Risk of myocardial infarction (MI) and death following MI in people with chronic obstructive pulmonary disease (COPD): a systematic review and meta-analysis

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ABSTRACT

Objectives: Cardiovascular disease is an important comorbidity in patients with chronic obstructive pulmonary disease (COPD). We aimed to systematically review the evidence for: (1) risk of myocardial infarction (MI) in people with COPD; (2) risk of MI associated with acute exacerbation of COPD (AECOPD); (3) risk of death after MI in people with COPD.

Design: Systematic review and meta-analysis.

Methods: MEDLINE, EMBASE and SCI were searched up to January 2015. Two reviewers screened abstracts and full text records, extracted data and assessed studies for risk of bias. We used the generic inverse variance method to pool effect estimates, where possible. Evidence was synthesised in a narrative review where meta-analysis was not possible.

Results: Searches yielded 8362 records, and 24 observational studies were included. Meta-analysis showed increased risk of MI associated with COPD (HR 1.72, 95% CI 1.22 to 2.42) for cohort analyses, but not in case-control studies: OR 1.18 (0.80 to 1.76). Both included studies that investigated the risk of MI associated with AECOPD found an increased risk of MI after AECOPD (incidence rate ratios, IRR 2.27, 1.10 to 4.70, and IRR 13.04, 1.71 to 99.7). Meta-analysis showed weak evidence for increased risk of death for patients with COPD in hospital after MI (OR 1.13, 0.97 to 1.31). However, meta-analysis showed an increased risk of death after MI for patients with COPD during follow-up (HR 1.26, 1.13 to 1.40).

Conclusions: There is good evidence that COPD is associated with increased risk of MI; however, it is unclear to what extent this association is due to smoking status. There is some evidence that the risk of MI is higher during AECOPD than stable periods. There is poor evidence that COPD is associated with increased in hospital mortality after an MI, and good evidence that longer term mortality is higher for patients with COPD after an MI.

INTRODUCTION

Cardiovascular disease is a common comorbidity and cause of death in people with chronic obstructive pulmonary disease (COPD), with up to one-third dying of cardiovascular disease. Reducing the cardiovascular disease in this population is an important strategy for reducing the burden of COPD.

Several studies have shown that people with COPD have a higher risk of myocardial infarction (MI) than people without COPD. One of the reasons for the increased risk of MI in patients with COPD is the shared major risk factor of smoking. In addition, several other cardiovascular risk factors are increased in COPD in addition to smoking status. There is some evidence that the risk of MI associated with increased comorbidities in patients with chronic obstructive pulmonary disease.
factors, including hypertension, diabetes, inactivity, poor diet, and older age, are also prevalent in patients with COPD.\textsuperscript{3-7} In addition, several studies have found an association between reduced FEV\textsubscript{1} (forced expiratory volume 1) and cardiovascular mortality in the general population.\textsuperscript{8} However, COPD itself is also thought to be an independent risk factor for MI with increased risk of MI possibly being mediated through increased systemic inflammation or reduced FEV\textsubscript{1} in people with COPD.

Acute exacerbations of COPD are events in the natural history of COPD which are characterised by an increase in COPD symptoms such as breathlessness, cough, sputum volume, and sputum purulence. It has recently been suggested that acute exacerbations of COPD (AECOPD) represent a period of increased risk of MI for people with COPD.\textsuperscript{9} A subtype of patients with COPD appears to have more frequent exacerbations than others. Frequent exacerbators have been defined as individuals who have two or more treated exacerbations per year. Frequent exacerbators may be at higher risk of MI compared to infrequent exacerbators, even during stable periods.

Several investigators have found that patients with COPD have worse mortality in hospital and following discharge after an MI compared to patient without COPD.\textsuperscript{10-12} However, the finding that patients with COPD have greater in hospital and short-term mortality has not been found by all investigators.\textsuperscript{13-15}

We aimed to systematically review the literature reporting on: (1) the risk of MI in people with COPD; (2) the risk of MI associated with AECOPD, either during AECOPD or that associated with frequent exacerbator phenotype; and (3) the risk of death after MI in people with COPD. These questions represent the most salient aspects of current research into the relationship between COPD and cardiovascular disease, and no systematic reviews have been published on these topics to date.

METHODS

Literature search

MEDLINE, EMBASE, BIOSIS & Science Citation Index were searched up to January 2015. A search strategy was devised which would pick up articles relevant to all three research questions. All strategies were based on the MEDLINE search strategy, which is presented in the online supplementary material. In brief, the literature was searched for terms which relate to COPD and terms with relate to MI, and these searches were combined using the AND Boolean logic operator. MeSH terms were combined with natural language searching using truncation where appropriate.

Inclusion and exclusion criteria

Inclusion and exclusion criteria were applied for each of the three research questions. Studies were included if they met the population, exposure, comparator and outcome criteria. These are presented below for each research question. Studies were included from database start date and were not restricted by language.

Risk of MI in people with COPD

The population of interest was the general population. The exposure of interest was diagnosis of COPD. The un-exposed group was people without a diagnosis of COPD. The outcome of interest was acute MI.

Risk of MI associated with AECOPD

The population of interest was people with a diagnosis of COPD. The exposures of interest were either: (1) discrete episodes of AECOPD or periods within 8 weeks of an AECOPD; or (2) frequent exacerbator phenotype. The comparators of interest were either: (1) periods of stable COPD; or (2) infrequent exacerbator phenotype. Studies were included if these reported a relative risk of MI, or if this could be calculated.

Risk of death after MI in people with COPD

The population of interest was those presenting to a hospital with an MI. Studies were included if these compared patients with a diagnosis of COPD to those without a diagnosis of COPD. Outcomes of interest were death in hospital and at any reported time points post discharge. Studies investigating risk of death for patients with COPD after an interventional procedure following an MI (such as percutaneous coronary intervention or coronary artery bypass graft) were specifically excluded under the population criterion.

Selection of included studies

Titles and abstracts, where available, were initially screened for potential inclusion by one reviewer. Full text versions of potentially included studies were then obtained and were screened by two reviewers. Authors were contacted if the information provided in articles was not sufficient to assess whether inclusion criteria were satisfied.

Risk of bias assessment

All included studies, except for those only reported as conference abstracts, were assessed for risk of bias. The risk of bias tool was informed by the Newcastle-Ottawa scale;\textsuperscript{16} however, we did not make use of a summary score as this is not advisable.\textsuperscript{17, 18} Risk of bias was assessed across the key domains related to selection of participants, comparability of groups, and measurement of outcomes. Several items were included under each domain, and were adapted for different study types. Where reports of studies included more than one analysis (eg, a case control as well as a cohort analysis) the risk of bias for these analyses were conducted separately. Risk of bias assessment was completed by one reviewer and checked by another.
Evidence synthesis

Characteristics and findings of studies were tabulated and compared. Data on severity of COPD was extracted as GOLD stage or FEV$_1$% predicted, where available. Information was also extracted on smoking status and previous cardiovascular disease. Estimates of effect were extracted or calculated, and are presented as ORs, risk ratios (RR), incidence rate ratios (IRR) or HRs.

Where included studies were reasonably statistically and clinically similar, we pooled results using random effects meta-analysis. We used the generic inverse variance method to pool maximally adjusted effect estimates. Analysis was conducted in Review Manager 5.3. Where studies were too statistically (I$^2$ over 75%) or clinically heterogeneous, meta-analysis was not conducted, but study summary results were graphed on forest plots without pooling the results. Studies which were not adjusted at all were not included in forest plots. For the question on risk of MI associated with COPD, studies were stratified by adjustment for smoking status (yes or no) and study design (cohort or case-control). For the question on risk of death following MI in COPD compared to patient without COPD, studies were stratified by outcome time point (in-hospital mortality or follow-up mortality). For follow-up mortality, studies were further stratified by analysis.

RESULTS

Identified studies

Literature searches yielded 8362 records. After title and abstract screening, 49 records were selected for full-text assessment, which resulted in the inclusion of 24 studies. The inclusion and exclusion process is summarised in figure 1. Of the 24 included studies, 9 investigated the risk of MI in patients with COPD compared to patient without COPD; 2 investigated the risk of MI associated with AECOPD; no studies were found which investigated the risk of MI associated with the frequent exacerbator phenotype; and 12 investigated outcomes after MI for patients with COPD compared to patient without COPD.

Summary characteristics of included studies are presented in tables 1–3.

All of the included studies which investigated risk of MI in people with COPD used data from either routine clinical or administrative databases. COPD was defined using diagnostic codes; these varied between COPD diagnosis in primary care, outpatient departments, hospital admission or discharge codes, and cause of death codes. Three studies also required that patients with COPD had been prescribed COPD medicines. One of the studies, Rodriguez et al. included only patients with a recent diagnosis of COPD and followed up to 5 years after this diagnosis to identify MI. Only one study reported a summary of COPD severity, and only two reported prevalence of current smokers. Four studies reported a cohort analysis only. Two studies reported a cohort analysis as well as a case-control analysis. One study reported the results of a cohort analysis and an analysis of period prevalence. One study compared rates of MI in patients with COPD to standardised populations rates of MI.

Two studies were identified which investigated the risk of MI associated with AECOPD. Both studies defined risk periods after the onset of AECOPD and used within person designs to compare the risk to a baseline period.

Nine studies reported mortality for patients with COPD after an MI compared to patient without COPD. Five studies reported a comparison of in-hospital mortality after an MI between patients with COPD and patient without COPD. Eight studies used a time-to-event analysis to investigate death after discharge from a hospital admission for MI.

Risk of bias assessment

The proportion of studies (or analyses, where appropriate) which were assessed as having either lower, unclear or higher risk of bias for each of the research questions is presented in figure 2. Detailed results from the risk of bias assessment for individual studies are presented in the appendix.

Risk of MI in people with COPD

Of nine included studies, eight found a higher risk of MI in patients with COPD compared to patient without COPD. Six studies estimated the ratio of incidence rates of MI in patients with COPD compared to patient without COPD. Five studies estimated this for all MIs, this ranged from IRR 1.18 (95% CI 0.81 to 1.71) to 5.31 (4.54 to 6.21). One study estimated the IRR for hospitalisation due to MI (IRR 1.49, 95% CI 0.71 to 3.13) and fatal MIs (1.51, 1.14 to 2.01). Two studies estimated the ratio of hazard of MI in patients with COPD compared to patient without COPD one study...
## Table 1. Characteristics of included studies – risk of MI associated with COPD

<table>
<thead>
<tr>
<th>Study</th>
<th>Design and setting</th>
<th>Population</th>
<th>Characteristics of COPD patients</th>
<th>MI definition</th>
<th>Maximally adjusted estimate (95% CI)</th>
<th>Factors adjusted for</th>
</tr>
</thead>
<tbody>
<tr>
<td>Curkendall et al²⁰</td>
<td>Cohort in the Saskatchewan Health databases 1998–2001</td>
<td>11 493 COPD patients ≥40 years, identified by physician claim or hospital discharge COPD code and at least two prescriptions for COPD medicines within 6 months of the index COPD code. 22 986 age and sex matched non-COPD patients</td>
<td>Age, sex, COPD severity, current smokers</td>
<td>Any MI during follow up: any inpatient or outpatient diagnosis of MI Hospitalisation due to MI: IRR 1.49 (0.71–3.13) Fatal MI: IRR 1.51 (1.14–2.01)</td>
<td>Age, sex, history of cardiovascular disease, diabetes, hypertension, hypercholesterolaemia</td>
<td>Period prevalence of MI: Age, sex, history of cardiovascular events, diabetes, hypertension, hypercholesterolaemia Hospitalisation for MI: adjusted for history of cardiovascular events, diabetes, hypertension, and hypercholesterolaemia using Poisson regression, age and sex by matching. Fatal MI: age and sex by matching only.</td>
</tr>
<tr>
<td>Feary et al⁸</td>
<td>Cohort in The Health Improvement Network, 2005–2007</td>
<td>29 870 COPD patients &gt;35 years identified by COPD diagnostic code. 1 174 240 non-COPD patients</td>
<td>Age, sex, COPD severity, current smokers</td>
<td>Diagnostic code for MI in primary care record</td>
<td>35–44 years: HR 10.34 (3.28–32.6) 45–54 years: HR 1.22 (0.55–2.74) 55–64 years: HR 1.55 (1.07–2.26) 65–74 years: HR 1.78 (1.37–2.31) ≥75 years: 1.34 (1.03–1.73)</td>
<td>Age, sex and smoking status</td>
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<thead>
<tr>
<th>Study</th>
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<tr>
<td>Huiart <em>et al</em></td>
<td>Cohort in the Saskatchewan Health databases, 1990–1999</td>
<td>5,648 COPD patients ≥50 years, identified by prescription of three or more bronchodilators within the period of one year. Rates of MI compared to those of general Saskatchewan population</td>
<td>Age NR Sex NR COPD severity NR Current smokers NR History of CVD NR</td>
<td>Primary hospital discharge diagnosis of MI</td>
<td>Standardised IRR: 1.30 (1.15–1.44)</td>
<td>Age and sex by standardisation</td>
</tr>
<tr>
<td>Mapel <em>et al</em></td>
<td>Cohort in the Veterans Administration Medical System, 1991–1999</td>
<td>COPD patients identified by discharge codes (1991–1999) and/or outpatient codes (1997–1999) Age and sex matched controls without COPD</td>
<td>Age Median 60 (IQR, 49–62) Sex 95.7% male COPD severity NR Current smokers NR History of CVD Cardiovascular disease—71.2%</td>
<td>Specific ICD-9-CM code for MI during 1999 that was not present in 1998</td>
<td>COPD patients identified using discharge codes IRR: 1.28 (1.18–1.38) COPD patients identified using outpatient codes IRR: 5.31 (4.54–6.21)</td>
<td>Age and sex by matching</td>
</tr>
<tr>
<td>Rodriguez <em>et al</em></td>
<td>Cohort and case-control study in the General Practice Research Database, 1996–2001</td>
<td>1,532 patients with a first COPD diagnosis in 1996, and no history of cardiovascular disease 13,500 age and sex matched non-COPD patients, with no history of cardiovascular disease</td>
<td>Age NR Sex NR COPD severity NR Current smokers NR History of CVD NR</td>
<td>Diagnostic code for MI in primary care record</td>
<td>Cohort analysis: IRR 1.18 (0.81–1.71) Case-control analysis: OR 0.93 (0.62–1.39)</td>
<td>Age and sex Case control analysis: Age, sex, smoking and number of primary care physician visits</td>
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<tr>
<td>Schneider et al</td>
<td>Cohort and nested case-control study in the General Practice Research Database, 1995–2005</td>
<td>35 772 patients with a first COPD diagnosis between 1995–2005 35 772 non-COPD patients matched on age, sex and calendar time and general practice</td>
<td>Age 40–49–6.8% 50–59–19.9% 60–69–33.8% &gt;70–39.8%</td>
<td>Diagnostic code for MI along with death or hospitalisation within 30 days of the diagnosis; and/or start of new treatment with ACE antagonist, β-blocker, statin, vitamin K antagonist, platelet aggregation inhibitor or aspirin within 90 days of the diagnosis in primary care record</td>
<td>Cohort analysis: IRR 1.56 (1.43–1.75) Case control analysis: Any COPD: OR 1.40 (1.13–1.73) Mild COPD: OR 1.79 (1.12–2.86) Moderate COPD: OR 1.30 (1.04–1.62) Severe COPD: OR 3.00 (1.53–5.86)</td>
<td>Cohort analysis: matched on age, sex, calendar time and general practice Case-control analysis: Smoking status, BMI, hypertension, hyperlipidaemia, diabetes and NSAID use</td>
</tr>
<tr>
<td>Sidney et al</td>
<td>Cohort in health insurance database. North Carolina, 1996–1999</td>
<td>COPD defined as: hospitalisation or outpatient diagnosis of COPD, two or more prescriptions for COPD medicines, aged over 40 years. Non-COPD patients matched on age, sex and length of care plan membership.</td>
<td>Age 40–59–35% 60–79–55% &gt;80–10%</td>
<td>ICD code for acute MI</td>
<td>Overall: IRR 1.89 (1.71–2.09) Men: IRR 1.77 (1.56–2.01) Women: IRR 2.09 (1.78–2.46) 40–64 years: IRR 2.43 (1.98–2.98) ≥65 years: IRR 1.73 (1.54–1.94)</td>
<td>Age, sex and baseline cardiovascular risk profile.</td>
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<tr>
<td>Sode et al</td>
<td>Cohort study within the National Danish patient registry, 1980–2006</td>
<td>Entire Danish population. COPD identified through hospital admission codes or COPD as cause of death</td>
<td>Age &lt;30–7% 30–59–54% 60–79–35% &gt;80–3%</td>
<td>Discharge diagnosis of MI or cause of death from Danish Causes of Death Registry listed as MI</td>
<td>HR 1.26 (1.25–1.27)</td>
<td>Age, sex, Danish ancestry, geographical residency (rurality), and level of education</td>
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<tr>
<td>Study</td>
<td>Design and setting</td>
<td>Population</td>
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<td>Yin et al</td>
<td>Cohort of all residents of Sweden aged over 18, July 2005-December 2008.</td>
<td>51,348 COPD patients identified by diagnostic codes from patient records. 6,743,342 non-COPD patients.</td>
<td>Sex 55% male COPD severity NR Current smokers NR History of CVD NR</td>
<td>Those with no previous MI or stroke Age Mean 71.1 Sex 44.3% male COPD severity NR Current smokers NR History of CVD NR</td>
<td>Diagnostic code for MI, or primary cause of death listed as MI</td>
<td>No previous MI or stroke: HR 1.47 (1.41–1.55)* Previous MI: HR 1.33 (1.23–1.43)*</td>
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*Data from personal communication (Magnus Back. Email communication. 18/08/2014).

COPD, chronic obstructive pulmonary disease; CVD, cardiovascular disease; FEV1, forced expiratory volume in 1 s; ICD, International Classification of Diseases; IRR, incidence rate ratios; MI, myocardial infarction.
Table 2  Characteristics of included studies

<table>
<thead>
<tr>
<th>Study</th>
<th>Characteristics</th>
<th>Population</th>
<th>AECOPD definition</th>
<th>MI definition</th>
<th>Risk periods</th>
<th>Risk estimate (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Donaldson et al.</td>
<td>3 512 Patients with COPD</td>
<td>COPD defined using Quality and Outcomes Framework, 2003</td>
<td>Increase in or new onset of more than one of: cough, sputum, sputum purulence, wheezing or dyspnoea; lasting three or more days and requiring treatment with an antibiotic or oral steroid.</td>
<td>MI defined as age ≥ 40 years, smoking history ≥10 pack-years, FEV1 &lt;70% predicted, and FEV1/FVC &lt;50%</td>
<td>30 days after AECOPD</td>
<td>IRR 2.27 1.10 to 4.70</td>
</tr>
<tr>
<td>Feary et al.</td>
<td>490 patients with COPD and AECOPD</td>
<td>COPD severity: GOLD stage I 24%, GOLD stage II 43%, GOLD stage III 46%, GOLD stage IV 9%</td>
<td>Median FEV1 &lt;70% of predicted</td>
<td>MI defined as age ≥ 18 years, smoking or non-smoking history, and FEV1 &lt;50% predicted</td>
<td>1-5 days</td>
<td>IRR 1.18 1.05 to 1.31</td>
</tr>
<tr>
<td>Schneider et al.</td>
<td>150 patients with COPD with a primary outcome of reduction in FEV1</td>
<td>MI defined as age ≥ 40 years, smoking or non-smoking history, and FEV1 &lt;50% predicted</td>
<td>MI defined as age ≥ 45 years, smoking or non-smoking history, and FEV1 &lt;50% predicted</td>
<td>1-5 days</td>
<td>IRR 2.09 1.14 to 3.85</td>
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</table>

Risk of MI associated with AECOPD

Donaldson et al. conducted a self-controlled case series using data from The Health Improvement Network (THIN). They used prescription of antibiotics and steroids in patient with COPD to identify AECOPD, and report an increased risk of MI in the 1-5 days following the onset of AECOPD (IRR 2.27, 95% CI 1.10 to 4.70). No difference in the risk of MI was found for the period 6-49 days, or at any time point when the alternative definitions of AECOPD of prescription of steroids alone or antibiotics alone were used. Halpin et al. reported a secondary analysis of the UPLIFT trial, which was an RCT comparing inhaled tiotropium and placebo in patients with COPD with a primary outcome of reduction in FEV1 decline. Time to first AECOPD was a secondary outcome. AECOPD were identified using a symptom-based definition and were reported to trial staff at

estimated this to be HR 1.26 95%, CI 1.25 to 1.27, while the other study estimated this to be HR 1.47 (1.41 to 1.55) for those with no previous MI, and HR 1.33 (1.25 to 1.43) for those with a previous MI. One study estimated the ratio of odds of period prevalence over 5 years of acute MI in patients with COPD compared to patient without COPD (OR 1.61, 95% CI 1.43 to 1.81). Only one of the included cohort studies compared risk of MI in people with COPD and people without COPD adjusted for smoking status. This study reported results stratified by age groups. Meta-analysis of these results showed an increased risk of MI for people with COPD (HR 1.72, 95% CI 1.22 to 2.42) (figure 3). Two of the included case-control studies adjusted for smoking status. Meta-analysis of these results did not show an increased risk of MI for people with COPD (OR 1.18, 95% CI 0.80 to 1.76) (figure 4). Meta-analysis was not conducted for the studies which did not adjust for smoking as heterogeneity was too high (I²=93%). These results are graphically summarised in figure 5.

Some studies investigated whether the effect of COPD on the risk of MI was different in terms of age and severity of airflow obstruction. Feary et al. found that the effect of COPD on risk of MI was higher in the 35–44 year age group (HR 10.34, 95% CI 3.28 to 32.6) compared to older age groups (45–54 years: HR 1.22 (95% CI 0.55–2.74), 55–64 years: HR 1.55 (95% CI 1.07 to 2.26), 65–74 years: HR 1.78 (95% CI 1.37 to 2.31), ≥75 years: HR 1.34 (95% CI 1.03 to 1.73)). Sidney et al. reported similar findings; the effect of COPD on risk of MI was higher in those who were aged 40–64 years (HR 2.43, 95% CI 1.98 to 2.98) compared to those who were aged over 64 years (HR 1.73, 95% CI 1.54 to 1.94). Schneider et al. investigated the risk of MI by sub-group of COPD severity. They found that the effect of COPD on the risk of MI was greater in those with severe COPD (OR 3.00, 95% CI 1.53 to 5.86) compared to those with moderate COPD (OR 1.30, 95% CI 1.04 to 1.62) or mild COPD (OR 1.79, 95% CI 1.12 to 2.86).
<table>
<thead>
<tr>
<th>Study</th>
<th>Design and setting</th>
<th>Population</th>
<th>COPD patient characteristics</th>
<th>Maximally adjusted estimate for mortality (95% CI)</th>
<th>Factors adjusted for</th>
</tr>
</thead>
<tbody>
<tr>
<td>Andell et al&lt;sup&gt;10&lt;/sup&gt;</td>
<td>Cohort study within the Swedish SWEDHEART registry between 2005–2010.</td>
<td>Consecutive patients admitted to Swedish coronary care units. COPD diagnosis ascertained through linkage to the Swedish National Patient Registry.</td>
<td>Age: Mean 75 years (SD, 9)</td>
<td>Mortality at one year: HR 1.14 (1.07–1.21)</td>
<td>Age, sex, smoking, comorbidity (previous MI, previous stroke, heart failure, renal failure, hypertension, diabetes, peripheral artery disease, cancer and previous bleeding), in hospital treatment and discharge medications (heparin, fondaparinux, dalteparin, enoxaparin, glycoprotein IIb/IIa inhibitors, angioplasty, coronary stenting, β-blockers, aspirin, clopidogrel, prasugrel, calcium channel blockers, digoxin, diuretics, statins, nitrates and warfarin).</td>
</tr>
<tr>
<td>Behar et al&lt;sup&gt;22&lt;/sup&gt;</td>
<td>Cohort study in Israel between 1981–1983</td>
<td>2276 consecutive patients surviving an MI after admission to 13 coronary care units. Patients with a history of chronic bronchitis or chronic airways obstruction and clinical and/or radiographic findings compatible with COPD during hospitalisation for MI were included.</td>
<td>Age: Mean 66.8 years (SD, 9.7)</td>
<td>Unadjusted:*  In–hospital RR 1.39 (1.16–1.67)</td>
<td>1 year RR 1.34 (1.16–1.55) 5 years RR 1.28 (1.18–1.40)</td>
</tr>
<tr>
<td>Bursi et al&lt;sup&gt;23&lt;/sup&gt;</td>
<td>Cohort study of the population in the Rochester Epidemiology project involving residents in Olmsted County, Minnesota from 1979 to 2007</td>
<td>Local residents in Olmsted County. MI ascertained from medical records compatible with ICD criteria. Information on COPD was also obtained from ICD codes.</td>
<td>Age: Mean 73 years (SD, 11)</td>
<td>HR 1.30 (1.10 to 1.54), mean follow up 4.7 years.</td>
<td>Age, sex, smoking, hypertension, MI type (STEMI/non-STEMI), creatine kinase level,Killip class, reperfusion treatment in hospital, use of drugs on discharge (β-blockers, ACEI, diuretics)</td>
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</table>

*Unadjusted:*
## Table 3. Continued

<table>
<thead>
<tr>
<th>Study</th>
<th>Design and setting</th>
<th>Population</th>
<th>COPD patient characteristics</th>
<th>Maximally adjusted estimate for mortality (95% CI)</th>
<th>Factors adjusted for</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dziewierz et al 24</td>
<td>Cohort study within Krakow Registry of ACS in February 2005-March 2005 and December 2005-January 2006</td>
<td>1414 patients with MI admitted to hospital in Krakow, Poland. Those with a previous history of COPD and current treatment with a steroid or bronchodilator were classified as COPD patients.</td>
<td>Age Mean 71.8 years (SD, 11)</td>
<td>Sex 62% male COPD severity NR Current smokers 40.7% History of CVD MI 34.6% Angina 80.2% HF 30.9%</td>
<td>HR 2.15 (1.30–3.55) Age, sex, BMI, diabetes, hypertension, hyperlipidaemia, prior angina, prior MI, prior heart failure, left ventricular ejection fraction, prior PCI, prior CABG, prior stroke or transient ischaemic attack, smoking status, peripheral arterial disease, chronic renal insufficiency, parameters on admission (chest pain, cardiogenic shock, heart rate, systolic blood pressure, diastolic blood pressure), time from chest pain onset to admission and type of MI (STEMI or NSTEMI)</td>
</tr>
<tr>
<td>Enriquez et al 11</td>
<td>Cross sectional study of National Cardiovascular Data Registry in the USA between January 2008 and December 2010</td>
<td>158 890 patients admitted to one of 445 sites with an MI. COPD patients had a history of COPD or were using long term inhaled or oral β-agonists, inhaled anti-inflammatory agents, leukotriene receptor antagonists or inhaled steroids.</td>
<td>Age STEMI—median 66 years nSTEMI—median 70 years</td>
<td>Sex STEMI—60.4% male nSTEMI—57.5% male COPD severity NR Current smokers STEMI—57.0% nSTEMI—41.9% History of CVD STEMI Prior MI—29.7% Prior CHF—15.3% nSTEMI Prior MI—39% Prior CHF—33.3%</td>
<td>In-hospital mortality STEMI OR 1.05 (0.95–1.17) Non-STEMI OR 1.21 (1.11–1.33) Age, serum creatinine, systolic blood pressure, troponin elevation, heart failure or cardiogenic shock at presentation, ST-segment changes, heart rate and prior peripheral arterial disease.</td>
</tr>
<tr>
<td>Study</td>
<td>Design and setting</td>
<td>Population</td>
<td>COPD patient characteristics</td>
<td>Maximally adjusted estimate for mortality (95% CI)</td>
<td>Factors adjusted for</td>
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<tr>
<td>Hadi <em>et al</em>&lt;sup&gt;15&lt;/sup&gt;</td>
<td>Cross sectional study of patients hospitalised with ACS in May 2006 and January 2007 to June 2007 in six Middle Eastern countries</td>
<td>8169 consecutive patients in the Gulf RACE registry presenting with ACS at 65 centres across six countries. COPD patients were identified from 1) medical records or 2) use of COPD medicines.</td>
<td><strong>Age</strong>&lt;br&gt;Median 64 (IQR, 56–71)&lt;br&gt;<strong>Sex</strong>&lt;br&gt;NR&lt;br&gt;<strong>COPD severity</strong>&lt;br&gt;NR&lt;br&gt;<strong>Current smokers</strong>&lt;br&gt;38.7%&lt;br&gt;<strong>History of CVD</strong>&lt;br&gt;Prior MI—34.8%&lt;br&gt;Prior angina – 54.4%</td>
<td><em>In hospital mortality:</em>&lt;br&gt;OR 0.40 (0.20–1.24)</td>
<td>Age, sex, cardiogenic shock, use of thrombolysis, use of aspirin, use of β-blocker, use of ACEi</td>
</tr>
<tr>
<td>Hawkins <em>et al</em>&lt;sup&gt;25&lt;/sup&gt;</td>
<td>Cohort study of patients with acute MI enrolled in VALIANT trial</td>
<td>Patients with MI complicated by LVSD and HF. COPD was identified by a questionnaire completed by trial site investigators.</td>
<td><strong>Age</strong>&lt;br&gt;Mean 68.1 (SD, 9.9)&lt;br&gt;<strong>Sex</strong>&lt;br&gt;71.1% male&lt;br&gt;<strong>COPD severity</strong>&lt;br&gt;NR&lt;br&gt;<strong>Current smokers</strong>&lt;br&gt;42.0%&lt;br&gt;<strong>History of CVD</strong>&lt;br&gt;Prior MI—39.9%&lt;br&gt;Prior angina—46.1%&lt;br&gt;Prior HF—27.3%</td>
<td>HR 1.14 (1.02–1.28)</td>
<td>Age, heart rate, systolic and diastolic blood pressure, weight, baseline creatinine, smoking status, diabetes, dyslipidaemia, hypertension, killip classification, anterior MI, new lower bundle branch block, thrombolytic therapy, primary PCI, coronary artery bypass graft, history of heart failure, atrial fibrillation, previous MI, angina, previous stroke, peripheral arterial disease, renal insufficiency, alcohol abuse, country of enrolment, beta blocker use, randomised treatment</td>
</tr>
<tr>
<td>Kjoller <em>et al</em>&lt;sup&gt;13&lt;/sup&gt;</td>
<td>Cohort study of consecutive patients recruited 1–6 days after an MI</td>
<td>Danish hospitals between May 1990 and July 1992 as part of TRACE study. COPD was identified using either 1) medical records or 2) patient report in addition to use of COPD medicines.</td>
<td><strong>Age</strong>&lt;br&gt;Median 70.5 (5–95 percentiles, 50.7–83.5)&lt;br&gt;<strong>Sex</strong>&lt;br&gt;68.2% men&lt;br&gt;<strong>COPD severity</strong>&lt;br&gt;NR&lt;br&gt;<strong>Current smokers</strong>&lt;br&gt;60.0%&lt;br&gt;<strong>History of CVD</strong>&lt;br&gt;Prior MI—25.1%</td>
<td><strong>Cohort entry to 30 days:</strong>&lt;br&gt;HR 0.89 (0.68–1.11)&lt;br&gt;<strong>Cohort entry to 7 years:</strong>&lt;br&gt;HR 1.15 (1.04–1.28)</td>
<td>Age, sex, BMI, hypertension, diabetes, smoking status, previous angina, wall motion index, angina, history of CHF, new CHF, atrial fibrillation, bundle branch block, wall motion index, use of thrombolytic therapy</td>
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<tr>
<td>Study</td>
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<td>Quint et al(^{26}) (abstract)</td>
<td>Cohort study of patients admitted after a first MI using data from the UK CALIBER database</td>
<td>8,065 patients admitted to UK hospitals with a first MI between Jan 2003-Dec 2008. COPD was identified using primary care records.</td>
<td>Previous angina—43.9, Previous CHF—28.2%</td>
<td>Mortality up to 7 years: HR 1.37 (1.23–1.52)</td>
<td>Age and sex</td>
</tr>
<tr>
<td>Raposeiras et al(^{13}) (abstract)</td>
<td>Cross sectional and cohort study of patients with ACS</td>
<td>4,497 consecutive patients admitted to Spanish hospitals for ACS. The ascertainment method for COPD was unclear.</td>
<td>Age, NR, Sex, NR, COPD severity NR, Current smokers NR, History of CVD NR</td>
<td>In-hospital death OR 1.04 (1.03–1.04)</td>
<td>GRACE score, β-blocker therapy</td>
</tr>
<tr>
<td>Rha et al(^{33}) (abstract)</td>
<td>Case control study in Korea AMI registry from 2005 to 2007</td>
<td>AMI patients in KAMIR</td>
<td>Age, Mean 71.7 (SD 10.0), Sex, NR, COPD severity NR, Current smokers NR, History of CVD NR</td>
<td>Mortality at 8 months OR 2.69, 95% CI could not be calculated from reported information.</td>
<td>Unadjusted</td>
</tr>
<tr>
<td>Salisbury et al(^{27})</td>
<td>19 centre prospective study of patients presenting with MI in a cohort study</td>
<td>MI patients in PREMIER study restricted to patients discharged alive after MI. Patients were considered to have COPD if they had a documented history of</td>
<td>Mortality up to 1 year HR 2.00 (1.44–2.79)</td>
<td>Age, gender, race, avoidance of health care due to cost, smoking, diabetes, hypertension, CHF, ejection fraction, previous CVD, MI diagnosis type, new onset HF after</td>
<td>Unadjusted</td>
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</table>
### Study Design and Setting

Patients hospitalised with AMI in greater Worcester, Massachusetts in medical centres. COPD patients were identified by previous mention of clinical or radiographic evidence for COPD in their medical record.

### COPD Patient Characteristics

<table>
<thead>
<tr>
<th>Study</th>
<th>Design and setting</th>
<th>Population (COPD or asthma) or had therapy specific for obstructive pulmonary disease.</th>
<th>COPD severity</th>
<th>Factors adjusted for</th>
</tr>
</thead>
<tbody>
<tr>
<td>Stefan et al.</td>
<td>Cross sectional study with follow up of patients hospitalised with AMI at greater Worcester, Massachusetts between 1997-2007</td>
<td>NR</td>
<td>NR</td>
<td>Smoking status used in secondary analysis</td>
</tr>
</tbody>
</table>

### Maximaly adjusted estimate for mortality

<table>
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<th>Study</th>
<th>Design and setting</th>
<th>Population (COPD or asthma) or had therapy specific for obstructive pulmonary disease.</th>
<th>COPD severity</th>
<th>Factors adjusted for</th>
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<td></td>
<td></td>
<td>NR</td>
<td>NR</td>
<td>Smoking status used in secondary analysis</td>
</tr>
</tbody>
</table>

### Factors adjusted for

- In hospital: OR 1.25 (0.97 – 1.34)
- 30 day mortality: OR 1.31 (1.0 – 1.58)
- Age, sex, year of hospitalisation, history of CVD, history of renal failure, type of MI (STEMI/non-STEMI), length of stay, smoking status used in secondary analysis, indicators of the centre, treatment type

### Table 3. Continued

<table>
<thead>
<tr>
<th>Study</th>
<th>Design and setting</th>
<th>Population (COPD or asthma) or had therapy specific for obstructive pulmonary disease.</th>
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<tr>
<td></td>
<td></td>
<td>NR</td>
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</tbody>
</table>

**Note:** Calculated from reported data. LVD, left ventricular systolic dysfunction; MeSH, Medical Subject Headings; MI, myocardial infarction; NR, not reported; PCI, percutaneous coronary intervention; NSTEMI, non-ST-segment elevation myocardial infarction; STEMI, ST-segment elevation myocardial infarction.
regular study visits. Data on MI were collected as serious adverse events. This study found that compared to the 30 days prior to AECOPD, risk of MI in the 30 days following AECOPD was increased (IRR 13.04; 95% CI 1.71 to 99.7). These results are graphically summarised in figure 6. Owing to different exposure time periods, the results for within person studies investigating the risk of MI associated with AECOPD were not pooled in meta-analysis.

Risk of death after MI in people with COPD
Of the studies investigating differences in in-hospital mortality after an MI, two found an increased risk of mortality for patients with COPD (RR 1.39, 95% CI 1.16 to 1.67 (unadjusted); and OR 1.04, 95% CI 1.03 to 1.04). Two studies did not find evidence for increased in-hospital mortality for patients with COPD (OR 0.40, 95% CI 0.20 to 1.24; OR 1.25, 95% CI 0.97 to 1.34). One study reported results split by type of MI: it

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**Figure 2** Summary of risk of bias for risk assessments for: A studies investigating risk of MI associated with COPD; B studies investigating risk of MI associated with AECOPD; and C studies investigating risk of death following MI in people with COPD. AECOPD, acute exacerbation of COPD; COPD, chronic obstructive pulmonary disease; MI, myocardial infarction.

**Figure 3** Forest plot showing risk of MI associated with COPD in cohort studies which adjusted for smoking status. CIs may vary slightly from those quoted in tables due to transformation during meta-analysis. COPD, chronic obstructive pulmonary disease; MI, myocardial infarction.
did not find an increased in-hospital mortality for patients with COPD after a STEMI (OR 1.05, 95% CI 0.95 to 1.17), but did find increased in-hospital death for patients with COPD after a non-STEMI (OR 1.21, 95% CI 1.11 to 1.33). Meta-analysis of adjusted results showed weak evidence for an increased risk of in-hospital death for patients with COPD (OR 1.13, 95% CI 0.97 to 1.31) (figure 7).

One study reported mortality at 30 days for patients with COPD compared to patient without COPD. This study found increased mortality for patients with COPD (OR 1.31, 1.10 to 1.58). Another study reported mortality at 8 months, and in an unadjusted analysis, found increased mortality for COPD compared to patients without COPD (OR 2.69, 95% CI was not reported and could not be calculated). One study also found, on unadjusted analysis, that mortality was greater for patients with COPD at 1 year (RR 1.34, 95% CI 1.16 to 1.55) and 5 years (RR 1.28, 95% CI 1.18 to 1.40) after MI.

Eight studies reported results of survival analysis of mortality during follow-up after an MI. All of the studies reported higher mortality for patients with COPD compared to patients without COPD during follow-up after discharge following an MI. HRs ranged from 1.15 (95% CI 1.04 to 1.28) to 2.15 (95% CI 1.30 to 3.55). However, one of these studies found no evidence of a difference in mortality when restricting the time period to the first 30 days following discharge (HR 0.89, 95% CI 0.68 to 1.11). Meta-analysis of studies which reported adjusted results showed an increased risk of death after discharge following MI for patients with COPD compared to patients without COPD (HR 1.26, 1.13 to 1.40) (figure 8). Four of the studies included under this question were excluded from meta-analysis for methodological or clinical heterogeneity.

**DISCUSSION**

**Main findings**

Most studies which investigated the risk of MI in people with COPD found that those with COPD have higher risk of MI than people who do not have COPD; however,
it is unclear how much of this increased risk is due to smoking status. The included cohort study which adjusted for smoking status showed an increased risk of MI in people with COPD, but this was not apparent in pooled analysis of the case–control studies which adjusted for smoking status. Both of the included studies which investigated the risk of MI associated with AECOPD found an increased risk of MI in the weeks following AECOPD. Most studies which investigated mortality after an MI for patients with COPD as compared to patients without COPD found that mortality after discharge was greater for those with COPD, and an increased risk of death was found on pooled analysis. However, findings on in-hospital mortality after an MI were mixed, and there was only weak evidence for increased risk of death in hospital for patients with COPD on pooled analysis.

**Limitations of included studies and future work**

One common limitation among the included studies, particularly those which investigated the risk of MI associated with COPD, was missing information on smoking status. As smoking is very strongly associated with COPD and risk of MI, it is likely to be a major confounder in all studies investigating this association. All of the studies in this review which investigated this association used either clinical or administrative routine data sources. Routine data are a potentially rich source of information about huge numbers of patients. However, data on smoking are not routinely recorded in all administrative databases. Indeed, all of those studies which did not have data for smoking in this question used administrative databases. Future studies on the association between COPD and cardiovascular disease should use data sources which contain reliable information on smoking status.

Further studies should be carried out to confirm findings that AECOPD are periods of increased risk of MI for people with COPD. These studies should ensure they use validated exposure measures and are adequately powered. Possible reasons for an increased risk of MI during AECOPD include increased inflammation and the potential cardiovascular effects of the drugs used to treat AECOPD. If indeed the finding of increased risk during AECOPD is confirmed, future studies should attempt to disentangle the reasons for increased risk of MI. In addition, studies should investigate factors which might modify this relationship, such as drugs used for treatment of COPD and cardiovascular prevention. Another potential bias in studies which investigate the relationship between AECOPD and MI, which could explain some of the increased risk of MI after AECOPD, is differential misclassification of episodes of angina as AECOPD.

No studies were found which investigated the risk of MI associated with the frequent exacerbator phenotype. The frequent exacerbator phenotype may prove to be a useful characteristic for stratifying cardiovascular risk among patients with COPD. Future cohort studies of cardiovascular disease in people with COPD should, where possible, phenotype participants and investigate the relationship between exacerbator phenotype and risk of MI. Few included studies assessed the influence of severity of COPD on risk of MI; further research should investigate
this relationship as well as the influence of severity of COPD on risk of death following MI.

A further limitation of several of the included studies on death following MI was availability of information on cause of death. Collection of information on cause of death in future studies would allow investigators to draw more confident conclusions about the reasons for increased risk of death following MI for people with COPD.

Strengths and limitations of this review

This review benefitted from using a comprehensive search strategy which covered several bibliographic databases. As the relationship between AECOPD and MI has not been extensively studied, the inclusion criteria for this research question were kept purposively broad. This allowed all information pertaining to this relationship to be included in the evidence synthesis. One potential limitation of systematic reviews is publication bias. The potential for publication bias was highest for the review of outcomes after MI. In order to reduce the risk of this bias, we only included studies which specifically investigated the risk of COPD on MI rather than several different potential prognostic factors, as studies which investigated several factors which did not find an association between COPD and MI may not have reported this in the abstract or even in the text. Owing to clinical and statistical heterogeneity, meta-analysis could only be conducted for some of the research questions. Where meta-analysis was conducted, statistical heterogeneity was generally high, and this may limit the generalisability of pooled estimates.

CONCLUSIONS

There is good evidence of an increased risk of MI in people with COPD; however, it is unclear to what extent this association is due to smoking status.

There is some evidence that among people with COPD, AECOPD represent periods of increased risk of MI. However, further larger studies using validated exposure methods are needed to support this finding.

There is weak evidence that in-hospital mortality is higher for people with COPD after an MI. There is good evidence that postdischarge mortality after an MI is higher for people with COPD.

Contributors

KJR, LS and JKQ conceived and designed the study. KJR and RY screened abstracts and full text articles for inclusion, extracted data, and assessed the risk of bias of included studies. KJR conducted the meta-analysis. KJR, RY, LS and JKQ interpreted findings. KJR wrote the first draft. KJR, RY, LS and JKQ revised the article critically for important intellectual content.

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Competing interests

JKQ reports grants from Medical Research Council, grants from GSK—MRC and the British Lung Foundation during the conduct of the study; personal fees from GSK, Almirall and AstraZeneca, outside the submitted work. LS reports grants from Wellcome Trust during the conduct of the study; grants from Wellcome Trust, grants from MRC, grants from NIHR, personal fees from GSK, outside the submitted work. KJR has no conflicts of interest to disclose. RY has no conflicts of interest to disclose.

Provenance and peer review

Not commissioned; externally peer reviewed.

Data sharing statement

This is a systematic review of previously published studies.

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REFERENCES


