The role of quantitative computed tomography in the diagnosis of chronic obstructive pulmonary disease

Abstract

In the last 25 years, there have been important improvements in computed tomography (CT) that may give more details about the lung structure in chronic obstructive pulmonary disease (COPD). The clinical exam and “classic” radiology (chest X-ray, conventional CT) have important roles: they raise the suspicion of hyperinflation, they highlight aspects of pulmonary hypertension, they may detect the triggers of exacerbations, they rule out some COPD complications and other lung diseases that can cause dyspnea (pneumothorax, tumors, bronchiectasis, and fibrosis). The spirometry may confirm the obstructive ventilatory disorder pattern of the disease. The modern CT scan technique - High Resolution CT (HRCT) with Multi-Detector CT procedure (MDCT) gives additional information about morphological details of parenchyma, bronchi, pulmonary vessels or lung function (ventilation/perfusion disorders) without significant lung irradiation. The new techniques provide quantifiable parameters that characterize the emphysema, the main COPD phenotypes and the risk of disease progression. Quantitative volumetric analysis of emphysema provides an early diagnosis of the disease in patients exposed to smoking and pollution. An early personalized diagnostic in COPD offers stronger reasons to prophylaxis (inhaled bronchodilators, anti-inflammatory medication, pulmonary rehabilitation, education for lifestyle changes).

Keywords: COPD, emphysema, multidetector CT

Introduction

In the last 25 years, there have been several improvements in computed tomography (CT) that might give more details on the structure of lung parenchyma in chronic obstructive pulmonary disease (COPD), on the main phenotypes or even on lung function. The clinical exam and “classic” radiology (chest X-ray, conventional CT) have important roles: to raise the suspicion of hyperinflation, highlight aspects of pulmonary hypertension, they may detect the triggers of exacerbations, and rule out some complications and other diseases with dyspnea (pneumothorax, tumors, bronchiectasis, tuberculosis, fibrosis, or cardiomegaly). The classical confirmation of the COPD relies on ventilatory obstructive disorder pattern on respiratory function tests (RFTs).

RFTs cannot cover the morphological analysis of the disease and detail the correlation between lesions and lung function. In the same time, RFTs become abnormal late in the disease evolution and cannot distinguish the very early COPD stage (when the patients are asymptomatic) to provide a targeted prophylaxis.

The primary evaluation of COPD in the pulmonology clinic has to be comprehensive and performed according to several combined criteria (symptoms, COPD Assessment Test, lung function by spirometry/plethysmography, number of previous year exacerbations, comorbidities, phenotypes characterization). The modern CT scan technique – High Resolution CT scan (HRCT) with Multi-Detector CT procedure (MDCT) – can offer important additional information to the clinical investigation and RFTs about anatomy of lung parenchyma, bronchi, pulmonary vessels and even about ventilation/perfusion function, without significant lung irradiation.
Conventional computed tomography (CT)

Chest X-ray is the first-line imaging technique for pulmonary disease assessment. It is widespread, cheap and everywhere accessible but it is not good enough for emphysema analysis. Chest X-ray has a low resolution and produces images summation effect that gives it a low sensitivity and specificity. A normal chest X-ray cannot rule out emphysema or characterize well enough the different phenotypes of emphysema. Chest X-ray is not considered to be the most accurate tool for identifying lung cancer in COPD patients. Thoracic CT scan became the most efficient imaging method that largely characterizes COPD. CT is not a routine investigation during initial diagnosis assessment. First, diagnosis in COPD relies on clinical features and spirometry.

Thoracic CT will be performed in selected cases upon indication of the pulmonology specialist. CT becomes necessary for confirming a specific cause for an exacerbation suspected on a standard chest X-ray or an associated disease complication (pneumonia, pneumothorax, interstitial lung diseases that produce shortness of breath or cough (pneumothorax, bronchiolitis, interstitial lung disease, bronchiectasis)).

The advantages of conventional CT scan compared with standard chest X-ray are multiple in terms of high sensitivity and specificity:

- CT highlights diffuse or localized emphysema (bullae).
- CT can detect small structures/details and analyze the tissue density in the lung parenchyma, bronchi, vessels, chest wall, and mediastinum.
- CT allows detection of aggravating factors (neoplastic diseases, fibrosis, bronchiectasis, cardiovascular comorbidities), triggers of COPD exacerbations or disease complication (pneumonia, pneumothorax, pleural effusion, pulmonary infarction).
- CT can detect some other respiratory diseases that associate the same symptoms as COPD: dyspnea, cough (sarcoidosis, fibrosis, tuberculosis, silicosis, lung tumors).

Kurashima et al. showed in 2005 in a study conducted on 516 patients with FEV1 below 70% that 12.7% of patients had other diseases than COPD.

High Resolution CT/ Multidetector CT

Multidetector CT (MDCT) is an advanced form of computed tomography (CT) technology for diagnostic imaging. In MDCT, a two-dimensional array of detector elements replaces the linear array of detector elements used in typical conventional and helical CT scanners. CT technique is capable to perform quantitative analysis of emphysema, air trapping, airways and vessels. It uses thin slices (0.625 and 1 mm), with three-dimensional (3D) reconstruction, low dose radiation (effective tube current, 30-60 mAs), analysis of lung volumes and attenuation in inspiration and expiration. 3D HRCT is nowadays considered the gold standard technique for noninvasive airspace evaluation.

Newer HRCT techniques provide an opportunity for a comprehensive depiction of COPD in term of lung parenchyma and bronchi morphology, as well as lung functionality.

Advantages of MDCT in COPD:

- Early detection of emphysema in asymptomatic chronic smokers (equivalent to COPD stage 0) in time of normal lung function.
- Quantitative evaluation and topographic disposition of emphysema (bullae).
- Characterization of phenotypes of emphysema (centrlobular, panacinar, paraseptal emphysema, localized bubbles) and COPD.
- Significant correlation between MDCT issues and clinical parameters: age, smoking history, St. George’s Questionnaire, BMI, spirometry, CPR.
- Differential diagnosis of emphysema with other diseases that produce shortness of breath or cough (pneumothorax, bronchiolitis, interstitial lung disease, bronchiectasis).
- Noninvasive quantitative evaluation of airway dimensions (bronchi and bronchioles, small air spaces) and virtual bronchoscopy.
- Functional assessment of pulmonary reserve before thoracic surgery (including lung transplantation) and before lung volume reduction techniques (by surgery or by unidirectional valves insertion).
- 3D reconstruction of the pulmonary vessels and evaluation of different pathologies: shunts, malformations, fistulas, tumors.
- Monitoring of progression of the disease under therapy.

Emphysema detection and characterization

In radiological terms, emphysema consists of parenchymal areas with low attenuation without definable walls. The attenuation coefficient of emphysematous lung usually decreases to a value below minus 950 HU. On expiratory CT, air trapping may be quantified by evaluating the percentage of lung volume less than a given threshold (e.g. -856 HU) comparing lung volumes and attenuation between expiration and inspiration.

For emphysema extension, MDCT elaborates a “voxels density mask model” = quantitative analysis of the region with abnormally low attenuation (threshold -950 UH) and calculating the emphysema score (ES). An overall “lung density map” will be created that offers a global picture of the affected lung. MDCT-derived ES seems to be the best predictor of all-cause mortality in a cohort of stable COPD outpatients along with age.

In addition, MDCT may highlight several parameters to assess emphysema:

- The thoracic cross-sectional area (measurement of the thoracic cage hyperinflation).
- The sterno-aortic distance at the carinal level (increased in hyperinflation).
- The “saber-sheath trachea configuration” (static modeling of the trachea by coronal narrowing and sagittal widening of the thoracic cage in emphysema).
MDCT is able to indicate also the emphysema distribution. This parameter may be a predictor of mortality (large emphysema in the upper areas associates an increased survival in COPD patients compared to basal localization)\(^{(13)}\).

Different phenotypes of emphysema and of COPD can be differentiated by MDCT with a high degree of sensitivity and specificity\(^{(18,19)}\):

- **Early centrilobular emphysema** (related to smoking and exposure to gases) is characterized by round low-attenuation in the central area of the secondary pulmonary nodule around the lobular artery\(^{(20)}\). The secondary pulmonary lobules are not destroyed. As emphysema progresses, the hypertransparent areas become confluent (Figure 1).
- **Panlobular emphysema** commonly involves the lung bases. It affects the entire secondary pulmonary lobule. Panlobular emphysema typically occurs as a result of \(\alpha_1\) antiprotease deficiency, but may also be determined by chronic smoking in the advanced stage of common COPD. Genetic evaluation of emphysema is useful to identify specific groups of patients who may benefit from specific enzymatic treatment. MDCT is much more sensitive for progression evolution and response to specific therapy than conventional CT or RFTs\(^{(21, 22)}\) (Figure 2).
- **Paraseptal emphysema** is defined as subpleural, and subfissural gas collection located in the periphery of the secondary pulmonary lobules. It is considered a panlobular type and a precursor of bullae (Figure 3).
- **Pericicatricial emphysema** is found near the scars and is accompanied by parenchymal and bronchial distortion.
- **Association of bronchiolitis or bronchiectasis to emphysema** includes a poor prognosis, an increased risk for exacerbation, but a better response to inhaled corticoids and bronchodilators\(^{(23, 24)}\) (Figure 4).
- Morphological characterization of emphysema (centri-/panlobular/extralobular emphysema) by MDCT correlates with histological analysis\(^{(17)}\).

**Imaging of the airways and pulmonary vessels**

HRCT/MDCT may characterize both large and small airways by constructing a 3D model of the airways. It can visualize the external and internal diameters of bronchi, the airway wall thickness and the inner design – “virtual bronchoscopy”.

Measurement of airway parameters correlates with the severity of airflow obstruction and with history of COPD exacerbation. Gas trapping was associated with...

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**Figure 1.** Thoracic CT scan (sagittal section). Diffuse emphysema, multiple apical bullae, bronchiectasis

**Figure 2.** Thoracic CT scan (axial sections): paraseptal emphysema, “mosaic” images of emphysema (with normal and decreased attenuation), panlobular emphysema in right upper lobe (posterior).
smaller airway lumen diameters less than 2 mm, greater dyspnea and chronic bronchitis\(^{(15)}\).

The obliteration of the small bronchioles determines patchy high-density areas (under-ventilated areas) of the parenchyma which alternate with low attenuation areas (with air trapping and poor perfusion). It will result a “mosaic attenuation pattern” with a lack of homogeneity of the lung density strongly specific for COPD especially on the end-expiratory scans\(^{(12)}\). Mosaic perfusion pattern has a good correlation with a lower FEV\(_1\) and FEV\(_1\)/FVC values\(^{(13)}\).

In the same time, MDCT may accurately characterize the disorders of the large bronchi or trachea (malacia, dilation, stenosis or diverticula, fistulae, tumors)\(^{(25)}\) (Figure 5). The precise location of the obliterated bronchi that determine the retrostenotic air trapping is important for the lung reduction intervention (by surgery or valves insertion).

Pulmonary vessels can also be characterized by MDCT using several parameters:
- The diameter of the pulmonary artery trunk.
- The diameters of the main pulmonary arteries.
- Wall arteries thickening.
- Vascular distortion (increased angles of arterial branches separation)\(^{(20)}\).
- Evaluation of “vascular attenuation” (thinning of pulmonary vessels and reduction in their number)\(^{(12)}\). Low “density” of the vascular images occurs in advanced emphysema\(^{(6,11,12)}\).

MDCT/HRCT has shown several benefits for a comprehensive morphological and functional characterization of COPD. It stands as a noninvasive method with multiple valences for an early valuable diagnostic which may indicate a targeted therapy of the disease.

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**Figure 3.** Thoracic CT scan (axial section). Panlobular basal emphysema (CT density -1024 UH)

**Figure 4.** Thoracic CT scan (axial section). Diffuse emphysema. Bronchiectasis and bronchiolitis phenotype (image “tree-in-bud” right lower lobe, posterior)

**Figure 5.** HRCT (3D Reconstruction). Lung bilobar right hypoplasia. Left mediastinal lung hernia.
Further studies are needed on the use of algorithms for precise quantification of changes in different lung structures in emphysema. The risk of irradiation remains the most important disadvantage of the technique which restricts indications for repeated usage of the HRCT. The clinical benefit of MDCT is higher than the risk of radiation. Despite this fact, reduction and optimisation of radiation are strongly recommended in accordance with the ALARA principle. The dose cannot be higher than the standard dose of 10-20% recommended by scanner vendors (26).

Morphologic and quantitative assessment of the lung parenchyma and airways in practice may be obtained at a radiation exposure value of 300-350 mGy·cm. This leads to an effective dose of 5-6 mSv. The COPD patients are usually more than 40 years old, this allowing repeated follow-up CT studies, that might be acceptable (27).