Micro-computed tomography of pulmonary fibrosis in mice induced by adenoviral gene transfer of biologically active transforming growth factor-β1

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Abstract

Background: Micro-computed tomography (micro-CT) is a novel tool for monitoring acute and chronic disease states in small laboratory animals. Its value for assessing progressive lung fibrosis in mice has not been reported so far. Here we examined the importance of in vivo micro-CT as non-invasive tool to assess progression of pulmonary fibrosis in mice over time.

Methods: Pulmonary fibrosis was induced in mice by intratracheal delivery of an adenoviral gene vector encoding biologically active TGF-β1 (AdTGF-β1). Respiratory gated and ungated micro-CT scans were performed at 1, 2, 3, and 4 weeks post pulmonary adenoviral gene or control vector delivery, and were then correlated with respective histopathology-based Ashcroft scoring of pulmonary fibrosis in mice. Visual assessment of image quality and consolidation was performed by 3 observers and a semi-automated quantification algorithm was applied to quantify aerated pulmonary volume as an inverse surrogate marker for pulmonary fibrosis.

Results: We found a significant correlation between classical Ashcroft scoring and micro-CT assessment using both visual assessment and the semi-automated quantification algorithm. Pulmonary fibrosis could be clearly detected in micro-CT, image quality values were higher for respiratory gated exams, although differences were not significant. For assessment of fibrosis no significant difference between respiratory gated and ungated exams was observed.

Conclusions: Together, we show that micro-CT is a powerful tool to assess pulmonary fibrosis in mice, using both visual assessment and semi-automated quantification algorithms. These data may be important in view of pre-clinical pharmacologic interventions for the treatment of lung fibrosis in small laboratory animals.

Background

Pulmonary fibrosis is a severe, chronic lung disease, which is associated with high mortality rates, afflicting more than five million patients worldwide [1]. Its most severe form, idiopathic pulmonary fibrosis (IPF) is typically associated with an average survival of 2-5 years from time of diagnosis [2,3]. Depending on the underlying cause, progression of pulmonary fibrosis is associated with alveolar epithelial cell injury and hyperplasia, inflammatory leukocyte accumulations within the bronchoalveolar space and lung parenchymal tissue, as well as fibroblast hyperplasia and exuberant deposition of extracellular matrix (ECM) components such as collagen and fibronectin in the lungs, eventually leading to formation of honey comb cysts and scar tissue [2,4]. Specifically in IPF, immunosuppressive and anti-inflammatory therapies have proved to be ineffective in preventing the fatal consequences of lung tissue remodelling characterizing pulmonary fibrosis.

Over the past few years, it has become evident that animal model systems to study the pathogenesis of lung fibrosis in rodents need to be more carefully evaluated...
with respect to their applicability to human lung fibrosis [4]. As an example, application of the anti-neoplastic
drug bleomycin into the lungs of mice or rats is the
most frequently employed model system to study the
pathogenesis of this devastating lung disease in small
laboratory animals. However, in contrast to human lung
fibrosis frequently developing over years and decades,
bleomycin treatment of mice is characterized by early
alveolar epithelial cell injury and acute lung inflamma-
tion followed by a transient lung fibrotic response,
thereby not adequately reflecting human disease pro-
gression [4]. Another previously described model system
to initiate fibrosis in small laboratory animals makes use
of somatic cell gene transfer of biologically active trans-
forming growth factor-β1, which, due to its unique
fibrogenic characteristics, has been shown to trigger
slowly progressing lung fibrosis in rats, which is irrever-
sible and lacks acute inflammatory episodes [5].

Diagnostic tools to assess the degree of lung fibrosis in
humans currently include lung CT scans, lung function
tests and histological examinations of lung tissue biops-
ies. In mice, where complete lungs can be removed,
particularly histological and biochemical approaches are
employed to assess the degree of fibrosis [6,7]. In recent
years, small animal imaging has emerged as an innova-
tive and powerful, non-invasive tool to perform follow
up of both acute and chronic disease processes in
laboratory animals over time [8-11]. Various non-inva-
sive techniques such as micro-CT, micro-positron emis-
tion tomography (micro-PET), and magnetic resonance
imaging (MRI) are innovative tools that may be useful
for evaluation of novel anti-fibrogenic therapeutic strate-
gies in mouse models in vivo. However, systematic
micro-computed tomography-based analysis of develop-
lung fibrosis induced by gene transfer of biologically
active transforming growth factor-β1 as compared to
classic histopathological approaches to monitor fibrotic
disease progression in mice using both visual assessment
and semi-automated quantification algorithms has not
been reported so far.

Therefore, in the current study, pulmonary fibrosis
was induced in mice by gene transfer of biologically
active transforming growth factor-β1, and fibrotic dis-
ease progression was evaluated by micro-computed
tomography and then correlated to histological scoring
of lung fibrosis as pioneered by Ashcroft [12].

**Methods**

**Adenoviral vectors**

Two adenoviral vectors were employed, which were
constructed as outlined in detail previously [5]. AdTGF-
β1[223-225] contains the cDNA of the coding region
of full-length porcine TGF-β1, while containing a mutation
of cysteine to serine at positions 223 and 225, rendering
expressed TGF-β1 biologically active [5]. Empty viral
vector AdDL70-3 constructed as previously described
was used as control [13].

**Induction of pulmonary fibrosis**

Groups of female C57BL/6 mice (30 mice, age 8-12
weeks, weight 18-20 g) were either infected with AdTGF-
β1[223-225] (herewith referred to as AdTGF-β1, 18 mice) or
control vector AdDL70-3 (12 mice) via intratracheal
instillation, as described recently in detail [6,14,15].
Briefly, anaesthetized mice were orotracheally intubated
with an Abbocath (Abbott, Wiesbaden, Germany), fol-
lowed by careful intratracheal instillation of adenoviral
vectors (~1 × 10⁸ plaque-forming units [PFU] per
mouse) suspended in a total volume of 50 μl of phos-
phate-buffered saline (PBS). At 1, 2, 3, and 4 weeks after
adenoviral transfer, mice (AdTGF-β1 for week 1-4: n = 3,
n = 6, n = 6, n = 3; AdDL70-3 for week 1-4: n = 3 for
each time point) were subjected to micro-computed
tomography (Figure 1), followed by immediate sacrifice
and isolation of lungs for histopathological evaluation of
the degree of pulmonary fibrosis, using the scoring sys-
tem described by Ashcroft et al. [12].

**Lung histology and assessment of pulmonary fibrosis**

Complete lungs were inflated in situ with PBS-buffered
(4.5%, pH 7.0) formalin (Roth, Darmstadt, Germany), and
were then carefully removed and immersed in PBS-buf-
fered formalin. Lung tissue samples were paraffin-
embedded, and sections of 5 μm were prepared and
stained with haematoxylin/eosin (H/E) and Masson’s or
Elastica-van-Giesen staining. The degree of lung fibrosis
was determined according to the method described by
Ashcroft and colleagues [12], employing a numerical scal-
ing system in lung samples ranging from 0 (normal lung)
to 8 (total fibrous obliteration of the field). The mean
degree of lung fibrosis according to the definitions by Ash-
croft et al. [12] was calculated from individual scores of
~15 microscopic fields analyzed per mouse lung.

**Micro-CT scans of mice**

A GE Explore Locus cone-beam micro-CT (GE Health-
care, Chalfont St Giles, GB) was used for data-acquisi-
tion in prone position under isoflurane inhalation
anaesthesia (tube voltage 80 kV, tube current 450 μA,
0.094 mm effective pixel size) with and without respira-
tory gating (i.e. synchronization of acquisition of micro-
CT projections with a timepoint in the respiratory cycle
of the individual mouse). Scanning took approximately
10 and 20 min without and with respiratory gating,
respectively. Respiratory monitoring was performed
using a pressure transducer pad under the animal’s
chest. Images were reconstructed and assessed at a con-
stant window width/window level (5000/2000).
Assessment of fibrosis by observer rating scale
Three radiologists blinded to the experimental groups and respiratory gating strategies, independently assessed image qualities of all individual entire micro CT scans, using a rating scale ranging from 1 (very good) to 5 (insufficient) and the amount of lung consolidations using a rating scale ranging from 1 (no consolidation) to 5 (very pronounced consolidation).

Post-processing
Due to the diffuse nature of fibrotic consolidations, direct quantification of the pathology of lung fibrosis by measurement of focal fibrotic lesions was not feasible. Therefore, quantification of aerated lung areas known to be compromised by developing lung fibrosis was used as a surrogate marker for progressive lung fibrosis. A region-growing segmentation algorithm was applied on each individual entire micro CT scan using the MevisLab software package (Mevis Research, Bremen, Germany) [16]. The algorithm uses the seed points as starting points. From these points the segmentation volume “grows”, with the structures included in the volume depending on the mathematical parameters. The segmentation algorithm was originally developed for quantification of diffuse lung tumors in transgenic mice, and was slightly modified for its use in the current study. Segmentation was carried out using 20-40 seed points, the segmentation threshold tolerance of the region growing algorithm was set to 2%.

Statistical analysis
All statistical computations have been performed using the statistical program SPSS 17 (SPSS Inc., Chicago, Ill., USA) The significance level of all tests has been set to 0.05 (5%). Because most of the observed variables were ordinal scaled we used non-parametric tests for statistical analysis. Significance of differences in image quality values were tested for all three observers in respiratory gated and respiratory ungated exams using a Wilcoxon signed ranks test. Furthermore, significance of differences in consolidation assessment values for respiratory gated and respiratory ungated exams was tested using a Wilcoxon signed ranks test for all three observers. Differences between the assessment of the 3 observers were evaluated comparing the 95% confidence interval for the mean for assessment values of the image quality for respiratory gated/ungated exams and the assessment values of the consolidation assessment for respiratory gated/ungated exams.

Correlation of the Ashcroft histology scores (assessing the degree of fibrosis) and the lung consolidation values (obtained by assessment of the images by three observers) was calculated for respiratory ungated/respiratory
gated exams and all three observers using Spearman’s rank correlation coefficient for ordinary scaled variables.

Correlation of the Ashcroft histology scores and the lung segmentation values (obtained by the semiautomated segmentation routine described above) was calculated for respiratory ungated and respiratory gated exams using a Pearson correlation for continuous variables. To assess differences between the 4 time points for histology scores, visual consolidation assessment and segmentation of aerated lung areas 95% confidence intervals were plotted against the 4 time points and differences were tested for significance.

**Results**

**In vivo micro-CT imaging of mouse lungs after intrapulmonary delivery of control vector or AdTGFβ1**

As shown in Figure 1 micro-CT imaging of mouse lungs subjected to treatment with control vector or application of AdTGFβ1 for up to 4 weeks revealed that no major consolidations were detectable in control vector treated mice, whereas AdTGFβ1 treatment of mice provoked strong consolidations easily detectable particularly on weeks 2 and 3 post-treatment (Figure 1).

**Assessment of image quality and consolidations in respiratory gated as compared to ungated exams of fibrosing lungs**

We next questioned whether detection of consolidations in fibrosing mouse lungs would differ between observers. However, initial comparative image quality assessment between three different observers revealed only slightly increased mean image quality values in respiratory gated (MVSD 2.70, MV CVF 2.33, MV TR 2.27) as compared to ungated exams of fibrosing lungs (MVSD 2.33, MV CVF 2.10, MV TR 1.97). Moreover, all three observers determined similar consolidation assessment values for respiratory gated (MV SD 2.13, MV CVF 2.40, MV TR 2.30) as compared to ungated exams (MV SD 2.13, MV CVF 2.43, MV TR 2.40).

Regarding the assessment variability between the 3 observers, we found that the lower and upper bound of the 95% confidence interval for the mean for the 3 observers showed no relevant differences except for the image quality assessment in gated exams. Here, lower and upper bound of the 95% confidence interval for the mean were distinctly higher for observer SD. The Cohen’s kappa statistic also showed significant agreement between nearly all observer scores, but no significant agreement was found for the image quality evaluation scores by observers SD and TR for gated (p = 0.051) and ungated exams (p = 0.699). Overall, these data illustrate that visual assessment of micro-CT based lung fibrosis in mice was independent of the observer.

**Correlation of histological grading of lung fibrosis with micro-CT based fibrosis grading**

We next compared classical Ashcroft scoring with micro-CT based evaluation of TGFβ-1 induced lung fibrosis in mice. Ashcroft histology scores and lung consolidation values showed significant correlations for all observers and respiratory gated as well as respiratory ungated exams using Spearman’s rank correlation coefficient (p < 0.001, Figure 2). Moreover, we also assessed progression of pulmonary fibrosis in mice by making use of a semi-automated region growing segmentation algorithm, where aerated lung volume was used as a surrogate marker for pulmonary fibrosis, which is illustrated in Figure 3. Ashcroft histology scores and lung segmentation values showed significant negative correlation using a Pearson correlation test for respiratory gated (p = 0.004) and respiratory ungated exams (p = 0.006) (Figure 4).

Due to the limited number of study objects within each group (AdTGF-β1 for weeks 1-4: n = 3, n = 6, n = 6, n = 3), we were not able to detect significant differences between different time points for visual consolidation assessment and segmentation of aerated lung areas. However, to better exemplify the time course of fibrosis in mice on days 7, 14, 21, and 28 post treatment, mean values and standard deviations for Ashcroft histology scores and visual assessment of all three observers for ungated exams are given (histology for control group: 0.15 ± 0.03; 0.14 ± 0.01; 0.17 ± 0.07; 0.15 ± 0.05/histology for fibrosis group: 1.79 ± 0.58; 2.13 ± 0.5; 1.67 ± 0.84; 1.64 ± 0.45/visual consolidation assessment scores for control group: 1.11 ± 0.19; 1.11 ± 0.19; 1.22 ± 0.38; 1.11 ± 0.19/visual consolidation assessment scores for fibrosis group: 3.56 ± 0.58; 4.06 ± 1.11; 2.50 ± 0.89; 2.00 ± 0.72).

**Discussion**

In this study we evaluated the use of micro-CT for assessment of pulmonary fibrosis induced by adenoviral gene transfer of biologically active TGF-β1 in mice. We found micro-CT imaging to be a highly valuable tool to study pulmonary fibrosis progression in mice using both visual inspection of lung images and a semi-automated segmentation algorithm that estimates aerated lung areas as a surrogate marker inversely correlating with developing lung fibrosis in mice.

Micro-CT allows a detailed assessment of lung morphology due to high density differences and well defined borders between air and lung tissue. However, one limitation of this method is that the activity of inflammatory processes can only be assessed by secondary characteristics by micro-CT, such as affection of adjacent structure or pleural effusion. Other imaging techniques such as magnetic resonance imaging (MRI),
micro-positron emission tomography (PET) and optical imaging could provide functional data, with micro-PET being an especially promising candidate to robustly assess inflammatory activity by increase of glucose metabolism as detected by $^{18}$F-fluoro-deoxyglucose micro-PET [17-19], which might be particularly relevant in terms of monitoring acute exacerbations of pulmonary fibrosis in small laboratory animals.

Image quality is of equal importance in small animal imaging as in human imaging. However, data on imaging protocols are still not extensive. One major issue addressed in previous studies relates to respiratory gating. Various gating techniques, including prospective and retrospective as well as intrinsic and extrinsic techniques have revealed that mostly, gated imaging resulted in increased image quality [20,21]. In the current study, we found that mean values for image quality were better in gated exams for all observers, however differences were minor and not significant. Both gated and ungated exams resulted in a highly significant correlation with histology-based Ashcroft scores. Therefore, we conclude that gating should be applied whenever possible, but is not an absolutely critical issue when this setup is not available.

A further important aspect of micro CT imaging of normal or fibrosing mouse lungs is the time-point in the respiratory cycle when images are acquired. Due to the typical respiratory pattern in anesthesia, with a short breath followed by a long expiratory plateau, imaging is mostly performed in expiration. Under those experimental conditions, where intratracheal intubation and ventilation is performed, micro-CT imaging can also be performed in inspiration, as a different and consistent respiratory pattern can be created [22]. Without gating, imaging is effectively performed during expiration, due to the longer period of time as compared to inspiration that only represents a small fraction contributing to the average imaging time-point.

The semi-automated segmentation routine described in this study was found to assess consolidations in a quantitative way, as proven by the significant correlation of segmentation values with Ashcroft-based histological grading. To a minor degree, observer interaction is possible, as the observer has to place its own seed points for the region growing algorithm. As seed points have to be placed carefully within aerated lung areas, inadvertent positioning of seed points within non-aerated lung areas may increase the likelihood that consolidated aeras will be included in the segmentation volume and thus may cause assessment bias, which however, according to the presented data, appears to be very low.

In the current study, segmentation volumes were not correlated with total lung volume or body weight of
Figure 3 Semiautomated region growing segmentation for assessment of pulmonary consolidation 3 weeks after instillation of control vector AdDL70-3 (A-C) or AdTGF-β1 (D-E). Aerated lung volume is used as an inverse surrogate marker for pulmonary fibrosis. (A, D) Axial micro-CT. (B, E) Placement of seed points for the region growing segmentation in the aerated lung. Seed points are enlarged for better detection. (C, F) Segmentation result shows aerated lung volume in white.

Figure 4 Correlation of aerated lung volume determined by region growing segmentation with histological assessment of lung fibrosis by Ashcroft scoring in mice treated with control vector (A, showing very low Ashcroft histology scores (i.e. no fibrosis) and normal aerated lung volume) or AdTGF-β1 (B). Aerated lung volumes are plotted against Ashcroft scores for each animal in respiratory gated and respiratory ungated exams, as indicated. Linear interpolation is given for respiratory gated and ungated exams, respectively. Note the different scaling.
individual mice. However, when considering that differences in expiratory total lung volumes between individual mice of the same age and strain are negligible, it may be largely excluded that this aspect may have confounded the reported correlations. Future algorithms might take segmentation assessments of the entire lung into account, thereby allowing the correlation of measured consolidations with total lung volumes, or simply by employing a quotient taking the body weight into account.

Additionally, different techniques of fibrosis assessment could be applied. A promising approach is the measurement of CT densities expressed by Hounsfield Units and assessing fibrosis not only by consolidation but also by evaluation of density distribution [23]. Morphological pattern assessment as applied in human fibrosis assessment has also been described for imaging of lung fibrosis in small laboratory animals [24,25]. However, such approaches would be dependent on an image quality that is consistently comparable to human imaging, which is technically still difficult to achieve in in vivo imaging of small laboratory animals.

In another study we evaluated the applied radiation dose for the protocols used in this study applying phantom and cadaver thermoluminescence dosimetric measurements (own unpublished observations). The expected mean doses in an average C57BL/6 mouse for the respiratory gated and ungated protocols used in this study were 201 mGy and 194 mGy respectively. The expected mean value for an estimated time of 1 minute fluoroscopy to determine the scan field of view was 22 mGy. Day et al. reported that high radiation doses of 7 - 9 Gy did not result in significant lung fibrosis in mice after 90 days, just few microscopically small collagen-rich foci were detected subpleurally [26]. Therefore, it appears that radiation-induced fibrosis can be largely excluded even when serial examinations are performed.

Human lung fibrosis differs from the animal model reported here, as it is irreversible and shows typical morphological patterns in computed tomography representing definite fibrotic changes of the lung. The application of the methods reported for imaging in human lung fibrosis therefore is not intended. We believe that the currently applied technique will primarily be helpful to better assess e.g. novel therapeutic strategies for the treatment of pulmonary fibrosis in rodent model systems.

**Conclusion**

In conclusion, the current study shows that respiratory gated/ungated micro-CT allows valid in vivo evaluation of the degree of pulmonary fibrosis in mice using both visual assessment and semi-automated quantification algorithms. Micro-CT thus offers a non-invasive imaging tool for evaluation of novel therapeutic strategies for the treatment of pulmonary fibrosis in mice.

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**Authors’ contributions**

TR, CvF and UAM designed the study. TR, RM, RH and UAM carried out the preparation of the animals and micro-CT exams. TR, CvF and SD read and assessed the imaging studies. LF performed the histopathological workup. LH performed the statistical analysis. KA, MK, JG, TW, MG and JB participated in its design and coordination and helped to draft the manuscript. TR and UAM wrote the manuscript. All authors read and approved the final manuscript.

**Competing interests**

The authors declare that they have no competing interests.

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