Pulmonary perfusion imaging using MRI: clinical application

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Abstract

Background Lung perfusion is one of the key components of oxygenation. It is hampered in pulmonary arterial diseases and secondary due to parenchymal diseases.

Methods Assessment is frequently required during the workup of a patient for either of these disease categories.

Results This review provides insight into imaging techniques, qualitative and quantitative evaluation, and focuses on clinical application of MR perfusion.

Conclusion The two major techniques, non-contrast-enhanced (arterial spin labeling) and contrast-enhanced perfusion techniques, are discussed.

Keywords Magnetic resonance imaging · Perfusion · Lung · Application · Arterial spin labeling perfusion · Contrast-enhanced perfusion

Introduction

Perfusion of the lung is one of the main components required for pulmonary gas exchange. Pulmonary perfusion is altered in physiological and pathological conditions such as pulmonary embolism, pulmonary hypertension and chronic obstructive disease. Therefore, assessment of the pulmonary perfusion is of physiological and clinical importance. As the lung is a large, diffuse and inhomogeneous organ it can only be studied in detail with techniques providing good to excellent spatial resolution. One of these techniques is magnetic resonance imaging (MRI). For more than 20 years MRI has been available as a technique for assessment of pulmonary perfusion [1]. Two principal techniques are established: arterial spin-labeling and contrast-enhanced perfusion [2]. Especially the tremendous technical developments of the recent years regarding hardware and new sequences make MR-based perfusion imaging broadly available. Therefore, this review article will elucidate the current state of the art sequences, clinical applications and future developments of each technique.

Non-contrast-enhanced pulmonary perfusion

Technique

MR offers the possibility to mark a specific part of spins magnetically by selective radiofrequency (RF) excitation, leading to a specific magnetisation to a selected fraction of blood. This technique is called arterial spin labeling (ASL) and requires no intravenous application of contrast material and is therefore considered to be completely non invasive [3].

It has the advantage that measurements can be made repeatedly over very short time periods (seconds). The absence of contrast also means that the measurements can be repeated indefinitely as it has not to be cleared out from the blood before reimaging is possible (also, the absolute number of measurements is limited by the total amount of
contrast material that can be applied in 1 day). There are some limitations of these techniques: (1) more difficult to implement compared with dynamic contrast-enhanced imaging; (2) requires a regular cardiac cycle; (3) single slice technique and 5–8 slices have to be acquired to cover the whole lung; (4) the limited voxel resolution (i.e. 5×5×30 mm³) [4].

In essence, two ECG-gated images of a selected slice are taken 5–8 s apart during a single apnea. The two images differ only in the way that tagging radiofrequency (RF) pulses change the signal of blood flowing into the imaged section while keeping the signal from the stationary structures unchanged between images. Subtraction of the two images leaves an image in which the signal from a volume element (voxel) is proportional to the amount of pulmonary arterial blood delivered during the previous heart cycle [3, 5–11]. In principle, ASL preparation can be combined with many fast data recording techniques. True fast imaging with steady precession (true FISP) sequences in particular has been found to be ideally suited for assessment of pulmonary perfusion.

Specific types of ASL techniques that allow for quantification of regional pulmonary perfusion during a single breath hold are called “Flow-sensitive Alternating Inversion Recovery” (FAIR) and “Flow-sensitive Alternating Inversion Recovery with an Extra Radio frequency (RF) pulse” (FAIER). Detailed descriptions of the different principles are given elsewhere [4, 12]. Perfusion-weighted images are obtained from the subtraction of a control and tag images, which are interleaved and can be acquired within one breathholding period of 20–27 s (for each slice) [13]. The potential sources of artefacts are cardiac and respiratory motion. Cardiac motion and pulsatile flow of the descending aorta can generate smearing and flow artefacts, respectively. Spatial variation between the control images can lead to ghosting of the blood vessels, and spatial movement between the control and tag images causes the appearance of the adjacent dark-bright pairs of the blood vessels.

A closely related technique, called the T1 method, has also been used to measure pulmonary perfusion (Fig. 1) [14, 15]. Methodologically, this technique is similar to the ASL techniques. It differs primarily in how the experiment is conceptualised—as the delivery of a bolus of tagged spins or as an apparent alteration of the longitudinal relaxation time. Perfusion-weighted images are acquired in a single shot by completely saturating the static background tissue before image acquisition. The decrease of the lung T1 values in the same slice position from the global to the selective experiment is statistically significant and results from the wash-in effect of the noninverted blood water ¹H spins outside the imaging slice. Because the perfusion-weighted image is acquired in a single shot, artefacts arising from patient motion or different respiratory levels can be avoided.

Furthermore, only very short acquisition times (≤ 5 s per slice) are required for the perfusion-weighted image, facilitating rapid and comfortable examinations. This is especially important for lung imaging, as some patients are not able to hold their breath for more than a few seconds.

Quantification

ASL-based techniques allow for calculation of relative pulmonary blood flow (rPBF) in ml/100 g/min [12, 16, 17]. For conversion of rPBF into PBF the lung density is required, which is usually assumed to be constant with 0.33 g/ml (at functional residual capacity) [18]. However, this might not be the case in lung diseases like fibrosis or emphysema. Normal values are reported to be in the range of 5.5 ml/g/min.

Using a rapid IR Snapshot FLASH technique, a spin labeling method within the imaging slice was used in combination with a simple two-compartment lung tissue model. The measured pulmonary perfusion rates of the healthy lung parenchyma were on the order of 400 to 600 ml/100 g/min, depending on the subject investigated and the actual slice position in the lung [19].

In healthy volunteers (n=6) a quantitative comparison between non contrast-enhanced and contrast-enhanced techniques was performed [20]. Two coronal slices (middle and dorsal part of the lung) were assessed and maps of relative pulmonary blood flow (rPBF) were calculated—no absolute perfusion measures were calculated. By placing ten regions of interest in each lung, a right-to-left perfusion ratio analysis was performed. For the dorsal slice the differences between both techniques were not statistically different. The middle position however showed a 16–43% underestimation of FAIR-derived rPBF in the right lung. This was explained by tracer saturation for the right lung as the right pulmonary artery is in the plane of both pulses needed for tagging and
readout. In an internal control this difference was not present if slices were acquired in a sagittal orientation.

Application

Based on the strength and limitations noted above, the clinical application so far has been limited. However, the strength of the techniques lies in the ability to study physiological aspects of the lung perfusion on a regional level [21–24]. Important for all perfusion studies is to notice that the SNR of ASL images is low, particularly in the anterior portions of the lung. Therefore, positioning the patient so that the area of interest is down-gravity may increase SNR and improve visibility of perfusion defects. Also, signal intensity and subsequently perfusion increases during examination on end-expiratory breath-hold compared to end-inspiratory breath-hold [25, 26].

Normal aging is associated with a decline in pulmonary function and efficiency of gas exchange. The effect of aging on spatial heterogeneity of pulmonary perfusion was studied on 56 healthy, non smoking volunteers with an age range of 21–76 years [27]. Relative dispersion increased significantly with increasing age by 0.1/decade until age 50–59 years, and a significant positive relationship between relative dispersion and age was found. No regional lung density measurements were performed. Therefore, it was not possible to investigate if the age-related parenchymal changes were the cause for this phenomenon. In an early study Keilholz et al. demonstrated the ability of ASL to depict the gravity-dependent perfusion effects [28]. Healthy volunteers were placed inside the scanner on their right and left lateral side and coronal images were acquired within a 10-s breath-hold using a FAIRER HASTE sequence. In both positions, an increase in the intensity of the dependent lung was found (229% for left lateral, 40% for right lateral). No change was seen along an isogravitational plane. These results were confirmed for volunteers lying on their back [15].

In six rabbits a pulmonary emboli model was introduced using a balloon catheter that was placed in the left pulmonary artery [29]. After occlusion of the left pulmonary artery a set of FAIRER images followed by a contrast-enhanced angiography was performed. The huge perfusion defect (whole left lung) was depicted by both imaging techniques. The huge perfusion defect (whole left lung) was depicted by both imaging techniques. It has to be noted again, that 8 s were required for a single slice of ASL, while the 3D acquisition of the contrast-enhanced perfusion needed 30 s. A similar experiment was done in 5 rabbits were ASL perfusion and ventilation images using hyperpolarised 3helium were assessed [30]. After occlusion of the pulmonary artery by a balloon catheter the ventilation was unchanged and the ASL perfusion images demonstrated the large perfusion defect.

In patients susceptible to high altitude pulmonary oedema (HAPE) it was possible to demonstrate the uneven hypoxic pulmonary vasoconstriction compared to non-susceptible patients by assessing the relative dispersion of blood flow [31].

Future developments

Up to now, only few reports exist on the clinical application of non contrast-enhanced perfusion techniques. Due to the lengthy acquisition times and relatively low signal increase, the clinical usage will be limited in the future. It may be the imaging technique of choice for pregnant women to rule out pulmonary emboli. However, in this setting other native imaging techniques, like steady-state free precession sequences, also showed a good clinical potential [32]. Therefore, the main application will be in exploration of physiological processes where the possibility of unrestricted measurements outweighs the prolonged acquisition time [3].

Contrast-enhanced MR perfusion

Technique

MRI of contrast-enhanced lung perfusion is realised by rapid imaging of the first pass of contrast material through the lungs after intravenous bolus injection. Two-dimensional approaches can be acquired in a fast fashion such as 0.3 s for each slice. On the other hand, they lack sufficient spatial resolution and anatomic coverage to characterise most of the perfusion defects or inhomogeneities. For this purpose 3D techniques are mandatory. To visualise the peak enhancement of the lungs, the temporal resolution of contrast-enhanced MRI of lung perfusion has to be below the transit time of a contrast bolus through the lungs, being in the range of 3 to 4 s [33]. Additionally, for clinical usage a reasonable spatial resolution and anatomic coverage are required to allow for visualisation of perfusion changes on a segmental level. The most frequently used technique is a 3D gradient echo pulse sequence (FLASH) with a short TE and TR with a resulting temporal resolution of 1.0 to 1.5 s for each 3D data set, dependent on the used slice thickness of 10 mm down to 5 mm [34, 35]. Improvements in k-space sampling techniques such as the use of parallel imaging techniques allow for increased spatial resolution without reduction of temporal resolution. A second strategy is to use echo sharing (like TREAT), where the low frequency k-space data are updated more often than the high frequency k-space data, which is interpolated between consecutive time frames, thus leading to an effective shortening of the total acquisition time [36]. Image quality of both techniques was assessed in nine patients [37]. Using the same temporal resolution the voxel size was decreased by 24% in the TREAT data sets compared to the standard perfusion
technique (both techniques used parallel imaging with an acceleration factor of 2). For central and peripheral vascular perception the TREAT perfusion was rated significantly superior to the standard technique. Basic evaluation of MR perfusion was performed qualitatively in comparison to perfusion scintigraphy. Intermodality comparison showed a good agreement (kappa = 0.74) in 14 patients with various diseases (COPD, lung cancer, etc.) [38]. A direct comparison between different k-space acquisition techniques was performed recently [39]. Three k-space acquisition techniques were compared for image quality and SNR: (1) generalised autocalibrating partial parallel acquisition (GRAPPA) with an internal acquisition of reference lines, (2) GRAPPA with a single “external” acquisition of reference lines before the measurement, and (3) a combination of GRAPPA with an internal acquisition of reference lines and view sharing. All examinations were performed in ten healthy volunteers (5 ml Gadobutrol with an injection speed of 2.5 ml/s) with a constant voxel size: 3.9 × 3.9 × 5 mm³. At 1.5 T, view sharing resulted in the best image quality and SNR. Furthermore, the temporal resolution was the shortest with 1.07 s.

So far, most of the studies used 0.1 mmol Gd-DTPA/kg bodyweight for visualisation of pulmonary perfusion. Different amounts of contrast media (0.05 mmol/kg bodyweight and 0.1 mmol/kg bodyweight), contrast media concentrations (0.5 mmol/l and 1 mmol/l), and injection protocols (2.5 ml/s and 5 ml/s) were compared in 10 healthy volunteers [40]. By analysing the signal-to-noise ratio no difference was found between the different contrast media concentrations giving the same injection speed. For both contrast agents a dose of 0.1 mmol/kg bodyweight was superior to the half dose protocol. In combination with a rapid injection (5 ml/s) the highest SNR in all vascular segments were achieved. Regarding the contrast media injection speed, a small bolus profile and fast attenuation of the lung parenchyma is optimal. It was found that with increased speed, the mean transit time (MTT) for the pulmonary arteries decreased significantly [41]. However, this decrease was not linear: a four-fold increase in injection rate led to a two-fold decrease in MTT. In the same study it was found, that the cardiac function parameters had no significant influence on the bolus profile. Overall, an injection speed of 3 ml/s was recommended for pulmonary perfusion [42].

Basic post-processing for improvement of visualisation of perfusion is done by subtraction of the data set with the highest signal intensity from a baseline data set (Fig. 2). In most cases this allows for a rapid analysis of the clinical situation of the patient. However, for any detailed analysis, therapy monitoring or presurgical estimation of the regional pulmonary function, a quantitative analysis is important. A summary of frequently used contrast media injection regimens are provided in Table 1.

Quantification

Quantification of pulmonary perfusion is based on the indicator dilution theory where the maximum of signal intensity and the temporal course of the signal change are used. For quantification an arterial input function (AIF) has to be defined by placing a region of interest (ROI) in the main pulmonary artery. The tissue response function of the lung parenchyma can be either determined by multiple single small ROIs or, better, for the whole lung.

Assuming a linear relation between the signal and the concentration of contrast agent, signal-time curves are converted to concentration-time curves [43]. Hypothesising a negligible amount of extravasating contrast agent during the first pass of the bolus in the lungs, the principles of the indicator dilution theory [44] were applicable. The central volume theorem states the relationship between the perfusion parameters regional pulmonary blood flow (PBF ml/100 ml lung tissue/min), regional pulmonary blood volume (PBV ml/100 ml lung tissue), and the mean transit time (MTT, s):

$$MTT = \frac{PBV}{PBF}$$

Regional PBV can be calculated by normalising the area under the tissue concentration-time curve to the integral of the arterial input function AIF:

$$PBV = \frac{\int C(t)dt}{\int AIF(t)dt}$$

In consideration of the finite length of the bolus and its dispersion during its way to the tissue volume, the relationship between the AIF and the tissue concentration-time curve is described by the convolution of the AIF and a residue function R(t):

$$C(t) = PBF \int AIF(\tau) \cdot R(t - \tau) d\tau$$

where R(t) is the amount of contrast agent remaining in the tissue at time t. PBF is therefore the initial height of R(t 0), respectively, the maximum of R(t) in case of a delay between the AIF(t) and C(t), and can be assessed by deconvolving AIF(t) and C(t). The deconvolution of the last equation is performed using singular value decomposition (SVD) [45], a model-independent nonparametric deconvolution analysis with the potentiality of reducing the noise contribution by applying a suitable threshold to a singular value matrix. This method has shown reliable results for first-pass bolus tracking in clinical studies [46].

Inclusion of the central pulmonary arteries leads to a systematic overestimation of pulmonary perfusion. Therefore, the use of small ROIs in the lung periphery can be
beneficial. However, if the spatial distribution of the underlying lung disease is unknown, or if lung perfusion quantification is required for an entire lung (e.g. to assess side differences of lung perfusion) a measurement of a limited lung sample in a small ROI will only reflect a poor estimate of lung perfusion. One study used a cross-correlation analysis (CCA) of perfusion data sets for suppression of central pulmonary vessels [47]. CCA has previously been used to improve the SNR and arteriovenous separation in time-resolved MR angiography [48, 49]. It was shown that it is feasible to identify vessels with a similar signal-intensity time curve as a reference vessel using CCA. For this purpose reference ROIs have to be drawn in the pulmonary artery and pulmonary vein. The approach was evaluated in

**Table 1** Summary of used contrast media injection protocols and spatial resolution of perfusion-weighted MR

| Injection speed [ml/s] | Dose               | Spatial resolution [mm³] | Reference |
|-----------------------|--------------------|--------------------------|-----------|
| 2.5                   | 5 ml Gadobutrol    | 3.9×3.9×5                | [39]      |
| 5                     | 5 ml Magnevist     | 1.9×1.8×10               | [56]      |
| 3                     | 0.05 mmol/kg BW Magnevist | 3.9×3.9×6.3         | [59]      |
| 4                     | 0.125 mmol/kg BW Magnevist | 2.9×1.6×10           | [76]      |
| 5                     | 5 ml Omniscan      | 1.9×1.8×12               | [63]      |
| 5                     | 0.1 mmol/kg BW Magnevist | 3.5×1.9×4              | [65]      |
| 5                     | 0.1 mmol/kg BW Magnevist | 1.9×3.8×3.6           | [66]      |
| 5                     | 0.1 mmol/kg BW Multihance | 1.9×3.8×4           | [69]      |
| 3-5                   | 3-5 ml Magnevist   | 1.9×1.8×10               | [34]      |
| 5                     | 0.1 mmol/kg BW Omniscan | 1.9×3.6×4             | [73]      |
seven healthy volunteers. The quantification of PBV and PBF resulted in an average reduction of 25% for PBV and 15% for PBF compared with the segmentations without vessel exclusion. A comparison of the manual exclusion of vessel structures and the CCA revealed an average reduction of 13% for the PBV and 8% for PBF. The reduction by applying the CCA was significant (P ≤ 0.001) compared with the results with manual and without blood vessel exclusion for all calculated parameters.

Basically, during quantitative perfusion evaluation the signal-time curves are converted into concentration-time curves. Therefore, a linear relationship between signal intensity and concentration is mandatory. This is especially important for the arterial input function where all contrast media passes by in a compact bolus [50]. A linear relationship, however, is only seen for a small range of concentrations of 3, 4 or 5 mmol/l as seen in in-vitro experiments [51–53]. The linearity between signal and concentration is dependent on the sequence and parameters used (like TE). Thus, the use of a saturation recovery prepared FLASH sequence was proposed [53], which is often used in myocardial perfusion imaging, to improve the linearity of the measurements and thus the contrast-to-noise ratio (CNR) at higher doses. Applied to volunteers the usage of 0.057 mmol/kg bodyweight was feasible [53], while in an animal experiment it was found that already 0.05 mmol Gd-DTPA/kg bodyweight resulted in in-vivo concentrations outside the linearity range [52]. However, comparing the quantitative measures of the 0.05 mmol/kg bodyweight the values were too high compared to results from PET where PBF was found to be 121 ml/min/100 ml [54] and PBV 17–20 ml/100 ml [55]. In a prospective trial different injection protocols and contrast media concentrations were investigated and compared to quantitative analysis of perfusion SPECT [56]. The amount of the MR contrast bolus was fixed with 5 ml and an injection speed of 5 ml/s, followed by a 20 ml saline flush. This approach ensured similar bolus widths in all applications. It was found that for patients weighing less than 70 kg a concentration of 0.3 mmol/ml showed the highest accordance to SPECT and 0.5 mmol/ml for patients weighing more than 70 kg. However, this study used 10-mm-thick slice partitions. Therefore, the small amount of 5 ml of contrast media resulted in an appropriate contrast-to-noise ratio.

Another solution for this complex problem was the usage of a dual bolus approach [51]. In this experiment, the arterial input function is determined by a low volume application of contrast media (i.e. 0.01 mmol/kg bodyweight) to achieve a linear relationship between contrast-agent concentration and signal intensity in the large pulmonary vessels. This is followed by boluses with higher doses to obtain sufficient CNR in lung tissue. This may allow absolute quantification as well as the visual analysis of perfusion in lung tissue, which is often not possible when a single low dose is used for absolute perfusion quantification. The drawback of this approach is the small and therefore short bolus (i.e. 1.5 ml) for the arterial input function, which requires an extremely high temporal resolution of the perfusion sequence to assess the peak signal. The combination of a small prebolus (0.01 mmol/kg bodyweight) with a main bolus of 0.04 mmol/kg bodyweight resulted in perfusion parameters best comparable with literature data. A combination of the prebolus with a main bolus of 0.08 mmol/kg bodyweight resulted in an underestimation of PBV [51].

For perfusion imaging of the lung, the optimal combination of temporal resolution and contrast-to-noise ratio is essential. This complex interaction was studied using a Monte-Carlo simulation and volunteer examinations [57]. It was found that a temporal resolution of approximately 2 s has to be achieved for correct application of the deconvolution algorithm. The influence of noise in the data is especially relevant for the estimates of PBF: high noise levels (i.e. low CNR values) lead to a substantial underestimation and a significant loss of accuracy in the PBF estimates. It was concluded that the assessment of PBF might benefit from less aggressive acceleration, for example, by a lower acceleration factor of 2. The CNR can be notably increased if larger voxel sizes are employed; the voxel size should thus be chosen as large as clinically acceptable.

Repeatability of quantitative perfusion measurements was tested in two patient groups and one large volunteer study. In one study ten patients with bronchioalveolar carcinoma were scanned twice within 3 days in inspiratory breath-hold [56]. Evaluation was performed by manually placing three regions of interest in the data set. An excellent agreement between both measurements was found, without any significant difference. Ten patients with pulmonary hypertension were re-scanned 3 weeks after the initial scan [58]. Three-dimensional perfusion data sets were acquired in inspiratory breath-hold and evaluation comprised the whole lung. No significant difference was found for PBF, PBV, and MTT. Fourteen volunteers were examined twice after 24 h in inspiratory and expiratory breath-hold [59]. Evaluation was done comprising quantitative whole lung evaluation and was performed by two observers. No significant difference was found for the pair of measurements performed in expiration and evaluated by the same observer. The inspiratory data sets however showed significant differences for PBF. Significant differences were found in inspiration and expiration for the interobserver variability. These three studies demonstrate the significant influence of the observer on the quantitative results. Therefore, for a broad usage of the technique an automatic segmentation approach assisting evaluation should be developed. Especially in data sets with pathologically changed perfusion (i.e. peripheral perfusion defects), automatic segmentation algorithms will
be difficult to be implemented. An initial study used morphological HASTE images for segmentation of the lung borders [60]. These contours were applied to the perfusion data sets after image registration of the HASTE and perfusion images. This reduced the time needed for evaluation of the perfusion data sets to approximately 10 min. However, this technique was only applied to data sets of healthy volunteers and no reproducibility experiments were performed.

So far MR perfusion of the lung parenchyma has been performed at 1.5 T. Given the benefits of higher field strengths (like 3 T) a higher SNR can be expected, which should be beneficial regarding quantification or which allows an increase in spatial resolution.

Application

In clinical routine, contrast-enhanced perfusion can be easily applied for visual assessment of disease characteristics. For a qualitative approach the discussion of temporal resolution and contrast media dose is obsolete and it can be performed at any MR scanner. Perfusion defects due to vascular obstruction show a typical wedge shape including the lung periphery (Fig. 3) [61]. In 41 patients with acute PE the agreement of MR perfusion with single-photon emission computed tomography (SPECT) perfusion for perfusion defects down to the subsegmental level was assessed [62]. MR perfusion showed a very high agreement with SPECT (kappa value per examination 0.98, and 0.98, 0.83, and 0.69 for lobar, segmental, and subsegmental perfusion defects, respectively). By a quantitative segmental perfusion analysis in patients with acute pulmonary embolism it was possible to determine the obstruction index [63]. This perfusion-based index showed a higher correlation with clinical severity and also was a significant predictor of outcome. This initial report brings MR perfusion in the pole position for being discussed as new modality of choice for PE assessment. Furthermore, this radiation-free technique is optimally suited for follow-up studies and treatment monitoring [64]. Visual evaluation is also sufficient to differentiate between different causes of pulmonary hypertension (PH) like PAH and chronic thromboembolic PH (CTEPH) [65]. Perfusion was diffusely reduced in PAH and focal defects occurred in CTEPH (Fig. 4, Fig. 5). In 45 patients with moderate to severe COPD, perfusion was matched to parenchymal alterations demonstrated by MDCT [66]. A high agreement on a lobar level was found between parenchymal destruction and reduction of perfusion in patients with severe emphysema (kappa of 0.8). In patients with cystic fibrosis the functional score, based on the visual evaluation of the perfusion defects, showed an acceptable intra- and interreader agreement (concordance correlation coefficient 0.89 and 0.8, respectively) [67].

Semi-quantitative analysis of 3D perfusion data sets in eight healthy volunteers allowed for calculation of the left-to-right perfusion ratio (ratio 0.9), which was confirmed by phase-contrast flow measurements [68]. Furthermore, a shorter transit time and higher peak signal were found in the dorsal lung regions. Quantitative evaluation of perfusion was compared to perfusion scintigraphy for assessment of left-to-right perfusion ratios in 23 patients with various lung diseases [69]. The MR estimations of left-to-right perfusion ratios correlated significantly with those of perfusion scintigraphy scans ($P<0.01$). The MR ratios computed from PBF showed the highest accuracy, followed by those from peak concentration and PBV.

One important clinical field for perfusion assessment is estimation of post-surgical lung function, for example, after lung cancer resection. So far, perfusion scintigraphy is routinely used for this purpose. In 60 patients with lung cancer, regional blood flow was assessed by scintigraphy and MR [34]. MR outperformed scintigraphy for assessment of postoperative FEV1 ($r=0.93$ and $r=0.89$, respectively). Therefore, it was concluded that perfusion MR is a feasible alternative to pulmonary perfusion scintigraphy for predicting postoperative lung function in patients with lung cancer.

Quantitative evaluation of different physiological effects was performed in nine healthy volunteers [70]. Examinations were performed in inspiratory and expiratory breath-holds. A significantly higher perfusion was observed at expiratory breath-hold. Also a higher PBF was observed for the right lung. However, no ventro-dorsal perfusion gradient was observed, which was explained by the inclusion of central pulmonary vessels in the evaluation. After application of a vasodilatative agent (inhalation of 100% oxygen) a significant increase and redistribution of the pulmonary perfusion were found in ten healthy volunteers [71].

![Fig. 3](image-url) Coronal 10-mm maximum intensity projection (MIP) image in a patient in a follow-up examination after acute pulmonary emboli. There is a residual occlusion of the middle lobe leading to a wedge-shaped perfusion defect on the right. There are also residual subtotal obstructions noted in the left upper and lower lobe characterised by a reduced perfusion.
Therefore, MR perfusion seems to be suited for estimation of therapy response in patients with PH after administration of vasodilative drugs.

In a group of 13 COPD patients, the pulmonary perfusion was found to be heterogeneously altered—PBF and PBV were significantly reduced and MTT prolonged [72].

Despite the pure quantification, another aspect of perfusion, although harder to establish, is the degree of homogeneity. In patients susceptible to high-altitude pulmonary oedema (HAPE-S) the perfusion inhomogeneity was compared between HAPE resistant volunteers and HAPE-S patients after 2 h of hypoxia (being equivalent to an altitude of 4,500 m) [73]. Parameters indicating perfusion inhomogeneity increased during hypoxia in both populations, particularly in HAPE-S subjects where they increased significantly for almost all evaluations. These results were in accordance with those obtained by ASL technique [31]. Therefore, this kind of analysis, although not yet broadly used, may be important for future applications in diseases with peripheral vasoconstriction like pulmonary hypertension. So far, only conventional perfusion parameters have been assessed in patients with pulmonary hypertension. Compared to healthy, aged-matched volunteers, perfusion parameters are different in patients with pulmonary hypertension, with reduced PBF and PBV and increased MTT [74, 75]. Only the MTT showed a moderate linear relationship with MPAP ($r=0.54$...
and \( r=0.56 \), respectively). Therefore, a severity assessment for pulmonary hypertension is currently not possible. It remains to be determined if perfusion MR might have a role for follow-up after treatment [58].

Future developments

Contrast-enhanced pulmonary perfusion can be routinely acquired on the present hardware. However the large data sets are difficult to handle and even visual evaluation might be challenging. Therefore, future developments needs to aim at providing easy-to-use post-processing utilities for visual and quantitative evaluation.

Conclusion

Non contrast-enhanced techniques for assessment of pulmonary perfusion are an important tool for understanding of lung physiology. Given the low spatial resolution and limited signal-to-noise ratio, a broad clinical application appears to be difficult. Contrast-enhanced perfusion techniques are broadly available and are easily and quickly performed, even in critical ill patients. Perfusion MR has shown good agreement with perfusion scintigraphy and SPECT. Therefore, this technique is ready to be used in a broader clinical area.

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