The relationship between lung function impairment and quantitative computed tomography in chronic obstructive pulmonary disease

O. M. Mets · K. Murphy · P. Zanen · H. A. Gietema · J. W. Lammers · B. van Ginneken · M. Prokop · P. A. de Jong

Received: 5 April 2011 / Revised: 1 June 2011 / Accepted: 2 July 2011 / Published online: 12 August 2011 © The Author(s) 2011. This article is published with open access at Springerlink.com

Abstract

Objectives To determine the relationship between lung function impairment and quantitative computed tomography (CT) measurements of air trapping and emphysema in a population of current and former heavy smokers with and without airflow limitation.

Methods In 248 subjects (50 normal smokers; 50 mild obstruction; 50 moderate obstruction; 50 severe obstruction; 48 very severe obstruction) CT emphysema and CT air trapping were quantified on paired inspiratory and end-expiratory CT examinations using several available quantification methods. CT measurements were related to lung function (FEV₁, FEV₁/FVC, RV/TLC, Kco) by univariate and multivariate linear regression analysis.

Results Quantitative CT measurements of emphysema and air trapping were strongly correlated to airflow limitation (univariate r-squared up to 0.72, \( p < 0.001 \)). In multivariate analysis, the combination of CT emphysema and CT air trapping explained 68-83% of the variability in airflow limitation in subjects covering the total range of airflow limitation (\( p < 0.001 \)).

Conclusions The combination of quantitative CT air trapping and emphysema measurements is strongly associated with lung function impairment in current and former heavy smokers with a wide range of airflow limitation.

Key Points
- CT helps to automatically assess lung disease in heavy smokers
- CT quantitatively measures emphysema and small airways disease in heavy smokers
- CT air trapping and CT emphysema are associated with lung function impairment

Keywords Pulmonary emphysema · Respiratory bronchioles · Quantitative computed tomography · Pulmonary function test · Chronic obstructive pulmonary disease
Introduction

Chronic Obstructive Pulmonary Disease (COPD) is characterized by chronic airflow limitation, and its severity is graded according to the Global Initiative for Chronic Obstructive Lung Disease (GOLD) criteria [1]. The airflow limitation in COPD patients is caused by parenchymal destruction (i.e. emphysema) and small airways disease with airway inflammation and airway remodeling, which leads to expiratory airway obstruction and air trapping [2, 3]. Emphysema has been quantified in vivo by computed tomography (CT) for decades [4], but in vivo detection of small airways disease has been difficult since these small airways (diameter <2 mm) cannot be visualized directly with current CT techniques. Nakano et al suggested that measurements of large airway wall thickening can be used as an indirect sign of small airways disease [5]. However, quantification of airway wall dimensions remains technically challenging. Alternatively, air trapping on end-expiratory CT may be easier to quantify, and is possibly a more direct measurement of small airways dysfunction. For this, the main challenge is to separate air trapping due to small airways disease from air that remains in emphysematous spaces after expiration [6].

Separate quantification of air trapping and emphysema is relevant for two reasons. First, COPD is a disease with a wide heterogeneity [7–10], and COPD phenotyping may be possible when quantitative CT is combined with clinical and physiological characteristics. Second, quantitative CT of COPD components might gain a role in (early) diagnosis of the disease; a risk estimate for COPD may be calculated from CT data, providing additional information in subjects who undergo CT. This strategy may be especially interesting in lung cancer screening, where large numbers of CT examinations are obtained in heavy smokers. Here, CT may reveal ‘hidden’ disease in undiagnosed COPD patients, and might reveal disease in patients with lung function still within the normal range [11].

The purpose of this study was to determine the relationship between lung function impairment and quantitative CT measures of air trapping and emphysema in heavy smokers with and without COPD.

Methods

To study COPD in a lung cancer screening trial [12], pulmonary function tests (PFT) and an expiratory CT data acquisition had previously been added to the screening protocol in our center. In the present study, we automatically assessed emphysema and air trapping in CT examinations obtained in this lung cancer screening population. Since limited subjects with severe airflow limitation were present in the screening trial, we added clinical subjects to be able to study the total range of COPD.

This retrospective study was approved by our institutional ethics board; written informed consent was obtained in the screening trial participants, and waived for the clinical subjects.

Subjects

Within the lung cancer screening trial participants, random samples of 50 subjects were drawn for each GOLD stage [1], resulting in 50 GOLD 1, 50 GOLD 2 and 24 GOLD 3 subjects. The clinical subjects underwent outpatient inspiratory and expiratory CT and PFT examination for evaluation of COPD and/or lung transplantation screening. Cases with GOLD 3 of GOLD 4 were selected. No CTs contained major breathing artifacts, and scans with signs of acute pneumonia were excluded. The GOLD 3 study population was extended to 50 subjects with a random sample of clinical subjects. The GOLD 4 group consisted of all available clinical subjects (N=48).

Additionally, we selected a group of normal smokers from the screening trial participants. We defined ‘normal smoker’ as two normal lung function tests with at least a two-year interval, plus a slow rate of lung function decline. Normal lung function was defined as FEV1 (forced expiratory volume in one second) >80% predicted and FEV1/FVC (forced expiratory volume in one second / forced vital capacity) >70%. A slow rate of decline was defined as an FEV1 decline less than 29 ml per year [13]. Seventy subjects fulfilled these criteria, of whom 50 were randomly selected.

Pulmonary function testing

Spirometry, body plethysmography and diffusion capacity testing was obtained according to ERS guidelines [14]. PFT was performed with ZAN, Oberthulba, Germany. Spirometry was obtained in all subjects and included forced expiratory volume in 1 second (FEV1, forced vital capacity (FVC)) and the ratio of forced expiratory volume in 1 second over forced vital capacity (FEV1/FVC). Data on bronchodilator use on the day of examination was not available, and reversibility with bronchodilation was not assessed. Due to restrictions in testing capacity during screening rounds, extensive pulmonary function testing was only performed in a subgroup of subjects. For our study population, RV/TLC (ratio of residual volume to total lung capacity) was available in 121 subjects, and Kco (ratio of diffusion capacity over alveolar volume) in 103 subjects. For lung cancer screening subjects, PFT was performed on the same day as CT. For clinical subjects, the PFT
performed closest to the CT date was selected (median 1 day, interquartile range 0–4 days).

Computed tomography

Volumetric inspiratory CT and volumetric end-expiratory CT after standardized breathing instructions were obtained between July 2007 and September 2008. All CT examinations were performed without intravenous contrast medium, using 16 slice CT (Brilliance 16P, Philips Medical Systems, Cleveland, OH) and 0.75 mm collimation. Settings were adjusted to body weight: 120 kVp (≤80 kg) or 140 kVp (>80 kg), both with 30 mAs, for inspiratory CT, and 90 kVp (≤80 kg) or 120 kVp (>80 kg), both with 20 mAs, for expiratory CT. A smooth reconstruction filter (B filter, Philips, Cleveland, OH) was used, according to the protocol. Images with slice thickness of 1.0 mm at 0.7 mm increment were reconstructed from lung bases to lung apices.

The clinically evaluated patients that were added to the screening population, underwent volumetric inspiratory CT and volumetric end-expiratory CT after standardized breathing instructions between October 2003 and May 2010. CT was performed without any intravenous contrast medium. These CT examinations were acquired on various generations of CT equipment from the same vendor (Philips Medical Systems, Cleveland, OH) with 16x0.75 mm collimation on Brilliance 16P (N=16) and Mx8000 IDT (N=12), 40x0.625 mm collimation on Brilliance 40 (N=8), 64x0.625 mm collimation on Brilliance 64 (N=32) or 128x0.625 mm collimation on Brilliance iCT (N=6). Exposure settings were: 100kVp or 120kVp, both with 130mAs, for inspiratory CT, and 80-120kVp with 10-30mAs for expiratory CT. A sharp reconstruction filter (C kernel, Philips) was used, according to the clinical protocol. Images with slice-thickness of either 0.9 or 1.0 mm at 0.45 or 0.7 mm slice-spacing were reconstructed from lung bases to lung apices.

Emphysema and air trapping quantification

The lung parenchyma on inspiratory and expiratory CT data sets was automatically segmented [15]. Additionally, lung segmentation was checked visually, and those with major errors (see Appendix 1) were excluded. Subjects that were excluded were replaced by another random subject with a corresponding GOLD stage (N=16). A noise reduction filter [16] was applied to decrease the influence of noise on the quantitative CT measurements.

The extent of emphysema was assessed by two widely used quantitative CT measures:

1) The percentage of lung voxels below -950HU in inspiration (IN -950) [17, 18].

2) The HU value at the 15th percentile of the attenuation distribution histogram in inspiration (Perc 15) [19].

The extent of air trapping was quantified by four different methods:

1) The percentage of lung voxels below -850HU in expiration (EXP -850) [20, 21].

2) The percentage of lung voxels between -850HU and -910HU in expiration (EXP -850 to -910). To correct for air in emphysematous areas, method 1 was adjusted by excluding the voxels below -910HU, based on reference [22].

3) The change in relative lung volume with attenuation values from -860HU to -950HU (RVC -860 to -950) between paired inspiratory and expiratory examinations [6]. Practically, the RVC -860 to -950 is calculated using the formula: expiratory relative lung volume -860 to -950HU - inspiratory relative lung volume -860 to -950HU. In this, the relative lung volume -860 to -950HU is defined as the volume of voxels with attenuation values between -860 to -950HU divided by the volume of voxels with attenuation values above -950HU (i.e. limited lung volume, without emphysema) [6].

4) The expiration to inspiration ratio of mean lung density (E/I-ratioMLD) [23–25].

Data analysis

CT measurements were compared between the five groups (i.e. normal smokers and the four COPD stages) with analysis of covariance (ANCOVA), with age, height and sex as covariates. Univariate and multivariate linear regression analysis were performed to evaluate the relationship between lung function parameters (dependent variables) and the CT measurements (independent variables). Multivariate analysis incorporated CT emphysema (2 methods) and CT air trapping (4 methods), while age, sex and height were introduced as correction factors for PFT. In order to meet linear regression requirements, we used absolute values instead of the percentage of predicted value, as the latter often elicits a heteroscedastic distribution of the residuals to one side [26]. Furthermore, multicollinearity and the distribution of residuals was assessed for all models; FEV1 and IN -950 were log-transformed (logFEV1 and logIN -950) as to obtain a normal distribution, and warrant symmetrical variance of the errors around zero (i.e. homoscedasticity). Linear regression analyses were performed in the total cohort of 248 subjects, as well as in the 174 screening trial participants separately.

All statistical analyses were performed using SPSS software v15.0 (SPSS Inc, Chicago, Illinois, USA). A p-value below 0.05 was considered statistical significant. Continuous data are given as mean±SD, unless indicated otherwise.
Results

Study population characteristics

The majority were male and the average age was around 60 years, except for the subjects in the GOLD 4 group who were younger and more often female (Table 1). CT emphysema and CT air trapping results were worse in the COPD cases when compared to the smokers with normal lung function, and the results worsened in the subsequent COPD stages. Differences were significant except for IN<sub>-950</sub> between normal smokers to GOLD2, Perc15 between GOLD1 and GOLD2, EXP<sub>-850</sub> to EXP<sub>-910</sub> and E/I-ratio<sub>MLD</sub> between normal smokers and GOLD1, RVC<sub>-860</sub> to RVC<sub>-950</sub> between normal smokers and GOLD2, and EXP<sub>-850</sub> to EXP<sub>-910</sub> between GOLD3 and GOLD4.

Univariate regression analysis

Quantitative CT measurements showed moderate to strong association with lung function parameters of airflow limitation (R-squared values ranging from 0.25 to 0.72, p < 0.001 (Table 2, Fig. 1). CT emphysema measurements were better correlated to Kco compared to RV/TLC. Also, CT air trapping measurements were better correlated to RV/
TLC compared to Kco. These findings did not change when only the screening trial participants were included, though the associations were weaker after exclusion of the more severely affected subjects (Appendix 2).

Multivariate regression analysis

Testing of collinearity and residuals showed a variance inflation factor below 10 [27] in all models (max: 3.85), and a normal, homoscedastic distribution of residuals. The multivariate CT models largely explained the variability in airflow limitation; R-squared value 0.68 to 0.83 (Table 3). This pattern did not change when only the screening trial participants were included, though the explained variance was lower after exclusion of the more severely affected subjects; R-squared value 0.49 to 0.59 (Table 3).

Discussion

In a population that covered the range of airflow limitation from mild to very severe obstruction, we showed that the combination of quantitative CT measurements of emphysema and air trapping is strongly associated with lung function impairment, and explains a large proportion of the variability in airflow limitation.

Previous studies have addressed the small airways disease component in COPD by quantitative CT, but used large airway wall dimensions as a surrogate for small airways disease [28]. Nakano et al reported on the combination of large airway wall thickening and quantitative emphysema to explain lung function variability and to detect COPD subgroups. They evaluated 114 smokers, and showed that quantitative CT measurements of emphysema and airways wall thickness explained 23% to 49% of lung function variability [28]. In the present study we found a higher R-squared value; 68% to 83% for the whole study population, and 49% to 58% in the screening participants only. This difference in explained variance compared to the study of Nakano et al., who found an R-squared of 23% to 49%, is likely due to the fact that air trapping is a more direct measure of small airways dysfunction compared to airway wall thickness. Additionally, part of the increase may be due to differences between the study populations. Previously, Nakano et al showed that airway wall thickening in CT closely correlates with small airway dimensions in histological specimens, and thus can be used as an indirect sign of small airways disease [5]. Other groups have shown that the correlation between large airway measurements and lung function parameters increases with subsequent airway generations [29–32]. This supports our view that the signal is strongest in the more peripheral airways. Additionally, quantification of airway wall dimensions remains technically challenging, especially in the more distal generations (i.e. beyond the fifth generation). Therefore, assessment of air trapping with quantitative

### Table 3 Results of multivariate linear regression for quantitative CT measurements of emphysema and air trapping and pulmonary function tests

| CT Model | All subjects (N=248) | Subanalysis (N=174) |
|----------|----------------------|---------------------|
|          | logFEV1 | FEV1/FVC | logFEV1 | FEV1/FVC |
| logIN\_950 +EXP-850 | 0.81 | 0.76 | 0.54 | 0.58 |
| logIN\_950 +EXP-850 to -910 | 0.75 | 0.72 | 0.51 | 0.56 |
| logIN\_950 +RVC-860 to -950 | 0.83 | 0.75 | 0.53 | 0.55 |
| logIN\_950 +E/I-ratioMLD | 0.81 | 0.76 | 0.54 | 0.57 |
| Perc15 +EXP-850 | 0.81 | 0.75 | 0.54 | 0.55 |
| Perc15 +EXP-850 to -910 | 0.71 | 0.68 | 0.49 | 0.49 |
| Perc15 +RVC-860 to -950 | 0.83 | 0.75 | 0.55 | 0.56 |
| Perc15 +E/I-ratioMLD | 0.79 | 0.73 | 0.51 | 0.51 |

Data given are r-squared (R²) values

* Lung cancer screening trial participants only, controls to GOLD III; logIN\_950 = log-transformed emphysema score below -950HU in inspiratory scan; Perc15 = emphysema score as 15th percentile of attenuation distribution curve in inspiratory scan; EXP-850 = air trapping score below -850HU in inspiratory scan; EXP-850 to -910 = air trapping score between -850HU and -910HU in inspiratory scan; RVC-860 to -950 = air trapping score as relative volume change between -860HU and -950HU; E/I-ratioMLD = expiration to inspiration ratio of mean lung density
measures in expiratory CT may also have important technical benefits. Although there is an increase in radiation dose for an additional expiratory CT acquisition, this is limited; in our screening trial protocol the effective radiation dose was less than 0.65 mSv for the additional expiratory CT acquisition.

Previously, the univariate relationship between quantitative CT measures and lung function in COPD is reported in a few studies [23–25, 33, 34]. In these studies, correlation coefficients between E/I-ratioMLD and FEV1 of 0.45 and 0.64 ($p<0.001$) are reported, which is lower than what we found. Reported correlation coefficients with RV/TLC are between 0.54 and 0.71, which is also slightly lower than our results. These differences may well be due to differences in study populations, since our study population was larger and equally covered all the stages of COPD. Further, RVC−860 to −950 has been previously studied by Matsuoka and colleagues in a cohort of 36 COPD subjects with GOLD stage 1–4 [6]. They found correlation coefficients of 0.80 and 0.78 between this CT air trapping measure and FEV1 and FEV1/FVC, respectively [6]. Despite the considerably smaller study population their results are comparable to ours, both for the whole population, and the less severely affected trial participants only. Last, the EXP−850 method has been published only in abstract format [20], and was previously used in asthma research for quantification of air trapping [21].

One has to be careful with the conclusion that one method is better than the other, given the absence of a clear reference standard for either emphysema or air trapping in our study. In fact, the main purpose of our study was to assess how well the quantitative CT measurements can explain the lung function impairment, instead of testing for the optimal method. Nevertheless, we feel that air trapping measurements that combine inspiratory and expiratory CT data may be preferable when ones purpose is to quantify the individual disease components in COPD (i.e. emphysema and small airways disease). This radiologic separation of disease components may be a next step in further defining morphologic phenotypes within COPD. The solely expiratory measurement of air trapping (i.e. EXP−850) is not useful for this purpose, because it does not correct for air that remains in emphysematous spaces on end-expiratory CT; it thus combines quantification of emphysema and small airways disease in one measure. Therefore, we intended to eliminate emphysema from this measurement by calculating the EXP−850−910−90 since Gevenois et al demonstrated that voxels with a density below −910HU on expiratory CT represent emphysema [22]. However, the performance of the models including this measure were worse, which makes us believe that the air trapping measurements that combine inspiration and expiration CT data will outperform the EXP−850−910 when confronted with a proper reference standard.

Since only a moderate number of GOLD3 and no GOLD4 subjects were present in the screening trial, we added clinical subjects with advanced airflow limitation to our study population. Unfortunately, the inspiratory radiation exposure and the reconstruction kernel used were not identical in the clinical and screening subjects. Despite the differences, the selection of clinical subjects was reasonably the best achievable option to investigate the total spectrum of COPD. The use of two different reconstruction filters and a difference in inspiratory radiation dose may have had some influence on the quantitative CT measurements [35–38]. However, it has also been suggested that dose-reduction to as low as 20mAs can be applied without clinical importance [39–41]. Moreover, after analyzing only the screening trial participants, examined with a single CT system and protocol, the pattern of univariate- and multivariate regression analysis remained the same as in the total population, though was less strong after exclusion of the 74 more severely affected cases. These findings support the validity of our conclusions.

Our study has some limitations. First, there is no good reference standard for air trapping, which is an in vivo phenomenon. This limitation makes it difficult to prove that the combined inspiratory-expiratory air trapping measurements are the best methods to quantify air trapping outside emphysematous spaces. Second, CT data acquisitions were not spirometrically controlled, which would enable a standardized inspiration and expiration level. Nevertheless, its absence strengthens the generalizability of our results since spirometer-gated CT is not widely applied. Despite these limitations, the combination of CT emphysema and CT air trapping explains a large part of the lung function variability. Third, we acknowledge that spirometry was obtained before bronchodilator administration, and that some asthma, whether or not coexisting with COPD, can be present. However, this likely concerns only a small amount of heavy smoking subjects. Last, we used a retrospective design and added a separate patient group due to the paucity of severely obstructed subjects in the screening group. However, this was the best achievable option to investigate the total spectrum of COPD. Moreover, although the explained variance decreased, the pattern found did not change after exclusion of the severely affected subjects.

In summary, quantitative CT measures of emphysema and air trapping are strongly associated with airflow limitation in a population that covers the total range of airflow limitation; the combination of CT emphysema and air trapping, as simple density measurements on paired inspiratory and expiratory CT images, explains around three-quarters of the lung function variability in such a population of COPD subjects. We suggest that these measures might gain a role in the identification of subjects with suspected lung function impairment, based on radiological data, and might prove useful in phenotyping COPD. This may be especially interesting in lung cancer.
screening, where large numbers of examinations are obtained in heavy smokers.

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Appendix 1

To score lung segmentation the observer was able to scroll through the entire scan with a semi-translucent colour overlay of the segmentation. This overlay could be removed and replaced at any time. The guidelines for scoring, as provided to the observers, was as follows:

**Excellent:** over 95% of the segmentation is correct

**Minor errors:** 80-95% of the segmentation is correct

**Major errors:** 50-80% of the segmentation is correct

**Totally wrong:** less than 50% of the segmentation is correct

Appendix 2

**Table 4** Results of univariate linear regression analysis for quantitative CT emphysema and air trapping measurements and pulmonary function tests in a subset of 174 subjects

|                         | logFEV₁ | FEV₁/FVC | RV/TLC | Kco  |
|-------------------------|---------|----------|--------|------|
| CT emphysema            |         |          |        |      |
| logIN₉₅₀                | 0.25    | 0.46     | 0.07   | 0.25 |
| Perc₁₅                  | 0.14    | 0.35     | 0.03    | 0.16 |
| CT air trapping         |         |          |        |      |
| EX₉₅₀                   | 0.41    | 0.54     | 0.24   | 0.15 |
| EX₉₅₀₋₉₁₀              | 0.36    | 0.44     | 0.24   | 0.11 |
| RVC₋₉₅₀₋₉₅₀           | 0.28    | 0.21     | 0.08   | 0.01 |
| E/I-ratioM₉₅₀          | 0.34    | 0.34     | 0.27   | 0.09 |

Data given are r-squared (R²) values

- Lung cancer screening trial participants only, controls to GOLD3; a 71 subjects with body box measurements; b 70 subjects with diffusion capacity measurements
- p<0.001; † p<0.01; ‡ p<0.05; NS not significant; logIN₉₅₀ = log-transformed emphysema score below -950HU in inspiratory scan; Perc₁₅ = emphysema score as 15th percentile of attenuation distribution curve in inspiratory scan; EXP₉₅₀ = air trapping score below -850HU in expiratory scan; EXP₉₅₀₋₉₁₀ = air trapping score between -850HU and -910HU in expiratory scan; RVC₋₉₅₀₋₉₅₀ = air trapping score as relative volume change between -860HU and -950HU; E/I-ratioM₉₅₀ = expiration to inspiration ratio of mean lung density

References

1. Global Initiative for Chronic Obstructive Pulmonary Disease. Global strategy for the diagnosis, management, and prevention of chronic obstructive pulmonary disease (2009) Available at http://www.goldcopd.com. Accessed March 2011
2. Pauwels RA, Buist AS, Calverley PM, Jenkins CR, Hurd SS (2001) Global strategy for the diagnosis, management, and prevention of chronic obstructive pulmonary disease. NHLBI/WHO Global Initiative for Chronic Obstructive Lung Disease (GOLD) Workshop summary. Am J Respir Crit Care Med 163:1256–1276
3. Hogg JC (2004) Pathophysiology of airflow limitation in chronic obstructive pulmonary disease. Lancet 364:709–721
4. Muller NL, Staples CA, Miller RR, Abboud RT (1988) “Density mask”. An objective method to quantitate emphysema using computed tomography. Chest 94:782–787
5. Nakano Y, Wong JC, de Jong PA, Buzatu L, Nagao T, Coxson HO, Elliott WM, Hogg JC, Pare PD (2005) The prediction of small airway dimensions using computed tomography. Am J Respir Crit Care Med 171:142–146
6. Matsuoka S, Kurihara Y, Yagihashi K, Hoshino M, Watanabe N, Nakajima Y (2008) Quantitative assessment of air trapping in chronic obstructive pulmonary disease using inspiratory and expiratory volumetric MDCT. AJR Am J Roentgenol 190:762–769
7. Reilly JJ (2008) COPD and declining FEV₁–time to divide and conquer? N Engl J Med 359:1616–1618
8. Rennard SI, Vestbo J (2008) The many "small COPDs": COPD should be an orphan disease. Chest 134:623–627
9. Han MK, Agusti A, Calverley PM, Celli BR, Criner G, Curtis JL, Fabbri LM, Goldin JG, Jones PW, MacNee W, Make B, Rabke KF, Rennard SI, Sciruba FC, Silverman EK, Vestbo J, Washko GR, Wouters EF, Martinez FJ (2010) Chronic obstructive pulmonary disease phenotypes: the future of COPD. Am J Respir Crit Care Med 182:598–604
10. Agusti A, Calverley PM, Celli B, Coxson HO, Edwards LD, Lomas DA, MacNee W, Miller BE, Rennard S, Silverman EK, Tal-Singer R, Wouters E, Yates JC, Vestbo J (2010) Characterisation of COPD heterogeneity in the ECLIPSE cohort. Respir Res 11:122
11. Hoop BJ, Mohamed Hoesfin F, Zanen P, Gietema HA, van Ginneken B, Isgum I, Mol C, van Klaveren R, Dijkstra A, Boezen H, Groen H, Postma D, Lammers JW, Prokop M (2011) CT-quantified emphysema in heavy smokers: predictive value for rate of lung function decline. Thorax. doi:10.1136/thx.2010.145995
12. van Iersel CA, de Koning HJ, Draisma G, Mali WP, Scholten ET, Ginneken B, Isgum I, Mol C, van Klaveren R, Dijkstra A, Boezen H, Groen H, Postma D, Lammers JW, Prokop M (2011) CT-quantified emphysema in heavy smokers: predictive value for rate of lung function decline. Thorax. doi:10.1136/thx.2010.145995
13. Quanjer PH, Tammeling GJ, Cotes JE, Pedersen OF, Peslin R, Yernault JC (1993) Lung volumes and forced ventilatory flows. Report Working Party Standardization of Lung Function Tests, European Community for Steel and Coal. Official Statement of the European Respiratory Society. Eur Respir J Suppl 16:5–76
14. Pauwels RA, Buist AS, Calverley PM, Jenkins CR, Hurd SS (2001) Global strategy for the diagnosis, management, and prevention of chronic obstructive pulmonary disease. NHLBI/WHO Global Initiative for Chronic Obstructive Lung Disease (GOLD) Workshop summary. Am J Respir Crit Care Med 163:1256–1276
15. van Rikxoort EM, de Hoop B, Vlieger MA, Prokop M, van Ginneken B (2009) Automatic lung segmentation from thoracic computed tomography scans using a hybrid approach with error detection. Med Phys 36:2934–2947
16. Schilham AM, van Ginneken B, Gietema H, Prokop M (2006) Local noise weighted filtering for emphysema scoring of low-dose CT images. IEEE Trans Med Imaging 25:451–463
17. Gevenois PA, De Maertelaer V, De Vuyst P, Zanen J, Yernault JC (1995) Comparison of computed density and macroscopic morphometry in pulmonary emphysema. Am J Respir Crit Care Med 152:653–657
18. Gevenois PA, De Vuyst P, De Maertelaer V, Zanen J, Jacobovitz D, Cosio MG, Yernault JC (1996) Comparison of computed density and macroscopic morphometry in pulmonary emphysema. Am J Respir Crit Care Med 154:187–192
19. Newell JD Jr, Hogg JC, Snider GL (2004) Report of a workshop: quantitative computed tomography scanning in longitudinal studies of emphysema. Eur Respir J 23:769–775
20. van Ginneken B, Murphy K, van Rikxoort EM, Isgum I, de Hoop B, Busacker A, Newell JD Jr, Keefe T, Hoffman EA, Granroth JC, Gevenois PA, De Vuyst P, De Maertelaer V, Zanen J, Yernault JC, Schilham AM, van Ginneken B, Gietema H, Prokop M (2006) Pulmonary emphysema: radiation dose and section thickness at multidetector CT for quantification of emphysema. AJR Am J Roentgenol 187:1281–1286
21. Boedeker KL, Nitt-Gray MF, Rogers SR, Truong DA, Brown MS, Gjertson DW, Goldin JG (2004) Emphysema: effect of reconstruction algorithm on CT imaging measures. Radiology 232:295–301
22. Yuan R, Mayo JR, Hogg JC, Pare PD, McWilliams AM, Lam S, Coxson HO (2007) The effects of radiation dose and CT manufacturer on measurements of lung densitometry. Chest 132:617–623
23. Madani A, De Maertelaer V, Zanen J, Gevenois PA (2007) Pulmonary emphysema: radiation dose and section thickness at multidetector CT quantification—comparison with macroscopic and microscopic morphometry. Radiology 243:250–257
24. Gierada DS, Pilgram TK, Whiting BR, Hogg JC, Pare PD, McWilliams AM, Lam S, Coxson HO (2007) The effects of radiation dose and CT manufacturer on measurements of lung densitometry. Chest 132:617–623
25. Madani A, De Maertelaer V, Zanen J, Gevenois PA (2007) Pulmonary emphysema: radiation dose and section thickness at multidetector CT quantification—comparison with macroscopic and microscopic morphometry. Radiology 243:250–257
26. Gierada DS, Pilgram TK, Whiting BR, Hogg JC, Bierhals AJ, Kim JH, Bae KT (2007) Comparison of standard- and low-radiation-dose CT for quantification of emphysema. AJR Am J Roentgenol 188:42–47
27. Zaporozhan J, Ley S, Weinheimer O, Eberhardt R, Tsakiris I, Noshi Y, Herth F, Kauzcor Hu (2006) Multi-detector CT of the chest: influence of dose onto quantitative evaluation of severe emphysema: a simulation study. J Comput Assist Tomogr 30:460–468