Functional MRI of the Lung

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Abstract: Imaging of the pulmonary parenchyma represents a unique challenge for MRI. Limited signal is caused by low proton density, susceptibility artefacts, and physiological motion (cardiac pulsation, respiration). Improvements in MRI techniques have extended the potential for investigations of pulmonary parenchymal disease with the evaluation of lung perfusion. More recently inhalation of non proton-MRI nuclei such as hyperpolarized gases (3He or 129Xe) can provide functional ventilation images.

Keywords: Lung diseases, pulmonary function, lung MRI, lung ventilation, lung perfusion.

INTRODUCTION

While proton-MRI has emerged in the past few years as a technique of choice for imaging many organs and diseases, lung parenchyma still remains one of the most difficult tissues to be imaged because of three main factors:

- The low density of air-filled lung parenchyma, resulting in low proton density and weak NMR signal intensity
- The important motion artefacts due to physiological displacements (cardiac and respiratory motions);
- Susceptibility effects induced by multiple air and tissue interfaces which cause many local gradients responsible for very low values of T2.

Several strategies have been developed to overcome these difficulties. Image acquisition during breathhold or using respiratory gating and/or EKG triggering procedures results in reduced motion artefacts and a fixed position of the diaphragm and the cardiac cavities during image reconstruction. Susceptibility artefacts can be minimized by the use of spin echo sequences and very short echo-time. Nonetheless, the weak NMR signal remains the major drawback for lung MRI. In this context, the development of new contrast agents is of crucial importance for increasing the NMR signal from the lung. Extra cellular or strictly intravascular contrast agents administrated intravenously (Gd chelates) have been used to enhance lung parenchyma and evaluate lung perfusion. Beside proton MRI, other nuclei can be used for MRI. They include Helium-3, Xenon-129 and Fluorine-19.3He and 129Xe are inert gases, and they can be applied easily by inhalation.

PROTON IMAGING

Low spin density is the major drawback for MRI of the lung, especially for the visualization of normal lung parenchyma and of lung diseases with loss of tissue such as emphysema. It can be approached by the use of gradient-echo sequences with short repetition times, low flip angle, a higher number of acquisitions and large voxels. In many lung diseases, the amount of tissue, fluid and/or cells is increased by the pathological process. Thus, a higher number of protons are accessible, and significant improvements of the SNR can be achieved. To make use of the higher spin density short echo times are important to avoid signal loss from T2 relaxation [1]. However, in various cases such a disease related increase of spin density is not sufficient for accurate delineation, characterization and diagnosis of the disease process. Paramagnetic contrast agents have the potential to increase the MR signal from the lungs when using very short echo times which reduce susceptibility effects arising from air-tissue interfaces. They are used for enhancement of nodules or consolidations, for MR angiography or for perfusion imaging.

Chronic Infiltrative Lung Disease

In Chronic infiltrative lung disease, the pathology itself induces an increase in proton density and a reduction of susceptibility artefacts [2]. Alveolitis demonstrates high signal intensity on T1w and T2w images and also shows manifest enhancement after contrast administration. Signal intensity correlated with clinical severity of disease and potential response to therapy. Successful anti-inflammatory treatment decreasing the activity of alveolitis or development of fibrosis in the natural course of the disease are associated with a decrease in signal intensity. King et al. have shown that, the administration of a contrast agent significantly improved the detection of honeycombing but not the detection of ground-glass opacities [3]. Although, extent and distribution of airspace disease is clearly depicted at MRI, it is still inferior to CT regarding anatomic assessment and demonstration of fibrosis, i.e. parenchymal bands and reticular abnormalities [5]. In animals, macromolecular contrast agents may improve the differentiation between alveolitis and fibrosis even further [6]. In the active alveolitic phase, leakage of macromolecules from the intravascular into the extravascular compartment was observed. In the fibrotic phase, enhancement was markedly diminished which
indicates a decrease in plasma volume within the fibrotic lung [6]. Macromolecular contrast agents will greatly improve the assessment of the inflammatory alveolitic processes in the future.

**Pulmonary Edema**

Magnetic resonance imaging has also been used successfully to evaluate the lung water content qualitatively and quantitatively [7]. These properties can be used to characterize the microvascular barrier function within the lung. In hydrostatic pulmonary edema in rats, the T1 relaxation time was significantly longer than in normal controls, whereas T2 relaxation time was not different. In permeability pulmonary edema, however, T1 and T2 relaxation times were significantly longer than in normal controls [8]. It could also be demonstrated that T1 and T2 are linearly related to extravascular lung water in an induced permeability edema in rats [9]. In normal excised pig lungs, a 95% correlation between MR measurements and gravimetric lung water content was found [10]. Using a three-dimensional gradient-echo sequence which is sensitive to water the entire lung can be covered with a spatial and temporal resolution sufficient to observe the development of pulmonary edema in an experimentally induced acute respiratory distress syndrome [11]. The results indicate that edema formation can be imaged regionally and quantified globally estimating the mismatch of transcapillary filtration flow and lymph clearance [12].

Lung water sequences were also successfully applied in humans. Measurements of average lung density were easily performed. Lung intensity decreased with full inspiration. It gives an estimate of regional lung expansion. Regional measurements of lung density are used to determine lung density gradients. These gradients did not show a significant change in the prone or the supine position. From the calculated lung density gradients, pleural pressure gradients can be calculated. They also decrease with full inspiration. In normal volunteers, MR measurements of these parameters were similar to those obtained with more invasive procedures [10]. Edema and infiltrative lung disease increase lung density and decrease the elastic properties of the lung, and MR measurements of lung water content may prove useful in the assessment of parenchymal lung disease. In animal models, using macromolecular contrast media, pulmonary edema of different etiologies (increased vascular pressure vs damaged capillary barriers) can be separated [13]. Measurements of the total intravascular volume using macromolecular contrast agents and the determination of total lung water using a multi-spin-echo sequence allow for an accurate calculation of extravascular lung water. There was a high correlation with gravimetric reference measurements in the quantitation of pulmonary edema.

Although some approaches are ready to use, MRI of pulmonary edema has not been established in clinical routine.

**Perfusion**

There are several different approaches to image lung perfusion. Some of them are based on spin tagging techniques, called Flow sensitive Alternating Inversion Recovery (FAIR) and FAIR with an extra radiofrequency pulse (FAIRER). The blood water is magnetically labeled and used as an endogenous, freely diffusible tracer. With inversion times of 1000-1500 ms enhancement of the lung parenchyma and the peripheral pulmonary vessels down to the subsegmental level was observed [14]. The classical method requires the administration of intravenous contrast agent (Gd-chelates) to evaluate lung perfusion. Gd-chelates will reduce the longitudinal relaxation (T1) of blood resulting in an increase in signal intensity. Usually, imaging is performed during the first pass of the contrast agent through the lung using a fast imaging sequence with a short TE to reduce susceptibility artefacts.

Changes of signal intensity during the first passage are detected with inversion recovery 2D turbo-FLASH with ultrashort TR or 3D FISP with short TR and short TE [15, 16]. For both techniques, the non-enhanced images are subtracted from the enhanced images, such as in digital subtraction angiography. The final image is a selective perfusion map of the lung. For the 2D sequence time-intensity curves were calculated to assess the time course of perfusion [17]. From the 3D FISP acquisition maximum intensity projection of perfusion were generated and assessed visually [18]. These techniques were successfully used to detect perfusion defects in patients with pulmonary embolism [19] (Fig. 1a).

In a prospective comparative study with perfusion lung scintigraphy, perfusion MRI had an average sensitivity of 69% and specificity of 91% for the detection of perfusion defects. The overall agreement between MR and scintigraphy appeared to be good (kappa 0.63) [20]. MR perfusion imaging was also capable to demonstrate physiological preferences of pulmonary perfusion. The gravity-dependence of lung perfusion was observed in volunteers who were investigated in supine and prone position [21]. In patients after single lung transplantation, preferred perfusion of the dorsal parts of the lung in supine position was clearly demonstrated by a higher increase in signal intensity [22]. These preliminary data in patients suggest that perfusion imaging might be quite helpful in the diagnosis of pulmonary embolism. It remains unclear whether perfusion imaging will provide clinically helpful information on perfusion defects and heterogeneity in emphysema or chronic infiltrative lung disease.

**Oxygen-Enhanced Lung MRI**

The use of 100% oxygen as a paramagnetic contrast agent in MR imaging of the lung was first proposed by Edelman, Chen and Stock [23-25]. A shortening of the T1 of the lung parenchyma on the order of 100-200 ms (10%) was reported during breathing 100% oxygen, resulting in a signal intensity variation on the order of 20% on optimized T1-weighted sequences (Fig. 1b). The optimum dosage for 100% O2 was shown to be 15 L/min [26]. Administration of 100% oxygen with this dosage has been shown to produce oxygen concentrations in the lung of 60-80% [27]. Above 15 L/min, a plateau in signal intensity increase was observed. Several investigators have suggested that prolonged hyperoxia due to 100% oxygen inhalation can cause diffuse alveolar damage and lung fibrosis [27]. Such noxious effects were observed on animal experiments and occurred during prolonged hyperoxia of several hours or days, whereas in O2-enhanced MRI lung studies, hyperoxia lasts less than 1 h in total and is alternated with periods of normoxia. MR images
are acquired with the subjects in a supine position alternately inhaling room air and 100% oxygen using a non-rebreathing ventilation mask or a mouthpiece associated with a nose clamp. A typical O₂ inhalation paradigm consists of subjects alternately breathing room air for about 1 min and 100% oxygen for about 2 min. The difference between the images of the two states yields the oxygen-enhanced ventilation images. Ventilation maps can be expressed as the percentage of relative enhancement. Image acquisition can be triggered either right after the air/O₂ switch for assessment of wash-in or washout or about 120 s after the switch to allow static imaging.

**Fig. (1).** Patient with a suspected pulmonary embolism. Gadolinium perfusion image demonstrates a perfusion defect in the left lower lobe (a). Oxygen coronal ventilation image demonstrates a homogeneous lung enhancement (b).

Simultaneous cardiac and respiratory synchronization has been shown to improve image quality [28] with a reduced variability of signal intensity and diaphragm mismatch. Image acquisition can be performed during quiet breathing, at the end of expiration when the signal intensity is higher (due to higher proton density and smaller susceptibility effects) as compared with acquisition at full inspiration. Only short suspensions of breathing of less than 3 s are required during acquisition, which is especially important for patients who could have difficulties in holding their breath at end-expiration.

Several studies aimed to compare signal intensity changes or signal intensity time course with DLCO, standard pulmonary function tests, HRCT or scintigraphy in healthy volunteers and patients with pathologic diffusive capacity such as idiopathic lung fibrosis or emphysema [27, 29]. In patients, mean enhancement ratio was significantly lower and signal intensity time course showed a delayed signal intensity increase [27, 29]. Weak and heterogeneous enhancement of 10-26% was found in patients with pulmonary emphysema [27]. Regional differences that correlated with findings on radiographs and CT scans were seen on the signal intensity slope and on the maps of signal intensity change. Enhancement was excellently correlated with DLCO [27] and strongly correlated with HRCT emphysema score. Mean upslope of enhancement calculated from dynamic acquisitions was strongly correlated with forced expiratory volume in 1 s (FEV1) and had a good to excellent correlation with DLCO [29]. A prospective study in 30 patients with lung cancer, showed that post-surgical FEV1 and predicted oxygen enhanced MRI FEV1 was excellently correlated [27].

**HYPERPOLARIZED GASES**

Compared with conventional water proton imaging, MRI of gases is inherently limited by the much lower spin density of the gas phase (typically three orders of magnitude drop in spin density).

One effective way to circumvent the low NMR sensitivity of gases is to acquire MR signal from the so-called hyperpolarized (HP) or laser-polarized gases.

Among other parameters, the NMR signal-to-noise ratio (SNR) per unit volume is depending linearly on the polarization level P, defined as the percentage of the maximum achievable nuclear macroscopic magnetization of the sample. For the nuclei of the atoms of interest in biomedical NMR (hydrogen, fluorine, phosphorus, sodium, etc), the value of the polarization level at thermal equilibrium, the so-called Boltzmann polarization level, is in the order of a few parts per million (ppm). However, nuclear polarization high above thermal equilibrium can be achieved by the use of appropriate polarization techniques. The polarization levels can be typically increased by 5 orders of magnitude compared with the Boltzmann equilibrium level, resulting in a similar increase in detectable NMR signal.

**MR Imaging Using HP 3He and 129Xe**

The large NMR signal offered by HP gases allows one to image their distribution in the pulmonary tree and the alveolar spaces. The first NMR biomedical application of HP 129Xe was reported in 1994 with intrapulmonary space imaging of excised mouse lungs [31]. The first in vivo animal lung ventilation images obtained using HP 3He were reported one year later [32] followed by the first human lung images [33, 34]. Imaging of the intrapulmonary distribution of HP gases can be used for the visualization of ventilated airspaces. The spatial resolution of lung ventilation images obtained using 3He exceed by an order of magnitude the spatial resolution that are routinely obtained with...
scintigraphy techniques using radioactive gases. In human studies, typical spatial resolution in the millimeter range are reported (Fig. 2), while in rodents studies sub-millimetric resolution (Fig. 3) is usually reached [36]. Spatial resolution obtained with 129Xe ventilation images is typically 3 to 5 times lower due to reduced available SNR. Physical and chemical interactions of the HP gases with their biological environment can be advantageously exploited for obtaining structural or functional information on the lungs. The most commonly used physical parameters are the diffusion length, the longitudinal relaxation times, the velocity, the flow and the chemical shift of the HP gases.

Fig. (2). Coronal hyperpolarized 3He gas MR image in a normal volunteer.

MR diffusion imaging is a well known technique applied essentially for measuring the so-called apparent diffusion coefficient (ADC) of water in tissues. The ADC values of HP gases in airspaces depend on gas composition (dilution of HP gases in pulmonary gases) and above all on the restriction of gas atoms diffusion by the broncho-alveolar walls. The diffusion length of helium atoms during typical diffusion sensitizing times (a few milliseconds) exceeds the diameter of alveolar sacks (a few hundreds of micrometers). Hence, in the time scale of MR acquisition, 3He diffusion in alveolar space takes place in a restricted regime. The dependence of HP gases ADC values upon the dimensions of the alveolar space can be used as a non-invasive tool for probing the lung architecture at a sub-pixel level. Indeed, 3He ADC values have been shown to significantly increase in patients with emphysema compared with healthy volunteers [37]. These 3He ADC changes in emphysematous lungs are attributed to morphological changes in alveolar structure and more specifically to airspace enlargements that characterize emphysema [37-40]. The longitudinal relaxation time, T1, of 3He in the lungs is approximately 20 s. The relaxation rate R1 of 3He is varying linearly with the partial pressure of oxygen due to dipolar interactions of 3He nucleus with paramagnetic molecular oxygen [41].

Fig. (3). Coronal 3D hyperpolarized 3He gas MR image in a normal rat.

By measuring the time variation of the relaxation time of HP 3He in the lungs, it is then possible to compute locally the alveolar oxygen concentration and the consumption rate of oxygen in vivo [42,43] (Fig. 4). The velocity of HP gas in the lungs can be measured from velocity-encoded MR imaging sequences and HP gases lung inflow or outflow can be derived from dynamic ventilation image series [44]. Among other applications, these dynamic gas MRI techniques have been used in animal models for the evaluation of bronchoconstrictive drug effects [45] and in clinical studies following lung transplant in patients [46]. Chemical shift imaging has been demonstrated using HP 129Xe. The high solubility and the large chemical shift (several hundreds of ppm) of xenon allow one to differentiate between xenon in alveoli and xenon dissolved in tissue. The so-called xenon polarization transfer contrast (XTC) technique aims to probe the xenon exchange between alveolar space and blood/tissue compartments [47]. The method is based on the selective destruction of xenon polarization in lung parenchyma. Owing to the rapid exchange of xenon between the gas and the tissue phase, the depolarization of xenon dissolved in tissue affects the xenon signal from the gaseous phase. Using an appropriate pixel-based signal analysis of this effect, the authors obtained so-called XTC lung images with a contrast related to the exchange rate of xenon between the tissue and the alveolar space. A similar approach, called CSSR (chemical shift saturation recovery) was recently demonstrated and applied in human studies to investigate the alveolar surface area per unit volume of gas [48]. Besides ventilation imaging, HP gas imaging has been applied for the assessment of tissue perfusion, the acquisition of
angiographic images and the characterization of biological tissues. These applications of HP gases have been restricted so far essentially to pre-clinical studies.

**Preclinical Examinations**

Emphysema disease in animal models has been extensively studied using 3He and 129Xe MRI. Elastase-induced emphysema has been investigated in rat [38, 40], mouse [49] and rabbit [50] using 3He or 129Xe diffusion MRI. Measurements performed at end-expiratory inflation volume demonstrated an increase by 20% of ADC values in the lungs of elastase-treated animals compared with ADC values in healthy control animals [38]. When measurements were carried out at total lung capacity, 3He ADC values increased from 0.15 cm²/s in normal rats to 0.18 cm²/s in elastase-challenged animals; moreover, a significant correlation was found between the 3He ADC values and the alveolar internal area assessed by histology in lungs fixed with formalin at an airway pressure corresponding to the total lung capacity [40]. Similarly, 3He ADC values averaged over the entire lungs were found to be approximately 25% higher in emphysema mice than in healthy animals [35].

As mentioned previously, it is possible to compute the alveolar partial pressure of oxygen, (pO₂) in vivo from the relaxation time of the HP 3He in the lungs. As the intrapulmonary oxygen partial pressure distribution is governed by local ventilation, perfusion and O₂ uptake, pO₂ assessment can be used to evaluate lung function. As a matter of fact, the determination of pO₂ values and of their

![Fig. (4).](image_url)
time evolution during breathhold represents an indirect measure of the ventilation/perfusion ratio. The potential of 3He imaging for detecting perfusion abnormalities based on pO2 measurement was demonstrated in an experimental pig model [51]. After isolated pulmonary arterial occlusion using a balloon catheter, a focal T1 reduction corresponding to an abnormally high pO2 (related to the absence of perfusion) was observed, which normalized upon deflation of the balloon. More recently, pO2 imaging and oxygen depletion rate imaging was extended to small animal studies in rats and mice [52,53]. The potential of 3He MRI for assessing airways constriction has been demonstrated in methacholine-induced bronchoconstriction in rat models [45]. Using a Cine-MRI approach in which image acquisition was synchronized with the inhalation of the gas mixture (3He with oxygen and nitrogen), heterogeneously distributed airways constriction was observed, resulting in a partition of the lung between ventilated and non-ventilated regions. The diameter of the main airways decreased by approximately 11% following methacholine injection. In a methacholine-induced broncho-constriction rat model [45], dynamic ventilation image series obtained from a single breath were used to generate parametric pixel-by-pixel maps of gas arrival time, filling time constant, inflation rate and gas volume and to assess these parameters in distal areas of the lung. Quantitative and regional analysis of gas flow, volume and arrival time demonstrated statistically significant differences between the baseline and constricted states. These differences were attributed to constricted areas of peripheral airways.

Clinical Applications

Longitudinal studies or follow-up examinations for treatment or therapy monitoring are limited by the ionizing nature of the standard clinical imaging techniques (radiography, CT or scintigraphy). HP gases imaging represents a relevant diagnostic modality when multiple examinations are required for improving the patient management care. Up to now, the clinical applications of HP gases have been focused mainly on the investigation of three lung diseases: emphysema, cystic fibrosis and asthma.

In clinical studies, the regional measurement of 3He ADC values has been used to evaluate the severity of emphysema in patients [54-56]. This approach is supported by the good correlation observed between the 3He ADC values and alveolar dimensions [40]. Additionally, the 3He ADC values have been shown to correlate with spirometric indexes such as FEV1 (forced expiratory volume in 1 second) in healthy volunteers and in emphysema patients [57]. 3He imaging has been also applied for the investigation of young patients (children and adolescent) with cystic fibrosis. Since the radiation exposure experienced in case of ionizing imaging modalities is problematic for children, 3He MRI represents an interesting alternative to standard imaging technique. 3He MRI has been demonstrated to be well suited for the visualization of ventilation defects (Fig. 5) and for evaluation of the efficacy of therapies such as physical therapies or bronchodilator inhalation [58-60]. Asthma disease is characterized by the inflammation and the obstruction of small airways. Several studies have investigated and demonstrated the potential of HP 3He in clinical studies with asthmatic patients [61-63]. It has shown that small and reversible ventilation defects that characterize asthma can be detected using HP 3He and that the efficiency of bronchodilator and physical challenge can be locally assessed. Furthermore, the variations in airflow obstructions as depicted with 3He MRI correlate with the clinical evaluation of asthma severity and spirometry measurements.

CONCLUSION

CT plays a major role in the imaging approach to structure and function in ARDS. CT has an impact on the optimization of the ventilatory strategies. However, great improvement has been made in MRI of the pulmonary parenchyma during the past years. The lung will not remain the forgotten organ for MR imaging in the future. Promising research activities are under way to introduce functional MRI of the lung. MRI will offer visualization of lung morphology as well as functional assessment of ventilation and pulmonary perfusion in a single examination.

REFERENCES

Lung MRI

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