Chronic obstructive pulmonary disease: inhale deeply and start to exercise

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Dual bronchodilation results in better lung deflation but similar exercise capacity compared to LAMA monotherapy http://ow.ly/7zL1309M6LK

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Chronic obstructive pulmonary disease (COPD) is defined by partially reversible airflow limitation as a result of either small airway obstruction or loss of elastic recoil due to emphysematous lung destruction, or both [1]. Airflow limitation resulting from small airway obstruction is treated with bronchodilators. Resulting improvements in airflow limitation are in turn quantitatively assessed by measuring the forced expiratory volume in 1 s (FEV1), which is the most widely used spirometric measure in COPD. In fact, most of our knowledge about the magnitude of improvements from bronchodilator therapy comes from this standard measurement of lung function. So we know that the simultaneous inhalation of long-acting muscarinic antagonists and long-acting beta-2 agonists from one inhaler once or twice daily results in improvements in FEV1 that are superior to the inhalation of the individual components [2–5] and clearly exceed the threshold for clinical importance of ~100 mL [6].

However, lessons learned from respiratory physiology clearly indicate that it is more important to measure the volumes of air that remain in the lungs of patients with COPD rather than to measure the volume that can be exhaled during a forced breathing manoeuvre [7]. These functional consequences of COPD are called lung hyperinflation and air trapping, i.e. an increase in the functional residual capacity and residual volume and decrease in the inspiratory capacity [7]. Lung hyperinflation has been shown to be closely related to exertional dyspnoea, exercise intolerance, physical inactivity and mortality [7]. Hence, pulmonologists began measuring the improvements in lung hyperinflation by bronchodilator therapy and were able to demonstrate that bronchodilators are also effective lung deflators, as they improve functional residual capacity, residual volume and inspiratory capacity [8, 9]. Furthermore, these improvements in lung hyperinflation lead to improved exercise tolerance, exertional dyspnoea and physical activity in daily life [8–11].

Given the strong link between lung hyperinflation and resulting exercise intolerance, the question arises whether better lung deflation will physiologically translate into better exercise tolerance. In this regard, the pooled MORACTO studies published in the current issue of the European Respiratory Journal by O’DONNELL et al. [12] provide an impressively large dataset to study in detail the effects of fixed dual bronchodilator therapy compared to monobronchodilator therapy on exercise intolerance. Besides the strong statistical power to analyse the primary end-points, this dataset permits the analysis of specific subgroups that might benefit from dual bronchodilator compared to monotherapy. Being asked, as a pulmonologist, whether it would be relevant for our patients to have an increase in inspiratory capacity of 0.254 L, which is 0.100 L better than the improvements observed by monotherapy, it is tempting to answer “yes”. Now we see the results of the MORACTO studies, which on average do not show a statistically
significant additional benefit of maximised lung deflation due to the simultaneous inhalation of the approved doses of tiotropium/olodaterol on exercise endurance time compared to tiotropium monotherapy. However, our clinical impression that better lung deflation is conducive to better exercise tolerance is at least partly confirmed by the MORACTO data, as exercise endurance time is numerically better with the approved doses of tiotropium/olodaterol than with monotherapy. Accordingly, numerically more patients experienced improvement in exercise endurance time, as shown by additional analyses summarised in the supplementary material of the article. So how should we interpret the data? First, we need to remember that the MORACTO studies are placebo-controlled cross-over trials with long wash-out periods without any long-acting bronchodilator therapy. Thus, highly symptomatic patients in need of effective lung deflation are likely to be underrepresented in these studies. Secondly, muscle deconditioning is highly prevalent in COPD [13]. To make use of the improved respiratory breathing physiology, exercise training is needed to further improve the exercise endurance time. This approach is subject to a different study protocol combining pharmacological and nonpharmacological treatments [14].

References


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