Quantitative and Qualitative Assessment of Pulmonary Emphysema with T2-Weighted PROPELLER MRI in a High-Risk Population Compared to Low-Dose CT

Quantitative and qualitative Bewertung von Lungenemphysemen mit T2-gewichteter PROPELLER-MRT in einer Hochrisiko-Population verglichen mit Niedrigdosis-CT

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ZUSAMMENFASSUNG
Ziel Erkennen und Schätzen von Lungenemphysemen mittels T2-gewichteter PROPELLER-MRT für die Bewertung von Lungenemphysemen.


Ergebnisse In allen 30 Fällen war das Vorliegen von Lungenemphysem korrekt erkannt. Eine signifikante Korrelation wurde sowohl für die klinischen Scores als auch für die computergestützte Evaluation festgestellt.

Schlussfolgerung Die T2-gewichtete PROPELLER-MRT kann als hilfreich für die Diagnostik von Lungenemphysemen angesehen werden.

Kernaussagen:
- Die T2-gewichtete PROPELLER-MRT kann für die Bewertung von Lungenemphysemen genutzt werden.
- Es zeigt sich, dass die klinischen Scores mit den computergestützten Evaluationen in der T2-gewichteten PROPELLER-MRT korrelieren.

ABSTRACT
Purpose To determine the suitability of T2-weighted PROPELLER MRI for the assessment of pulmonary emphysema.

Materials and Methods A retrospective analysis of 60 participants in a lung cancer screening program (30 subjects with pulmonary emphysema, and 30 control subjects without emphysema) was performed. The T2-weighted PROPELLER MRI sequence was used for the assessment.
Pulmonary emphysema was analyzed and correlated with the results of LDCT. The presence and the extent of pulmonary emphysema were first assessed qualitatively using a three-point score, and then quantitatively with a semi-automated software program to obtain emphysema indices.

**Results** All 30 cases with pulmonary emphysema were accurately detected by MRI. There were 3 cases with emphysema according to MRI without emphysematous changes on LDCT (false-positive results). The qualitative scores as well as the emphysema indices were significantly higher in the emphysema group compared to the control group for MRI and LDCT (p < 0.001). Both the scores and the indices correlated significantly between MRI and LDCT (qualitative score of severity: r = 0.912/p < 0.001 in the emphysema group and r = 0.668/p < 0.001 in the control group; emphysema index: r = 0.960/p < 0.001 in the emphysema group and r = 0.746/p < 0.001 in the control group).

**Conclusion** The presence and the extent of pulmonary emphysema may be assessed qualitatively and quantitatively by T2-weighted PROPELLER MRI with very good correlation to LDCT.

### Introduction

Pulmonary emphysema, as a common finding in long-term smokers and a leading cause of mortality worldwide, is characterized by irreversible destruction of the lung parenchyma [1, 2]. Computed tomography (CT) plays an important role in the visualization and characterization of pulmonary emphysema, and it has previously been shown that CT findings can predict mortality in COPD [1, 3]. Moreover, the degree of emphysema on CT images can be quantified by computing low attenuation areas under a specific threshold [4 – 6].

Magnetic resonance imaging (MRI) allows for radiation-free lung imaging, but the low proton density of the lung parenchyma, susceptibility artifacts at air-tissue interfaces, and the vulnerability to respiratory and cardiovascular motion artifacts makes lung imaging with MRI still challenging [7 – 9].

Even though a decrease in lung density further complicates image acquisition, several studies have shown the feasibility of MRI for the assessment of pulmonary ventilation and emphysema. Most studies report on the use of ultrashort echo time (UTE) imaging, as well as functional MRI with Fourier decomposition or inhalation of hyperpolarized noble gases [10 – 17]. However, especially the latter approach is highly technically demanding.

Besides the aforementioned MRI techniques, it has been shown that conventional structural T2-weighted MRI using the Periodically Rotated Overlapping ParallEl Lines with Enhanced Reconstruction (PROPELLER) technique can depict pulmonary nodules and changes in lung tissue in general [18 – 21]. Hence, the aim of this study was to evaluate whether T2-weighted PROPELLER imaging might also be used for the assessment of pulmonary emphysema.

### Materials and methods

#### Study participants

The study population consisted of 224 participants in a lung cancer screening study comparing low-dose CT (LDCT) and MRI. To be included for the screening, the participants had to be 50 – 70 years old with a long history of cigarette smoking (at least 15 cigarettes per day for at least 25 years, or at least 10 cigarettes per day for at least 30 years). Study participants were active smokers or had quit for not more than 10 years. The screening study was approved by the institutional review board and by the federal agency for radiation protection. Written informed consent was obtained from all study subjects. For the present study, we retrospectively included 30 participants with pulmonary emphysema according to LDCT, and 30 participants without emphysema.

#### Technique

For this study, a transverse T2-weighted sequence using the PROPELLER technique (MultiVane XD, Philips Healthcare, Best, The Netherlands) as part of our MRI screening protocol was evaluated. The sequence was acquired on a clinical 1.5 Tesla scanner (Ingenia, Philips Healthcare, Best, The Netherlands) with an anterior phased-array body coil. The imaging parameters were as follows: repetition time 2200 – 2500 ms, echo time 60 ms, flip angle 90°, FOV 400 mm, matrix 432 x 432 mm, slice thickness 6 mm, acquisition time 3:18 min with respiratory gating. Other sequences of the MRI protocol were transverse T2-weighted STIR (short tau inversion recovery) MVXD, coronal T2-weighted MVXD, transverse balanced steady-state free precession, and coronal 3D T1-weighted gradient echo, yet these sequences were not evaluated for this present study.
LDCT was performed on a clinical 128-slice spiral CT scanner (iCT, Philips Healthcare, Best, The Netherlands) in inspiratory breath-hold with a reconstructed slice thickness of 2 mm. The tube current-time product was 25 mAs, the tube voltage was 120 kV and the volume CT dose index was 1.8 mGy, leading to a dose length product of 70–90 mGy*cm. All participants underwent LDCT and MRI within the same day or week.

**Image analysis**

LDCT images of 224 participants of our lung cancer screening program were retrospectively evaluated for the presence of pulmonary emphysema by a radiologist with 4.5 years of experience. The window settings of the LDCT datasets for this analysis were window width 1500 Hounsfield Units (HU) and window level –700 HU. First, the presence and severity of emphysema were assessed qualitatively using a three-point scale: 0 = no emphysema, 1 = moderate emphysema (centrilobular lucencies occupying approximately > 5 % of a lung zone OR scattered small juxtapleural lucencies), 3 = advanced emphysema (large juxtapleural lucencies). Second, morphological patterns of emphysema were evaluated: 1 = predominantly centrilobular emphysema with scattered, multiple or coalescent centrilobular lucencies (few paraseptal lucencies may be present), 2 = predominantly paraseptal emphysema with scattered or multiple paraseptal lucencies (few centrilobular lucencies may be present), 3 = mixed or advanced destructive emphysema. These definitions followed the statements of the Fleischner Society published in 2015 [4]. Third, automated emphysema analysis was performed using a commercially available software program (IntelliSpace Portal, Philips Healthcare, Best, The Netherlands) in order to obtain emphysema indices (EI).

Lung parenchyma was considered emphysematous when it showed attenuation values of below –930 HU at inspiration. An emphysema index (EI) was calculated for each LDCT dataset, defined as the percentage of lung volume with emphysema divided by the total lung volume. The presence of pulmonary emphysema was defined as an EI of ≥ 6 % or when multiple lucencies were clearly visible on LDCT. LDCT datasets in which extensive atelectasis and/or pulmonary infiltrates led to an opacification of approximately one third of a pulmonary lobe were excluded (n = 10). Subjects for which the automated analysis of LDCT images did not work appropriately due to noise overlay were also excluded (n = 27). This was seen in adipose individuals in particular. There are three reasons for the malfunction of the automated analysis: 1) Automated lung segmentation from trachea and surrounding tissue failed. 2) Quantitative results showed excessive emphysema indices that were apparently false positive. 3) Emphysema assessment included several parts of soft tissue and bones also leading to false-positive results.

This led to 30 subjects with pulmonary emphysema. 30 of the remaining 157 participants of our lung cancer screening program who did not show pulmonary emphysema were randomly selected as control subjects.

MR images of the 30 subjects with pulmonary emphysema according to the LDCT definition mentioned above (mean age: 60.3 ± 6.4 years) were presented to two radiologists with 6.5 years and 16 years of experience, respectively, together with the 30 control subjects (mean age: 58.3 ± 5.8 years). The datasets were anonymized and presented in random order. The MR images were evaluated in consensus. Again, the presence of emphysema was first assessed qualitatively with predefined image windowing using the same three-point scales as for LDCT. Then, automated quantitative analysis was performed using a software program custom written in MATLAB (The MathWorks, Inc., Natick, Massachusetts, USA). With the help of this program, the lungs were segmented from soft tissue, bones and large vessels using a region growing algorithm with a threshold value of < 50 % of the mean muscle signal obtained at 3 different positions across the imaging volume. Voxels erroneously classified as lung tissue (e.g. trachea) were manually removed from the lung segmentation. A second threshold value of < 15 % of the muscle signal was used to define emphysematous lung parenchyma. Both thresholds followed the study of Roach et al. on the performance of ultrashort echo time (UTE) MRI for evaluating pulmonary emphysema [13], but were adjusted empirically to the T2-weighted sequence used in this present study corresponding well to the –930 HU selected for emphysema assessment on LDCT. The chosen MRI thresholds were determined by one author of the manuscript who was not taking part in qualitative and quantitative emphysema analysis to avoid a bias. The thresholds were chosen after investigating other different thresholds (e.g. < 70 % for lung segmentation and < 10 % for definition of emphysema).

**Statistical analysis**

Statistical analysis was performed with SPSS 24 (IBM, Armonk, New York, USA). Spearman coefficient was applied for correlation of qualitative scores between MRI and LDCT (presence/severity and morphological patterns). Pearson coefficient, linear regression analysis and Bland-Altman plot were applied for comparison of emphysema indices as calculated by MRI and LDCT. The Mann-Whitney U-Test was used to define differences of qualitative scores and emphysema indices between MRI and LDCT.

**Results**

All 30 cases with pulmonary emphysema according to low-dose CT (LDCT) were accurately detected by MRI. There were 3 subjects who seemed to have emphysema on MRI according to qualitative assessment, yet they did not show emphysematous changes on LDCT (false-positive results). One of these subjects showed multiple centrilobular lucencies and a slightly elevated emphysema index (EI) of 5.4 % according to MRI, while the LDCT did not show emphysema and an EI of 0.9 %. LDCT and corresponding MR images of this case are shown in [Fig. 1](#). In the other two cases, quantitative measurement on MRI did not correspond to emphysema with indices of 2.7 % and 2.1 %, respectively. The mean qualitative emphysema score was significantly higher in the emphysema group compared to the control group for MRI (1.47 vs. 0.10, p < 0.001) and for LDCT (1.50 vs. 0.0, p < 0.001). The scores regarding severity and morphological patterns of emphysema correlated significantly between MRI and
LDCT ($r = 0.912$ and $p < 0.001$ for severity in the emphysema group; $r = 0.668$ and $p < 0.001$ for severity in the control group; $r = 0.843$ and $p < 0.001$ for emphysema pattern in the emphysema group; $r = 1.000$ and $p < 0.001$ for emphysema pattern in the control group).

▶ Fig. 2 shows an example of the qualitative assessment of emphysema.

▶ Table 1 shows the number of morphological patterns of emphysema as detected by MRI and CT.

<table>
<thead>
<tr>
<th>Pattern</th>
<th>MRI</th>
<th>CT</th>
</tr>
</thead>
<tbody>
<tr>
<td>centrilobular</td>
<td>13</td>
<td>13</td>
</tr>
<tr>
<td>paraseptal</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>mixed or advanced destructive</td>
<td>16</td>
<td>17</td>
</tr>
</tbody>
</table>

The slight discrepancies between the images are due to different breathing positions (CT images were acquired in inspiratory breath-hold while acquisition of MR images was gated to the expiratory phase of the respiratory cycle).

▶ Fig. 3 shows an example of the software-based quantitative analysis of emphysema.

The semi-automated software-based segmentation of the lung from the surrounding soft tissue, bones and vessels on MRI was technically successful in all 60 cases. The manual effort for the correction of lung segmentation was less than 3 minutes per case. Representative images of quantitative emphysema analysis are shown in ▶ Fig. 3.

The emphysema index was significantly higher in the emphysema group for MRI and LDCT ($p < 0.001$) (Table 2) with significant correlation between MRI and LDCT ($r = 0.960$ and $p < 0.001$ for emphysema group; $r = 0.746$ and $p < 0.001$ for control group). The Bland-Altman plot and linear regression analysis are shown in ▶ Fig. 4.
Discussion

The main finding of this study is that T2-weighted PROPELLER MRI, which has been shown to be suitable for lung imaging [18 – 21], may also be used for the assessment of pulmonary emphysema despite its lower spatial resolution compared to CT. The results are comparable to previously published studies using UTE imaging and functional MRI in subjects with emphysema [10 – 13, 15], while being technically more easy to implement.

The software-based quantitative assessment of pulmonary emphysema on CT images is well established. Regarding normal-

| Table 2 | Mean emphysema index (EI) as calculated on MR and CT images. |
|-----------------|-----------------|-----------------|-----------------|
| **Emphysema group** | **EI MRI** | **EI LDCT** |
| 11.6 ± 11.3 | 11.6 ± 10.3 |
| **Control group** | 1.4 ± 1.2 | 0.7 ± 0.7 |

Tab. 2 Durchschnittlicher Emphysem-Index (EI) berechnet in MRT- und CT-Aufnahmen.

Fig. 4 Linear regression of emphysema indices (EI) for emphysema group a and control group b, as well as Bland-Altman plot of emphysema indices for emphysema group c and control group d.

Abb. 4 Lineare Regression der Emphysem-Indices (EI) für die Emphysemgruppe a und die Kontrollgruppe b, sowie Bland-Altman-Diagramm der Emphysem-Indices für die Emphysemgruppe c und die Kontrollgruppe d.
dose CT with a slice thickness of 1 mm, a threshold of –950 HU seems to be optimal for CT densitometry analysis of emphysema [4–6, 22]. Yet, in a statement of the Fleischner Society, Lynch et al. pointed out that excessive image noise with a reduced CT dose can simulate emphysema, particularly on quantitative CT [4]. Moreover, image quality and noise level with simulated mAs levels below 60 mAs were significantly inferior to images with higher simulated mAs levels in a study by Ley-Zaporozhan et al. [23]. They concluded that imaging dose could be lowered to 60 mAs in thin-slice CT without a diagnostically relevant increase in noise impairing image quality. Hence, the optimal threshold for emphysema quantification with low-dose CT using mAs levels below 30 mAs as in our study has yet to be determined. After testing different thresholds from –910 to –950 HU, we decided to use a midway threshold of –930 HU in our study, since this value correlated best with the results of the qualitative emphysema analysis (in many cases with a threshold of –950 HU; lucencies that were apparently related to emphysema were not indicated as such by the CT software tool, while a threshold of –910 HU led to an obvious overestimation in visual analysis). This is in contrast with a study by Gierada et al., who showed that there were no significant differences between normal-dose and LDCT for emphysema analysis [24]. However, the reconstructed slice thickness in their study was 5 mm and the tube current-exposure time product was 30–60 mAs, while slice thickness and radiation dose of LDCT were much lower in our study (2 mm and 25 mAs). In addition, the absolute CT threshold was not particularly important for this study, since our intention was the correlation with MRI, and not to find the most suitable threshold for densitometry analysis on LDCT.

Based on our study results, pulmonary emphysema may be assessed quantitatively with structural T2-weighted PROPELLER MRI. Still, it should be mentioned that the correlation was excellent when emphysema was present, but a little less valid in subjects without emphysema.

To the best of our knowledge, there is only one study by Roach et al. [13] including software-based emphysema quantification using structural MRI (UTE). However, high-resolution UTE imaging of the lung might not be feasible in the clinical routine, since it is technically demanding and might take up to 20 minutes for image acquisition [12, 13]. In comparison, most clinical MRI scanners should be able to yield a T2-weighted PROPELLER sequence similar to the one being used in our study. Thus, the presented approach may be more transferable to clinical routine. Our study results suggest that T2-weighted PROPELLER MRI may have the potential to be used for the quantification and phenotyping of severe pulmonary emphysema, and subsequently for the identification of progression in follow-up examinations to avoid the radiation exposure of repeated CT scans. At the same time, this approach would allow for the detection of other relevant findings, such as pulmonary nodules or inflammatory changes as previously shown [10, 12, 21].

Our study has several limitations. First, this is a retrospective study, since the images were not primarily acquired for the analysis of pulmonary emphysema. However, the intention of the present study was to evaluate whether emphysema could be assessed with a conventional T2-weighted PROPELLER sequence that can be easily applied in the clinical routine. Second, the results of emphysema analysis were not correlated with spirometric parameters such as the forced expiratory volume in the first second/forced vital capacity ratio (FEV1/FVC). This is desirable in future studies. The third limitation is the low sample size of our study population. A fourth limitation is that some LDCT datasets with distinct noise overlay (especially seen in adipose individuals) or extensive atelectasis/infiltrates were excluded from further analysis with MRI, which may have led to a preselection bias. And fifth, subjects with mild emphysema were not part of the analysis. Even though it was not evaluated in this study, we have to assume that the current T2-weighted PROPELLER sequence would probably not be capable of detecting slight emphysematous changes as reliably as severe emphysema due to the much lower spatial resolution compared to LDCT. Still, this did not have a major influence on the visual scoring.

In conclusion, the presence and extent of pulmonary emphysema may be assessed qualitatively and quantitatively using T2-weighted PROPELLER MRI with very good correlation to LDCT according to the present study. However, our study results should be validated in larger prospective studies.

**Conflict of Interest**

The authors declare that they have no conflict of interest.

**References**


