Review article

Is community acquired pneumonia an independent risk factor for cardiovascular disease?

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Abstract

Community-acquired pneumonia (CAP) is the most frequent infectious cause of death in Western countries. The high mortality rate in CAP is commonly related to comorbid conditions including cardiovascular disease.

Clinical studies in both primary and secondary care settings have identified an increase in short and long term risk of cardiovascular events and death from vascular events following acute respiratory infections. The mechanism remains to be fully established, but it has been suggested that the inflammatory state in patients affected by CAP acts to promote platelet activation, thrombosis and to narrow coronary arteries through vasoconstriction. Acute infections destabilise vascular endothelium and create an imbalance between myocardial oxygen supply and demand, leading to an increased risk of cardiovascular events. Acute infections have been shown to have both systemic effects, and local effects on coronary vessels. These effects are mediated through both the host response to infection and, in some cases, direct effects of bacterial infection or bacterial products.

In this review, we discuss the link between CAP and increased risk of cardiovascular events, drawing on existing evidence from clinical and mechanistic studies. Further studies and increased awareness into this association is warranted to promote novel ways of protecting high risk patients.
**Introduction**

Community acquired pneumonia (CAP) is the leading cause of death from infectious diseases in Western countries [1]. The mortality rate from CAP remains high at around 5-15% [2-6] and this figure has not changed significantly since the development of antibiotics in the 1950’s [2, 7]. Initial survivors of CAP have been shown to be at ongoing risk of mortality in the months to years following infection [8-12].

In patients with pneumonia, more than a quarter of deaths within 30 days and nearly 50% of deaths overall are not directly related to pneumonia but are related to co-morbidities [2, 13]. The elderly population, who are the most frequently affected by CAP, are also at high risk of mortality from cardiovascular disease [14].

Accumulating evidence points to a direct link between acute respiratory infections and an increased risk of cardiovascular events. Whether or not CAP acts as an acute trigger for cardiovascular events is important to ascertain as preventative strategies may aid in reducing mortality from this common infection.

The aim of this review is to discuss the role of community acquired pneumonia as a trigger for acute vascular events, drawing on evidence from experimental and clinical studies in the literature.

**Search strategy**

The current narrative review was based on a search of Pubmed for articles on major MeSH terms ‘myocardial infarction’ or ‘acute coronary syndrome’ or ‘cardiovascular diseases’ with ‘pneumonia’ or ‘respiratory tract infections’ or ‘sepsis’ between 1950
and December 2010. Only articles in English or English translation were reviewed. The articles were selected on the basis of originality and relevance.

**Atherosclerosis, thrombosis and plaque vulnerability**

Central to the pathogenesis of acute coronary syndromes and stroke are the rupture of the fibrous cap overlying a previously stable atherosclerotic plaque. The risk of plaque rupture is determined by the size and consistency of the lipid-rich core, the thickness of the fibrous cap and the extent of ongoing inflammation and repair within the cap [15]. Rupture exposes the underlying lipid-rich core which is highly pro-thrombotic. Formation of thrombus in-situ leads to occlusion of the coronary (or cerebral) vessel with the subsequent clinical presentation of acute coronary syndrome or stroke [16]. The severity of disease is determined by a number of factors including extent of thrombosis, the degree of arterial narrowing, the degree of vasoconstriction and ultimately the coronary perfusion pressure relative to the myocardial oxygen demand. A detailed discussion of the pathogenesis of atherosclerosis and acute coronary syndrome is beyond the scope of this review but is summarised in figure 1.

The process of plaque rupture, in-situ thrombus formation and the balance of arterial perfusion vs. myocardial (or cerebral) oxygen demand may be significantly affected during acute pneumonia.

Figure 1.

*Title:* Atherosclerosis.

*Legend:*
Figure 1A shows a stable atherosclerotic plaque. Oxidised lipoproteins are taken up by macrophages to produce lipid laden foam cells. Release of cytokines promote further infiltration by macrophages and promote smooth muscle proliferation. The fibrous cap surrounding the lipid core is dominated by vascular smooth muscle but also contains inflammatory cells. Figure 1B shows a ruptured atherosclerotic plaque. Rupture of the endothelial covering of the plaque exposes the highly thrombogenic lipid-rich core, leading to platelet aggregation and thrombus formation which may occlude the coronary vessel and give rise to the clinical presentation of acute coronary syndrome [16-21].

The Proposed Mechanism of Increased Cardiovascular Events following Respiratory Infections

A large body of evidence from clinical, in vitro and in vivo mechanistic studies provide a biologically plausible link between acute respiratory infections and atherosclerosis and thrombosis.

Systemic inflammation

Atherosclerosis is an inflammatory disorder [16]. Patients with acute coronary syndromes have evidence of systemic inflammation, with elevated pro-inflammatory cytokines and coagulation markers [22, 23]. In stable patients with atherosclerosis, raised inflammatory markers correlate with increased 10-year cardiovascular risk [24, 25]. C-reactive protein (CRP) in particular has been identified as an independent predictor of a risk of cardiovascular events [26-31] and has been shown to correlate with the extent of coronary atherosclerosis seen on angiography [32-34].
Inflammatory cytokines (interleukin (IL)-6, tumour necrosis factor-alpha (TNFα)) and markers of coagulation such as fibrinogen are elevated in patients with coronary artery disease [22, 23, 35-37].

Recent evidence has provided new information about the profound acute inflammatory response in CAP patients. CAP leads to a rapid elevation in pro-inflammatory cytokines such as CRP, IL-6, TNF-α and IL-8. These cytokines correlate to some extent with severity and decline with treatment. However, long-term follow up studies have shown that ongoing subclinical inflammation persists in CAP patients even after clinical recovery. Raised levels of IL-6 and IL-10 at hospital discharge have shown an association with an increased risk of death at 1 year [38]. Unfortunately it is not yet known how long this inflammation persists after hospital discharge or how changes in cytokine levels over time in CAP correlate with vascular events.

Patients with CAP therefore have systematic inflammation which places them at risk of acute cardiovascular events both during the acute infection, but also persisting after recovery and potentially increasing their risk over the longer term.

**Coronary artery inflammation**

Evidence also suggests that acute infections may trigger inflammation directly in the coronary vasculature in addition to causing systemic inflammation. A post-mortem study suggested that patients with acute infections had increased infiltration of established atherosclerotic plaques with macrophages and increased infiltration of the adventitia with macrophages, T cells and dendritic cells [39].
Platelet activation and thrombosis

As discussed above, platelet aggregation on the surface of a ruptured plaque is the initiating event in acute coronary and cerebral syndromes. Acute infections may stimulate platelet activation directly through pro-inflammatory cytokines or by bacterial products such as lipopolysaccharide [40]. Indirect platelet activation may also result during infection from mechanical stress induced by changes blood volume and vascular tone [41, 42].

This activation is an essential part of innate immunity, as activated platelets participate in host defence through reactive oxygen species generation, phagocytosis, cytokine release and interaction with neutrophils and monocytes [42]. This process may, however, also predispose to the initiation of thrombosis and a acute ischaemic events.

CAP also induces a marked pro-thrombotic state associated with elevation of thrombotic markers such as fibrinogen, factor IX, thrombin-anti-thrombin complex and D-dimer [43-45]. Septic patients demonstrate elevated levels of tissue factor without a corresponding increase in tissue factor pathway inhibitor [46]. Milbrandt et al. [47] identified that significant pro-thrombotic effects and elevation of coagulation markers persisted throughout hospital stay and occurred even in patients with clinically very mild pneumonia. Data from the same group also suggests that elevation of coagulation markers is greatest in elderly patients with pneumonia, the group at greatest cardiovascular risk [48].
**Endothelial dysfunction**

The vascular endothelium is a dynamic autocrine and paracrine organ that regulates vascular tone and secretes vasoactive and inflammatory mediators. In atherosclerosis, there is endothelial dysfunction with impaired vascular tone and proliferation of vascular smooth muscle. Dysfunctional endothelium promotes platelet aggregation and adhesion of leukocytes [49]. In sepsis, activation of the endothelium occurs early and may be beneficial facilitating the passage of leukocytes to infected areas [50]. However, activated endothelium releases nitric oxide, a key mediator in sepsis, resulting in vasodilation through relaxation of vascular smooth muscle. Acute infections lead to increased expression of inducible nitric oxide synthase (iNOS) with elevated nitric oxide (NO) production [51-54]. High levels of NO are toxic to the myocardium and cause systemic vasodilation [54-56] leading to increased leakiness of the systemic vascular bed and multiple organ failure [57]. The crucial role of iNOS is demonstrated by the finding that iNOS deficient mice are relatively resistant to lipopolysaccharide (LPS) induced sepsis [58]. Sepsis also induces increased expression of cyclooxygenase-2 [59] leading to increased production of prostanoid mediators from the arachidonic acid pathway including thromboxane and prostacyclin [60]. These mediators along with endothelin-1 [61] are increased in sepsis and impair the ability of the endothelium to regulate vascular tone, prevent platelet activation and regulate inflammatory cell adhesion [54]. Furthermore plasma von Willebrand factor levels are increased with endothelial damage, and may contribute to an overall thrombotic tendency [62].

**Effects of community acquired pneumonia on the heart**
Acute respiratory infections are associated with the damaging combination of reduced myocardial contractility, increased myocardial oxygen demand and reduced myocardial oxygen delivery.

Cytokines (IL-1, TNF-α, IL-6), prostanoids, endothelin-1 and nitric oxide produced in sepsis are all known to depress myocardial contractility (reviewed extensively by Merx et al [54]). Decreased systolic and diastolic ventricular contractility in septic patients has been evidenced by numerous scientific and clinical studies, and results in reduced coronary perfusion pressure [54, 63, 64].

Infection also results in increased myocardial oxygen demand secondary to systemic vasodilation, tachycardia and catecholamine release. Catecholamines cause coronary vasoconstriction and endothelial dysfunction [65]. In addition, patients with respiratory infections are frequently hypoxic. Consequently, despite the increased oxygen demands described in sepsis, there may be significantly reduced myocardial oxygen delivery. Reduced myocardial oxygen delivery may lead to myocyte necrosis [66]. Cardiac troponin levels are a sensitive marker of myocyte necrosis and are frequently elevated in sepsis and are related to sepsis severity, as well as left ventricular dysfunction in sepsis [67-69].

Figure 2 summarises the proposed mechanisms of how CAP may lead to acute cardiovascular events.

Figure 2.
Title: Summary of the proposed mechanisms by which community acquired pneumonia may provoke acute coronary syndrome/acute vascular events.

Direct effects of respiratory pathogens

Experiments on apolipoprotein deficient mice (a well established mouse model of atherosclerosis) have shown that infection with influenza virus promotes inflammatory cell infiltration, smooth muscle cell proliferation and fibrin deposition in atherosclerotic plaques. [70]. Unfortunately, no such study has been done using Streptococcus pneumoniae, the most common cause of CAP. Much of the experimental work into the link between respiratory infection and atherosclerosis has been directed at Chlamydia pneumoniae. Pathological studies have identified C pneumoniae in atherosclerotic plaques using numerous methods (polymerase chain reaction, electron microscopy, immunocytochemistry) whereas C pneumoniae antigens were not detectable in normal or non-atherosclerotic segments of arterial wall [71, 72]. Animal models have shown the atherosclerotic process to be induced and accelerated by infection with C. pneumoniae [71, 72]. Whether these findings in animal models translate to the atherosclerotic process in humans remains to be seen.

A number of comprehensive reviews have been published summarising the evidence for and against the link between C pneumoniae and atherosclerosis [73, 74].
Evidence from Clinical Studies

Epidemiological Studies

Numerous surveillance studies have reported seasonal patterns of cardiovascular deaths with a peak during winter months that cannot otherwise be explained by the conventionally recognised cardiovascular risk factors [75, 76]. Respiratory tract infections also peak in the winter, but no epidemiological studies have clearly established a link between the two disease entities.

Crawford et al. [77] identified a seasonal increase in fibrinogen level and mean platelet size that corresponded to an increased incidence of acute cardiovascular events and postulated that this was linked to a seasonal pro-thrombotic state. In another study, Woodhouse et al. [78] suggested that the seasonal variation in fibrinogen might be induced by winter respiratory infections: fibrinogen level was strongly related to neutrophil count, C-reactive protein, self-reported cough and coryza.

Population based studies

Large cohort studies utilising primary care databases have investigated a link between respiratory tract infection and the incidence of acute vascular events in the weeks to months following infection.

Primary care research databases use computer-recorded diagnostic codes entered by general practitioners [79]. Validation studies have verified the accuracy of computer entries for acute myocardial infarction (AMI) and stroke at >89% [80]. Respiratory tract infections (RTI) are entered into the database using a number of standard coding
terms including “respiratory tract infection”, “bronchitis”, “pneumonia”, “chesty productive cough” [81].

There have been 3 major primary care studies (summarised in Table 1) comprising a total of 33,563 patients with first episode of AMI and 28,271 with first episode of stroke [82-84]. All have documented an increased risk of acute MI or stroke following RTI which persists for up to 90 days after infection. In all 3 studies, the increased risk was maximal shortly after infection and decreased with time. Interestingly, these studies have shown minimal or no association between urinary tract infection and AMI suggesting the increased cardiovascular risk may be specific to respiratory infections.
<table>
<thead>
<tr>
<th>Study</th>
<th>Time (days)</th>
<th>Myocardial infarction</th>
<th>Stroke</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>OR</td>
<td>CI</td>
<td>OR</td>
</tr>
<tr>
<td>Meier</td>
<td>1-5</td>
<td>3.6</td>
<td>2.2-5.7</td>
</tr>
<tr>
<td>[82]</td>
<td>6-10</td>
<td>2.3</td>
<td>1.3-4.2</td>
</tr>
<tr>
<td></td>
<td>11-30</td>
<td>1.2</td>
<td>0.9-1.7</td>
</tr>
<tr>
<td></td>
<td>31-90</td>
<td>1.1</td>
<td>0.9-1.4</td>
</tr>
<tr>
<td></td>
<td>91-365</td>
<td>1.0</td>
<td>0.8-1.0</td>
</tr>
<tr>
<td>Smeeth</td>
<td>1-3</td>
<td>4.95</td>
<td>4.43-5.53</td>
</tr>
<tr>
<td>[83]</td>
<td>4-7</td>
<td>3.2</td>
<td>2.84-3.6</td>
</tr>
<tr>
<td></td>
<td>8-14</td>
<td>2.81</td>
<td>2.54-3.09</td>
</tr>
<tr>
<td></td>
<td>15-28</td>
<td>1.95</td>
<td>1.79-2.12</td>
</tr>
<tr>
<td></td>
<td>29-91</td>
<td>1.4</td>
<td>1.33-1.48</td>
</tr>
<tr>
<td>Clayton</td>
<td>1-3</td>
<td>3.75</td>
<td>1.86-7.56</td>
</tr>
<tr>
<td>[84]</td>
<td>1-7</td>
<td>2.1</td>
<td>1.38-3.21</td>
</tr>
<tr>
<td></td>
<td>8-28</td>
<td>1.93</td>
<td>1.42-2.63</td>
</tr>
<tr>
<td></td>
<td>29-91</td>
<td>1.16</td>
<td>0.92-1.47</td>
</tr>
<tr>
<td></td>
<td>92-365</td>
<td>1.08</td>
<td>0.94-1.23</td>
</tr>
</tbody>
</table>

Abbreviations; RTI = respiratory tract infection; OR = odds ratio; CI = confidence interval
These primary care based studies are large and uniformly give similar conclusions but there are a number of inherent limitations related to the study design. One major problem is the classification of respiratory infection within general practice databases. Diagnosing CAP in the community is challenging as GPs have to rely on clinical diagnoses without chest radiographic confirmation. Previous reviews have highlighted clinical findings alone to be inadequate in diagnosing CAP [85]. Pneumonia accounts for only 5-12% of all cases of adult lower respiratory tract infection treated with antibiotics by GPs in the community [86]. In addition, there may be some overlap between the less acute symptoms of coronary artery disease, left ventricular failure and acute respiratory infection. In many of these studies it is not possible to exclude the possibility of an inaccurate initial diagnosis of infection. This is also a consideration in some of the hospital based studies described below.

Another major limitation of current clinical studies is that it is not possible to delineate whether the association between CAP and cardiovascular disease is because of the high burden of mutual risk factors (age, diabetes, smoking) in these patients or whether a direct link is being observed.

**Hospital based and Short term studies**

Questionnaire studies of patients admitted to hospital following an acute myocardial infarction or ischaemic stroke have shown a frequent reporting of symptoms of respiratory infection preceding admission [87, 88]. In addition, one small study identified that positive serology for bacterial infection (including that of pathogens causing CAP e.g. *Staphylococcus aureus, Streptococcus pneumoniae, Mycoplasma*)
*pneumoniae, Chlamydophila pneumoniae*, was 5 times more likely in patients presenting with ischaemic stroke than in controls [89].

A multi-centre study by Lichtman et al. [90] looking at non-cardiac conditions in patients admitted with AMI found that 7.2% had concurrent pneumonia. A further study by the same investigators [91] identified severe pneumonia at admission in 18.4% of patients. The study by Zurru et al. [92] on patients admitted with atherothrombotic ischaemic stroke noted respiratory infections in the past year to be more common, with radiologically confirmed CAP being the most prevalent type of infection (19% in cases versus 6% in controls (odds ratio (OR)=3.9, 95% confidence interval (CI) 1.9 to 8, p<0.0001)).

Overwhelmingly therefore, in studies looking at prevalence of infection before ischaemic events, infections of the respiratory tract are consistently the most common cause. The potential overlap between respiratory and cardiac symptoms, as discussed above, remains a concern in some of these studies.

Further hospital based studies have looked at patients admitted with CAP for the incidence of concurrent acute cardiac events and whether clinical outcome/mortality from CAP might be attributable to acute MI. These are observational studies which typically do not utilise a control population.

Musher et al. [93] studied 170 patients admitted with community-acquired pneumococcal pneumonia and identified concurrent AMI occurring in 7.1%. 50% of patients with concurrent MI were bacteraemic. Mortality in patients with concurrent
pneumococcal pneumonia and acute cardiac events was higher than in those with pneumonia alone (OR 3.9, P<0.008).

These findings have been extended by 2 studies from the CAPO international database suggesting an important role for acute MI in clinical outcomes for pneumonia patients. Ramirez et al. [94] found AMI to be present in 5.8% of 500 consecutive patients admitted with CAP. 15% of those with severe CAP (i.e. requiring ITU admission) had concurrent AMI and an association between pneumonia severity index (PSI) score and AMI was noted. Of those patients transferred to the ITU with the first 24 hours after hospital admission, 50% had concurrent AMI. Aliberti et al. [95] subsequently reported in the same cohort that acute myocardial infarction was the second most common cause of clinical failure (28% of cases of treatment failure).

In addition, Corrales-Medina et al. [96] followed patients with CAP caused by *S. pneumoniae* or *H. influenzae* for 15 days after hospital admission. In their study, they found that CAP patients were eight times more likely than controls to have a cardiac event. The risk of AMI increased to more than 40-fold when comparing the risk 1 year before the episode of CAP to the risk during the 15 day follow up period [97].

**Long term studies**

A number of studies have sought to study the long term sequelae of CAP by following patients after discharge from hospital. Overwhelmingly, these studies suggest that survivors of CAP have increased long term mortality compared to the healthy population and that cardiovascular disease comprises a substantial proportion of long term deaths in CAP patients. Mortensen et al. investigated causes of mortality within
90 days of presentation with CAP. After neurological conditions and malignancy, cardiac ischaemia was identified as the 3rd most common underlying cause of death in this cohort of patients [2].

Two prospective multi-centre cohort studies suggest that cardiovascular events are a major cause of mortality following CAP. Yende et al. [10] followed patients for 1 year and Johnstone et al. [9] followed patients for an average of 5.4 years after discharge. Both identified that one third of deaths following CAP episode were due to cardiovascular causes. Recently, Bruns et al. [11] reported an increased risk of death at 1, 5 and 7 years following an episode of CAP compared to the general Dutch population. Overall, cardiovascular events contributed to 16% of deaths in this cohort. A population based study from Finland [12] followed elderly patients for a mean of 9.2 years and identified a significantly increased risk of long term mortality among CAP patients and an independent association with cardiovascular mortality (relative risk (RR) = 2.4, 95% CI 1.7-3.4, p<0.001).

Studies assessing cardiovascular risk in patients admitted to hospital with CAP are summarised in table 2.
Table 2: Evidence from studies on patients admitted to hospital with pneumonia.

<table>
<thead>
<tr>
<th>Study</th>
<th>No. of cases</th>
<th>Study Design</th>
<th>Follow up period post CAP</th>
<th>Outcome</th>
<th>Incidence (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Musher [93]</td>
<td>170</td>
<td>Retrospective cohort study</td>
<td>Duration of hospital stay</td>
<td>AMI</td>
<td>7.1</td>
</tr>
<tr>
<td>Ramirez [94]</td>
<td>500</td>
<td>Retrospective cohort study</td>
<td>Duration of hospital stay</td>
<td>AMI</td>
<td>5.8</td>
</tr>
<tr>
<td>Corrales-Medina [96]</td>
<td>206</td>
<td>Retrospective case control study and case-crossover design</td>
<td>15 days</td>
<td>AMI</td>
<td>10.7</td>
</tr>
<tr>
<td>Mortenson [2]</td>
<td>208</td>
<td>Prospective observational study</td>
<td>90 days</td>
<td>Cardiovascular mortality</td>
<td>13</td>
</tr>
<tr>
<td>Yende [10]</td>
<td>1799</td>
<td>Prospective cohort study</td>
<td>1 year</td>
<td>Cardiovascular mortality</td>
<td>33</td>
</tr>
<tr>
<td>Johnstone [9]</td>
<td>3415</td>
<td>Population based cohort study</td>
<td>5.4 years</td>
<td>Cardiovascular mortality</td>
<td>31</td>
</tr>
<tr>
<td>Bruns [11]</td>
<td>356</td>
<td>Prospective observational study</td>
<td>7 years</td>
<td>Cardiovascular mortality</td>
<td>16</td>
</tr>
</tbody>
</table>
Preventative measures and the impact on the relationship between CAP and vascular events

i) Pneumococcal and Influenza vaccination

If respiratory infections predispose to cardiovascular events, it would be expected that prevention of respiratory infections with vaccination should also prevent vascular events. Previous studies have shown that influenza vaccine reduces the risk of recurrent MI, stroke and cardiac hospital admissions [98, 99] and it has been hypothesised that pneumococcal vaccine might provide a similar protective role. To date, such studies have produced conflicting results. It is acknowledged that the current available polysaccharide pneumococcal vaccine is likely less effective than the influenza vaccine, which may explain the incongruent results [100].

A case-control study by Lamontagne et al. [101] assessing 999 patients admitted to hospital with first MI showed that cases were less likely than controls to have been vaccinated (OR 0.53). A protective role of pneumococcal vaccination was dependent on time since vaccination, with increased protection offered if the vaccine was received more than 2 years ago (OR 0.33) compared with no protection if the vaccine was received less than one year ago. Conversely, in a cohort study by Tseng et al.
involving 84,170 patients there was no significant association between pneumococcal vaccination and risk of acute MI or stroke. A further case-control study from the UK [103] on 16,012 patients with first MI showed a significant protective benefit from influenza vaccine (OR 0.81) but not from pneumococcal vaccine.

A recent study from China [104] evaluating the dual influenza/pneumococcal vaccine has added to this debate by showing a marked reduction in mortality (hazard ratio (HR) 0.65; p<0.001), ischaemic stroke (HR 0.67; p<0.001), and acute myocardial infarction (HR 0.52; p<0.001) in vaccinated patients compared to unvaccinated subjects. In this study, both influenza vaccine and pneumococcal vaccine alone did not show a significant protective effect.

It is important to note that none of these studies were randomised controlled trials and cannot conclusively answer the question as to whether pneumococcal vaccination truly prevents acute arteriovascular events. Evidence from vaccination studies to date is summarised in table 3.

<table>
<thead>
<tr>
<th>First author, year</th>
<th>Study design</th>
<th>Vaccine</th>
<th>Cardiovascular event</th>
<th>OR/HR</th>
<th>CI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lamontagne, 2008</td>
<td>Case-control</td>
<td>Pneumococcal</td>
<td>AMI</td>
<td>0.53</td>
<td>0.40-0.70</td>
</tr>
<tr>
<td>Tseng, 2010</td>
<td>Prospective cohort</td>
<td>Pneumococcal</td>
<td>AMI</td>
<td>1.09</td>
<td>0.98-1.21</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Stroke</td>
<td>1.14</td>
<td>1.00-1.31</td>
</tr>
</tbody>
</table>
Siriwardena, Case-control Pneumococcal AMI 0.96 0.91-1.02
2010 [103] Influenza AMI 0.81 0.77-0.85
Hung, 2010 Prospective Dual AMI 0.52 0.38-0.71
[104] cohort Stroke 0.67 0.54-0.83

Abbreviations; HR=hazard ratio,

ii) Antimicrobial therapy

Multiple large-scale randomised placebo-controlled trials of antibiotic regimes against *C. pneumoniae* have collectively shown no major cardioprotective effect. Meta-analysis data on anti-chlamydial antibiotics and cardiovascular disease from 13 studies comprising 12,491 treated patients and 12, 518 controls, did not show a significant benefit (RR = 0.96; 95% CI 0.90-1.04) [105].

iii) Pharmacotherapy

Several observational studies have shown a potential benefit for drugs used to treat cardiovascular disease on outcome for patients with community acquired pneumonia or sepsis. Significant reduction in short term mortality have been found with prior statin therapy [106] and prior ACE-inhibitor therapy in patients with CAP [106]. Statins have established immunomodulatory effects in addition to their role in preventing cardiovascular disease.[99] A recent study also suggested a possible benefit associated with anti-platelet agents [107]. A detailed discussion of the potential role of these drugs are beyond the scope of this review. These data raise the question as to whether these drugs may reduce deaths from CAP by preventing cardiovascular events.
Clinical significance

Current evidence suggests that patients with CAP are at increased short and long term risk of acute arteriovascular events. Cardiovascular disease is implicated in a significant proportion of deaths in CAP patients. Therapies, such as anti-platelet agents, statins, angiotensin II inhibitors/angiotensin receptor blockers are established in the secondary prevention of vascular events and may become relevant treatments in the long term management of CAP patients. The evidence so far collated (see table 4) meets the Bradford Hill criteria [108] for defining causal association to some extent, but because pneumonia and cardiovascular disease share common risk factors (e.g. smoking, diabetes, age), a true cause-and-effect relationship is difficult to confirm.
Table 4: Bradford Hill’s criteria for causality and the evidence so far

<table>
<thead>
<tr>
<th>Criterion</th>
<th>Evidence</th>
</tr>
</thead>
<tbody>
<tr>
<td>Strength of association</td>
<td>Strong association depicted in the study by Corrales-Medina et al. [96]. Patients were eight times more likely than controls to have a cardiac event 15 days after hospital admission with CAP.</td>
</tr>
<tr>
<td>Consistency of results in a variety of settings</td>
<td>Studies in both primary and secondary care of a variety of study design including case-control and cohort have shown an increased risk of CV events following acute respiratory infection.</td>
</tr>
<tr>
<td>Specificity</td>
<td>Both pneumonia and cardiovascular disease are multifactorial and share common risk factors therefore this is difficult to confirm.</td>
</tr>
<tr>
<td>Temporality</td>
<td>Primary care database studies showed risk of MI decreasing with time following infection [82-84].</td>
</tr>
<tr>
<td>Biological gradient</td>
<td>In the study by Ramirez et al. [94], a positive association between pneumonia severity score and AMI was noted.</td>
</tr>
<tr>
<td>Plausibility</td>
<td>Experimental evidence as highlighted in this review supports a role for infection in cardiovascular events.</td>
</tr>
</tbody>
</table>
Coherence Laboratory, epidemiological and clinical evidence is consistent in supporting a role for infection in cardiovascular events.

Experimental evidence In vitro/animal model studies specific to pneumonia are awaited. Vaccination studies to date are not consistent enough to satisfy this criterion[101-104].

Analogy The effect of other established cardiovascular risk factors could be considered analogous to that of pneumonia.

**Limitations of existing studies and future direction**

While great progress has been made in understanding the link between respiratory infections and cardiovascular events, many areas require further study. While much is known on the effects of severe sepsis on the heart and endothelium, many of the patients described in clinical studies had mild pneumonia and many did not require hospitalization. These patients are unlikely to have had sepsis and less is known about how relatively mild infections affect the heart, vasculature and inflammatory pathways. Furthermore, there remains a lack of experimental models into specific CAP causing organisms and cardiovascular disease. It may be that pneumococcal pneumonia, for example, has a more potent effect on cardiovascular disease than other organisms, as has been shown in studies of influenza virus and cardiovascular disease, compared with other viral infections.
Conclusion

Current evidence suggests that community acquired pneumonia is associated with significant increase in short and long term risk of cardiovascular events and death from vascular events.

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Conflict of interest statement

The authors confirm they have no conflicts of interest regarding the present manuscript.
References


Figure 1

[Diagram showing the relationship between platelet activation, pro-coagulant state, systemic inflammation, local inflammation within plaques, endothelial dysfunction, hypoxemia, systemic vasodilation, increased circulating catecholamines, and coronary vasoconstriction.

Figure 2

[Diagram illustrating acute coronary syndrome with atheroma instability, thrombogenesis, plaque rupture, increased myocardial oxygen demand, and reduced myocardial oxygen delivery.]