

ORIGINAL RESEARCH

Quantitative CT Analysis in Patients with Pulmonary Emphysema: Do Calculated Differences Between Full Inspiration and Expiration Correlate with Lung Function?

This article was published in the following Dove Press journal: International Journal of Chronic Obstructive Pulmonary Disease

Lan Song^{1,*} Jonas A Leppig^{2,*} Ralf H Hubner³ Bianca C Lassen-Schmidt⁴ Konrad Neumann 1005 Dorothea C Theilig² Felix W Feldhaus 1002 Ute L Fahlenkamp² Bernd Hamm² Wei Song Zhengyu Jin (b) Felix Doellinger 102

¹Department of Radiology, Peking Union Medical College Hospital, Chinese Academy of Medical Sciences and Peking Union Medical College, Beijing, People's Republic of China; ²Department of Radiology, Charité Universitätsmedizin Berlin, Berlin, Germany; ³Department of Internal Medicine/Infectious Diseases and Respiratory Medicine, Charité Universitätsmedizin Berlin, Berlin, Germany; ⁴Fraunhofer Institute for Digital Medicine MEVIS, Bremen, Germany; 5 Institute of Biometrics and Clinical Epidemiology, Charité Universitätsmedizin Berlin, Berlin, Germany

*These authors contributed equally to this work

Correspondence: Zhengyu Jin Department of Radiology, Peking Union Medical College Hospital, Chinese Academy of Medical Sciences and Peking Union Medical College, Shuaifuyuan No. I, Dongcheng District, Beijing, People's Republic of China Tel +86 10 69 155 439 Fax +86 10 69 155 058 Email jinzy@pumch.cn

Purpose: The aim of this retrospective study was to evaluate correlations between parameters of quantitative computed tomography (QCT) analysis, especially the 15th percentile of lung attenuation (P15), and parameters of clinical tests in a large group of patients with pulmonary emphysema.

Patients and Methods: One hundred and seventy-two patients with pulmonary emphysema and chronic obstructive pulmonary disease (COPD) global initiative for chronic obstructive lung disease (GOLD) stage 3 or 4 were assessed by nonenhanced thin-section CT scans in full inspiratory and expiratory breath-hold, pulmonary function test (PFT), a 6-minute walk test (6MWT), and quality of life questionnaires (SGRQ and CAT). QCT parameters included total lung volume (TLV), total emphysema score (TES), and P15, all measured at inspiration (IN) and expiration (EX). Differences between inspiration and expiration were calculated for TLV (TLV_{Diff}), TES (TES_{Diff}), and P15 (P15_{Diff}). Spearman correlation analysis was performed.

Results: CT-measured lung volume in inspiration (TLV_{IN}) correlated strongly with spirometry-measured total lung capacity (TLC) (r=0.81, p<0.001) and moderately to strongly with residual volume (RV), forced vital capacity (FVC), and forced expiratory volume in 1 second (FEV₁)/FVC (r=0.60, 0.56, and -0.49, each p<0.001). Lung volume in expiration (TLV_{EX}) correlated moderately to strongly with TLC, RV and FEV₁/FVC ratio (r=0.75, 0.66, and -0.43, each p < 0.001). TES and P15 showed stronger correlations with the carbon monoxide transfer coefficient (KCO%) (r= -0.42, 0.44, both p<0.001), when measured during expiration. P15_{Diff} correlated moderately with KCO% and carbon monoxide diffusing capacity (DLCO%) (r= 0.41, 0.40, both p<0.001). The 6MWT and most QCT parameters showed significant differences between COPD GOLD 3 and 4 groups.

Conclusion: Our results suggest that QCT can help predict the severity of lung function decrease in patients with pulmonary emphysema and COPD GOLD 3 or 4. Some QCT parameters, including $P15_{EX}$ and $P15_{Diff}$, correlated moderately to strongly with parameters of pulmonary function tests.

Keywords: chronic obstructive pulmonary disease, pulmonary emphysema, quantitative CT, pulmonary function test

Introduction

Chronic obstructive pulmonary disease (COPD) is the third leading cause of mortality in industrialized countries.¹ Pulmonary emphysema is characterized by irreversible hyperinflation of lung parenchyma and is a key contributor to impaired

Song et al Dovepress

lung function in COPD. Patients with pulmonary emphysema have a higher lung volume than healthy people, whereas the difference in volume between inspiration and expiration is smaller than in healthy nonsmokers.^{2,3} The pulmonary function test (PFT) is the gold standard for diagnosing COPD, typically showing a reduced forced expiratory volume in 1 second (FEV₁), an increased total lung capacity (TLC), and a decreased FEV₁/forced vital capacity (FVC) ratio. However, the PFT results depend on how well a patient can cooperate, and the number of unacceptable and nonrepeatable maneuvers is relatively higher among older subjects and patients in poor health.⁴ The latest COPD guidelines emphasize the feasibility and necessity of various methods for evaluating COPD. While PFT continues to be the mainstay of diagnosis, quantitative computed tomography (QCT) analysis has gained importance in the evaluation of COPD and is increasingly used to identify and classify emphysema. 5-8 Emphysematous lung destruction results in the replacement of normal lung tissue, which has a typical attenuation of approximately -850 Hounsfield units (HU) on inspiratory CT scans in healthy humans. Hyperinflation reduces lung attenuation towards -1000 HU. The normal increase in lung density from inspiration to expiration is reduced in COPD due to air trapping/hyperinflation. 5 CT attenuation values can quantify the extent of emphysema. Low attenuation volumes (LAVs) are considered to represent emphysematous lung parenchyma and reportedly correlate with the degree of lung function impairment.⁶

The diagnostic value of QCT for the objective evaluation of pulmonary emphysema has been described for total lung volume (TLV), the extent of emphysema (thresholdbased identification of LAV), total emphysema score (TES, i.e. the emphysema index), and mean lung density (MLD). 6-10 MLD is the most basic parameter for estimating emphysema extent. Previous studies have shown that the MLD derived from expiratory CT scans is a good predictor of pulmonary ventilation. Alternatively, indices of lung attenuation at a given percentile along the HU histogram, such as the first percentile (P1) or 15th percentile (P15), can be used for the evaluation of emphysema.⁸ As the emphysema extent increases, the density histogram shifts towards lower HU values, and the P index decreases. There is some evidence that the percentile approach is more robust for longitudinal monitoring of emphysema and less susceptible to changes in lung volume or concomitant high-attenuation disease (e.g. fibrosis) than MLD.¹¹ Several percentile indices derived from inspiratory scans were reported to significantly correlate with microscopic and macroscopic emphysema extent. Most long-term studies have used P15, as it is less severely degraded by image noise and truncation effects than P1. Net we expect the percentage difference of P15 values measured at inspiration and expiration (P15_{Diff}) to be a useful parameter for evaluating pulmonary emphysema severity. P15_{Diff} might be affected less by patient size than absolute values acquired in only one respiratory phase and it should be able to reflect dynamic changes between full inspiration and expiration. However, P15_{Diff} has not yet been investigated in COPD patients.

We conducted a systematic retrospective analysis to correlate QCT parameters measured at inspiration and expiration and the differences between inspiration and expiration with clinical test results in a large number of patients with severe COPD. CT-based inspiratory and expiratory TLV, TES, and P15 and the calculated parameters of the corresponding inspiratory/expiratory differences were compared with PFT results and scores of the 6-minute walk test (6MWT), St. George's Respiratory Questionnaire (SGRQ), and COPD assessment test (CAT).

Patients and Methods

This study was approved by the local ethics committee/institutional review board (IRB; Charité Ethikkommission; Entscheid EA1/213/16). Patient consent to anonymous and retrospective review of their medical records was not required by IRB for this retrospective study. The confidential and anonymous evaluation of the patient data was guaranteed. The study was conducted in accordance with the Declaration of Helsinki.

Patients

In this retrospective study, we analyzed 172 pulmonary emphysema patients that were clinically diagnosed with severe COPD, Global Initiative for Chronic Obstructive Lung Disease (GOLD) stage 3 or 4. All patients were evaluated for surgical or bronchoscopic lung volume reduction and underwent noncontrast multislice CT (MSCT) scanning and a PFT within a span of 30 days in our department from May 2014 to May 2019 (Table 1). The inclusion criteria for patients with severe COPD were FEV₁/FVC ratio <70% and FEV₁ < 50% of the predicted value after bronchodilator inhalation. COPD was graded according to the 2019 GOLD guidelines: GOLD 3 (severe): $30\% \le \text{FEV}_1 < 50\%$ of the predicted value, GOLD 4 (very severe): FEV₁ < 30% of the predicted value, both based on postbronchodilator use. ¹ The

Table I Demographics and Quantitative CT Parameters, Pulmonary Function Test Data, and 6-Minute Walk Test and Clinical Scores in 172 Patients with COPD GOLD 3 or 4

Parameters	Overall	GOLD 3	GOLD 4	p-value
Number (%)	172 (100%)	71 (41.3%)	101 (58.7%)	_
Age (years)	65.9±6.7 (47~80)	67.1±6.9 (50~80)	65.0±6.4 (47~80)	0.054
Sex (male/female)	99 (57.6%)/73 (42.4%)	35 (49.3%)/36 (50.7%)	64 (63.4%)/37 (36.6%)	0.067
Quantitative CT measurements				
TLV _{IN} (mL)	6751±1383 (3875~10,710)	6566±1364 (3875~10,710)	6880±1388 (4043~10,305)	0.144
TLV _{EX} (mL)	5631±1227 (2529~9465)	5316±1156 (2529~8716)	5853±1232 (3543~9465)	0.004
TLV _{Diff} (%)	16.3±9.0 (0.3~39.7)	18.7±9.5 (0.3~39.7)	14.6±8.3 (0.3~37.1)	0.003
TES _{IN} (%)	28.0±9.5 (6.5~51.6)	25.8±9.6 (6.5~48.4)	29.5±9.2 (7.2~51.6)	0.013
TES _{EX} (%)	21.6±9.3 (2.0~45.9)	19.1±9.4 (2.0~44.7)	23.3±9.2 (3.5~45.9)	0.004
TES _{Diff} (%)	24.7±16.4 (-2.7~82.3)	28.3±17.7 (-1.3~82.3)	22.1±15.1 (-2.7~67.3)	0.015
PI5 _{IN} (HU)	-969±14 (-1003~ -932)	-967±15 (-1003~ -932)	-971±13 (-993~ -935)	0.057
PI5 _{EX} (HU)	-960±18 (-997~ -889)	-956±21 (-997~ -889)	-964±16 (-991~ -921)	0.009
PI5 _{Diff} (%)	0.9±0.8(-0.3~5.7)	1.1±1.0 (0.1~5.7)	0.8±0.6 (-0.3~3.2)	0.006
Data of clinical pulmonary function tests				
FEV ₁ (mL)	767±213 (260~1630)	975±227 (520~1630)	640±190 (260~1030)	<0.001
FEV ₁ % (% of predicted)	27.9±4.6 (12~49)	36.3±4.7 (30~49)	23.2±4.5 (12~29)	<0.001
FVC (mL)	2188±713 (890~4390)	2663±711 (1430~4240)	2144±719 (890~4390)	<0.001
FVC% (% of predicted)	62.1±13.7 (33~120)	75.1±13.4 (53~120)	59.5±14.3 (33~102)	<0.001
TLC (mL)	7361±1374 (4490~11,880)	7337±1523 (4960~11,880)	7378±1293 (4490~11,130)	0.888
TLC% (% of predicted)	121.7±14.7 (98~159)	121.4±12.5 (98~152)	121.8±15.9 (99~159)	0.609
RV (mL)	5019±1136 (2120~8300)	4708±1129 (2120~8130)	5194±1141 (2290~8300)	0.009
RV% (% of predicted)	217.6±42.9 (104~331)	201.2±36.6 (104~295)	226.9±43.7 (128~331)	<0.001
FEV ₁ /FVC (%)	12.7±3.9 (6~25)	14.6±4.3 (8~25)	11.6±3.3 (6~22)	0.001
RV/TLC (%)	67.8±8.0 (43~85)	63.2±7.5 (43~75)	70.4±8.8 (43~85)	<0.001
KCO% (% of predicted, n=95)	34.5±20.3 (25~45.3)	39±18 (33~51)	30±21 (23~44)	0.093
DLCO% (% of predicted, n=110)	28.5±14.3 (21.8~36)	33±14 (29~43)	25±12 (17~29)	0.001
Exercise capacity test and scores of				
quality of life questionnaires				
6MWT (m, n=103)	243.8±107.5 (0~570)	280.8±114.1 (0~570)	219.3±96.3 (0~535)	0.004
SGRQ (score, n=74)	64.6±12.8 (38~98)	61.3±14.1 (38~88)	66.5±11.8 (42~98)	0.093
CAT (score, n=116)	23.6±6.2 (7~40)	22.5±6.2 (10~36)	24.3±6.1 (7~40)	0.126

Notes: All PFT parameters recorded here were obtained without short-acting bronchodilators. KCO% and DLCO% are presented as median ± interquartile interval (interquartile range). Other values are presented as n (%) or mean ± SD (range).

Abbreviations: COPD, chronic obstructive pulmonary disease; GOLD, global initiative for chronic obstructive lung disease; TLV, total lung volume; TES, total emphysema score; P15, 15th percentile of lung attenuation distribution curve; IN, inspiration; EX, expiration; Diff, percentage difference between inspiration and expiration; FEV₁, forced expiratory volume in I second; FVC, forced vital capacity; TLC, total lung capacity; RV, residual volume; FEV₁/FVC, ratio of FEV₁ to FVC; RV/TLC, ratio of RV to TLC; DLCO, carbon monoxide diffusing capacity; KCO, carbon monoxide transfer coefficient; 6MWT, 6-minute walk test; SGRQ, St. George's Respiratory Questionnaire; CAT, COPD assessment test.

exclusion criteria (based on a retrospective, blinded review of chest CT datasets) were active lung diseases such as acute exacerbation of COPD, pneumothorax, pleural effusion, extensive ground-glass opacities, suspicious nodules, lung tumors, relevant atelectasis, consolidation, major scars, thoracic deformity, postoperative lung, and poor respiratory coordination. All subjects had a long-term history of cigarette smoking with a smoking index of more than 20 pack-years. The results of the 6MWT, SGRQ, and CAT were retrieved from medical records.

Quantitative Computed Tomography

Patients were randomly assigned to undergo inspiratory and expiratory nonenhanced CT for emphysema evaluation on one of two CT scanners (Light Speed Ultra 8 or Revolution EVO, both General Electric Healthcare, Chicago, IL, USA) with the same protocol (supine position, two scans of the whole lung in full inspiratory and expiratory breath-hold) including standardized scanning parameters (0.625 or 1.25 mm primary slice thickness, slice interval the same as slice thickness, 120 kV voltage, 100 mA tube current,

Song et al Dovepress

standard (soft) kernel). All patients were thoroughly coached in breathing techniques prior to the CT scan. Quantitative analysis of chest CT scans was performed using the MeVis PULMO3D software (v3.7.1, Fraunhofer MEVIS, Bremen, Germany) to obtain TLV, TES, and P15 (i.e. the 15th percentile of the lung attenuation distribution curve) using the density mask technique. TES was defined as the percentage of voxels with density values below the threshold of –950 HU out of the total number of voxels in the whole lung as described in many previous studies. The full datasets of axial slices without interscan gaps were used for analysis.

Differences in the QCT parameters between inspiration and expiration were calculated as follows:

$$TLV_{Diff} = (TLV_{IN} - TLV_{EX})/TLV_{IN}$$

$$TES_{Diff} = (TES_{IN} - TES_{EX}) / TES_{IN}$$

$$P15_{Diff} = (P15_{IN} - P15_{EX})/P15_{IN}$$

Examples of QCT analysis of the total lung during full inspiratory and full expiratory breath-hold are presented in Figures 1 and 2.

Pulmonary Function Test

Dynamic spirometry (including FEV₁ and FVC) and static lung volumes (including TLC and residual volume (RV)) were measured according to American Thoracic Society/ European Respiratory Society standards. 17,18 All spirometric maneuvers were performed before and 15 min after inhalation of 400 mg of salbutamol. A Ganshorn PowerCube (Ganshorn Medizin Electronic GmbH, Niederlauer, Germany) was used for all measurements. Diffusing capacities, including carbon monoxide diffusing capacity (DLCO) and carbon monoxide transfer coefficient (KCO), were measured according to single-breath carbon monoxide uptake. All lung function parameters except FEV₁/FVC ratio and RV/TLC ratio are expressed as absolute values and as the percentage of the predicted value. FEV₁/FVC was calculated as the percentage of FEV₁ to FVC, and RV/TLC was calculated as the percentage of RV to TLC. Diffusing capacity parameters DLCO% and KCO% were calculated as the percentage of the predicted value. The PFT results are summarized in Table 1.

Statistics

Continuous variables with nonnormal distribution, such as KCO% and DLCO%, are expressed as median \pm interquartile interval (interquartile range). Continuous variables with normal distribution are presented as mean \pm SD (range).

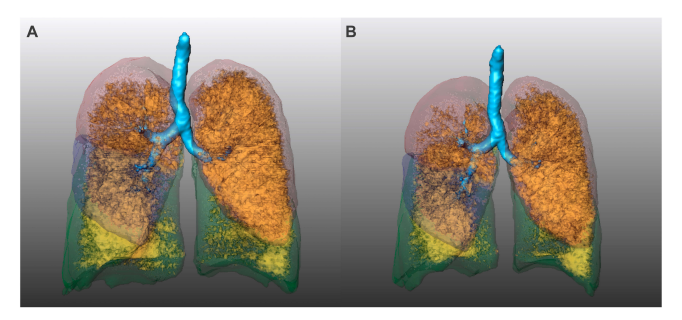


Figure 1 Coronal 3D surface views of the lungs of a 79-year-old female smoker with COPD GOLD 3 (**A**, **B**). These surface views were generated with the MeVis PULMO3D software by processing CT scans acquired at full inspiration (**A**) and full expiration (**B**). Lung lobes are depicted semitransparent green (lower lobes), red (upper lobes) and blue (right middle lobe). Emphysematous lung parenchyma, demonstrated as a LAV (i.e., voxels with a density ≤ −950 HU), is depicted in orange and yellow. In this patient, LAV in the right upper lobe significantly decreased in expiration, while LAV in the left upper lobe did not show a significant change between the two respiratory phases.

Abbreviations: 3D, three dimensional; COPD, chronic obstructive pulmonary disease; GOLD, global initiative for chronic obstructive lung disease; LAV, low attenuation volumes.

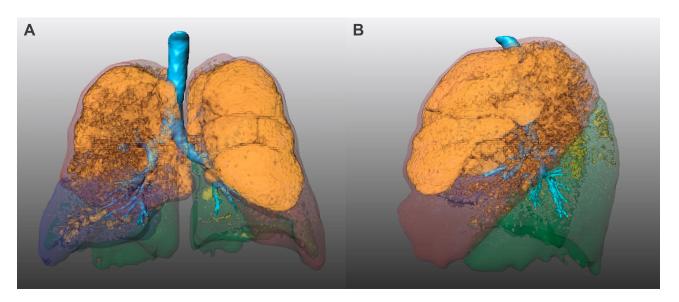


Figure 2 3D surface views of the lung of a 59-year-old male smoker with COPD GOLD 3 (**A**, **B**). Coronal (**A**) and sagittal (**B**) reconstructions of a CT scan acquired at full inspiration demonstrates a predominant emphysema distribution in the upper lobe. **Abbreviations:** 3D, three dimensional; COPD, chronic obstructive pulmonary disease; GOLD, global initiative for chronic obstructive lung disease.

Categorical variables are expressed as frequencies (percentages). For the comparison of QCT parameters and clinical parameters, Spearman's rank correlation analysis was used. Differences between COPD GOLD 3 and GOLD 4 were assessed using the independent samples *t*-test or Mann–Whitney *U*-test for continuous variables as appropriate, and the chi-square test was used for categorical variables (sex). A two-tailed *p*-value <0.05 was considered statistically significant. The SPSS Statistics 26 software package (IBM Corporation, Armonk, NY, USA) was used for all statistical analyses. In this observational study, no adjustment for multiple testing was performed. Hence, all *p*-values are exploratory.

Results

Whole-Lung Analysis and Comparison Based on GOLD Stages

The demographics, QCT data, PFT data, and scores of the 6MWT, SGRQ and CAT of 172 severe COPD patients are shown in Table 1. No significant differences were found for age, sex, TLV_{IN} , $P15_{IN}$, TLC, TLC%, KCO%, SGRQ and CAT scores between GOLD 3 and GOLD 4. All other QCT and PFT parameters and 6MWT showed significant differences between both GOLD stages.

Correlation Analysis of Different Measurement Methods

The correlation analyses for all tested QCT and PFT parameters are shown in Table 2 and Figure 3. TLV_{IN} showed

moderate to very strong correlations with TLC, RV, FVC and FEV₁/FVC (r=0.81, 0.60, 0.56, and -0.49; each p<0.001). TLV_{EX} showed a moderate to strong correlation with TLC, RV and FEV₁/FVC (r=0.75, 0.66, and -0.43; each p<0.001). TLV_{Diff} showed a moderate, negative correlation with RV/TLC (r=-0.43; p<0.001). TES_{EX} showed a moderate, negative correlation with KCO% (r=-0.42; p<0.001). It is worth noting that both P15_{EX} and P15_{Diff} showed moderate correlations with KCO% (r=0.44 and 0.41; both p<0.001), and P15_{Diff} showed a moderate correlation with DLCO% (r=0.40; p<0.001). There were only weak but measurable correlations in some other pairings. The strongest correlation was found for CT-measured TLV_{IN} and PFT-measured TLC, but paired t-test showed that the mean TLC (7361 mL prebronchodilator) was significantly higher than that of TLV_{IN} (6751 mL), with a mean difference of 610 mL in all 172 patients (p<0.001).

The correlations of the QCT parameters and the 6MWT, SGRQ, and CAT scores are shown in Table 3. The SGRQ and 6MWT scores only weakly correlated with TLV_{Diff}, TES_{EX}, TES_{Diff}, P15_{EX} and P15_{Diff} (SGRQ: r=-0.27, 0.32, -0.30, -0.29, and -0.28; 6MWT: r=0.27, -0.21, 0.29, 0.23, and 0.32). The CAT score showed no significant correlations with any of the QCT parameters, although it was strongly correlated with the SGRQ score (r=0.65; p<0.001). The 6MWT showed a moderate, negative correlation with the SGRQ score (r=-0.47; p<0.001) and a weak, negative correlation with the CAT score (r=-0.23; p<0.05).

Song et al Dovepress

Table 2 Spearman's Rank Correlation Coefficients Describing the Statistical Correlations Between QCT Parameters and the Results of Pulmonary Function Tests in the Evaluation of 172 Patients with COPD GOLD 3 or 4

	FEV ₁ (mL)	FEV _I % (% Pred)	TLC (mL)	TLC% (% Pred)	RV (mL)	RV (% Pred)	FVC (mL)	FVC% (% Pred)	FEV _I / FVC (%)	RV/TLC (%)	KCO% (% Pred)	DLCO% (% Pred)
TLV _{IN} (mL)	0.36***	-0.10	0.81***	0.20**	0.60***	0.30***	0.56***	0.04	-0.49***	-0.14	0.12	0.10
TLV _{EX} (mL)	0.17**	-0.24**	0.75***	0.22**	0.66***	0.37***	0.37***	-0.09	-0.43***	0.08	-0.0I	-0.04
TLV _{Diff} (%)	0.34***	0.32***	-0.05	-0.12	-0.23**	-0.20**	0.29***	0.25**	-0.04	-0.43***	0.22*	0.25**
TES _{IN} (%)	-0.24**	-0.26***	0.22*	0.36***	0.22**	0.27***	0.05	0.07	-0.21**	0.10	-0.36***	-0.26**
TES _{EX} (%)	-0.27**	-0.31***	0.24**	0.34***	0.26**	0.29***	-0.01	0.01	-0.18**	0.19*	-0.42***	-0.34***
TES _{Diff} (%)	0.24**	0.29***	-0.21*	-0.23**	-0.28**	-0.21**	0.13	0.11	0.02	-0.28**	0.37***	0.36***
PI5 _{IN} (HU)	0.20**	0.23**	-0.23*	-0.35***	-0.20**	-0.24**	-0.11	-0.12	0.23**	-0.06	0.36***	0.27**
PI5 _{EX} (HU)	0.28***	0.33***	-0.26**	-0.35***	-0.27**	-0.29***	-0.01	-0.04	0.18*	-0.19*	0.44***	0.37***
PI5 _{Diff} (%)	0.31***	0.37***	-0.22*	-0.23**	-0.31**	-0.25**	0.18*	0.15	0.00	-0.34***	0.41***	0.40***

Notes: At least moderate positive or negative statistical correlations with coefficients $r \ge 0.40$ or ≤ -0.40 are highlighted in **bold** ($r=0\sim0.19$, no or very weak correlation; $r=0.20\sim0.39$, weak correlation; $r=0.40\sim0.59$, moderate correlation; $r=0.60\sim0.79$, strong correlation; $r=0.80\sim1$, very strong correlation; r=0.0.05, **r=0.00.] Abbreviations: TLV, total lung volume; TES, total emphysema score; P15, 15th percentile of lung attenuation distribution curve; IN, inspiration; EX, expiration; Diff, percentage difference between inspiration and expiration; FEV₁, forced expiratory volume in 1 second; TLC, total lung capacity; RV, residual volume; FVC, forced vital capacity; DLCO, carbon monoxide diffusing capacity; KCO, carbon monoxide transfer coefficient.

None of the calculated differences between the CT measurements in inspiration and expiration showed strong correlations with the clinical tests, including the questionnaires.

Discussion

In this study, we measured TLV, TES and P15 in inspiration and expiration and calculated TLV_{Diff}, TES_{Diff}, and P15_{Diff} from paired inspiratory/expiratory chest CT scans for two reasons: PFTs also provide dynamic breathing information, and normal lung volumes vary with patient size.

Comparison of Our results and Comparable Studies

A LAV threshold of -950 HU, as used in our study, is most commonly used for CT-based emphysema quantification, yielding a good balance between sensitivity and specificity.^{8,14,16} As expected, our patients with severe COPD (FEV₁% <50%) had a much higher mean TES_{IN} of 28.0% (SD 9.5) than the reported range for healthy individuals aged 30–70 years (2.6–4.5%). 19 The mean P15_{IN} in GOLD 3 and GOLD 4 was -967HU and -971HU, respectively, in our study, compared with a mean of -919 HU in young men with normal lung function.²⁰ In 36 severe COPD patients, the means of TLV_{Diff} , $P15_{IN}$ and $P15_{EX}$ were similar to ours: 0.19±0.10 vs. 0.16±0.09, -961±12 HU vs. -969±14 HU, and -951±21 HU vs. -960±18 HU.²¹ Heussel et al found median TLV_{IN}, TES_{IN} and P15_{IN} of 7200mL, 45%, and -985 HU, respectively, in 102 COPD patients GOLD stages 3 and 4, versus the corresponding mean values of 6751 mL, 28% and -969 HU in 172 COPD patients GOLD stages 3 and 4 in our study. This discrepancy may be due to differences in data presentation and different emphysema extents in our study population from those in the study from Heussel et al. In another study, TLV_{IN} was on average 670 mL smaller than PFT-derived TLC, consistent with our result of 610 mL.²² There are two potential reasons for this discrepancy. First, QCT-measured TLV_{IN} does not include upper and lower airway gas volumes. Second, the clinical PFT is examined in the sitting position, while the CT scans needed for QCT evaluation are acquired in the supine position. The inspiratory capacity may be affected by the different positions. However, a very strong correlation between TLV_{IN} and TLC (r=0.81; p<0.001) was shown in our study, comparable to the previous result (r=0.90).

Many publications on the correlation between other QCT and PFT parameters are inconsistent. Our study found the strongest correlations for TLC and TLV_{IN} (r=0.81; p<0.001), RV and TLV_{EX} (r=0.66; p<0.001), and RV/TLC and TLV_{Diff} (r=-0.43; p<0.001). TLV_{IN} had moderate to strong correlations with RV, FVC and FEV_I/FVC, and TLV_{EX} had a strong correlation with TLC. Functionally, hyperinflation of the entire lung is characterized by high RV and low FVC. Therefore, we used CT scans in full inspiration and full expiration to measure vital capacity (i.e. TLV_{IN}-TLV_{EX}) and total lung capacity (i.e. TLV_{IN}). The lung with the largest amount of air trapping was considered to be the lung with the lowest VC/TLC ratio (i.e. TLV_{Diff}). Therefore, TLV_{Diff} indicated lung collapse.

Frequently, a negative correlation of TES with FEV_1 or FEV_1/FVC is expected.^{7,9} Our study also shows weak, negative correlations between FEV_1 , $FEV_1\%$, FEV_1/FVC , and TES in both inspiration and expiration. Schroeder et al reported that TES_{IN} (r=-0.67, -0.76),

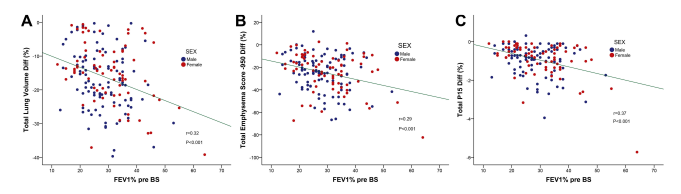


Figure 3 Correlation between FEV₁% pre-BS and parameters of quantitative CT analysis in 172 patients with COPD GOLD 3 or 4 (**A–C**). These three scattered plot diagrams show only weak statistical correlations between FEV₁% pre-BS and TLV_{Diff} (**A**, r=0.32, p<0.001), TES_{Diff} (**B**, r=0.29, p<0.001) and P15_{Diff} (**C**, r=0.37, p<0.001). **Abbreviations:** FEV₁%, percentage predicted of forced expiratory volume in 1 second; pre-BS, before application of short-acting bronchodilators; TLV_{Diff}, percentage difference of total lung volume between inspiration and expiration; TES_{Diff}, percentage difference of total emphysema score between inspiration and expiration; P15_{Diff}, percentage difference of the 15th percentile of lung attenuation distribution curve between inspiration and expiration.

 $P15_{IN}$ (r=0.55, 0.69) and $P15_{EX}$ (r=0.71, 0.80) had moderate to strong correlations with FEV_1 and FEV_1/FVC in COPD patients. Consistent with this result, $P15_{IN}$ and $P15_{EX}$ also had significant positive correlations with FEV_1 and FEV_1/FVC in our study. Overall, most of the correlation coefficients between TLV, TES and P15 and the results of PFT and 6MWT in our study were similar to those of the previous study, while some correlations were weaker in our study. However, we found better correlations with KCO% and DLCO%. The correlation of

Table 3 Spearman's Rank Correlation Coefficients Describing Statistical Correlations Between QCT Parameters and the Results of Clinical Tests in the Evaluation of 172 Patients with COPD GOLD 3 or 4

	6MWT(m)	SGRQ (Score)	CAT (Score)			
TLV _{IN} (mL)	0.21*	-0.13	0.01			
TLV _{EX} (mL)	0.07	0.08	0.05			
TLV _{Diff} (%)	0.27**	-0.27*	-0.07			
TES _{IN} (%)	-0.14	0.27*	-0.08			
TES _{EX} (%)	-0.21*	0.32**	-0.06			
TES _{Diff} (%)	0.29**	-0.30*	0.01			
PI5 _{IN} (HU)	0.12	-0.27*	0.10			
PI5 _{EX} (HU)	0.23*	-0.29*	0.09			
PI5 _{Diff} (%)	0.32**	-0.28*	0.02			
SGRQ (score)	-0.47***	_	0.65***			
CAT (score)	-0.23*	0.65***	-			
I	I	1	I			

Notes: At least moderate positive or negative statistical correlations with coefficients $r \ge 0.40$ or ≤ -0.40 are highlighted in **bold** ($r=0\sim0.19$, no or very weak correlation; $r=0.20\sim0.39$, weak correlation; $r=0.40\sim0.59$, moderate correlation; $r=0.60\sim0.79$, strong correlation; $r=0.80\sim1$, very strong correlation; *p<0.05, **p<0.01, ***p<0.001).

Abbreviations: TLV, total lung volume; TES, total emphysema score; P15, 15th percentile of lung attenuation distribution curve; IN, inspiration; EX, expiration; Diff, percentage difference between inspiration and expiration; 6MWT, 6-minute walk test; SGRQ, St. George's Respiratory Questionnaire; CAT, COPD assessment test.

FEV₁% with TLV, TES and P15 was not strong in our study. This might be explained by the fact that TLV, TES, and P15 are morphological measures mainly influenced by parenchymal destruction, while FEV₁ is a bronchial functional parameter. Emphysema extent is only partially predicted by the severity of changes revealed by PFTs.²³ In our study, TES correlated best with KCO% and TLC% in both inspiration (r=-0.36, 0.36) and expiration (r=-0.42, 0.34). Furthermore, we found that FEV₁/FVC correlated best with TLV_{IN} and TLV_{EX} (r=-0.49, -0.43; both p<0.001), KCO% correlated best with TES_{EX}, P15_{EX} and P15_{Diff} (r=-0.42, 0.44, 0.41; each p<0.001), and DLCO% correlated best with P15_{Diff} (r=0.40; p<0.001). TLV_{Diff}, TES_{Diff} and P15_{Diff} correlated better with RV/TLC than TLV_{IN} or TLV_{EX}, TES_{IN} or TES_{EX}, P15_{IN} or P15_{EX} do, suggesting that the calculated differences might be better indicators of air trapping than the corresponding absolute values measured in inspiration and expiration. In 30 COPD patients, Cavigli et al found stronger correlations of DLCO% with TES_{IN} and P15_{IN} (r=-0.63, 0.50, respectively) than we did (r=-0.26, 0.27, p<0.01).²⁴ Gelb et al found strong negative correlations between DLCO%, KCO% and TES only in patients with an absolute FEV₁ ≥ 1000 mL.²⁵ Our study population had a mean FEV₁ of 770 mL, which may explain why we found weaker correlations than Cavigli et al.²⁴ Akira et al reported that P15_{IN} and P15_{EX} to correlate well with FEV₁/FVC, RV/TLC and DLCO%, but not with FEV1%, and most of the corresponding correlations were stronger in patients with $\text{FEV}_1\% \ge 50\%$. They also found that TES_{EX} correlated better with FEV₁% and RV/TLC than with TES_{IN}, when FEV₁% was less than 50%, consistent with our result.²⁶ Our study shows that $P15_{\rm IN}$ and $P15_{\rm EX}$ correlated best

with KCO%, relatively well with DLCO% and TLC%, and weakly with FEV₁/FVC and FEV₁%. P15_{EX} and P15_{Diff} only had a weak negative correlation with RV/TLC. TES_{EX}. P15_{EX} and P15_{Diff} had a relatively good correlation with KCO% and DLCO% in our study. From our point of view, these results suggest that P15_{EX}, P15_{Diff} and TESEX are related to lung diffusion capacity.

Staging COPD Patients According to **QCT Parameters**

Except for TLV_{IN} and P15_{IN}, our study shows significant differences in lung volume (TLV_{EX}, TLV_{Diff}), emphysema index (TES_{IN}, TES_{EX} and TES_{Diff}), and 15th percentile of lung density (P15_{EX}, P15_{Diff}) between GOLD 3 and GOLD 4 patients. TLV_{EX}, TES_{IN} and TES_{EX} were significantly larger in GOLD 4 than in GOLD 3, while TLV_{Diff} , TES_{Diff}, P15_{Diff}, P15_{EX} and 6MWT were significantly smaller in GOLD 4 than in GOLD 3. There were no significant differences in TLV_{IN}, TLC or TLC% between both GOLD stages in our study. These results suggest that QCT, especially the parameters measured in expiration, can help to evaluate the severity of COPD in patients with GOLD 3 and 4. We also found better correlations with FEV₁, FEV₁%, TLC, RV, RV%, RV/TLC, KCO%, and DLCO% for TES_{EX} and P15_{EX} than for TES_{IN} and P15_{IN}. Zaporozhan et al reported no differences in TES_{IN}, TES_{EX}, TES_{Diff} and TLV_{Diff} between GOLD 3 and GOLD 4 among 31 patients, probably due to the small number of patients.³

Relationship Between QCT and Qualityof-Life Questionnaires

The GOLD guidelines group COPD patients by symptom and risk assessment. Quality of life assessments, such as the SGRQ and CAT, have been widely used to evaluate the condition of COPD patients and to predict their mortality risk.27,28 While the GOLD classification is an index reflecting the severity of COPD, CAT reflects the patients' overall quality of life, allowing patients to monitor their disease status and treatment effects easily.²⁸ Previous studies showed that the CAT score could reflect disease impact on exercise ability.²⁹ Our results show that the CAT score was weakly and negatively correlated with the 6MWT (r= -0.23; p<0.05), while the SGRQ score had a moderate to strong correlation with the 6MWT and CAT score (r= -0.47, 0.65; both p<0.001). Except for TLV_{IN} and TLV_{EX}, the QCT parameters only correlated weakly

with the SGRO score. No OCT parameter correlated with the CAT score, suggesting that the SGRQ may be superior in evaluating emphysematous lung destruction. The 6MWT and most QCT and PFT results were significantly different between COPD GOLD stage 3 and 4, while there was no significant difference in the SGRQ and CAT scores between the two groups. This finding might be related to the different ages and educational levels of the patients with respect to the recognition of acute exacerbations of COPD. QCT may assess the severity of COPD more objectively.

Drawbacks of the Study

Differences in study populations and CT slice thickness, hardware, protocols, and postprocessing software may have contributed to differences between our results and previous studies. The correlation between QCT parameters and clinical tests was not as strong as expected, but some important factors hampered our results. First, we conducted a retrospective study, and most CT scans were not performed on the exact same day as the PFTs, although we excluded patients with long intervals (>30 days). Second, CT scans at breath-hold, PFTs, the 6MWT, and quality of life questionnaires all depend on patient cooperation. Despite breathing training before the CT scans and PFTs, reaching the desired expiratory volumes is impaired in advanced emphysema (GOLD 3 and 4) with hyperinflated lungs, which was a confounder of lung density and volume in expiratory scans in these patients. Third, we did not use the mass-conserving volume change method for estimating voxel volume changes from the HU values within an inhale/exhale CT image pair as proposed by Castillo et al.³⁰ We plan to apply this method in further studies. Finally, this study focused on the CT assessment of lung parenchyma and did not consider airway abnormalities, which also affect lung ventilation function significantly. Therefore, the relationship between airway abnormalities and PFT results needs to be addressed in future studies.

Conclusion

The present study provides reference values for several QCT parameters in both inspiration and expiration compared with clinical tests in a large group of severe COPD patients. As expected, lung volumetry from CT scans works well and strongly correlates with spirometry. In particular, P15, including P15_{EX} and P15_{Diff}, correlates with the severity of emphysema and parameters of lung diffusion capacity. QCT can help to evaluate the severity

of COPD in GOLD 3 and 4 patients. Many studies, including ours, indicate that expiratory CT is superior for assessing COPD severity and better reflects airflow limitation and air trapping.

Abbreviations

6MWT, 6-minute walk test; CAT, COPD assessment test; COPD, chronic obstructive pulmonary disease; DLCO, carbon monoxide diffusing capacity; FEV₁, forced expiratory volume in 1 second; FVC, forced vital capacity; GOLD, global initiative for chronic obstructive lung disease; KCO, carbon monoxide transfer coefficient; P15, 15th percentile of lung attenuation distribution curve; PFT, pulmonary function test; pre BS, before application of short-acting bronchodilators; QCT, quantitative computed tomography; RV, residual volume; SGRQ, St. George's Respiratory Questionnaire; TES, total emphysema score; TLC, total lung capacity; TLV, total lung volume.

Acknowledgments

This work was supported by the non-profit Central Research Institute Fund of the Chinese Academy of Medical Sciences (2019PT320008, 2018PT32003). The authors thank Mrs. Bettina Herwig for English language editing.

Disclosure

Author Bianca C Lassen-Schmidt is an employee of Fraunhofer MEVIS, the developer of the MeVis PULMO3D software. Ralf H Hubner reports personal fees from Pulmonx during the conduct of the study. All other authors report no conflicts of interest in this work.

References

- Singh D, Agusti A, Anzueto A, et al. Global strategy for the diagnosis, management, and prevention of chronic obstructive lung disease: the GOLD science committee report 2019. Eur Respir J. 2019;53(5). doi:10.1183/13993003.00164-2019
- Cho HB, Chae KJ, Jin GY, et al. Structural and functional features on quantitative chest computed tomography in the Korean Asian versus the White American healthy non-smokers. *Korean J Radiol*. 2019;20 (7):1236–1245. doi:10.3348/kjr.2019.0083
- Zaporozhan J, Ley S, Eberhardt R, et al. Paired inspiratory/expiratory volumetric thin-slice CT scan for emphysema analysis: comparison of different quantitative evaluations and pulmonary function test. *Chest*. 2005;128(5):3212–3220. doi:10.1378/chest.128.5.3212
- Belzer RB, Lewis RJ. The practical significance of measurement error in pulmonary function testing conducted in research settings. *Risk Anal.* 2019;39(10):2316–2328. doi:10.1111/risa.13315
- Nishio M, Matsumoto S, Tsubakimoto M, et al. Paired inspiratory/ expiratory volumetric CT and deformable image registration for quantitative and qualitative evaluation of airflow limitation in smokers with or without COPD. *Acad Radiol*. 2015;22(3):330–336. doi:10.1016/j. acra.2014.09.011

 Madani A, De Maertelaer V, Zanen J, Gevenois PA. Pulmonary emphysema: radiation dose and section thickness at multidetector CT quantification - Comparison with macroscopic and microscopic morphometry. *Radiology*. 2007;243(1):250–257. doi:10.1148/radiol.2431060194

- Lee YK, Oh Y-M, Lee J-H, et al. Quantitative assessment of emphysema, air trapping, and airway thickening on computed tomography. Lung. 2008;186(3):157–165. doi:10.1007/s00408-008-9071-0
- Heussel CP, Herth FJ, Kappes J, et al. Fully automatic quantitative assessment of emphysema in computed tomography: comparison with pulmonary function testing and normal values. *Eur Radiol*. 2009;19(10):2391–2402. doi:10.1007/s00330-009-1437-z
- Koyama H, Ohno Y, Nishio M, et al. Three-dimensional airway lumen volumetry: comparison with bronchial wall area and parenchymal densitometry in assessment of airway obstruction in pulmonary emphysema. *Br J Radiol*. 2012;85(1020):1525–1532. doi:10.1259/bjr/22602417
- Schroeder JD, McKenzie AS, Zach JA, et al. Relationships between airflow obstruction and quantitative CT measurements of emphysema, air trapping, and airways in subjects with and without chronic obstructive pulmonary disease. AJR Am J Roentgenol. 2013;201(3): W460–W470. doi:10.2214/AJR.12.10102
- Dirksen A. Monitoring the progress of emphysema by repeat computed tomography scans with focus on noise reduction. *Proc Am Thorac Soc.* 2008;5(9):925–928. doi:10.1513/pats.200804-033QC
- Ashraf H, Lo P, Shaker SB, et al. Short-term effect of changes in smoking behaviour on emphysema quantification by CT. *Thorax*. 2011;66(1):55–60. doi:10.1136/thx.2009.132688
- Kuhnigk J-M, Dicken V, Zidowitz S, et al. Informatics in radiology (infoRAD): new tools for computer assistance in thoracic CT. Part 1. Functional analysis of lungs, lung lobes, and bronchopulmonary segments. *Radiographics*. 2005;25(2):525–536. doi:10.1148/rg.252045070
- Lynch DA. Progress in imaging COPD, 2004–2014. Chronic Obstr Pulm Dis. 2014;1(1):73–82. doi:10.15326/jcopdf.1.1.2014.0125
- Wang ZM, Gu SC, Leader JK, et al. Optimal threshold in CT quantification of emphysema. Eur Radiol. 2013;23(4):975–984. doi:10.1007/s00330-012-2683-z
- Lynch DA, Austin JHM, Hogg JC, et al. CT-definable subtypes of chronic obstructive pulmonary disease: a statement of the fleischner society. *Radiology*. 2015;277(1):192–205. doi:10.1148/radiol.2015141579
- Wanger J, Clausen JL, Coates A, et al. Standardisation of the measurement of lung volumes. *Eur Respir J.* 2005;26(3):511–522. doi:10.1183/09031936.05.00035005
- Miller MR, Crapo R, Hankinson J, et al. General considerations for lung function testing. Eur Respir J. 2005;26(1):153–161. doi:10.1183/09031936.05.00034505
- Hochhegger B, Alves GRT, Irion KL, Moreira JD, Marchiori ED. Emphysema index in a cohort of patients with no recognizable lung disease: influence of age. *J Bras Pneumol*. 2012;38(4):494–502. doi:10.1590/S1806-37132012000400012
- Mets OM, van Hulst RA, Jacobs C, van Ginneken B, de Jong PA. Normal range of emphysema and air trapping on CT in young men. AJR Am J Roentgenol. 2012;199(2):336–340. doi:10.2214/AJR.11.7808
- Hetzel J, Boeckeler M, Horger M, et al. A new functional method to choose the target lobe for lung volume reduction in emphysema - comparison with the conventional densitometric method. *Int J Chron Obstruct Pulmon Dis.* 2017;12:2621–2628. doi:10.2147/COPD.S139304
- Kauczor HU, Heussel CP, Fischer B, Klamm R, Mildenberger P, Thelen M. Assessment of lung volumes using helical CT at inspiration and expiration: comparison with pulmonary function tests. *Am J Roentgenol*. 1998;171(4):1091–1095. doi:10.2214/ajr.171.4.9763003
- Hersh CP, Jacobson FL, Gill R, Silverman EK. Computed tomography phenotypes in severe, early-onset chronic obstructive pulmonary disease. COPD. 2007;4(4):331–337. doi:10.1080/15412550701601 274
- 24. Cavigli E, Camiciottoli G, Diciotti S, et al. Whole-lung densitometry versus visual assessment of emphysema. *Eur Radiol*. 2009;19 (7):1686–1692. doi:10.1007/s00330-009-1320-y

25. Gelb AF, Schein M, Kuei J, et al. Limited contribution of emphysema in advanced chronic obstructive pulmonary disease. Am Rev Respir Dis. 1993;147(5):1157-1161. doi:10.1164/ajrccm/147.5.1157

- 26. Akira M, Toyokawa K, Inoue Y, Arai T. Quantitative CT in chronic obstructive pulmonary disease: inspiratory and expiratory assessment. AJR Am J Roentgenol. 2009;192(1):267-272. doi:10.2214/ AJR.07.3953
- 27. Sciriha A, Lungaro-Mifsud S, Scerri J, Magro R, Camilleri L, Montefort S. Health status of COPD patients undergoing pulmonary rehabilitation: a comparative responsiveness of the CAT and SGRQ. Chron Respir Dis. 2017;14(4):352-359. doi:10.1177/1479972317694622
- 28. Jones PW, Harding G, Berry P, Wiklund I, Chen WH, Kline Leidy N. Development and first validation of the COPD assessment test. Eur Respir J. 2009;34(3):648-654. doi:10.1183/09031936.00102509
- 29. Finch S, Laska IF, Abo-Leyah H, Fardon TC, Chalmers JD. Validation of the COPD Assessment Test (CAT) as an outcome measure in bronchiectasis. Chest. 2019.
- 30. Castillo E, Vinogradskiy Y, Castillo R. Robust HU-based CT ventilation from an integrated mass conservation formulation. Med Phys. 2019;46(11):5036-5046. doi:10.1002/mp.13817

International Journal of Chronic Obstructive Pulmonary Disease

Dovepress

Publish your work in this journal

The International Journal of COPD is an international, peer-reviewed journal of therapeutics and pharmacology focusing on concise rapid reporting of clinical studies and reviews in COPD. Special focus is given to the pathophysiological processes underlying the disease, intervention programs, patient focused education, and self management protocols. This journal is indexed on PubMed Central, MedLine and CAS. The manuscript management system is completely online and includes a very quick and fair peer-review system, which is all easy to use. Visit http://www.dovepress.com/testimonials.php to read real quotes from published authors.

Submit your manuscript here: https://www.dovepress.com/international-journal-of-chronic-obstructive-pulmonary-disease-journal