The Relevance of Targeting Treatment to Small Airways in Asthma and COPD

Pierachille Santus, Dejan Radovanovic, Matteo Pecchiari, Matteo Ferrando, Francesco Tursi, Vincenzo Patella, and Fulvio Braido

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Summary

COPD and asthma have different risk factors and pathogenesis, but they share a pathophysiologic hallmark characterized by small airways disease. Although difficult to explore and measure, modifications of distal airways’ pathophysiology and biology represent an early sign of obstructive disease and should be researched and assessed in everyday clinical practice. In the last 15 years, computed microtomography scans have shed light on the anatomy and physiology of the so-called silent zone, and research devoted to investigate the effect of inhaled treatments on small airway pathophysiology has been increasing. This narrative review offers a historical summary of researchers and landmark studies that reported, defined, and advanced the research on small airways. We then discuss the latest findings on the role and characteristics of the small airways’ inflammatory and cellular structure, and we describe the assessment tools available to detect small airways dysfunction in COPD and asthma and the effect of bronchodilators and inhaled steroids on functional and biological biomarkers. Finally, we analyze the newest technological therapeutic advances aimed at small airways treatment in terms of inhalation devices and small particle size molecules. Key words: small airways; COPD; asthma; severe asthma; small particles; ventilation inhomogeneity; closing volume. [Respir Care 2020;65(9):1392–1412. © 2020 Daedalus Enterprises]
**Introduction**

Asthma and COPD represent 2 of the most common non-communicable diseases worldwide. Globally, asthma is the most prevalent chronic respiratory disease, with around 339 million people affected (http://globalasthmareport.org, Accessed February 24, 2019); COPD prevalence is continuously increasing, estimates being around 251 million cases in 2016 (https://www.who.int/respiratory/copd/burden/en, Accessed February 24, 2019). Despite advances in therapy and prevention, it is estimated that 1,000 people die each day from asthma (~400,000 each year); deaths from COPD are 8 times more common, and COPD is projected to be the third leading cause of death worldwide by 2030 (https://www.who.int/respiratory/copd/burden/en, Accessed February 24, 2019). Asthma and COPD both represent an unbearable social and economic burden on health care systems all over the world. Although these diseases differ regarding risk factors and natural history, they both present with alterations of the small airways. The importance of the peripheral airways in the pathophysiology and clinical manifestations of asthma and COPD make them the intuitive target for long-term pharmacologic approaches. However, the so-called silent zone of the lung has been always difficult to access, and the lack of standardized and unanimously accepted methods of measurement has often left the assessment of the small airways to the experimental and investigational level, never becoming accessible to clinical practice. In the last 20 years, however, due to better, more reliable, and more standardized techniques, the study of small airways has gained renewed interest. Bronchodilators and inhaled steroids represent the treatment mainstay for COPD and asthma, respectively. Technological progress has allowed the development of new delivery systems and drug formulations designed to increase drug deposition and to improve therapeutic efficiency, effectiveness, and drug safety.

This narrative review introduces the concept of small airways disease pathophysiology and biology, then summarizes the latest findings and advances in the assessment of small airways, and finally provides an overview of the pharmacology and mechanics behind molecule and delivery systems that are currently available for the treatment of asthma and COPD.

**Pathophysiology of Small Airways in Asthma and COPD**

**Defining Small Airways**

The section provides a brief description of the evolution of airway physiology that led to the current notion that the airways with a diameter < 2 mm offer very low resistance in normal lungs. A schematic representation of the main steps that led to the discovery of small airway pathophysiology is reported in Figure 1.

Since 1915, following the studies by Fritz Rohrer, it was thought that the main contribution to airflow resistance within the respiratory system came from the small conducting airways. In fact, Rohrer studying 1 excised undistended lung, noted that the cross-sectional area at every successive branching did not increase as he traced the trachea-bronchial tree, which supports the idea that there is a significant pressure drop along the bronchial generations, ie, an increase in resistance for every daughter branch almost twice that of the parent bronchial branch (Fig. 1). What Rohrer was missing in his aerodynamic calculations was that, in inflated lungs, total cross-sectional area of the conducting airways increased exponentially as the gas-exchanging surface was approached, a piece of information added only in 1963 by Ewald R Weibel. In 1965, Malcolm Greer calculated for the first time the airway resistance, unifying 2 paramount concepts into lung pathophysiology, ie, the Hagen-Poiseuille equation and the Weibel quantitative morphometric studies (Fig. 1) and demonstrating that the peripheral airways contributed to < 10% of total airway resistance, much less than what was estimated by Rohrer. The explanation for such a phenomenon was that the small
airways’ cross-sectional area was much larger compared with conductive airways. In fact, starting from the fifth or sixth airway bronchial generation, the total airway cross-sectional area increases exponentially due to a semi-dichotomous branching system and a greater cross-sectional area of the 2 daughter branches compared with their parent branch.\(^9,11\) As a result of this anatomical structure, the closer the exchange surface, the shorter and more compacted the airways, allowing the high velocity and turbulent flow from the proximal airways to become laminar and independent of density in lung periphery.\(^12\)

Soon after, directly measuring resistance in airways < 2 mm by means of a retrograde catheter, Macklem and Mead\(^13\) confirmed Green’s calculations and demonstrated how peripheral airways offered very low resistance in normal lungs (Fig. 1). In 1967, Peter T Macklem and Jere Mead directly measured the resistance in airways < 2 mm in diameter,\(^15\) confirming the estimation made by Green. Soon after, Hogg, Macklem, and Thurlbeck reported that the small airways are the major site of airway resistance in patients with emphysema. The gray area in the airway generation-resistance diagrams represents airways < 2 mm.

\[ \Delta p = \frac{8\mu LQ}{\pi R^4} \]

The Hagen-Poiseuille equation

1915
Rohrer estimates airway resistance in undistended lungs

1963
Weibel publishes his morphometric studies

1838-1839
Green demonstrates that peripheral airways contribute to < 10% of total airway resistance

1965
Hogg, Macklem and Thurlbeck introduce the term small airways, the major site of airflow obstruction in patients with emphysema

1968
Macklem and Mead directly measure the resistance in airways < 2 mm with a retrograde catheter

Fig. 1. Brief historical overview showing the milestones that lead to the discovery of the nature of airway resistance\(^8\) and to small airways mechanics. In 1915, Fritz Rohrer erroneously posited that the peripheral airways were the major site of airway resistance; integrating the lung morphometric studies performed by Weibel\(^9\) with the Hagen-Poiseuille equation, Green estimated that peripheral airways contribute to < 10% of total airway resistance.\(^10\) In 1967, Peter T Macklem and Jere Mead directly measured the resistance in airways < 2 mm in diameter,\(^15\) confirming the estimation made by Green. Soon after, Hogg, Macklem, and Thurlbeck reported that the small airways are the major site of airway resistance in patients with emphysema. The gray area in the airway generation-resistance diagrams represents airways < 2 mm.

Although this only minimally affects the mechanical properties of the lung. This was later supported by Hogg and
colleagues\textsuperscript{16} by means of the single-breath nitrogen washout test (SBN2).

The Silent Zone

The redundancy of the lung obscures small airways disease, which becomes detectable with traditional lung function tests only when the lung periphery is consistently damaged. This section describes the pathophysiology of small airways and the need for effective pulmonary function tests.

The core investigations on pathophysiology and mechanics of small airways were predominantly conducted during the 1960s and 1970s, and the model that was most frequently adopted was chronic bronchitis and emphysema. At that time, it was clear that COPD was a condition related to cigarette smoking, with an insidious onset and an as-yet unclear epidemiology and natural history. Researchers were aware, however, that the earliest pathological changes in young tobacco smokers occurred in denuded epithelium and intramural inflammatory cells in the respiratory and membranous bronchioles.\textsuperscript{17} The latter, which occurred mainly in small airways, was known as peripheral airway disease.\textsuperscript{18} However, it was also clear that, before the damage became detectable with standard spirometric parameters such as FE\textsubscript{V\textsubscript{1}}, or FVC, the number of diseased airways had to be huge.\textsuperscript{5} Due to this particular feature, the small airways were referred as the silent zone of the lung.\textsuperscript{5,19} To identify early signs of disease, a series of new tests were developed. One of the earliest markers of precocious small airways damage was the identification of the frequency dependence of lung dynamic compliance in the case of small airway disease. Briefly, Woolcock and colleagues\textsuperscript{20} noted that, in contrast to healthy lungs, subjects with chronic bronchitis and asthma experienced a significant reduction of dynamic compliance during a wide range of breathing frequencies due to prolonged regional time constants, an observation that was not modified by the administration of bronchodilators.\textsuperscript{20} Together with the radioactive gas technique\textsuperscript{21} and the SBN2 test,\textsuperscript{22} the dynamic compliance assessment was characterized by poor standardization and availability.\textsuperscript{18,20} Moreover, these tests were considered markers of early smoke exposure rather than having a significant clinical prognostic meaning.\textsuperscript{18}

Pathobiology of Small Airways in COPD

The section deals with the immunology and inflammatory cells that characterize the airways of patients with COPD, with a peculiar focus to small airways. Inhalation of noxious particles, in western countries mainly represented by cigarette smoke, represents the most important risk factor for the development of COPD.\textsuperscript{23} In low-income countries, however, COPD can be also found in nonsmokers exposed to smoke from burning wood or biomass.\textsuperscript{24,25} The inflammation in COPD is localized predominantly to peripheral airways and lung parenchyma.\textsuperscript{26} Airways \(< 2\) mm in diameter are more prone to undergo these pathological changes and thus to be occluded by mucus, inflammatory infiltrates, smooth muscle and mucus glands hypertrophy, and airway wall thickening, which involves the epithelium, the smooth muscle, and the adventitia.\textsuperscript{27}

Compared with healthy individuals, the airway lumen of patients with COPD is rich in neutrophils and macrophages, and greater concentrations of macrophages, T-lymphocytes, and B-lymphocytes have been reported in the airway wall and parenchyma.\textsuperscript{23,27,28} The inflammatory response promoted by macrophages is upregulated by the nuclear transcription factor-\(\kappa\)B and by p38 mitogen-activated protein kinase. In turn, macrophages produce chemotactic agents for CD8\textsuperscript{+} and CD4\textsuperscript{+} lymphocytes.\textsuperscript{23,29} CD8\textsuperscript{+} cells induce apoptosis and necrosis of epithelial and endothelial cells, which likely contribute to the lung damage observed in patients with COPD.\textsuperscript{30} In COPD, activated neutrophils and lymphocytes infiltrate the small airways and are correlated with disease severity.\textsuperscript{31-33} Neutrophils contribute to alveolar destruction by producing myeloperoxidase, neutrophil elastase, matrix metalloproteinase-8, and matrix metalloproteinase-9,\textsuperscript{23} with the latter mediators also being responsible for increased mucus production.\textsuperscript{34} Inconsistent results have been published on the presence of eosinophils in the bronchoalveolar lavage of subjects with COPD. Higher eosinophil concentrations may be regulated by epithelial mediators such as interleukin-33 (IL-33) in response to epithelial cell injury.\textsuperscript{35}

Upregulation of the histone deacetylase (HDAC) activity is one of the mechanisms by which corticosteroids inhibit inflammation. The neutrophil- and macrophage-driven reactive oxygen species released secondary to cigarette smoke exposure is responsible for the inactivation of HDAC-2. In fact, the activities of HDAC-2, -3, -5, and -8 have all been demonstrated to be reduced in COPD patients, suggesting a role for the inactivation of HDACs in steroid insensitivity in COPD.\textsuperscript{23,26,36} Isajevs and coworkers\textsuperscript{37} reported that cigarette smoke increased pro-inflammatory cytokines through the activation of nuclear transcription factor-\(\kappa\)B and by inhibiting HDAC-2. Nuclear transcription factor-\(\kappa\)B p65 expression was greater in large airways, correlating with an increased number of macrophages, whereas the HDAC-2 expression was mainly decreased in small airways, suggesting its potential role in the development of bronchiolitis.\textsuperscript{37}

Pathobiology of Small Airways in Asthma

The section deals with the immunology and inflammatory cells that characterize the airways of patients with asthma, with a particular focus on small airways. The role of the
Peripheral airways in asthma is increasingly recognized as a relevant target for asthma treatment. Previous research conducted on ex vivo lung specimens and transbronchial biopsies indicates that inflammatory and structural changes occur in both small airways and lung parenchyma of subjects with asthma. Thus the contribution of small airways to the pathobiology of asthma, combined with the presence of the relevant receptors in these airways, suggest that this site should not be neglected in the clinical monitoring of asthma or for the development of new treatments for the disease. Due to the anatomical site, procedural limitations, and the relatively poor availability of histological samples, the evaluation of the cellularity of small airways in asthma has always been difficult, all the more so if we consider that the majority of specimens are obtained during asthma exacerbations leading to a patient’s death, which do not reflect the disease in a stable state.

Inflammatory changes of the proximal airways of subjects with asthma also occur in the distal airways. Small airways of patients with asthma are thickened and the airway wall is infiltrated by a variable distribution of eosinophils, T-lymphocytes, neutrophils, and macrophages. Moreover, small airways often present histological signs of obstruction secondary to mucus plugging and epithelial damage. An increased amount of activated eosinophils have been described in peripheral airways compared with large conductive airways in subjects with asthma, suggesting a significant inflammatory process localized in lung periphery. In fact, in subjects who died from severe asthma, eosinophils and CD45+ leukocytes were found mainly in the outer wall of small airways. Accordingly, Hamid and colleagues reported a higher number of lymphocytes, eosinophils, and neutrophils associated with an increased mRNA expression of IL-4, IL-5, and eotaxin. This finding was confirmed with increased IL-4 and IL-5 mRNA-positive cells found in small airways in patients with mild and moderate asthma. These modifications may contribute to an uncoupling of the small airways from the surrounding lung parenchyma, promoting airway closure and collapse.

Small Airways and Severe Asthma

The section focuses on the role of small airways in the pathophysiology and inflammatory patterns in fatal and severe asthma. Studies performed in subjects with asthma have confirmed that the inflammatory changes to the peripheral airways parallel the severity of the disease. In a study by Carroll et al., the increased expression of mast cell chymase in the outer area of the distal airways was significantly associated with impaired lung function in individuals with severe asthma. Compared to subjects with mild to moderate asthma, subjects with severe asthma exhibited an increased production of alveolar nitric oxide (NO) but not bronchial NO, implyng that a more intense degree of inflammation occurs in the periphery of the bronchial tree in the most severe forms of asthma. In patients with severe or fatal asthma, there is increased thickness of the airway smooth muscle layer in the proximal and distal airways and increased smooth muscle cell size.

In patients with severe asthma, the progressive loss of lung function is caused by structural alterations and remodeling of the proximal and distal airways. Specific abnormalities of the distal airways and lung parenchyma have also been described in nocturnal asthma, which occurs in 30–75% of the asthmatic population. In nocturnal asthma, but not in non-nocturnal asthma, a more pronounced inflammation was described in the peripheral airways rather than in the large central airways, with a significant increase in lymphocytes, macrophages, and eosinophils during the night. Autopsy studies have reported increased numbers of lymphocytes and eosinophils uniformly distributed throughout the large and small airways in lung tissue from individuals with fatal cases of asthma and from subjects with mild and severe asthma who died of other causes, compared to control subjects with no history of asthma.

The percentage of eosinophils and macrophages was also significantly increased in the alveolar tissue and was related to a nocturnal reduction in lung function. The inflammatory response observed in the context of the alveolar tissue could explain the loss of airways-parenchyma coupling detected in individuals with nocturnal asthma. Reduced macrophage phagocytosis of bacteria has also been described in subjects with severe asthma.

Small Airways Assessment

The section discusses the latest evidence in terms of small airway disease assessment and quantification with lung function tests and inflammatory markers, with a special focus on lung ventilation heterogeneity and closing volume, as well as the radiological evaluation of small airways and the development of new technologies in small airways imaging.

Plethysmography

Mid-Exspiratory Flows. The mean forced expiratory flow (FEF) between 25% and 75% of FVC (FEF25-75) is one of the most popular indices of peripheral airways obstruction, resembling the concavity of the flow-volume curve. However, its use in clinical practice is limited by physiologic variability and measurement inconsistency. The repeatability of FEF25-75 is affected by its volume dependence and might also be evidence of air-flow inhomogeneity because it is changed by the presence of severe air-flow obstruction. Its value as a predictor of peripheral obstruction has also been
questioned following findings from the Severe Asthma Research Program of the National Heart, Lung and Blood Institute, which reported a lack of correlation between FEF25-75 and other indices of air trapping.58

**Lung Volume and Airway Resistance.** Commonly employed spirometric tests, such as FEV1 and FEV1/FVC, generally reflect large airways function. Vital capacity (VC), residual volume (RV), and FVC are, to some extent, correlated to the functional state of the peripheral airways. VC is reduced in patients with peripheral airways obstruction, and this reduction may occur before that of the FEV1/VC ratio.60 RV is increased in patients with air trapping, a phenomenon that is secondary to alterations that occur mainly at the level of peripheral airways.38 FVC is reduced at any level of airway obstruction, but it may be also a reflection of air trapping.38 In all normal lungs, the flow-volume curve depends on the density of the inhaled gas.61 Heliox is a mixture of 80% oxygen and 20% helium, and a decrease in pulmonary resistance is associated with an increase in maximum expiratory flow.61 In patients with irreversible airway obstruction, breathing heliox can lead to an increase in resistance because this mixture has similar viscosity but a lower density compared with ambient air.63 Laminar flow resistance is viscosity-dependent, so in the presence of fully developed laminar flow there is a small increase in resistance.19 Despas and colleagues61 reported that adults with asthma fall into 2 groups: those who respond to heliox and those who do not respond. The greatest prevalence of nonresponders was represented by smokers with asthma who had recurrent infections with chronic cough and sputum production,64 implicating a possible role of small airways disease in the response to heliox. Recently, Sood and coworkers65 noted that methacholine-induced cough in subjects with otherwise preserved airway sensitivity was reflected by signs of small airways obstruction such as air trapping and hyperinflation.65

A spirometric pattern characterized by a decrease in VC and FEV1, an increase in RV and RV/total lung capacity, and a normal total lung capacity and FEV1/FVC was described by Stanescu and colleagues.66 This pattern has been suggested as an early sign of peripheral airways disease and has been observed in early emphysema stages, in elderly people and asymptomatic asthmatics.67,68 Although the RV represents a reliable and reproducible parameter, it is considered a poor mirror of small airways dysfunction and mainly reflects premature airway closure and air trapping. More recently, Santos and coworkers69 reported that dyspnea at rest in subjects with COPD was significantly related to specific airway resistance (Raw) rather than FEV1, and that acute changes in specific Raw induced by bronchodilators are significantly associated with improvements in dyspnea, whereas no correlation was observed with dynamic volumes. Indeed, the assessment and extent of bronchodilator responsiveness in terms of dynamic volumes is influenced by the slow or rapid inhalation maneuver preceding the evaluation of FVC.70

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**Fig. 2.** Single-breath nitrogen washout test (SNB2) in a healthy young subject and a subject with severe COPD. The different phases that distinguish the SNB2 test are shown. The closing volume is the point at which the phase IV departs from the phase III plateau. The phase III slope is increased in subjects with COPD compared with normal subjects. After bronchodilator administration in the COPD subject, the phase III slope is reduced and the vital capacity is increased, with a consequent reduction in closing volume. The irregularities visible primarily on the tracing obtained in a healthy young subject represent the cardiac artifact.
Lung Inhomogeneity and Closing Volume. Inhaled bronchodilators represent the treatment mainstay in COPD, and their functional target has always been the small airways. However, the evaluation of bronchodilator efficacy has often been limited to dynamic volume response, which poorly reflect small airway modifications. On the other hand, peripheral lung injury and poor distribution of ventilation can be assessed readily with the SBN2 test, reflected in a steeper slope of phase III and an increased closing volume. The SBN2 test is a noninvasive tool to assess small airways dysfunction and ventilation heterogeneity, and it has been used since the 1960s to detect early closure of peripheral airways. It is based on the detection of regional differences in ventilation distribution, represented by the phase III slope of the flow-volume curve following the washout test. Toward the end of expiration, an abrupt increase in N2 concentration is observed, which corresponds to the beginning of phase IV. This is due to the preferential emptying of the apex after the closure of airways at the base of the lung. The volume at which phase IV begins is termed closing volume (CV), and it reflects small airways dysfunction. The CV identifies the point at which some regions of the lung stop contributing to lung deflation and trap air beyond functionally closed airways. An increased CV is responsible for 2 major abnormalities: poor distribution of ventilation with impaired gas exchange, and peripheral lung injury. Experiments in normal animals have shown that mechanical ventilation at low lung volume induces peripheral airway injury with concurrent increase in airway resistance. Further damage and a substantial inflammatory reaction occur in animal models when the collapse of small airways is promoted by increasing the surface tension, thus supporting the idea that cyclical opening and closing of the small airways during tidal breathing may also cause lung injury in humans. This should occur whenever CV exceeds the expiratory reserve volume, or when the closing capacity (ie, the sum of CV and RV) exceeds the functional residual capacity (Fig. 2). Despite being an easily performed and sensible technique, SBN2 is limited by poor reproducibility and predictability.

The assessment of CV by means of the SBN2 test has been limited by the observation that, especially in patients with severe air-flow obstruction, phase IV is not present in a significant percentage of cases. Another proposed operational indicator of the onset of airway closure is represented by the inflection of the transpulmonary pressure-volume curve obtained during slow exhalation from total lung capacity (Fig. 3). Technical difficulties to obtain reliable transpulmonary pressure-volume curves and controversies concerning the criteria for the identification of the inflection point have limited the use of this indicator in humans. Recently, Pecchiari and coworkers simultaneously recorded the SBN2 test and the slow deflation transpulmonary pressure-volume curve in 40 healthy subjects and 43 subjects with COPD. The onset of phase IV identified the CV in SBN2 test, whereas in the transpulmonary pressure-volume curve the CV was identified by the deviation from the exponential fit and the inflection point.
of the interpolating sigmoid function. The authors found that the CV evaluated with the SBN2 test always corresponded to the inflection point at the transpulmonary pressure-volume curve. However, in almost 50% of subjects with COPD, phase IV was not present during the SBN2 test, yet a clear inflection point at the transpulmonary pressure-volume curve was discernable. This evidence suggests that the absence of phase IV does not correspond to the absence of a CV, probably reflecting a sign of increased ventilation heterogeneity (Fig. 4). The study also reported that almost 50% of subjects with COPD experience cyclic opening and closing of the airways during tidal breathing, a phenomenon that implies the presence of continuous stresses at the level of peripheral airways because such stress is a source of mechanically-driven inflammation and a plausible mechanism sustaining damage to the airways and disease progression. Soon after, the same group demonstrated that both muscarinic antagonists and β-adrenergic agonists were similarly able to improve the heterogeneity of lung ventilation, positively affecting gas exchange and the extent of small airway closure in subjects with moderate to severe COPD, thus reducing the percentage who experienced cyclic opening and closure of the airways during tidal breathing (Fig. 5). Surprisingly, the administration of bronchodilators did not change the transpulmonary pressure at which the CV started, indicating that the effects were due to the reduced smooth muscle tone and improved airway compliance. The latter is the first-ever evidence of the potential effect of bronchodilator therapy as a key factor to reduce the risk of small airway damage in patients with COPD (Fig. 6). These hypotheses are complemented by the observations of Bhatt and colleagues, who reported how emphysema can mechanically influence surrounding areas of normal lung, thus increasing the risk of lung function decline.

The multiple-breath nitrogen washout test measures functional residual capacity. Patients with high airway resistance take longer to clear the nitrogen, and the test can distinguish between obstructions in the large airways or the small airways. Dutrieue et al and Verbanck et al used the multiple-breath nitrogen washout test to derive 2 variables, the index of conductive ventilation heterogeneity and the index of acinar ventilation heterogeneity. In subjects with asthma, the clinical application of the multiple-breath nitrogen washout test includes evaluation of ventilation heterogeneity after an airway challenge and analysis of response differences to inhaled steroids when comparing nonextrafine and extrafine particles. The influence of aging on ventilation heterogeneity needs to be taken into consideration. In this scenario, Verbanck and colleagues recently reported that, with respect to clinical response, age has a small but consistent effect that needs to be factored in when using the multiple-breath nitrogen washout test indices for the detection of small airways abnormality in disease.

**Forced Oscillation Technique and Impulse Oscillometry**

The forced oscillation technique works by superimposing pressure oscillatory waves on the airway during tidal
breathing. The forced oscillation technique was first described by DuBois in 1956 as a method to characterize respiratory impedance. Both the forced oscillation technique and impulse oscillometry are easy to perform and require minimal patient cooperation. Impulse oscillometry operates by delivering a continuous spectrum of frequencies, favoring the characterization of lung mechanics, both in adults and in preschool children with asthma. This method can reflect regional lung inhomogeneity and therefore represents a useful tool during bronchoprovocation challenges. Takeda and coworkers reported a strong correlation between impulse oscillometry parameters and clinical symptoms and asthma control in subjects with asthma. Impulse oscillometry allows a separate assessment of the peripheral and proximal airways, and it appears to be a sensitive screening tool for the early detection of airway obstruction. Oppenheimer and colleagues proposed that impulse oscillometry can be used as a noninvasive tool for assessment of distal airway function when spirometry is normal. More recently, the Assessment of Small Airways Involvement in Asthma study, a multinational prospective cohort study, explored the effects of small airway dysfunction in subjects with asthma. Small airway dysfunction was noted across all asthma severity stages; more importantly, however, subjects with milder asthma had an index of acinar ventilation heterogeneity within normal values, whereas impulse oscillometry parameters (ie, the fall in resistance from 5 Hz to 20 Hz [R5–R20]) and spirometric variables...
were directly correlated with more severe involvement of the small airways.\textsuperscript{99}

Peripheral Exhaled NO

The fraction of exhaled NO (F\textsubscript{ENO}) is a reliable marker of airway eosinophilia, and it is measured in clinical practice and used in the follow-up of patients with asthma. In both asthma and COPD, F\textsubscript{ENO} levels are influenced by airway bronchoconstriction.\textsuperscript{100} F\textsubscript{ENO} and alveolar NO can be elevated in patients with COPD, independent of smoking status and disease severity,\textsuperscript{101,102} and levels $\geq 20$ parts per billion have been associated with a higher risk of exacerbation.\textsuperscript{103}

More recently, Santus and coworkers\textsuperscript{104} reported that peripheral NO, in particular the alveolar fraction of F\textsubscript{ENO}, is associated with the amount of air trapping and diffusion capacity derangement in subjects with COPD. Moreover, after administration of long-acting $\beta_2$-agonists in acute conditions, F\textsubscript{ENO} was reduced and the decrease in peripheral NO was correlated with the degree of improvement of lung ventilation inhomogeneity and air trapping.\textsuperscript{104} These data may indicate that NO in patients with COPD may reflect the amount of mechanical stress and strain-induced production of inflammatory mediators such as NO, sustaining the hypothesis that peripheral airway injury is one of the major determinants of COPD pathogenesis and its role in the disease’s natural history.\textsuperscript{108} Although F\textsubscript{ENO} or alveolar NO evaluation seem to be potentially useful tools for COPD management, more data are needed to support their routine clinical use.

Imaging

Computed tomography (CT) has emerged as a useful tool to assess peripheral airways disease noninvasively in patients with asthma. The cardinal CT sign of peripheral airways disease in asthma is the presence of pulmonary decreased attenuation areas, which are more consistent on expiratory CT scans.\textsuperscript{105-107} Decreased attenuation areas reflect air trapping, which may be either secondary to transient small airway obstruction or spasm or may be caused by fixed peripheral remodeling.\textsuperscript{108,109}

Two different systems are currently used to quantify air trapping on CT: the visual score and the densitometric analysis. The former is limited by the subjective approach that causes sometimes significant interobserver variability; thus the densitometric analysis has been the technique of choice in recent scientific investigations. Specifically, either mean lung density or 850 HU cut-off CT density has been used to quantify air trapping in subjects with asthma.\textsuperscript{110,111}

Quantification of air trapping in expiratory CTs may provide additional information on the effects of inhaled therapy targeted to small airways disease, dysfunction which may not be readily revealed with pathophysiological testing.\textsuperscript{109,112} The latter technique has been used to compare the relative efficacy of 2 different inhaled medications on small airway hyper-reactivity; small-size inhaled particles prompted a significant improvement in air trapping compared with traditional-size particles.\textsuperscript{109,111}

Air-trapping quantification has also been used to define asthmatic phenotypes. A recent investigation by Busacker and colleagues\textsuperscript{111} reported that quantitative CT scan identified subjects with a high risk of severe disease. Subjects with air trapping above the median were significantly more likely to have a history of asthma-related hospitalizations, ICU visits, or mechanical ventilation.\textsuperscript{111} CT imaging of pulmonary ventilation patterns in patients with asthma has evolved over the last decade with the introduction of more sophisticated techniques. The direct assessment of peripheral remodeling with CT measurements is still limited due to the difficulty to obtain accurate measures of the distal airways. Computer-assisted 3-dimensional analysis of the airway tree structure is currently advancing our knowledge on the anatomy and pathology of small airway disease. With this technique, researchers have reported that airway wall measurements of the distal airways have the strongest correlation with air-flow obstruction parameters in subjects with asthma.\textsuperscript{113}

The spatial resolution of currently available high-resolution CT does not allow for an accurate assessment of small airways in patients with either asthma or COPD. Recent developments carried out by Tanabe and co-workers\textsuperscript{114} tested the possibility of using ultra-high-resolution CT to measure airway walls of bronchioles down to 2 mm in diameter with a significant reduction of the error rate compared to traditional high-resolution CT algorithms.

Although current evidence indicates that small airway obstruction could be routinely assessed with CT, this technique does have some limitations, including a lack of standardization of technical parameters for the CT scanner, a lack of consensus on the best index for small airways disease assessment, and exposure of subjects to ionizing radiation. Indeed, the regular assessment of small airway disease via CT imaging is still not part of routine clinical practice for young asthmatic patients.

To date, magnetic resonance imaging (MRI) of the lung has been cumbersome for two primary reasons: the density of lung parenchyma, which returns a weak magnetic resonance signal, and the many air–tissue interfaces, which introduce magnetic susceptibility effects and cause a rapid decay of the magnetic resonance signal in the lung.\textsuperscript{115} The development of new contrast agents such as hyperpolarized helium-3 ($^3$He) and xenon-19 ($^{129}$Xe) have allowed a much higher spatial and temporal resolution of MRI scans (eg, hyperpolarized gas lung MRI ventilation imaging). Helium is an inert gas that does not diffuse into pulmonary circulation, thus allowing the assessment of ventilation defects and heterogeneity, whereas xenon diffuses more rapidly to
lung parenchyma and further in the capillary bed, allowing the assessment of both the gas phase and the dissolved phase for the simultaneous assessment of ventilation and perfusion. This technique has been used successfully in subjects with asthma to evaluate regional ventilation defects as well as air-flow obstruction and its reversibility after methacholine challenge after exposure to bronchodilators.\textsuperscript{116-118} Hyperpolarized \textsuperscript{3}He-diffusion MRI at multiple diffusion timescales and quantitative CT scan densitometry was compared to the multiple-breath nitrogen washout test by Gonem and co-workers\textsuperscript{119} to assess the acinar airway involvement in subjects with asthma. Gonem et al.\textsuperscript{119} found that an increased index of acinar ventilation heterogeneity was associated with worse asthma severity, worse CT expiratory air-trapping indexes, and with the apparent diffusion coefficient of \textsuperscript{3}He at 1 s, a sign associated with alterations in long-range diffusion within the acinar airways.\textsuperscript{119} The objective of future research should be to integrate CT and MRI acquisitions with physiological measures of small airway dysfunction to provide both anatomico-radiological measures of peripheral airways and dynamic regional ventilation-perfusion defects.

According to recent evidence from pathoradiological studies, compared with control smokers, the number of terminal bronchioles in subjects with mild and moderate COPD was decreased by 40% and 43%, respectively, and the number of transitional bronchioles decreased by 56% and 59%, respectively.\textsuperscript{120} Moreover, subjects with mild COPD and a preserved FEV\textsubscript{1}, compared with at-risk smokers, exhibited a 19% reduction in total airway count measured with CT scan (ie, the total airway count represents the major determinant for disease progression).\textsuperscript{85} A recent study from Vasilescu and co-workers\textsuperscript{121} offered for the first-ever evidence that the analysis of functional small airways disease obtained by means of parametric response mapping (PRM) on subjects with severe COPD is correlated with loss of terminal bronchioles, luminal narrowing, and obstruction of ex-vivo eclipsed lungs analyzed with micro-CT. The latter is one of the first examples of a radiomorphological marker able to identify small airways damage in patients with COPD.\textsuperscript{121} Indirect evidence that may support the importance of the progressive loss of treatable airways is reported in a post hoc analysis of the 4-y Understanding Potential Long-Term Impacts on Function with Tiotropium (UPLIFT) trial, which reported a progressive reduction in the extent of the acute bronchodilator response, a reduction that was increasingly pronounced in subjects with severe to very severe COPD.\textsuperscript{122} The decline in lung function was generally larger in subjects with severe to very severe than with mild to moderate air-flow obstruction/MID EXP. FLOWS

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<td>LUNG VOLUMES AND RESISTANCES</td>
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| RV, RV/TV, TLC, 
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| Closing volume |
| CLOSING PRESSURES | Esophageal Balloon |
| PL-V curve (Paw) |
| CLOSING VOLUME | Bronchoscopy |
| BAL |
| Induced sputum |
| FLAMMATION/HISTOLOGY | BAL biopsy |
| Inflammatory cells |
| IMMAGING/VENTILATION HETEROGENEITY | CT/microCT |
| MRI |
| Nuclear Medicine |
| PRM |
| Xe vent CT |

\textbf{Fig. 7.} Available noninvasive and invasive assessment tools for small airway assessment. Diagnostic tests are ordered based on complexity and availability from the simplest (upper level) to the most complex (lower level). FVC = forced vital capacity; VC = vital capacity; RV = residual capacity; IC = inspiratory capacity; sRawtot = total specific airway resistance; IOS = impulse oscillometry; SB2N2 = single-breath nitrogen washout test; CV = closing volume; CC = closing capacity; MBW = multiple breath washout test; FRC = functional residual capacity; FENO = fractional exhaled nitric oxide; Calv = alveolar nitric oxide concentration; PL-V = transpulmonary pressure-volume curve; BAL = bronchoalveolar lavage; TB = transbronchial; CT = computed tomography; QCT = quantitative computed tomography; PRM = parametric response mapping; Xe vent = Xenon ventilation; MRI = magnetic resonance imaging; V/P SPECT = ventilation-perfusion single-photon emission CT.
obstruction, in older subjects (≥ 65 y), and in former rather than current smokers.123

In a study of small airway pathology performed on lungs from subjects with COPD, multidetector-row CT revealed a decreased number of airways that were 2–2.5 mm in diameter and the loss of lumen starting from fifth-generation airways; accordingly, the assessment with micro-CT revealed decreases in the number of terminal bronchioles and the lumen, wall volumes, and alveolar attachments of pre-terminal bronchioles (TB-1) and pre-preterminal bronchioles (TB-2).124 Micro-CT and histology also noted increased B cell infiltration of TB-1 and TB-2 walls, which correlated with a decreased number of alveolar attachments. These findings suggest that changes to the small airway related to disease extends from 2-mm diameter airways to the terminal bronchioles, and that a B cell–mediated immune response may lead to the loss of alveolar attachments.125

In the last decade, ventilation-perfusion single-photon-emission CT (V/P SPECT) has been recognized as a powerful tool to assess small airway dysfunction and early disease and to evaluate the presence of ventilation defects and comorbidities in patients with COPD.126-128 Airway obstruction, previously representing a limitation to nuclear imaging studies with hydrosoluble aerosols in patients with COPD, no longer represents an issue since the introduction of ultrafine aerosols such as Technegas, thus allowing the expansion of V/P SPECT application, which exhibited a higher sensitivity than chest CT and FEV1 in detecting early airway changes.128 In fact, Technegas penetration depends on small airway function, and bronchial deposition increases in small airway disease, thus allowing detection of airway obstruction earlier with Technegas ventilation image than with spirometry.129 Recently, Bajc and colleagues127 reported the possibility of using this technique to distinguish between healthy subjects and those with COPD, not only to grade disease severity but also to add important information on concomitant comorbidities in subjects with COPD, such as pulmonary embolism and heart failure with vascular or ventilation defects. Although promising, the technique requires standardization and refinement in terms of spatial resolution before it can be used to provide clinically useful data on small airway dysfunction and early disease in at-risk patients with normal lung function tests.

Cytology of Induced Sputum

Sputum induction after inhalation of hypertonic saline is a noninvasive technique that is valid for studying inflammatory cells in central airways. Modified protocols have been proposed to evaluate whether this method can also assess inflammation in small airways. In particular, it has been suggested that sequential inductions, performed after short intervals of time (ie, 20–30 min), may provide useful information on distal airway inflammation.130-132 However, this procedure may be associated with issues regarding reproducibility and standardization, and no studies have been performed to directly compare the inflammatory profile in sputum with small airways assessment through transbronchial biopsies. A schematic representation of the currently available assessment tools for small airways in asthma and COPD are reported in Figure 7.

Treating Small Airways in Asthma and COPD

This section summarizes the rationale for the development of devices and drug formulations that target the small airways and describes the physics behind the concept of small-particle delivery. This section also presents the latest bench and clinical evidence in the treatment of small airways in subjects with asthma or COPD.

Delivery Systems Targeting Small Airways

Small airway inflammation and dysfunction characterize specific clinical asthma phenotypes, such as nocturnal asthma, exercise-induced asthma, and allergic asthma.7 The transversal involvement of distal airways in different obstructive diseases such as asthma and COPD have justified research efforts to create pharmacologic treatments and technologies that can reach and target the peripheral airways.

Most inhaled therapies, however, do not sufficiently reach the small airways.133-135 Although the drug particle size pattern may vary markedly between inhalers, conventional pressurized metered dose inhalers (pMDIs) and most dry powder inhalers (DPIs) emit drug particles that are too large to effectively target the small airways. In fact, a high proportion of the drug deposits in the oropharynx, causing local and systemic effects through gastrointestinal absorption.136

Particle size is a critical factor determining the therapeutic efficacy of inhaled drugs. The mass median aerodynamic diameter (MMAD) is used to measure the heterogeneity of particles in an emitted dose. MMAD predicts the deposition and distribution of particles in the bronchial tree. Using radiolabeled salbutamol, Usmani and colleagues137 noted an inverse relationship between particle size and lung deposition. Lower MMAD reduces oral deposition, thereby attenuating the consequences of an incorrect inhalation technique. However, the optimal particle size for an inhaled drug to effectively reach and deposit in the small airways has not clearly been defined, as reflected by the different terms and sizes: sub-micron particles (< 1 μm); ultrafine particles (1.1 μm), and extrafine particles (1.5 μm).138-140 Small particles are defined those with a MMAD < 2 μm, and inhaled formulations with MMAD < 2 μm have been termed extrafine.139
The proportion of particles within the aerosol that are < 5 \mu m is often referred to as the fine-particle fraction, or the fine particle dose if expressed in absolute mass of drug.\textsuperscript{138} Both the particle size and the fine-particle fraction of an aerosol play a significant role for the deposition of an inhaled drug and its relative distribution within the large and small airways.\textsuperscript{7} The geometric standard deviation (GSD) is defined as the ratio of the median diameter to the diameter at \pm 1 SD from the median diameter, and it measures the dispersion of the particle diameter.\textsuperscript{141} If the particle size varies over a wide range (ie, GSD > 1.2 \mu m), the aerosol is described as having a polydisperse particle distribution; if the particles are of similar size (ie, GSD < 1.2 \mu m), the aerosol particle distribution is described as monodisperse.\textsuperscript{141} It has been reported in lung deposition studies that small-particle aerosols achieve better pulmonary deposition as well as effective penetration into the peripheral lung. Importantly, small-particle aerosols are not exhaled to any significantly greater level compared to large-particle aerosols when assessed using in vivo lung deposition studies.\textsuperscript{7}

The available data suggest that the more distal deposition of small-particle formulations of inhaled corticosteroids (ICS) is safe in patients with asthma, and, for some drugs, distal deposition may even result in a reduced effect on the hypothalamic–pituitary–adrenal axis, most probably through less oropharyngeal deposition and hence decreased gastrointestinal bioavailability.\textsuperscript{142-145}

**Pharmacologic Targeting of Small Airways in Asthma**

Recent technological advances in device engineering and drug formulation have led to a new generation of inhalers emitting small-particle drug aerosol at slower velocities, with enhanced lung deposition and more effective aerosol penetration into peripheral lung. The novel devices include solution-based hydrofluoralkane-propelled pMDIs delivering ICS aerosols (eg, beclomethasone dipropionate, ciclesonide, and flunisolide), long-acting \beta\textsubscript{2}-adrenergic bronchodilators (LABA) (eg, formoterol), and an ICS/LABA fixed-dose combination (eg, beclomethasone dipropionate/formoterol). Furthermore, a new multi-dose DPI (NextHaler, Chiesi, Italy), which delivers a small-particle ICS/LABA combination, and the novel Soft-Mist inhaler (Boehringer Ingelheim, Germany), which delivers small-particle aerosols of long-acting bronchodilators, have been developed recently. The extrafine fixed-dose ICS/LABA (beclomethasone dipropionate/formoterol, 100/6 \mu g) is delivered using both a pMDI and a DPI. Extrafine beclomethasone dipropionate/formoterol achieves satisfactory central and peripheral lung deposition, leading to an increased potential for synergistic interaction and improved efficacy.\textsuperscript{146-148}

Increased resistances and inflammation-driven structural changes of peripheral airways in patients with asthma are involved in the lack of symptom control and enhanced exposure to sudden attacks across all disease severity stages.\textsuperscript{149-151} The development of extrafine formulations and the ability to reach the distal airways more effectively prioritized the pharmacologic management of small airways in asthma to reduce the future risk of exacerbations and side effects as well as to achieve better symptom control.\textsuperscript{4}

The significant increase in peripheral airway deposition of the inhaled drug with respect to the delivered dose provided by extrafine formulations allows for better equivalence ratios when compared with nonextrafine formulations, thus reducing drug exposure and potentially minimizing the efficacy/safety ratio with comparable clinical efficacy.\textsuperscript{145,152} Extrafine ICS formulations have been reported to be superior to nonextrafine ICS in improving functional and inflammatory parameters of small airways in subjects with asthma.\textsuperscript{89,109}

Papi and coworkers\textsuperscript{153} reported that the extrafine beclomethasone dipropionate/formoterol DPI formulation, assessed in subjects with moderate to severe asthma, was not inferior to a budesonide/formoterol DPI formulation in improving lung function and disease control, and with a comparable safety profile. This was also demonstrated when extrafine beclomethasone dipropionate/formoterol DPI was compared with salmeterol/fluticasone.\textsuperscript{154} In the latter case, subjects treated with beclomethasone dipropionate/formoterol achieved more rapid bronchodilation, with a significant reduction in air-trapping indexes, which is an indirect sign of improvement of small airway parameters.\textsuperscript{154} Real-life studies have also noted that a higher proportion of subjects treated with the extrafine beclomethasone dipropionate/formoterol pMDI achieved asthma control with a lower cumulative exposure to daily ICS dose compared with nonextrafine formulations.\textsuperscript{155,156}

More recently, Marti et al\textsuperscript{157} confirmed previous randomized reports showing significant improvements in asthma control in subjects treated with extrafine beclomethasone dipropionate/formoterol, independent of smoking status, disease duration, or the presence of air trapping. Subjects with and without indirect signs of air trapping experienced a mean FVC increase of 700 and 400 mL, respectively, confirming the effects of the study treatment on small airways.\textsuperscript{157} Contoli and colleagues\textsuperscript{158} noted significant improvements in phase III slope and R5-R20 values in smokers but not in never-smokers treated with beclomethasone dipropionate/formoterol. Conversely, Cox and co-workers\textsuperscript{159} showed no difference in the adenosine provocation dose and peripheral airway resistance when comparing an extrafine formulation of beclomethasone dipropionate with nonextrafine beclomethasone dipropionate and fluticasone pMDIs. The findings of the latter studies support the notion that both ICS alone or in combination with a LABA in an extrafine formulation can
result in real-life improvements in lung function and disease control, independent of disease severity or smoking status.160

Vos and colleagues161 used functional imaging methods to study the effect of extrafine beclomethasone dipropionate/formoterol in steroid-naïve, partially and well-controlled subjects with asthma. The investigators reported significant improvements in airway resistance and asthma control scores, along with traditional lung function parameters and FENO levels. Well-controlled subjects experienced an improvement in small airway volume even though traditional lung function parameters did not change significantly.161

The effects of bronchodilators on the SBN2 test have been extensively investigated in normal subjects and in subjects with asthma and COPD, but with contrasting results. No effects were observed in normal lungs,162-167 whereas in subjects with asthma, adrenergic agents have been shown to not only lower the slope of phase III but also to reduce the closing volume by increasing the vital capacity.168-173 Scichilone and colleagues174 compared the effects of extrafine beclomethasone dipropionate/formoterol on the SBN2 test and on bronchial hyper-reactivity with the effects on nonextrafine salmeterol/fluticasone in 30 subjects with asthma. The investigators reported that only beclomethasone dipropionate/formoterol achieved a significant improvement in the cumulative methacholine provocation dose and a trend toward a reduction in closing volume and in air trapping indexes and closing volume and to improve regional lung ventilation in patients with moderate to severe asthma, either alone or when associated with salmeterol/fluticasone.178,179 Additionally, 24 weeks of treatment with montelukast reduced alveolar inflammation estimated with peripheral exhaled NO and reactance at low frequencies compared with placebo in subjects with stable asthma with background ICS treatment.180

In subjects with mild asthma, adding montelukast to fluticasone 250 μg for 3 weeks reduced FENO by 60% compared to baseline (P < .001, P = .08 for alveolar NO).181

Oral pharmacologic treatments have shown contrasting effects on small airways function in patients with asthma. The oral methylxanthine theophylline has long been used for chronic asthma treatment.177 However, the literature to support its effects on small airways in subjects with asthma is lacking. Conversely, the antileukotriene montelukast has been reported to reduce air-trapping indexes and closing volume and to improve regional lung ventilation in patients with moderate to severe asthma, either alone or when associated with salmeterol/fluticasone.178,179

**Pharmacologic Targeting of Small Airways in COPD**

Small airways represent a major site of interest in patients with COPD. In fact, the peripheral lung constitutes the initial site of airway damage and a challenging diagnostic and pharmacologic target due to its peculiar anatomy and physiology, as well as the current limited ability to identify early disease. A body of literature supports the idea that available bronchodilators can improve the pathophysiological parameters that reflect small airway damage. Indeed, simple spirometry has a limited sensitivity both in detecting small airway disease and in assessing the modifications induced by inhaled therapy.3,70,86,182

R5-R20 and X5 measured with impulse oscillometry in subjects with COPD have been associated with worse airway obstruction, hyperinflation, and worse symptom burden. In fact, the degree of volume response in terms of FVC has been associated with the impairment of peripheral airway resistance.182 To date, anticholinergics and β2-agonists have exhibited comparable effects on several lung function parameters in patients with COPD. In acute conditions, patients with moderate to severe air-flow obstruction experience a significant reduction in inspiratory capacity, RV, and specific R5 when randomly exposed to tiotropium bromide or indacaterol.70 Moreover, post-bronchodilator changes in specific R5 were more closely related to bronchodilator-induced improvements of lung mechanics and dyspnea, making the reduction of specific R5 a better criterion to study the acute functional effect of bronchodilators than the traditional criteria based on FEV1 and FVC.70

Small airway disease is not only associated with increases in static volumes and specific R5, but also with regional ventilation heterogeneity, which can be assessed with the SBN2 test or the multiple-breath nitrogen washout test. Indexes of ventilation heterogeneity are correlated with exercise capacity, exercise-induced desaturation, and
The acute administration of tiotropium and indacaterol produced similar improvements in phase III slope, reduced the extent of airway closure, and increased PaO2 levels in subjects with moderate and severe COPD. Similar results were reported for aclidinium bromide and glycopyrronium.

Compared with nonextrafine formulations, the intrapulmonary deposition of extrafine particles studied with the gamma scintigraphy technique, multi-slice CT scans, and computational fluid dynamics in subjects with moderate and severe COPD exhibited better regional lung deposition as well as significant improvements in static lung volumes and a reduction in extrathoracic drug deposition. This observation supports other results indicating that exposure to lower doses of ICS with the administration of an extrafine formulation of beclomethasone dipropionate yielded comparable results in preventing exacerbations and better odds of treatment stability at 2 y compared with nonextrafine fluticasone in subjects with moderate to very severe air-flow obstruction. In subjects with COPD, the administration of extrafine beclomethasone dipropionate/formoterol significantly improved FVC, air trapping, dyspnea and quality of life.

More recently, the extrafine triple combination therapy of beclomethasone, formoterol and glycopyrronium have been studied in three large randomized clinical trials – that is, TRILOGY, TRINITY, and TRIBUTE. In all trials, when coarse particles formulations were compared with the extrafine triple-combination therapy, the latter showed a statistically significant reduction in moderate and severe exacerbation risk.

Some studies have investigated the effect of theophylline on small airway function in subjects with COPD. In COPD subjects with nonreversible airflow obstruction, the administration of theophylline improved air trapping and airway resistance, with results comparable to those for salbutamol.

**Clinical Implications and Future Developments**

The role of small airways in COPD and asthma has suffered from a paradox in the last century. In the 1950s and 1960s, we witnessed the rapid development of tests able to assess small airway disease, thus enabling the early detection of peripheral lung dysfunction in patients at risk for chronic respiratory disease. This rapid evolution was slowed by the poor reliability and repeatability of some of these tests, which were due mainly to the limited technologization of such procedures. At that time, the idea that smoking habit was to be abandoned due to its undeniable negative effects on health and society, may have led to the progressive loss of interest in small airway dysfunction in COPD, but also conditioned research. The subsequent development of new molecules and delivery systems made the research on small airways appealing again. Radiomorphological findings from the COPD-gene and CANCOLD research projects have added the imaging piece to the work conducted first by Weibel and later by researchers Harvard University and McGill University. Landmark papers that were recently published have allowed us to link morpho-functional data to the natural history of COPD and to highlight the importance of early pharmacologic intervention. Moreover, recent studies conducted on pathophysiological parameters such as closing volume and ventilation heterogeneity by means of the SBN2 test and new interpretations of body plethysmography should help integrate these biomarkers of small airway dysfunction into the monitoring process of the pharmacologic response to therapy and the follow-up of patients with both asthma and COPD.

Finally, the standardization of exhaled biomarkers such as NO has added an important tool to the everyday clinical practice of respiratory physicians involved in the management of patients with asthma. In patients with COPD, in addition to a classic inflammatory pathway, this evidence has modeled the idea of a new role of small airways in the natural history of chronic obstructive diseases. The mechanical stress produced by the small airway damage in emphysema may mechanically influence surrounding areas of normal lung, thus increasing the risk of lung function decline. However, no large clinical trial has yet reported the effectiveness of COPD treatments in slowing disease progression. The potential role of treatments able to reach the lung periphery and increase lung deposition may become a valid option in patients with small airway involvement in asthma and with early signs of COPD. In asthma, the concept of “future risk” has gained a paramount importance in the management of the disease, as emphasized by recent guidelines (https://ginasthma.org, accessed May 24, 2019). Whether the detection of small airway abnormalities will identify patients at major risk of asthma exacerbations or more severe disease, or identifies those likely to benefit from a specific intervention, is unknown and should be pursued in randomized clinical trials.

**Summary**

Asthma and COPD are both characterized by small airway involvement and dysfunction. In most cases, the abnormalities of the peripheral lung can grow to a disproportional degree before the damage is identified and treated, thus limiting the effects of the pharmacologic intervention. Reliable and simple tests are now available, and are consistent both in terms of sensitivity and repeatability for the identification of small airway disease, making the so-called silent zone much more noisy. Although advanced and intriguing, data on the structure...
and function of small airways require clinical standardi-
zation and confirmatory results. Indeed, more effort
should be devoted to the global functional assessment of
patients at risk and patients with mild disease to achieve
the goal of changing the natural history of asthma and COPD.

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