudocontinuous ASL.
Figure 3 Example of an underestimation of relative CBF cerebral blood flow (relCBF) in ASL arterial spin labeling due to a proximal vessel stenosis. A, The estimated relative CBF cerebral blood flow based on DSC dynamic susceptibility contrast perfusion is within normal limits. B, In contrast, the relative CBF cerebral blood flow estimated by using a standard single–inflow-time ASL arterial spin labeling sequence demonstrates marked reduction in the left anterior and middle cerebral artery territories. The origin of this discrepancy is the increased perfusion delay, illustrated on C, DSC dynamic susceptibility contrast–derived delay map, which exactly matches the altered perfusion in ASL arterial spin labeling due to the presence of a high-grade stenosis of the left internal carotid artery on D, time-of-flight (TOF) image. The underlying principle is explained in E. The normal perfusion time series of DSC dynamic susceptibility contrast imaging (solid black line) is shifted to the right due to the presence of a proximal vessel stenosis and slower collateral flow (dotted line). DSC dynamic susceptibility contrast imaging acquires an entire time series, this shift in the bolus arrival simply causes a shift of the estimated curve, and the relative CBF cerebral blood flow can be accurately estimated in DSC dynamic susceptibility contrast. In contrast, a single–inflow-time ASL arterial spin labeling sequence with a standard inflow time (indicated by the vertical line) will underestimate the true perfusion simply because it is too early with respect to the peak of the perfusion curve.
Figure 4  Examples of dementia. Transverse FDG fluorodeoxyglucose and ASL arterial spin labeling images of, A, a healthy individual (male; age, 57 years; Mini-Mental State Examination [MMSE] score, 30), B, patient with AD Alzheimer disease (male; age, 52 years; MMSE score, 19), and, C, patient with frontotemporal lobar dementia (female; age, 53 years; MMSE score, 26). Functional images show predominant prefrontal abnormalities in FTD and parietal abnormalities in AD Alzheimer disease. Red color reflects normal metabolism and perfusion.
Figure 5 Images in 50-year-old woman presenting with stroke, with 14 hours of right hemiparesis and aphasia, National Institutes of Health Stroke Scale score of 9 at the time of imaging. Patient was later found to have extracranial left internal carotid artery dissection. A, Diffusion-weighted images demonstrate irreversibly damaged tissue within the left caudate and putamen. By using a multidelay ASL arterial spin labeling sequence capable of acquiring both, B, CBF cerebral blood flow and, C, arterial transit time images, a larger region of perfusion abnormality is identified. D, Conventional DSC dynamic susceptibility contrast images show the region of perfusion abnormality (time to the maximum of the residue function) is concordant with the findings on ASL arterial spin labeling images.
Figure 6  Example of multidelay ASL arterial spin labeling imaging. A fixed labeling duration of 2000 msec is used, but on subsequent images, different postlabel delays (PLD) ranging from 700 to 3000 msec are used, yielding ASL arterial spin labeling difference (control label) images as shown in A. From these data, and the use of general kinetic modeling, one can simultaneously measure, B, an arrival-time–corrected CBF cerebral blood flow and, C, the arterial arrival time itself. In this patient, there is near symmetric CBF cerebral blood flow but clear arterial arrival delay in the right hemisphere, as shown by the higher values on the arrival time map.
Figure 7 Images in 51-year-old man with exertional headaches and an arteriovenous fistula imaged at 1.5 T. A, T2-weighted images demonstrate very subtle flow voids in the inferior frontal lobe (arrow). B, ASLarterial spin labeling images demonstrate linear high signal intensity (arrows) in the region of the right inferior frontal lobe, which extends to the right cavernous sinus, indicative of an arteriovenous shunt lesion. C, Collapsed and, D, source MR angiographic images confirm the presence of an ethmoid dural fistula (arrow), which was recognized only after the observation of the abnormal ASLarterial spin labeling signal within the venous structures draining the fistula. This case also demonstrates that while 3 T is preferable to 1.5 T, relevant clinical information can be obtained with ASLarterial spin labeling at 1.5 T.
Figure 8  Comparison of ASL arterial spin labeling MR imaging and FDG fluorine 18 fluorodeoxyglucose PET in epilepsy. Each column represents imaging studies of a single patient. A, Preoperative axial T1-weighted MR images. B, Raw FDG fluorine 18 fluorodeoxyglucose PET images. C, Raw PASL pulsed ASL CBF cerebral blood flow maps. D, Image fusion of T1-weighted MR imaging and FDG fluorine 18 fluorodeoxyglucose PET. E, Image fusion of T1-weighted MR imaging and PASL pulsed ASL. Arrows indicate regions of focal cortical dysplasia that are epileptogenic. Numbers 1–5 refer to different patients. (Reprinted, with permission, from reference 80.)
Figure 9 Images in 78-year-old man with unresectable anaplastic astrocytoma (arrow) (World Health Organization Grade 3), seen on, A, postcontrast T1-weighted and, B, FLAIR images. C, ASLarterial spin labeling image demonstrates increased CBFcerebral blood flow (arrow) in the region of the tumor. D, Increased cerebral blood volume (arrow) is also visible on bolus DSCdynamic susceptibility contrast image, though it is more difficult to appreciate due to the extensive number of arterial and venous vessels that surround the tumor. The findings of high CBFcerebral blood flow are characteristic of high-grade (grades III and IV) glial neoplasms.
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**Figure 11** Images in 57-year-old man with lung cancer brain metastasis 15 months after resection, radiation, and chemotherapy. A, Axial T1-weighted gadolinium-enhanced image shows a new enhancing lesion, with, B, axial fluid-attenuated inversion recovery image demonstrating extensive vasogenic edema in the region of the resection cavity. C, ASL arterial spin labeling image demonstrates no increase in CBF cerebral blood flow in this region, similar to findings on, D, FDG fluorine 18 fluorodeoxyglucose PET image. These findings were considered to represent radiation necrosis and the patient is being followed serially rather than with re-resection.
Figure 12  Images in 29-year-old woman with bilateral Moyamoya disease. Both, A, fluid-attenuated inversion recovery and, B, diffusion-weighted images show evidence of prior infarcts in the deep white matter on the left side. C, Multidelay ASLarterial spin labeling image acquired at baseline shows normal CBFcerebral blood flow in the bilateral anterior circulation. D, However, image obtained 10 minutes after intravenous administration of 1 g of acetazolamide demonstrates the expected CBFcerebral blood flow increase in the posterior circulation but marked reduction in CBFcerebral blood flow in both anterior circulations, compatible with cerebrovascular steal. This finding has been associated with a high risk of subsequent cerebrovascular events and prompted bilateral direct superficial temporal artery-middle cerebral artery bypass.
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