Beta-blockers in COPD: time for reappraisal

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ABSTRACT The combined effects on the heart of smoking and hypoxaemia may contribute to an increased cardiovascular burden in chronic obstructive pulmonary disease (COPD). The use of beta-blockers in COPD has been proposed because of their known cardioprotective effects as well as reducing heart rate and improving systolic function. Despite the proven cardiac benefits of beta-blockers post-myocardial infarction and in heart failure they remain underused due to concerns regarding potential bronchoconstriction, even with cardioselective drugs. Initiating treatment with beta-blockers requires dose titration and monitoring over a period of weeks, and beta-blockers may be less well tolerated in older patients with COPD who have other comorbidities. Medium-term prospective placebo-controlled safety studies in COPD are warranted to reassure prescribers regarding the pulmonary and cardiac tolerability of beta-blockers as well as evaluating their potential interaction with concomitant inhaled long-acting bronchodilator therapy. Several retrospective observational studies have shown impressive reductions in mortality and exacerbations conferred by beta-blockers in COPD. However, this requires confirmation from long-term prospective placebo-controlled randomised controlled trials. The real challenge is to establish whether beta-blockers confer benefits on mortality and exacerbations in all patients with COPD, including those with silent cardiovascular disease where the situation is less clear.

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Introduction

Chronic obstructive pulmonary disease (COPD) is one of the world’s leading causes of morbidity and is now the third leading cause of mortality, amounting to 3 million deaths in 2010 [1, 2]. Exacerbations in particular account for up to three-quarters of the total costs due to COPD [3], with attributable costs exceeding USD 30 billion [4]. A recent COPD task force statement identified an unmet need in terms of finding drugs to treat common comorbidities specifically mentioning the putative effects of beta-blockers on the cardiovascular burden and its associated impact on mortality [5]. Cardiovascular comorbidity is common in patients with COPD due to smoking in addition to other shared risks including genetic susceptibility, systemic inflammation and ageing [6]. The prevalence of COPD in patients with heart failure ranges from 11% to 52% in North American patients and from 9% to 41% in European patients [7]. The purpose of this article is to critically reappraise current knowledge regarding beta-blockers in COPD, looking at the current evidence for their therapeutic index and how this relates to management guidelines.

We have not attempted a systematic review or meta-analysis as described elsewhere [8–10], but rather highlight the key areas of clinical relevance for physicians who treat patients with COPD. In this article we have: 1) considered the putative link between COPD and the heart in terms of potential targets for beta-blockers; 2) reviewed retrospective data linking the use of beta-blockers to reduced exacerbations and mortality; 3) examined the unmet need for use of beta-blockers in patients with COPD and both known, and potentially unknown, cardiovascular disease; 4) evaluated which beta-blocker to use based on their pharmacology and impact on pulmonary function; and 5) attempted to draw conclusions about the current clinical use of beta-blockers in COPD.

COPD and the heart

The main accepted clinical indications for the use of beta-blockers in COPD are for patients post-myocardial infarction and for patients with heart failure. However, the presence of untreated or unrecognised (i.e. silent) cardiovascular disease may contribute to mortality in COPD and may also be an underlying causative factor in exacerbations, which can be difficult to separate from respiratory aetiologies (figure 1 and box 1) [6, 7]. It is also possible, if not likely, that the burden of cardiovascular disease may be underrated by pulmonologists when treating COPD patients because symptoms are presumed to be primarily driven by airflow obstruction, especially during exacerbations.

The prevalence of left ventricular systolic dysfunction ranges between 10% and 46% in patients with COPD, and although the occurrence of heart failure with preserved left ventricular ejection fraction is less clear, estimates in patients with severe COPD are as high as 90% [7]. The benefits of beta-blockers in...
patients with heart failure due to left ventricular systolic dysfunction are well established from pivotal trials as well as meta-analysis [11–14]. The challenge in COPD may be more with respect to diagnosis of heart failure with echocardiography, where image acquisition is difficult due to lung hyperinflation [15].

Beta-blockers only have proven benefits in patients post-myocardial infarction but not in stable coronary arterial disease [16, 17]. Nevertheless, the presence of coronary calcium on chest computed tomography scans is associated with mortality in COPD [18], and known coronary arterial disease is also associated with longer exacerbations, more dyspnoea, and lower health status and exercise capacity in stable patients with COPD [19]. There is also an acute increase in arterial stiffness, particularly during infective exacerbations of COPD, along with increases in cardiac enzymes especially in patients with coronary arterial disease [20]; one particular study found that one in 12 patients admitted to hospital with an exacerbation of COPD met the criteria for a myocardial infarction [21]. The presence of coronary heart disease in COPD, along with the adverse effects of hypoxaemia [22], may be compounded by the positive chronotropic effects of concomitant inhaled beta-agonist therapy [23, 24], further compromising cardiac reserve. It has been shown that even a low dose of a beta-1 selective antagonist such as atenolol might protect against chronotropic, inotropic and electrocardiographic effects of inhaled beta-agonists, which are mediated by cardiac beta-2 receptor stimulation [25].

Another potential target is diastolic dysfunction, although a meta-analysis suggests that the beneficial effects of beta-blockers in such patients are less clear cut [26]. Several factors may contribute to the occurrence of impaired diastolic function in COPD. First, patients with COPD also appear to have a higher left ventricular mass (hypertrophy) even in the absence of left ventricular dilatation, which impacts upon survival [27]. Secondly, lung hyperinflation in COPD may cause cardiac compression reducing both left ventricular and atrial filling even in the absence of raised pulmonary arterial pressure [28–30]. These factors may also be compounded by the negative effects of hypoxaemia on diastolic filling [22, 31].

In addition to these COPD-related risks, patients with the disease commonly have other comorbidities such as coronary artery disease, hypertension and diabetes, which can all adversely affect diastolic function. This was addressed in a recent prospective longitudinal study of healthy young adults followed over 25 years, where a fall in the ratio of forced expiratory volume in 1 s (FEV1) to forced vital capacity (FVC) was associated with reduced left atrial size and cardiac output [32]. Left ventricular end diastolic and end systolic wall stress measured by magnetic resonance imaging is associated with increasing severity of airflow obstruction in patients with COPD and coexistent heart failure [33]. Impaired left ventricular filling is clinically important because it can eventually produce left atrial enlargement, which is a key risk factor for atrial fibrillation and associated mortality during exacerbations of COPD [34]. Furthermore, the presence of impaired diastolic filling in patients with COPD is also related to impaired walking distance [35]. Thus, the absence of benefits of beta-blockers in diastolic dysfunction may not apply in COPD and deserves re-evaluation in this patient group.

**Effects of beta-blockers on mortality and exacerbations**

Due to the high cardiovascular comorbidity in COPD from smoking along with increased sympathetic drive due to hypoxaemia [36], beta-blockers have been proposed as a cogent therapeutic intervention for...
their known cardioprotective effects in addition to reducing heart rate and improving systolic and diastolic dysfunction. One of the fundamental issues with regards to more widespread use of beta-blockers in COPD is the concern regarding beta-2 receptor antagonism and associated airway smooth muscle constriction, which may even occur with cardioselective agents that exhibit preferential beta-1 blockade, especially in more susceptible severe patients with impaired respiratory reserve. The risk-benefit equation in COPD becomes more favourable for patients who already have overt cardiac disease such as heart failure or post-myocardial infarction, where beta-blockers have proven protective effects [11, 16]. There are, however, no data as to the putative beneficial effects of beta-blockers in those COPD patients who may have concomitant silent coronary arterial disease or heart failure.

Retrospective observational data have shown beneficial effects of beta-blockers in a cohort of 5977 patients with COPD who were followed for a mean of 4.35 years [37], where their use was associated with an overall 22% (95% CI 8–33%) reduction in mortality. In a study of 825 patients admitted to hospital for an exacerbation of COPD, beta-blocker use among 142 patients was associated with a 61% (95% CI 1–86%) reduction in mortality [38]. RUTTEN et al. [39] showed 32% (95% CI 17–44%) and 29% (95% CI 17–40%) reductions in mortality and exacerbations, respectively, conferred by taking beta-blockers among 2230 patients with COPD followed up for a mean of 7.2 years. In a cohort study from Sweden of 4858 patients with COPD, those who were discharged on a beta-blocker (84%) post-myocardial infarction had 13% (95% CI 2–22%) lower mortality [40]. In a retrospective report of 256 patients with COPD who had either coronary heart disease or heart failure, 58% were taking beta-blockers associated with a 73% (95% CI 50–85%) reduction in the likelihood of being admitted to a hospital emergency room [41]. In contrast, in an observational study using time dependent analysis of 2249 severe oxygen-dependent COPD patients there was a 19% increase in mortality associated with taking beta-blockers [42]. However, in a prospectively followed cohort of 3464 patients with COPD who were followed for a mean of 7.2 years, BHATT et al. [43] found a 27% (95% CI 10–40%) reduction in total exacerbations, while in Global Initiative for Chronic Obstructive Lung Disease grade 3/4 patients on home oxygen there was a 67% reduction (95% CI 42–81%).

In a 2012 meta-analysis of nine retrospective cohort studies, the pooled estimate for mortality reduction with beta-blockers was reported to be 31% (95% CI 22–38%) [8]. In a subsequent 2014 meta-analysis of 15 retrospective studies of 21 596 patients with COPD, the pooled estimate for reduction in overall mortality conferred by beta-blockers was 28% (95% CI 17–37%) and for exacerbations was 38% (95% CI 18–58%) [9]. The reduction in mortality was 36% (95% CI 24–46%) among the subgroup of patients (five studies; 39% weighting) with known coronary heart disease and 26% (95% CI 7–42%) in the subgroup with known heart failure (three studies; 18% weighting).

The beneficial effects of beta-blockers on exacerbations may involve other potential noncardiac mechanisms whereby beta-blockers could reduce COPD exacerbations [44, 45]. In heart failure, use of cardioselective beta-blockers reduces systemic inflammatory cytokine release such as interleukin-6 and alters leukocyte distribution, which may also impact inflammation during respiratory infections [46]. Beta-blockers have also been reported to inhibit neutrophil chemotaxis and oxygen free radical production [47], while in human endothelial cells they have been reported to reduce the release of endothelin-1, a bronchoconstrictor peptide implicated in the pathogenesis of COPD exacerbations [48, 49].

It is not possible to eliminate the possibility of residual confounding in the observational studies suggesting beta-blockers may reduce exacerbations and mortality in COPD and thus definitive randomised trials are needed. There is now a planned placebo-controlled trial powered for a reduction in exacerbations using metoprolol over 1 year via the US COPD Clinical Research Network and funded by the Department of Defense (Clinicaltrials.gov identifier: NCT02587351).

This study will exclude those patients with an absolute indication for beta-blockers including an myocardial infarction or revascularisation procedure within 3 years or with an ejection fraction <40%. However, it remains possible that this and similar studies may run the risk of only including patients where beta-blockers are less efficacious.

The unmet cardiac need in COPD

Beta-blockers have an established position in the management of coronary artery disease while heart failure guidelines reinforce their use in patients with concomitant COPD [50]. Similarly, COPD management strategies also state that the benefits of selective beta-1 blocker treatment in heart failure clearly outweigh any potential risk associated with treatment even in patients with severe COPD [51]. Despite this guidance there is a reluctance to prescribe even cardioselective beta-blockers in COPD, even in the presence of known cardiac disease, because of persistent concerns regarding potential bronchoconstriction, especially in more severe patients. In a cohort from Scotland we found that only 14% of patients with COPD were taking beta-blockers for cardiovascular comorbidity [37]. Further evidence of a reluctance to prescribe beta-blockers
in COPD was documented by Quint et al. [52] where 55% of patients who had a myocardial infarction were not prescribed a beta-blocker, with only 22% being prescribed on admission. In the UK 64% of patients without COPD and acute coronary syndrome were prescribed beta-blockers as compared with 16% of similar patients with COPD who were prescribed beta-blockers [53]. Furthermore COPD was documented as a reason for withholding beta-blockers in 33% of patients who did not receive a beta-blocker, while noncardiologists were 40% less likely to prescribe beta-blockers. In the USA, Chen et al. [54] found that elderly patients after an acute myocardial infarction were 62% less likely to be given beta-blockers in the presence of a history of treated COPD or asthma. Initiating treatment with beta-blockers is not simple as it requires dose titration over a period of weeks along with monitoring of heart rate, blood pressure and perhaps spirometry, all of which take time, incurring extra healthcare costs. Moreover beta-blockers may be less well tolerated in older patients with coexisting comorbidities such as diabetes, peripheral vascular disease and renal impairment, who are more prone to postural hypotension.

Choice of beta-blocker and effects on pulmonary function

The mechanism of beta-blocker induced bronchoconstriction is thought to be due to the effects of pre- and post-junctional beta-2 receptor antagonism uncovering the prevailing cholinergic tone via post-junctional smooth muscle muscarinic type 3 receptors, resulting in airway smooth muscle constriction [55].

In a subgroup analysis of 2712 patients from a cohort who had serial spirometry measures over 4 years, there was no deleterious effect of long-term beta-blocker use (88% were cardioselective) on either FEV1 or FVC, even among the more severe patients taking triple inhaled therapy, who had the greatest reductions in exacerbations and mortality [37]. In a meta-analysis of randomised controlled trials with cardioselective beta-blockers there was no significant change in FEV1 compared with placebo, when given either as single −2.1% (95% CI −6.1–2.0%) or chronic dosing −2.6% (95% CI −5.9–0.8%), and also no significant effect on the FEV1 response to beta-2-agonists [10]. In a randomised controlled trial of 27 patients with heart failure who also had coexistent moderate-to-severe COPD, after 4 months of treatment there was a 190 mL significant fall in FEV1 between bisoprolol and placebo, while salbutamol reversibility, symptoms and quality of life were unchanged [56]. In a comparison of bisoprolol and placebo in patients with moderate-to-severe COPD, there was a significantly worsening of dynamic hyperinflation during cycle endurance while exercise duration was unaltered [57]. In a study comparing 24 COPD patients on beta-blockers matched to patients not taking beta-blockers there was no difference in exercise capacity or gas exchange despite lower heart rate and blood pressure, in turn suggesting great oxygen delivery per heart beat [58].

The beta-blockers currently licensed for heart failure are the beta1 selective bisoprolol, nebivolol, metoprolol and the non-selective carvedilol (box 2). As has already been shown in heart failure [59] and asthma [60] it is important to slowly titrate up the dose of beta-blocker to improve cardiovascular and pulmonary tolerability. Bisoprolol has a licensed indication for use in heart failure and coronary artery disease and has a beta-1/2 receptor selectivity ratio of 14:1, which is higher than either atenolol (5:1) or metoprolol (2:1) [61]. In a cross-over study of 51 patients with COPD and heart failure, directly comparing 6 weeks of bisoprolol, metoprolol and carvedilol [62], FEV1 was lowest with carvedilol and highest with bisoprolol with metoprolol in between. In a randomised controlled trial comparing bisoprolol (mean dose 6.4 mg) and carvedilol (mean dose 47 mg) in patients with heart failure and COPD, FEV1 significantly improved by 137 mL with bisoprolol, but not with carvedilol (30 mL improvement) [63]. In 15 mild-to-moderate COPD patients there

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**BOX 2** Prescribing of beta-blockers in chronic obstructive pulmonary disease for cardiovascular disease

- Beta-1 selective antagonists including metoprolol, bisoprolol and nebivolol exhibit dose related beta-2 receptor blockade
- Carvedilol is a nonselective beta-antagonist that is more likely to cause bronchoconstriction than beta-1 selective antagonists
- Slowly titrate the dose of beta-blockers at 1–2 weekly intervals up to the usual maintenance dose
- Monitor supine and erect blood pressure, heart rate and spirometry during dose titration
- Concomitant long-acting muscarinic antagonists may obviate potential bronchoconstriction
- Symptomatic bradycardia may occur if beta-blockers are used with other rate-limiting drugs such as calcium blockers (e.g. verapamil and diltiazem), ivabradine or anti-arrhythmic agents (e.g. digoxin, amiodarone and flecainide)
- Symptomatic hypotension may occur when beta-blockers are used with other vasodilatory drugs (e.g. angiotensin converting enzyme inhibitors, angiotensin receptor blockers, calcium channel blockers and alpha receptor blockers)
was a significant worsening in airway hyperreactivity to methacholine challenge with metoprolol and propranolol, but not nebivolol compared with placebo, while the acute bronchodilator response to fenoterol was only blunted by propranolol [64].

Nebivolol has been shown to exhibit greater in vitro beta-1/2 receptor selectivity than bisoprolol in human myocardium [65] and also suppresses endothelial nitric oxide [66]. In healthy volunteers attenuation of beta-2 receptor mediated terbutaline-induced hypokalaemia was significantly greater with bisoprolol 10 mg or atenolol 50 mg/100 mg versus nebivolol 5 mg, which in turn was not different from placebo [67]. Nebivolol produced significant blunting of terbutaline-induced glucose and insulin responses compared with placebo in keeping with beta-2 receptor antagonism at the 5 mg dose. However, the relative beta-1/2 selectivity cannot be inferred since this would require comparison of beta-blocker doses that exhibit the same degree of beta-1 antagonism as assessed by exercise heart rate reduction [68], which was not measured.

In a post hoc analysis of 2670 patients from the Organized Program to Initiate Lifesaving Treatment in Hospitalized Patients with Heart Failure (OPTIMIZE-HF), there were no differences between selective and non-selective beta-blockers in terms of lower mortality or re-hospitalisation in patients with and without COPD [69]. Carvedilol blocks cardiac beta-1 and beta-2 receptors as well as exhibiting peripheral vasodilatation due to alpha receptor blockade, which in addition to its antioxidant activity [70] may explain its superiority versus metoprolol in heart failure in one particular study, which may not have compared comparable doses [59]. Until there is more convincing evidence to support the superiority of carvedilol in heart failure, it would be prudent to choose a selective agent such as bisoprolol, nebivolol or metoprolol due to their superior safety profile in COPD.

Long-acting muscarinic antagonists such as tiotropium have been shown to obviate bronchoconstriction even when using nonselective beta-blockade with propranolol in asthmatic patients [71]. It is the more severe COPD patients who would, in theory, be most at risk of beta-blocker induced bronchoconstriction. These patients would usually already be taking concomitant long-acting muscarinic antagonists and hence be protected from bronchospasm. The relatively small degree of dose-related beta-2 receptor antagonism conferred, for example, by bisoprolol [72] would not be expected to result in worsening of pulmonary function. It is also important to consider the potential impact of beta-2 receptor genotype on the risk–benefit equation for beta-blockers in COPD. It has been shown that asthmatic patients who possess one or two copies of the arginine-16 beta-2 receptor polymorphism are more prone to propranolol-induced bronchoconstriction in terms of FEV1 and airway resistance [73]. While the arginine-16 polymorphism conferred a worse outcome on survival in patients receiving metoprolol after an acute coronary syndrome [74], it was not associated with survival in heart failure patients treated with metoprolol or carvedilol [75].

Conclusions and the way forward
There are compelling reasons to use cardioselective beta-blockers in patients with COPD who have coexistent heart failure or are post-myocardial infarction (box 3). Current evidence would suggest that

**BOX 3** Key messages

- Cardiovascular comorbidity, including coronary artery disease and heart failure, commonly coexists in chronic obstructive pulmonary disease (COPD) due to the effects of smoking, systemic inflammation, hypoxaemia and other shared risks.
- COPD may also be associated with impaired diastolic filling due to lung hyperinflation, which may be compounded by the negative lusitropic effects of hypoxaemia and left ventricular hypertrophy.
- The main indications for beta-blockers in patients with COPD are post-myocardial infarction and heart failure with reduced ejection fraction. Despite clear evidence beta-blockers improve outcomes in these COPD patients they remain significantly underused due to concerns about adverse respiratory effects, even with beta-1 selective antagonists.
- Meta-analyses of retrospective studies with beta-blockers in COPD have shown pooled estimates for reductions in mortality of 28% and exacerbations of 38%.
- Initiating treatment with beta-blockers requires careful dose titration and monitoring. This may be particularly relevant for patients with COPD who are often older and have other comorbidities that increase the risk of intolerance.
- Beta-1 selective antagonists such as bisoprolol, nebivolol and metoprolol are preferred to the nonselective carvedilol as they are less likely to produce bronchoconstriction in COPD.
- Long-acting muscarinic antagonists, which are commonly used in COPD, protect against the potential for bronchoconstriction due to dose related beta-2 receptor antagonism.
- The key unanswered question is whether beta-blockers may confer benefits on mortality and exacerbations in all patients with COPD including those with silent cardiovascular disease.
there remains a reticence to prescribe beta-blockers in such patients because of a fear of adverse events, particularly worsened lung function. Further prospective medium-term safety studies are therefore required to carefully follow the effects of cardioselective drugs on pulmonary function in patients with more severe COPD by employing slow initial dose titration as well as evaluating their interaction with long-acting bronchodilators (Clinicaltrials.gov identifier: NCT01656005).

There is currently not sufficient evidence at present to advocate treatment with beta-blockers for the prevention of exacerbations or exacerbation-related mortality. Long-term placebo-controlled multicentre trials in COPD are indicated to confirm the benefits of beta-blockers already seen on mortality and exacerbations in observational studies. The key question to answer is whether the potential benefits of beta-blockers are confined to those patients with known cardiovascular disease or are present in the wider population who may have silent cardiovascular disease. Likewise, beta-blockers are not currently indicated in COPD patients with diastolic dysfunction alone where controlled trials are also warranted.

Beta-blockers are likely to be part of a more complex therapeutic jigsaw in addressing the composite risk from different cardiovascular abnormalities in COPD, and as has already been shown with heart failure there may be additive effects from drugs acting on other neuro-hormonal pathways. This includes drugs which block the renin-angiotensin system that may be particularly effective at regressing left ventricular hypertrophy [76]. Dual angiotensin/neprilysin inhibition may also confer benefits by augmenting brain natriuretic peptide levels [77] and ameliorating the adverse effects of hypoxic pulmonary vasoconstriction [78, 79]. Anti-platelet drugs might also be beneficial for treating silent coronary artery disease in more severe COPD patients who are oxygen dependent [42]. Pulmonologists have tended to focus on drugs which act on the lung rather than the heart, because of the evidence supporting the former. Perhaps now is time to look at the lungs’ next door neighbour in the chest and begin to address the unmet need of cardiac disease in COPD.

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