INTRODUCTION

Chronic obstructive pulmonary disease (COPD) is defined as a common preventable and treatable disease characterized by persistent airflow limitation, which is usually progressive, and is associated with enhanced chronic inflammatory responses in the airways and lung (1). COPD is a major cause of morbidity and mortality in the general population, and the WHO estimates that it will become the third leading cause of death worldwide (2). It is a heterogeneous and complex condition, with a wide variety of clinical and pathologic characteristics. Even though pulmonary function test (PFT) parameters are currently used to diagnose and classify the severity of COPD, they cannot fully represent the type and range of the heterogeneous pathophysiologic abnormalities present in COPD. Additionally, the PFT tends to be relatively insensitive to early stages and subtle changes in COPD, before functional parameters begin to show changes (3). Furthermore, patients with similar PFT values may demonstrate completely different clinical and radiological phenotypes.

In the era of precision medicine, a more meticulous understanding of clinical disease manifestation and its relationship to COPD can be provided by advanced imaging techniques. In this review article, we focus on the current status and future challenges of CT imaging in COPD.
with endogenous disease pathogenesis will likely have significant implications on disease management and understanding of the disease process. Chest CT is the most commonly used imaging modality; it is essentially non-invasive and provides information regarding the structural and underlying pathophysiologic changes in the COPD patient. CT scans allow in vivo visual analysis of morphologic characteristics and the distribution of both emphysema and small airway disease, the two main components of COPD. However, qualitative visual CT assessment, which has been the mainstream method for acquiring information from CT scans, is prone to inter-reader, and sometimes even intra-reader, variability, limiting its application in broad clinical and experimental settings (4). Therefore, the need for more objective CT-based measures has grown significantly over the past decade. In this context, recent technical advances in quantitative CT imaging methods are of considerable interest for their ability to provide more precise and reproducible estimates of the severity and distribution of emphysema and airway disease (5). In the early days, research on the quantitation of CT in COPD was weighted towards quantification of emphysema. However, there have recently been an increasing number of publications targeting the airway component of COPD, and these can be further divided into direct airway parameter measurements and quantification of air trapping as the functional manifestation of small airway disease. In this review, we briefly address previous studies on COPD quantification, highlight the current concepts, and discuss the future direction of quantified CT imaging.

EMPHYSEMA QUANTIFICATION

Pulmonary emphysema, defined as an abnormal irreversible dilatation of air spaces and destruction of airway walls distal to terminal bronchioles, is a major finding in COPD. Due to the increased proportion of air compared with normal lung parenchyma, emphysema appears as a region of relatively lower CT attenuation, expressed as lower Hounsfield units (HU). The density mask method, first introduced in 1988, quantifies emphysema by calculating the voxels that are lower than a certain HU threshold, referring to such voxel regions as low attenuation areas (LAA) (6). The HU threshold of -910 was initially proposed in 1988 and various different threshold values have been tested afterwards. In 2006, Madani et al. (7) has shown the optimal HU threshold to be -960 HU or -970 HU according to macroscopic and microscopic pathologic correlations. However, because higher threshold values result in higher sensitivity to image noise, the most commonly accepted threshold value is -950 HU (8), with the percentage area of lung less than -950 HU (the emphysema index, or %LAA) being widely used to estimate the emphysema component in COPD patients (Fig. 1A, B) (9).

An alternative approach, based on the frequency histogram of lung attenuation, calculates the lung parenchymal HU at a given percentile along the histogram. This value is called the “percentile index,” and validation with pathologic specimens has shown that the optimal threshold is the first percentile (7). However, because of concerns regarding increased image artifacts and noise at the first percentile level, the 15th percentile threshold is most commonly used (8, 10). The percentile index has been reported to be more robust for evaluating longitudinal variations in emphysema, and as being less sensitive to lung volume changes (8, 11).

An innate limitation of CT histogram measurement approaches is that they do not take into account the heterogeneous patterns of distribution, size, or clusters of emphysema, which could be influential factors in the evaluation of a COPD patient. Since emphysema is a regionally distributed disease, determination of the anatomic zonal distribution of emphysema, rather than simple quantification of the total amount of emphysematous involvement, may be of importance. Most currently available quantitative CT methods are able to divide each lung into separate anatomical zones, allowing ratios of the extent of emphysema in different lung zones to be calculated. More recent methods also permit the segmentation of lobes to measure lobar volumes and the extent of emphysema objectively (Fig. 1C). To quantify the local size of lung parenchyma showing emphysematous changes, several investigators have used low attenuation cluster analysis by determining the power-law exponents (D) of emphysema holes; this represents the cumulative frequency-size distribution of the emphysema, and estimates how the emphysema areas are conglomerated to form small-to-large “clustered” regions of emphysema (12-14).

Quantification of emphysema is relatively simple, and quantitative CT measurements have been shown to correlate better with macroscopic measurements of emphysema than do mea-
sures of visual CT assessment (15). In patients with α1-antitrypsin deficiency, density-based emphysema measurement has shown a superior accuracy in detecting worsening of emphysema, and therefore has been accepted as the effective diagnostic method for detecting disease progression (16). The CT density mask method was found to have an association with COPD mortality, and is reported to be a stronger predictive parameter for cardiac and respiratory mortality than global initiative for chronic obstructive lung disease (GOLD) staging (17, 18). Patients with greater quantitative measures of emphysema were reported to have more rapid decline of PFT parameters (19). Furthermore, quantified measures of emphysema showed associations with frequent exacerbation and worse clinical outcomes from exacerbation events (20-22). When the heterogeneity of emphysema assessed using quantitative CT was taken into account, the basal distribution of emphysema was associated with greater impairment in forced expiratory volume in 1 second (FEV$_1$), with central areas of emphysema showing a stronger correlation with airflow limitation than those in the periphery of the lung (23, 24). In one report, the quantitatively measured heterogeneity of emphysema distribution was associated with a decrease in FEV$_1$ and FEV$_1$/forced vital capacity (FVC) (25). When com-

**Fig. 1.** Quantitative CT measurement of emphysema.


B. Using the density mask method with a threshold of -950 HU, areas with HU values lower than the threshold (low attenuation areas%) can be readily quantified, and overlaying of the density mask (shown in green on online figure) allows a more robust assessment of emphysema.

C. Most available CT quantification software provides reliable automatic segmentation of the lung, making regional quantification of emphysema possible.

HU = Hounsfield unit
pared with the %LAA value obtained from the whole lung, histogram-based measures of different patterns of emphysema involvement were more predictive of measures of PFT results, severity of dyspnea, and quality of life (26). Additionally, the regional distribution of emphysema has been shown to be clinically important in selecting optimal candidates for lung volume reduction surgery (27).

AIRWAY DISEASE: DIRECT MEASUREMENT

Small airway remodeling is an important factor in COPD, and is known to be the strongest determinant of airflow limitation. A small airway is defined as an airway with an internal diameter smaller than 2 mm, and because of the limited spatial resolution of even the most up-to-date CT scanners, the accurate direct measurement of small airway dimensions has proven to be a great challenge. Current quantitative CT techniques allow three dimensional reconstruction of the airways and adequate non-invasive measurement of segmental and sub-segmental airways, which are “large airways” by definition (Fig. 2A). However, a previous report showed that changes in large airways reflect changes in small airways on pathologic examination, and the measured CT parameters of large airways correlated with PFT results (28). Additionally, the measured CT parameters of more distal airways showed stronger correlations with spirometry (29). Therefore, it may be of value to obtain quantified airway parameters from measurable large airways on CT of COPD patients (Fig. 2B).

The most widely used methodology for obtaining quantitative airway measurements is the “full-width half-maximum (FWHM)” method. This uses projecting linear rays from the airway center to determine the inner and outer margin of the airway wall, using an HU threshold for differentiating the wall from

![Fig. 2. Quantitative assessment of airways.](image)

A. Recent technical advances allow more accurate and robust automatic extractions of airways, with three dimensional volumetric reconstructions.
B. Once airways are extracted and target points in the airways are selected, quantitative airway parameters are automatically measured (shown in blue on online figure).
Increased bronchial wall thickness and WA% is reported to be correlated with deterioration of FEV\textsubscript{1}, and this correlation is reported to be stronger with more distal airways (29, 31, 32, 36). Also, in subjects with a comparatively low emphysema component, quantitative CT measurements of airways showed stronger correlations with physiologic indices (37). Airway parameters also correlated with functional markers and exercise capacity of COPD patients (9, 38). It is hypothesized that increased bronchial wall thickness may be closely related to inflammation of the airway, and thus may have an association with aggravated symptoms of bronchitis or exacerbations. Accordingly, increased quantitative parameters of airway wall thickness were related to an increased frequency of, and mortality from, COPD exacerbations (20, 21). However, compared with emphysema quantification, there are still many technical difficulties and uncertainties regarding measurements of airway changes in COPD patients, and further investigation is warranted.

**AIRWAY DISEASE: AIR TRAPPING MEASUREMENT**

As previously mentioned, airways with a diameter less than 2 mm are the major sites of airflow obstruction, and small airway narrowing is known to occur in early stages of COPD, before major emphysematous changes occur (39). Small airway wall remodeling is known to be the most dominant factor for the air-

![Fig. 3. Full-width at half-maximum method and commonly measured airway parameters.](image)

A. In the attenuation profile along an outwards flowing ray from the luminal center-point through to the airway wall, the inner and outer airway wall boundaries are assumed halfway to the maximum on the lumen side, and halfway to the minimum on the parenchymal side, respectively.

B. Diverse airway parameters can be obtained using quantitative analysis, including wall thickness WA, WT, LA, LD, Pi, and WA%. HU = Hounsfield units, LA = lumen area, LD = lumen diameter, Pi = internal perimeter, WA = wall area, WA\% = wall area percentage, WT = wall thickness.
flow limitation observed in COPD patients, and small airway changes occur due to abnormal cellular proliferation and peribronchial fibrosis. As CT is unable to adequately image the small airways directly because of its limited resolution, many researchers have used different parameters of air trapping measured on expiratory phase CT as functional estimates of small airway disease.

One of the earliest and simplest ways to estimate air trapping using chest CT is to evaluate the percentage of low attenuation area at a threshold of -856 HU or -850 HU. The threshold is chosen because it is known to be the attenuation of normally inflated inspiratory lung, with the concept being that healthy expiratory lung should show higher attenuation than this value. Using this relatively straightforward method, several researchers have reported high correlations with spirometry results, including FEV₁/FVC and FEV₁ percent predicted (3, 40). Other methods for measuring air trapping have been addressed, using information from both inspiratory and expiratory CT, including the ratio of inspiratory to expiratory lung volume and the expiratory to inspiratory ratio of mean lung density.

However, the above methods have an innate limitation, in that it is impossible to separate trapped air from emphysematous lung or obstructed small airways. One method developed to quantify air trapping outside the emphysematous area is measurement of the relative volume change between -860 HU and -950 HU (41). This method excludes the emphysema portion of all voxels with attenuation lower than -950 HU from the inspiration and expiration scans, and calculates for the whole lung the relative volumes with attenuation values less than -860 HU on each inspiratory and expiratory CT. Results from this method revealed strong associations with physiologic parameters of gas trapping; however, this method was limited by the fact that pixel-by-pixel matching of inspiratory and expiratory scans was not performed.

To overcome such a limitation, a new quantitative method called parametric response mapping has recently been developed; this involves pixel-by-pixel co-registration of inspiratory and expiratory CT scans so that a more exact comparison can be

![Fig. 4. Air trapping measurement using a co-registration method. Using an image co-registration technique, expiratory CT images are modified and matched with inspiratory CT images. This technique allows voxel-by-voxel comparisons of attenuation changes between inspiration and expiration, with air trapping being defined as areas with less change in attenuation than the preset threshold (60 Hounsfield units in the example shown above).](image-url)
obtained (5). This method enables the classification of emphysema and functional small airway disease, and is reported to show an increased correlation with PFT results (42). Another method, called air trapping index (ATI), relies on the difference of each of the image voxels detected on co-registered inspiratory and expiratory CT scans (Fig. 4) (43, 44). Unlike other methods, both parametric response mapping and ATI methods allow additional assessment of the regional heterogeneous distribution of air trapping.

Quantified parameters of air trapping were found to be associated with changes in FEV₁ and other clinical parameters, and with aggravating GOLD classification (41, 45). Additionally, results from paired evaluations of inspiratory and expiratory CT images revealed that they were stronger predictors of spirometry results than the use of just the expiratory scan (46). Quantitatively measured air trapping parameters in a large cohort study were associated with FEV₁ deterioration, which was notably prominent in patients with a lesser severity of COPD (47). Although air trapping parameters provide assessment of only the consequence of pathological airway changes, such techniques contribute to the characterization of disease phenotypes, and considerable research is ongoing.

OTHER COMPONENTS OF COPD

Pulmonary vascular change is a prevalent feature of COPD and is reported to be a strong predictor of mortality. However, the actual pathogenesis of pulmonary vascular changes in COPD is still a vastly unknown territory, and new quantitative CT methods are being developed to better understand the relationship between vascular changes and other components of COPD. The pulmonary artery diameter and the ratio of the pulmonary artery diameter to aorta diameter were associated with pulmonary artery pressure and increased exacerbation rates (48). When the cross sectional area (CSA) of the pulmonary arteries was measured using quantitative volumetric CT, total CSA showed a strong correlation with the severity of emphysema, and a significant but lesser degree of correlation with air trapping in COPD patients (49). Using a technique to automatically segment pulmonary vasculature and measure the total blood volume, pruning of distal arteries was suggested as an early change in smoking-related COPD, with the extent of emphysema having a negative relationship with total pulmonary blood volume (50). The clinical significance of these findings still needs to be clarified, and continued investigation is much needed.

Ischemic heart disease is a common comorbidity in COPD. Patients with COPD are prone to more frequent and more fatal myocardial infarctions, even after adjusting for common risk factors such as smoking status and old age (51). Furthermore, reduced FEV₁ has been reported to be an increased risk factor for cardiovascular mortality (52). The quantitative measurement of coronary artery calcium (CAC) is an accurate and non-invasive parameter of coronary artery atherosclerosis. A high CAC score is reported to be an independent predictor of future cardiovascular events, and COPD patients were found to have higher CAC scores and increased cardiovascular mortality (53). In addition, calcium score used as a measurement of the degree of systemic atherosclerosis was reported to have a weak but significant correlation with the volume fraction of emphysema on quantitative CT, FEV₁/FVC, and diffusion capacity, with these correlations being independent of age, body mass index (BMI), and smoking amount (54).

Cachexia and skeletal muscle wasting are common comorbidities in COPD patients, and changes in body composition are reported to be associated with COPD severity, airflow obstruction, and mortality. Thoracic respiratory muscles in particular are unique and crucial for alveolar ventilation, and a weakness of respiratory muscle results in the dyspnea and respiratory failure associated with mortality in COPD patients. One group quantitatively measured the area of pectoralis muscle using axial CT images and revealed a significantly decreased muscle area in COPD patients compared with healthy individuals, with patients having a decreased muscle area generally having a worse disease stage. Decreased pectoralis muscle area was also associated with functional markers of disease, including BODE index, dyspnea score, and other parameters of patient activity (55). In a similar context, the measurements of intercostal muscle mass and attenuation using quantitative CT methods were significantly correlated with COPD severity and the quantitatively assessed extent of emphysema on CT (56). A decrease in thoracic muscle mass with increasing intercostal fat was associated with worsening of COPD severity.

COPD patients are at a higher risk of osteoporosis because the two entities share similar clinical factors such as smoking, de-
increased physical activity, low BMI, prolonged steroid use, and vitamin D deficiency. Furthermore, osteoporosis often results in vertebral compression fractures, which can deteriorate FEV\(_1\) and decrease vital capacity in COPD patients. Although dual-energy X-ray absorptiometry of the lumbar spine and hip is currently the gold standard for the diagnosis of osteoporosis, quantitative CT has recently emerged as a solid alternative for assessing bone mineral density (BMD), using a three-dimensional technique to provide volumetric assessment of bone. In a recent report, COPD, especially the emphysema dominant type, was associated with an increased severity of osteoporosis measured on chest CT, even after adjustment for conventional risk factors (57). Moreover, decreased thoracic vertebral BMD was an independent predictor of higher rates of exacerbations and other morbidity measures in COPD patients (58).

**OBSTACLES AND FUTURE PERSPECTIVES**

Although quantitative analysis of CT has shown promising results, further work is required to implement quantitative CT parameters as imaging biomarkers that can be used to practically manage COPD patients. First, methods for quantifying the various components of COPD need standardization, optimization, and simplification. Although various methods have shown good correlations with physiologic or clinical parameters, there is currently neither consensus nor a gold standard method for quantifying each component of COPD. Furthermore, quantitative parameters derived from CT images can be influenced by both patient and CT-related factors, including patient weight, inspiration adequacy, CT manufacturer, and calibration. Optimization of CT protocols and quality control in image acquisition are critical for large multicenter/multinational trials and comparisons of results from different subjects. Additionally, one very important practical point is that the time and effort required for quantitative assessment, including post-processing procedures, should be minimized, with procedures being automated to improve their clinical accessibility and utility.

The present GOLD strategy does not recommend routine use of CT scanning in COPD, and only advises that it may be helpful in excluding differential diagnosis or when surgical options, such as lung volume reduction surgery, are being considered. This is largely because of a relative lack of evidence that information gathered from CT can actually modify treatment plans, predict treatment response or prognosis, and eventually alter the mortality of COPD patients. A large number of prior studies have compared quantitative CT parameters with simple PFT results, and more analyses need to be assessed against disease outcome measures, effect of treatment, and other clinically significant markers of disease severity. Similarly, investigation of the relationship between genetic information and quantitative imaging parameters in COPD could provide crucial information on disease pathogenesis because individual manifestations of COPD may vary according to genetic variations. Additionally, more longitudinal cohort studies are required to track CT changes over time and obtain an accurate picture of disease progression and how it can be effectively captured by quantitative CT imaging. Further work should be performed to analyze the clinically important COPD phenotypes using quantitative CT techniques because such information may have an important influence on COPD trials and drug development, which are in great clinical demand.

Lastly, quantitative CT analysis should be augmented with the rapidly increasing number of new up-to-date technologies. Dose reduction technology, such as low dose protocols, noise lowering procedures, and new image reconstruction algorithms, are vital for longitudinal studies and the more widespread use of CT in clinical practice. Dual-energy CT technology allows functional imaging of the lung and may have further applications in quantitative evaluation of COPD patients. In the era of big data analysis, radiomics is a recent active field of study, in which high throughput data is extracted and a large amount of advanced quantitative imaging features are analyzed, often automatically or semi-automatically. Although most published radiomics studies have largely focused on cancer research, application of radiomics to COPD may reveal additional insights. Artificial intelligence, including deep learning and machine learning technology, although still a vastly unknown field of study, has also gained great worldwide attention in the past few years, and is another potential field of study that could be augmented by quantitative CT technology.

**CONCLUSION**

Quantitative CT imaging is a promising technique for objec-
tively measuring disease manifestation, and has already shown associations with traditional clinical and physiologic markers of COPD. Quantitative CT analysis provides reliable estimates of two dominant features of COPD, emphysema and airway disease, and various other features, including pulmonary vessel alterations, atherosclerosis, cachexia, and osteoporosis. These quantified parameters can be used as important research tools to understand the underlying heterogeneity of COPD, and to assist in revealing the fundamental pathogenesis in it. The translation of information derived from these approaches into actual clinical practice has the potential to guide individualized management strategies and improve disease outcomes in COPD patients.

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CT를 이용한 만성폐쇄성폐질환의 정량적 평가: 현 상태와 미래 과제에 대한 리뷰
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만성폐쇄성폐질환은 다양한 임상적, 병리학적 특징을 가지고 있는 복합적인 질병이다. CT는 만성폐쇄성폐질환을 평가하는 데 가장 널리 사용되는 영상도구이나, 지금까지의 주관적 평가 방법으로는 임상적으로 의미 있는 일관되고 객관적인 영상 지표를 얻는 데 한계점이 있었다. 이러한 문제점을 극복하기 위하여 만성폐쇄성폐질환 환자에게서 얻은 CT 정보를 정량적으로 분석하는 방법이 많은 연구자들의 주목을 받았고, 이러한 분야의 기술적 진보를 바탕으로 폐기종과 소기도 질환이라는 만성폐쇄성폐질환의 두 주요 요소뿐만 아니라 폐혈관의 변화, 심혈관 질환, 체 성분 변화, 그리고 골다공증을 포함하는 다양한 변화를 정량적으로 분석하고 측정할 수 있게 되었다. 그리고 이렇게 측정된 정량적인 수치들은 만성폐쇄성폐질환 환자의 폐 기능 검사 수치와 연관성 보고되었으며, 더 나아가 질병의 실제 임상적 경과와 환자의 예후와도 연관성이 있는 것으로 나타났다. 하지만 이러한 긍정적인 보고에도 불구하고 아직 CT를 이용한 만성폐쇄성폐질환의 정량적 평가 방법은 실제 진료와 환자의 치료에 널리 적용되고 있지 않다. 이에 저자들은 먼저 현재까지 이루어진 CT를 이용한 만성폐쇄성폐질환의 정량적 평가 방법의 기술적 진보에 대하여 리뷰하고, 이러한 방법이 질병과 환자의 폐기능에 갖는 임상적 의미에 대하여 서술하고자 한다. 또한, 이러한 기술이 실질적으로 임상적으로 유용한 도구로 자리잡기 위해서 극복해야 하는 한계점과 문제점에 대하여 논의하고, 마지막으로 앞으로 CT를 이용한 만성폐쇄성폐질환의 정량적 평가 방법에 대한 연구가 나아가야 하는 방향에 대하여 제안하고자 한다.

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