Prevalence of low-grade inflammation in depression: a systematic review and meta-analysis of CRP levels

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Abstract

Background. Peripheral low-grade inflammation in depression is increasingly seen as a therapeutic target. We aimed to establish the prevalence of low-grade inflammation in depression, using different C-reactive protein (CRP) levels, through a systematic literature review and meta-analysis.

Methods. We searched the PubMed database from its inception to July 2018, and selected studies that assessed depression using a validated tool/scale, and allowed the calculation of the proportion of patients with low-grade inflammation (CRP >3 mg/L) or elevated CRP (>1 mg/L).

Results. After quality assessment, 37 studies comprising 13,541 depressed patients and 155,728 controls were included. Based on the meta-analysis of 30 studies, the prevalence of low-grade inflammation (CRP >3 mg/L) in depression was 27% (95% CI 21–34%); this prevalence was not associated with sample source (inpatient, outpatient or population-based), antidepressant treatment, participant age, BMI or ethnicity. Based on the meta-analysis of 17 studies of depression and matched healthy controls, the odds ratio for low-grade inflammation in depression was 1.46 (95% CI 1.22–1.75). The prevalence of elevated CRP (>1 mg/L) in depression was 58% (95% CI 47–69%), and the meta-analytic odds ratio for elevated CRP in depression compared with controls was 1.47 (95% CI 1.18–1.82).

Conclusions. About a quarter of patients with depression show evidence of low-grade inflammation, and over half of patients show mildly elevated CRP levels. There are significant differences in the prevalence of low-grade inflammation between patients and matched healthy controls. These findings suggest that inflammation could be relevant to a large number of patients with depression.

Introduction

Depression is a common mental illness with a complex aetiology and is one of the leading causes of disability worldwide, affecting around 10–20% of the general population in their lifetime (Lim et al., 2018). There is now increasing evidence suggesting an association between depression and inflammation (Goldsmith et al., 2016). For instance, 'sickness behaviour' commonly seen following an acute infection, shares many characteristics with depression, such as fatigue, sleep disturbance and decreased motivation (Dantzer et al., 2008); early-life infection and autoimmune diseases are associated with a higher risk of depression in adulthood (Benros et al., 2013); people with chronic immune-mediated inflammatory diseases such as rheumatoid arthritis exhibit a higher prevalence of depression (Dickens et al., 2002). Depression is also associated with other conditions linked with elevated inflammatory markers, such as cardiovascular disease (CVD) (Ridker, 2003).

C-reactive protein (CRP) is a marker of acute phase response which has been used most extensively as a measure of low-grade inflammation in psychiatric (von Känel et al., 2007; Fernandes et al., 2016) and physical conditions (Visser et al., 1999; Danesh et al., 2000). CRP is associated with cardiovascular risk, including myocardial infarction, stroke, sudden cardiovascular death and peripheral vascular disease (Ridker, 2003). Meta-analyses of cross-sectional studies confirm that mean concentrations of circulating CRP and inflammatory cytokines such as interleukin 6 (IL-6) are higher in patients with acute depression compared with controls (Howren et al., 2009; Dowlati et al., 2010; Haapakoski et al., 2015; Goldsmith et al., 2016). Population-based longitudinal studies show that higher levels of CRP and IL-6 at baseline are associated with an increased risk of depression in subsequent follow-ups (Gimeno et al., 2009).
et al., 2009; Wiium-Andersen et al., 2013; Khandaker et al., 2014; Zalli et al., 2016), suggesting that inflammation could be a cause rather than simply a consequence of the illness.

The association between inflammation and depression is clinically relevant. Poor response to antidepressants is associated with the activation of inflammatory immune responses (Lanquillon et al., 2000; Benedetti et al., 2002; Carvalho et al., 2013; Chamberlain et al., 2018). It has been reported the mean CRP levels are higher in treatment-resistant compared with treatment-responsive patients with depression (Maes et al., 1997; Sluzewska et al., 1997). Anti-inflammatory treatment has antidepressant effects (Müller et al., 2006; Köhler et al., 2014; Kappelmann et al., 2018). Randomised controlled trials (RCTs) indicate that anti-inflammatory drugs are likely to be beneficial particularly for depressed patients who show evidence of inflammation (Raison et al., 2013; Kappelmann et al., 2018). Currently, a number of ongoing RCTs of anti-inflammatory treatments are recruiting specifically depressed patients with elevated CRP levels (e.g. >3 mg/L): NCT02473289; ISRCTN16942542 (Khandaker et al., 2018). Therefore, a better understanding of the prevalence of low-grade inflammation in depression, and of factors associated with inflammation could inform future research and clinical practice.

Inflammation is unlikely to be relevant for all patients with depression (Khandaker et al., 2017). While it is established that mean concentrations of peripheral inflammatory markers are higher in depressed patients compared with controls (Howren et al., 2009; Dowlati et al., 2010; Haapasoksi et al., 2015; Goldsmith et al., 2016), it is unclear what proportion of depressed patients show evidence of low-grade inflammation. Many studies have reported on the prevalence of inflammation in depressed patients using various CRP level thresholds to define inflammation, e.g. >3 or >1 mg/L. These studies have been conducted in different settings and populations, e.g. inpatient, outpatient, population-based (Raison et al., 2013; Wiium-Andersen et al., 2013; Shin et al., 2016). The reported prevalence of inflammation varies widely among these studies; for example, for low-grade inflammation (CRP >3 mg/L) it has been reported to vary between 0% and 60% in existing studies (Ma et al., 2011; Hannestad et al., 2013). However, as far as we are aware, a systematic review and meta-analysis of the prevalence of low-grade inflammation in patients with depression is currently lacking. While it is likely that the prevalence of low-grade inflammation is higher in patients with depression compared with controls, to our knowledge, no systematic review and meta-analysis has examined the odds ratio for inflammation in depressed patients compared with matched controls.

We conducted a systematic review of existing studies to: (1) quantify the prevalence of low-grade inflammation in patients with depression using meta-analysis; (2) calculate the odds ratio for low-grade inflammation in depressed patients compared with matched healthy controls using meta-analysis; (3) identify sociodemographic and other factors associated with inflammation prevalence in patients with depression using meta-regression analysis. We defined low-grade inflammation as serum CRP levels >3 mg/L. This cut-off has been chosen based on the American Heart Association and Center for Disease Control and Prevention recommendations, which defined CRP levels of >3 mg/L as high (Pearson et al., 2003). In addition, we carried out additional analyses using CRP levels >1 mg/L to define ‘elevated CRP’, and >10 mg/L to define ‘very high CRP’ indicative of current infection. We also carried out a number of sensitivity analyses; for example, meta-analyses using >1 and >3 mg/L thresholds for CRP were repeated using only studies that excluded patients with suspected infection (defined as CRP >10 mg/L); and after excluding poor quality studies.

Methods

Search strategy and study selection

This systematic review has been performed according to the Preferred Reporting Items for Systematic Reviews and Meta-analyses (PRISMA) guidelines. The search protocol was prospectively published on PROSPERO (see: http://www.crd.york.ac.uk/PROSPERO/display_record.asp?ID=CRD42018106640). The PubMed database was searched for published studies from its inception to 5 of July 2018 using the following keywords: 'CRP OR “C-reactive protein” OR “hs-CRP” OR hsCRP) OR (C-Reactive Protein[mesh] AND depression”). No language restriction was applied; we only selected studies based on human participants. The electronic search was complemented by hand-searching of meta-analyses and review articles. Abstracts were screened, and full texts of relevant studies were retrieved. Two authors applied the inclusion/exclusion criteria independently and selected the final studies for this review (LB and EFO).

Selection criteria

We included studies that: (1) examined CRP levels in people with depression; (2) assessed depression using clinical criteria (DSM or ICD) or a validated tool (e.g. Hamilton Depression Rating Scale), and reported it as a categorical variable (yes/no); (3) reported CRP levels allowing the calculation of the proportion of ‘infamed’ patients using cut-offs of either 3, 1 or 10 mg/L. One study used CRP cut-offs of 0.99 and 3.13 mg/L, which was included (Penninx et al., 2003), as these values are very close to the thresholds above. Exclusion criteria were (1) studies reporting measures of inflammation other than CRP, e.g. interleukins or genetic markers; (2) in vitro or animal studies; (3) non-original data, e.g. reviews; (4) studies exclusively based on patients with a medical condition, e.g. cancer.

Recorded variables

The main outcome measure was the proportion of subjects showing elevated CRP in patients and, where reported, in non-depressed controls. We also extracted the following data: author; year of publication; sampling criteria; diagnostic criteria for depression; age of participants; treatment status (antidepressant-free, treatment resistant); ethnicity; matching criteria for patients and controls (if present); study setting and sample source (e.g. community or inpatient); presence of comorbidities. If there were multiple publications from the same data set, we used the study with the largest sample.

Data synthesis

We performed meta-analyses of the prevalence of inflammation in depressed patients using three different CRP cut-offs to define inflammation: >3 (primary), >1 and >10 mg/L. The pooled prevalence of inflammation was calculated using quantitative random-effect meta-analysis, expressed as percentage and 95% CI. The use of random-effect meta-analysis, as opposed to fixed
effect, is appropriate when there is heterogeneity between studies. Pooling of studies was performed using the inverse variance method, so that studies with bigger samples were given greater weight. The Clopper-Pearson method was used to compute confidence interval for individual studies, and the logit transformation was used for the transformations of proportions, with a continuity correction of 0.5 in studies with zero cell frequencies. Heterogeneity between studies was measured using the $I^2$ statistic, which describes the percentage of the variability in effect estimates that is due to heterogeneity. Heterogeneity was tested using Cochrane’s Q-Test (Higgins and Thompson, 2002). Publication bias was assessed for each group of studies by visual inspection of funnel plots, and tested with an Egger’s regression test for funnel plot asymmetry (mixed-effects meta-regression model). $P$ values <0.05, two tailed, were considered statistically significant. We used meta-regression analyses to evaluate the association of inflammation prevalence with age, sex, body mass index (BMI), sample source, proportion of antidepressant-free patients and ethnicity. Seventeen studies reported CRP levels in matched non-depressed controls; these were used to calculate the meta-analytic odds ratio for inflammation in patients with depression vs. healthy controls using random-effects estimates for meta-analyses with binary outcome data; pooling of studies was performed using the inverse variance method and with a continuity correction of 0.5 in studies with zero cell frequencies. Study quality was assessed using the Newcastle–Ottawa Scale (Stang, 2010). Analyses were repeated with poor quality studies removed. Meta-analyses were carried out using the meta package (version 4.9 (Schwarzer, 2007)) in R 3.4 (R Core Team, 2017), and plotted using packages meta and Cairo v1.5 (Urbanek and Horner, 2015). Additional information on the methods can be found in the Supplementary Materials.

**Results**

The literature search yielded 1545 results, out of which 37 studies met the inclusion criteria for meta-analysis (Legros et al., 1985; Penninx et al., 2003; Ladwig et al., 2005; Liukkonen et al., 2006; O’Brien et al., 2006; Almeida et al., 2007; Kling et al., 2007; Danese et al., 2008; Nilsson et al., 2008; Cizza et al., 2009; Harley et al., 2010; Ma et al., 2011; N Gaslshpour et al., 2011; Hannestad et al., 2013; Raison et al., 2013; Shanahan et al., 2013; Park et al., 2014; Uher et al., 2014; Wium-Andersen et al., 2014; Courtet et al., 2015; Wysokiński et al., 2015; Cepeda et al., 2016; Haroon et al., 2016; Rapaport et al., 2016; Shin et al., 2016; Ekinici and Ekinici, 2017; Eutenweer et al., 2017; Gallagher et al., 2017; Horsdal et al., 2017; Iha et al., 2017; Cáceda et al., 2018; Chamberlain et al., 2018; Felger et al., 2018; Osimo et al., 2018b; Porcu et al., 2018; Shibata et al., 2018; Wei et al., 2018). Please see Supplementary Fig. S1 for the PRISMA diagram of study selection, and Table 1 for details of the included studies.

**Prevalence of low-grade inflammation (CRP >3 mg/L) in depressed patients**

**Results based on all available studies**

Thirty studies comprising 11 813 patients with depression were used for this analysis. The meta-analytic pooled prevalence of low-grade inflammation in depressed patients was 27% (95% CI 21–34%); see Fig. 1. There was evidence of heterogeneity among studies ($I^2 = 97.7%$; $95\%$ CI $97.3$–$98.1%$; Cochrane’s $Q = 1264; p < 0.01$). Further analyses after grouping studies by setting $I^2 = 91.9%$ (95% CI 21–42%); $I^2 = 91.9%$ (95% CI 21–42%); $I^2 = 95.9%$; Cochrane’s $Q = 338$; and in population-based samples ($N = 7020$) it was 24% (95% CI 17–34%; $I^2 = 98.3%$; Cochrane’s $Q = 483$).

**Analyses excluding poor quality studies or cases with past depression**

A sensitivity analysis excluding six poor quality studies, comprising 8778 patients, showed that the prevalence of inflammation was 27% (95% CI 22–33%); Supplementary Table S1 and Fig. S1. There was evidence of heterogeneity ($I^2 = 96.4%$; 95% CI 95.5–97.1%; Cochrane’s $Q = 644$; $p < 0.01$). A sensitivity analysis excluding two studies where depression was not active in all patients, comprising 11 763 patients, showed that the prevalence of inflammation was 26% (95% CI 20–34%); Supplementary Fig. S2. There was evidence of heterogeneity ($I^2 = 97.9%$; 95% CI 97.4–98.2%; Cochrane’s $Q = 1261$; $p < 0.01$).

**Analysis after excluding cases of suspected infection (CRP >10 mg/L)**

Nine studies reported the prevalence of low-grade inflammation after excluding participants with suspected infection, defined as CRP >10 mg/L. A separate meta-analysis based on these studies, comprising 6948 patients, showed that the prevalence of inflammation was 16% (95% CI 8–32%); Supplementary Fig. S4. There was evidence of heterogeneity ($I^2 = 98.8$%; 95% CI 98.5–99.1%; Cochrane’s $Q = 675$; $p < 0.01$).

**Association between prevalence of low-grade inflammation (CRP >3 mg/L) and characteristics of depressed patients**

Meta-regression was used on 19 studies comprising 7858 patients to test the association between the prevalence of inflammation and the proportion of patients who were antidepressant-free at the time of study. There was no association between these factors (estimate: $-0.007$; $z = -1.03$; $p = 0.30$). Similarly, sex, age, non-White ethnicity, BMI and sample source (inpatient, outpatient or population-based) were not associated with the prevalence of inflammation (see Supplementary Results).

**Assessment of publication bias**

A funnel plot of the 30 studies assessing the prevalence of low-grade inflammation (CRP >3 mg/L) in depression visually appeared symmetrical (Supplementary Fig. S5). Egger’s regression test for funnel plot asymmetry was non-significant ($t = -1.3$; df = 28; $p = 0.21$), suggesting there was no evidence of publication bias.

**Odds ratio for low-grade inflammation (>3 mg/L) in depressed patients**

Seventeen studies reported the prevalence of inflammation in 7761 depressed patients and 155 728 matched non-depressed controls (see Supplementary Table S2 for matching details). The meta-analytic OR for inflammation in depressed patients compared with matched controls was 1.46 (95% CI 1.22–1.75; $p < 0.0001$); see Fig. 2. There was evidence of heterogeneity ($I^2 = 71.9$%; 95% CI 54.3–82.7%; Cochrane’s $Q = 57$; $p < 0.01$).

Based on the same studies, we meta-analysed the prevalence of inflammation in depressed patients and matched non-depressed controls separately. The prevalence of inflammation in controls
<table>
<thead>
<tr>
<th>Study</th>
<th>Country</th>
<th>Setting</th>
<th>Depressed (N)</th>
<th>Controls (N)</th>
<th>Mean age of patients in years (SD)</th>
<th>Patient sex (% male)</th>
<th>Assessment of depression*</th>
<th>Quality rating*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Legros et al. (1985)</td>
<td>Belgium</td>
<td>Inpatient</td>
<td>34</td>
<td>NA</td>
<td>42 (NA)</td>
<td>NA</td>
<td>Feighner et al. (1972) criteria (Feighner et al., 1972)</td>
<td>Good</td>
</tr>
<tr>
<td>Penninx et al. (2003)</td>
<td>USA</td>
<td>Prospective /population-based</td>
<td>145</td>
<td>2879</td>
<td>74 (2.9)</td>
<td>38.62</td>
<td>CES-D</td>
<td>Good</td>
</tr>
<tr>
<td>Ladwig et al. (2005)</td>
<td>Germany</td>
<td>Prospective /population-based</td>
<td>986</td>
<td>2035</td>
<td>57.5 (7.8)</td>
<td>100</td>
<td>Subscale from the von Zerssen affective symptom checklist (von Zerssen and Cording, 1978); depression = score ⩾11</td>
<td>Good</td>
</tr>
<tr>
<td>Liukkonen et al. (2006)</td>
<td>Finland</td>
<td>Prospective /population-based</td>
<td>962</td>
<td>4097</td>
<td>31 (0)</td>
<td>39.4</td>
<td>Hopkins symptom checklist-25 (Parloff et al., 1954)</td>
<td>Good</td>
</tr>
<tr>
<td>O’Brien et al. (2006)</td>
<td>Ireland</td>
<td>Outpatient</td>
<td>32</td>
<td>20</td>
<td>44.05 (NA)</td>
<td>34.38</td>
<td>DSM-IV</td>
<td>Poor</td>
</tr>
<tr>
<td>Almeida et al. (2007)</td>
<td>Australia</td>
<td>Prospective /population-based</td>
<td>213</td>
<td>4000</td>
<td>76.6 (4.4)</td>
<td>100</td>
<td>GDS-15 score ⩾7</td>
<td>Good</td>
</tr>
<tr>
<td>Kling et al. (2007)</td>
<td>USA and Israel</td>
<td>Outpatient</td>
<td>18</td>
<td>18</td>
<td>41 (12)</td>
<td>0</td>
<td>DSM-IV</td>
<td>Good</td>
</tr>
<tr>
<td>Danese et al. (2008)</td>
<td>UK</td>
<td>Prospective /population-based</td>
<td>109</td>
<td>673</td>
<td>32 (0)</td>
<td>39.45</td>
<td>DSM-IV</td>
<td>Good</td>
</tr>
<tr>
<td>Nilsson et al. (2008)</td>
<td>Sweden</td>
<td>Outpatient</td>
<td>50</td>
<td>NA</td>
<td>Median age 71 years</td>
<td>NA</td>
<td>DSM IV</td>
<td>Good</td>
</tr>
<tr>
<td>Cizza et al. (2009)</td>
<td>USA</td>
<td>Outpatient</td>
<td>77</td>
<td>41</td>
<td>35.5 (7)</td>
<td>0</td>
<td>DSM-IV SCI</td>
<td>Good</td>
</tr>
<tr>
<td>Harley et al. (2010)</td>
<td>New Zealand</td>
<td>Outpatient</td>
<td>346</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>DSM-IV SCI, Montgomery Asberg Depression Rating Scale</td>
<td>Poor</td>
</tr>
<tr>
<td>Ma et al. (2011)</td>
<td>USA</td>
<td>Outpatient</td>
<td>10</td>
<td>498</td>
<td>NA</td>
<td>30</td>
<td>BDI score ⩾22</td>
<td>Poor</td>
</tr>
<tr>
<td>Naghashpour et al. (2011)</td>
<td>Iran</td>
<td>Outpatient</td>
<td>43</td>
<td>52</td>
<td>37.26 (6.5)</td>
<td>0</td>
<td>BDI score ⩾5</td>
<td>Poor</td>
</tr>
<tr>
<td>Hannestad et al. (2013)</td>
<td>USA</td>
<td>Outpatient</td>
<td>9</td>
<td>7</td>
<td>37 (14.3)</td>
<td>44.44</td>
<td>DSM-IV</td>
<td>Fair</td>
</tr>
<tr>
<td>Raison et al. (2013)</td>
<td>USA</td>
<td>Outpatient</td>
<td>60</td>
<td>NA</td>
<td>43.4 (8.8)</td>
<td>33.33</td>
<td>DSM-IV depression through SCID</td>
<td>Good</td>
</tr>
<tr>
<td>Shanahan et al. (2013)</td>
<td>USA</td>
<td>Prospective /population-based</td>
<td>61</td>
<td>NA</td>
<td>13.5 (1.9)</td>
<td>NA</td>
<td>DSM-IV criteria assessed through CAPA</td>
<td>Good</td>
</tr>
<tr>
<td>Park et al. (2014)</td>
<td>Korea</td>
<td>Outpatient</td>
<td>30</td>
<td>30</td>
<td>65.2 (4.8)</td>
<td>30</td>
<td>DSM-IV SCI</td>
<td>Good</td>
</tr>
<tr>
<td>Uher et al. (2014)</td>
<td>Canada, UK, Germany, Croatia, Denmark, Slovenia, Belgium</td>
<td>Outpatient</td>
<td>241</td>
<td>NA</td>
<td>40.7 (11.4)</td>
<td>37.34</td>
<td>Schedule for Clinical Assessment in Neuropsychiatry</td>
<td>Good</td>
</tr>
<tr>
<td>Wiium-Andersen et al. (2014)</td>
<td>Denmark</td>
<td>Prospective /population-based</td>
<td>1183</td>
<td>77 626</td>
<td>65 (NA)</td>
<td>36.35</td>
<td>ICD criteria</td>
<td>Poor</td>
</tr>
</tbody>
</table>

(Continued)
<table>
<thead>
<tr>
<th>Study</th>
<th>Country</th>
<th>Setting</th>
<th>Depressed (N)</th>
<th>Controls (N)</th>
<th>Mean age of patients in years (SD)</th>
<th>Patient sex (% male)</th>
<th>Assessment of depression</th>
<th>Quality rating</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wysokiński et al. (2015)</td>
<td>Poland</td>
<td>Inpatient</td>
<td>319</td>
<td>NA</td>
<td>59.7 (21)</td>
<td>23.82</td>
<td>ICD-10</td>
<td>Poor</td>
</tr>
<tr>
<td>Courtet et al. (2015)</td>
<td>France</td>
<td>Inpatient</td>
<td>600</td>
<td>NA</td>
<td>39.8 (13.4)</td>
<td>27.83</td>
<td>DSM-IV</td>
<td>Good</td>
</tr>
<tr>
<td>Cepeda et al. (2016)</td>
<td>USA</td>
<td>Prospective /population-based</td>
<td>1325</td>
<td>12 951</td>
<td>45.3 (12)</td>
<td>35.92</td>
<td>PHQ9 score &gt;9</td>
<td>Good</td>
</tr>
<tr>
<td>Haroon et al. (2016)</td>
<td>USA</td>
<td>Outpatient</td>
<td>50</td>
<td>NA</td>
<td>38.6 (10.8)</td>
<td>32</td>
<td>DSM-IV</td>
<td>Fair</td>
</tr>
<tr>
<td>Rapaport et al. (2016)</td>
<td>USA</td>
<td>Outpatient</td>
<td>155</td>
<td>NA</td>
<td>46.1(12.6)</td>
<td>41.29</td>
<td>DSM-IV SCID, Clinical global impressions severity score, HAMD-17</td>
<td>Fair</td>
</tr>
<tr>
<td>Shin et al. (2016)</td>
<td>Korea</td>
<td>Outpatient</td>
<td>2492</td>
<td>49 736</td>
<td>36.75 (6.52)</td>
<td>66.77</td>
<td>CES-D score ≥21</td>
<td>Poor</td>
</tr>
<tr>
<td>Ekinci and Ekinci (2017)</td>
<td>Turkey</td>
<td>Inpatient</td>
<td>139</td>
<td>50</td>
<td>42.2 (12.3)</td>
<td>30.21</td>
<td>DSM-IV</td>
<td>Poor</td>
</tr>
<tr>
<td>Euteneuer et al. (2017)</td>
<td>Germany</td>
<td>Outpatient</td>
<td>98</td>
<td>30</td>
<td>37.3 (12.2)</td>
<td>51.02</td>
<td>DSM-IV</td>
<td>Good</td>
</tr>
<tr>
<td>Gallagher et al. (2017)</td>
<td>Canada</td>
<td>Prospective /population-based</td>
<td>811</td>
<td>5084</td>
<td>67.3 (10.8)</td>
<td>32.18</td>
<td>CES-D ≥4</td>
<td>Poor</td>
</tr>
<tr>
<td>Horsdal et al. (2017)</td>
<td>Denmark</td>
<td>Prospective /population-based</td>
<td>2187</td>
<td>NA</td>
<td>Median age 35.7 years</td>
<td>37.4</td>
<td>ICD-8 and ICD-10</td>
<td>Good</td>
</tr>
<tr>
<td>Jha et al. (2017)</td>
<td>USA, Singapore</td>
<td>Outpatient</td>
<td>106</td>
<td>NA</td>
<td>46.64 (11.89)</td>
<td>30.19</td>
<td>International Neuropsychiatric Interview (MINI)</td>
<td>Good</td>
</tr>
<tr>
<td>Wei et al. (2018)</td>
<td>China</td>
<td>Inpatient</td>
<td>18</td>
<td>15</td>
<td>43.89 (20.78)</td>
<td>27.78</td>
<td>DSM-IV</td>
<td>Good</td>
</tr>
<tr>
<td>Cáceda et al. (2018)</td>
<td>USA</td>
<td>Inpatient</td>
<td>52</td>
<td>NA</td>
<td>36.8 (12.6)</td>
<td>36.54</td>
<td>DSM-IV</td>
<td>Good</td>
</tr>
<tr>
<td>Chamberlain et al. (2018)</td>
<td>UK</td>
<td>Outpatient</td>
<td>198</td>
<td>54</td>
<td>36.5 (NA)</td>
<td>15.15</td>
<td>DSM-5</td>
<td>Good</td>
</tr>
<tr>
<td>Felger et al. (2018)</td>
<td>USA</td>
<td>Outpatient</td>
<td>73</td>
<td>NA</td>
<td>42.1 (11.1)</td>
<td>42.47</td>
<td>SCID-IV</td>
<td>Good</td>
</tr>
<tr>
<td>Osimo et al. (2018b)</td>
<td>UK</td>
<td>Inpatient</td>
<td>137</td>
<td>NA</td>
<td>40(13)</td>
<td>47.44</td>
<td>ICD-10</td>
<td>Good</td>
</tr>
<tr>
<td>Porcu et al. (2018)</td>
<td>Brazil</td>
<td>Outpatient</td>
<td>67</td>
<td>NA</td>
<td>46 (NA)</td>
<td>NA</td>
<td>DSM-5, ICD-10</td>
<td>Good</td>
</tr>
<tr>
<td>Shibata et al. (2018)</td>
<td>Japan</td>
<td>Prospective /population-based</td>
<td>105</td>
<td>2424</td>
<td>65.5 (11.1)</td>
<td>NA</td>
<td>CESD score ≥16</td>
<td>Poor</td>
</tr>
</tbody>
</table>

*CES-D, The Center for Epidemiologic Studies Depression Scale; DSM-IV, Diagnostic and Statistical Manual of Mental Disorders, 4th. Edition; GDS, Geriatric Depression Scale; BDI, Beck’s Depression Inventory; SCID, Structured Clinical Interview for DSM; CAPA, The Child and Adolescent Psychiatric Assessment; ICD, World Health Organisation International Classification of Diseases; PHQ9, Patient Health Questionnaire-9; HAMD-17, Hamilton Depression Rating Scale (HDRS).

Studies evaluated using the Newcastle-Ottawa Scale (see Supplementary methods and Table S1), then converted to Agency for Healthcare Research and Quality - AHRQ - standards (good, fair and poor) using these thresholds:

- Good quality: ≥75% in Selection domain AND ≥50% in Comparability domain AND ≥50% in Outcome domain.
- Fair quality: 50% in Selection domain AND ≥50% in Comparability domain AND ≥50% in Outcome domain.
- Poor quality: ≤50% in Selection domain OR 0% in Comparability domain OR ≤50% in Outcome domain.

"Not available."
was 16% (95% CI 11–23%) and that in depressed patients was 24% (95% CI 17–34%); see Supplementary Figs S6 and S7.

We carried out sensitivity analyses based on five available studies of depressed patients and matched healthy controls that excluded subjects with very high levels of CRP (>10 mg/L). These studies, comprising 3868 patients and 63 212 controls, showed that the prevalence of inflammation in controls was 10% (95% CI 3–26%) and that in patients it was 13% (95% CI 4–36%); see Supplementary Figs S8 and S9. Based on these studies, the meta-analytic OR for inflammation in depressed patients compared with matched controls was 1.44 (95% CI 0.80–2.61; p = 0.23); see Supplementary Fig. S10.

A sensitivity analysis excluding poor quality studies, comprising 5045 patients and 105 372 controls, showed that the
Meta-analytic OR for inflammation in depressed patients compared with matched controls was 1.56 (95% CI 1.29–1.88; \( p < 0.0001 \)); see Supplementary Table S1 and Fig. S11. A further sensitivity analysis only including studies that matched patients and controls by BMI, comprising 2624 patients and 79 887 controls, showed that the meta-analytic OR for inflammation in depressed patients compared with matched controls was 1.59 (95% CI 1.08–2.34; \( p = 0.02 \)); see Supplementary Fig. S12. Finally, a sensitivity analysis excluding studies where depression was not active in all patients showed that the meta-analytic OR for inflammation in depressed patients compared with matched controls was 1.47 (95% CI 1.22–1.75; \( p = 0.02 \)); see Supplementary Fig. S13.

Prevalence of elevated CRP levels (>1 mg/L) in depressed patients

Results based on all available studies

Twenty-five studies comprising 8887 patients with depression were used for this analysis. The meta-analytic pooled prevalence of elevated CRP >1 mg/L in depressed patients was 58% (95% CI 47–69%); see Fig. 3. There was evidence of heterogeneity \( (I^2 = 98.7\%; 95\% \text{ CI } 98.5–98.9\%; \text{Cochrane’s } Q = 1862; p < 0.01) \). Further analyses after grouping studies by setting showed that the prevalence of elevated CRP in inpatient samples \( (N = 1023) \) was 56% (95% CI 46–66%; \( I^2 = 81.8\% \); Cochrane’s \( Q = 22 \)); in outpatient samples \( (N = 697) \) was 59% (95% CI 50–67%; \( I^2 = 74.9\% \); Cochrane’s \( Q = 40 \)); and in population-based samples \( (N = 7167) \) was 57% (95% CI 34–77%; \( I^2 = 99.3\% \); Cochrane’s \( Q = 1774 \)).

Analyses excluding poor quality studies or cases with past depression

A sensitivity analysis excluding four poor quality studies showed that the prevalence of elevated CRP >1 mg/L in depressed patients was 57% (95% CI 43–69%); Supplementary Table S1 and Fig. S14. There was evidence of heterogeneity \( (I^2 = 98.9\%; 95\% \text{ CI } 98.7–99.1\%; \text{Cochrane’s } Q = 1858; p < 0.01) \). A further sensitivity analysis excluding studies where depression was not active in all patients showed that the prevalence of elevated CRP in depressed patients was 58% (95% CI 45–69%); Supplementary Fig. S15. There was evidence of heterogeneity \( (I^2 = 98.8\%; 95\% \text{ CI } 98.6–99.0\%; \text{Cochrane’s } Q = 1861; p < 0.01) \).

Analysis after excluding cases of suspected infection (CRP>10 mg/L)

Eight studies also reported the prevalence of elevated CRP after excluding participants with suspected infection, defined as CRP >10 mg/L. A separate meta-analysis based on these studies, comprising 4456 patients that excluded patients with CRP levels >10 mg/L showed that the prevalence of elevated CRP >1 mg/L was 50% (95% CI 29–72%); see Supplementary Fig. S16. There was evidence of heterogeneity \( (I^2 = 99.1\%; 95\% \text{ CI } 98.9–99.3\%; \text{Cochrane’s } Q = 816; p < 0.01) \).

Association between prevalence of elevated CRP levels (>1 mg/L) and characteristics of depressed patients

Meta-regression analyses did not find any significant association of elevated CRP with sex, age, BMI, non-White ethnicity, being antidepressant-free or sample source (see Supplementary Results).
Assessment of publication bias

A funnel plot of the 25 studies assessing the prevalence of elevated CRP in depression appeared visually symmetrical. Egger’s regression test for funnel plot asymmetry was non-significant ($t = −0.43; df = 23; p = 0.67$), suggesting there was no evidence of publication bias (Supplementary Fig. S17).

Odds ratio for elevated CRP levels (>1 mg/L) in depressed patients

Fifteen studies reported the prevalence of elevated CRP >1 mg/L in 5177 depressed patients and 106 682 matched non-depressed controls (see Supplementary Table S2 for matching details). The meta-analytic OR for elevated CRP in depressed patients compared with matched controls was 1.47 (95% CI 1.18–1.82; $p = 0.0005$); see Fig. 4. There was evidence of heterogeneity ($I^2 = 75.6%; 95\%$ CI 59.8–85.2%; Cochran’s $Q = 57; p < 0.01$).

Based on the same studies, we meta-analysed the prevalence of elevated CRP in depressed patients and matched non-depressed controls separately. The prevalence of elevated CRP >1 mg/L in controls was 44% (95% CI 26–65%) and that in depressed patients was 59% (95% CI 41–75%); see Supplementary Figs S18 and S19.

A sensitivity analysis based on 12 studies after excluding poor quality studies showed that the meta-analytic OR for elevated CRP in depressed patients compared with matched controls was 1.51 (95% CI 1.22–1.88; $p = 0.0002$); see Supplementary Table S1 and Fig. S20. A sensitivity analysis only including the...
nine studies that matched the patients and controls by BMI showed that the meta-analytic OR for elevated CRP >1 mg/L in depressed patients compared with matched controls was 1.52 (95% CI 1.12–2.07; \( p = 0.01 \)); see Supplementary Fig. S21. A further sensitivity analysis excluding studies where depression was not active in all patients showed that the meta-analytic OR for elevated CRP in depressed patients compared with healthy controls was 1.47 (95% CI 1.19–1.81; \( p = 0.003 \)); see Supplementary Fig. S22. Finally, a sensitivity analysis of the four studies excluding subjects with very high levels of CRP (>10 mg/L) showed that the meta-analytic OR for elevated CRP in depressed patients compared with healthy controls was 1.29 (95% CI 0.38–4.30; \( p = 0.68 \)); see Supplementary Fig. S23.

**Very high CRP levels (>10 mg/L) in depressed patients and healthy controls**

We used data from four available studies comprising 3926 patients and 62 748 matched healthy controls. The meta-analytic pooled prevalence of very high CRP in depressed patients matched to healthy controls was 3% (95% CI 1–11%); in the same studies, prevalence of very high CRP in healthy controls matched to depressed patients was 1% (95% CI 0–4%); the meta-analytic OR for very high CRP in depressed patients compared with matched controls was 1.52 (95% CI 1.13–2.05; \( p = 0.006 \)); see Supplementary Figs S24–26.

**Discussion**

To our knowledge, this is one of the first systematic reviews and meta-analyses of the prevalence of low-grade inflammation in patients with depression. We report that a notable proportion of depressed patients show evidence of inflammation. Approximately one in four patients with depression show CRP levels >3 mg/L, a widely used threshold to define low-grade inflammation in the literature. The prevalence is unaltered after excluding poor quality studies, or after excluding studies where depression was not active. After excluding patients with suspected infection, the prevalence of low-grade inflammation is about one in six. We also report that approximately three patients out of five have mildly elevated CRP (>1 mg/L). The prevalence is unaltered after excluding poor quality studies, or after excluding studies where depression was not active. After excluding patients with suspected infection, the prevalence of elevated CRP is one in two. Meta-regression analyses show that the prevalence of inflammation in depression is not associated with sex, age, BMI, ethnicity or sample source.

Using matched non-depressed controls, we quantified the odds ratios of low-grade inflammation and of elevated CRP in depressed patients. We report that the proportion of patients with depression showing elevated inflammatory markers as compared to matched healthy controls is remarkably stable: the ORs were 1.46 for CRP levels >3 mg/L, 1.47 for CRP levels >1 mg/L, and 1.52 for CRP levels >10 mg/L. There was no evidence of publication bias within the included studies, but there was evidence of heterogeneity in all analyses.

Knowing inflammation levels in patients with depression could be important for several reasons, particularly for predicting the risk of physical illness and for predicting response to psychiatric treatment. Inflammation is a potentially causal risk factor for CVD (Pearson et al., 2003), because CVD is associated with circulating IL-6 and CRP levels (Pradhan et al., 2002; Danesh et al., 2003).
et al., 2004; Danesh et al., 2008) and with genetic variants regulating levels/activity of IL-6 (IL6R Genetics Consortium Emerging Risk Factors Collaboration, 2012; Interleukin-6 Receptor Mendelian Randomisation Analysis Consortium, 2012). Depression is co-morbid with CVD (Hare et al., 2013). Depression increases the risk of incident CVD, and is a marker of poor prognosis after myocardial infarction (Nicholson et al., 2006). Inflammation could be a shared mechanism for these conditions. Using Mendelian randomisation analysis of the UK Biobank sample, we previously found that out of all cardiovascular risk factors, IL-6, CRP and triglycerides are likely to be causally linked with depression (Khandaker et al., 2019). Therefore, cardiovascular risk screening in depressed patients who show evidence of inflammation could be useful. Our work suggests that such screening will be relevant for about a quarter of patients with depression.

We focussed on CRP levels as our preferred measure of inflammation because it has been widely used in different fields of medicine to measure inflammation, and standardised cut-offs for CRP exist in the literature. The American Heart Association and Center for Disease Control and Prevention have proposed clear CRP thresholds as indicators of inflammation levels (<1 = ‘low’, 1–3 = ‘medium’, >3 mg/L = ‘high’) (Pearson et al., 2003). Our findings are consistent with previous meta-analyses reporting higher mean concentrations of CRP, IL-6 and other inflammatory markers in depressed patients compared with controls (Howren et al., 2009; Dowlati et al., 2010; Haapakoski et al., 2015; Goldsmith et al., 2016). Our study adds to the literature by providing information on the proportion of depressed patients who have evidence of inflammation.

In addition to depression and CVD, inflammation is associated with other physical and psychiatric disorders including diabetes mellitus (Pradhan et al., 2001), schizophrenia (Miller et al., 2011; Khandaker et al., 2015) and dementias (Schmidt et al., 2006). Inflammation is also an important predictor of increased all-cause mortality (Zacho et al., 2010; Sung et al., 2014; Li et al., 2017). Therefore, routine CRP screening in patients with depression, and identification and treatment of the cause of inflammation could improve overall health-related mortality and morbidity. Public health interventions aimed at reducing inflammation could improve mortality and morbidity associated with a number of conditions.

It is unlikely that anti-inflammatory drugs will be useful for all patients with depression (Khandaker et al., 2017). Measurement of CRP levels could inform patient selection in RCTs of anti-inflammatory drugs for depression. We are aware of two studies that are testing novel anti-inflammatory drugs such as monoclonal antibodies (mAb) against the IL-6/IL-6R pathway. One study testing the efficacy and safety of sirukumab (anti-IL-6 mAb) for depression has completed recruitment (NCT02473289). We are conducting an RCT of tocilizumab (anti-IL-6R mAb) for patients with depression (Khandaker et al., 2018). Both of these studies are based on patients with CRP levels >3 mg/L. Secondary analysis of existing RCTs suggests mAb against specific inflammatory cytokines, such as IL-6/IL-6R, could be helpful for depression (Sun et al., 2017; Kappelmann et al., 2018). However, definitive efficacy trials need to be completed before anti-inflammatory drugs are approved for use in patients with depression.

Studies included in this review varied on setting, country and analytic methods, and the proportion of depressed patients with elevated CRP (>3 mg/L) in these studies ranged between 0% and 60%. In our analyses, the prevalence of inflammation was not associated with participant age and sex, antidepressant treatment, ethnicity or source of sample. This is the case despite the samples spanning all age groups [median age: 42.2 years; interquartile range (IQR): 37–59]. Both sexes were well represented (median proportion of males: 36%, IQR: 17–41%). The samples comprised both antidepressant-free and treated populations (antidepressant-free = 6 studies; 100% treated = 3 studies; mixed populations = 10 studies). Included studies covered samples collected from inpatient (N = 6), outpatient (N = 15) and general population (N = 9). One reason for not detecting an association between inflammation and sociodemographic factors could be that a number of studies matched patients and controls on these factors.

The meta-analytic prevalence of low-grade inflammation (CRP >3 mg/L) in non-depressed controls seen in our analysis is 16%, which is lower than the prevalence of inflammation reported in some general population studies. For instance, Ford et al. (2004) reported the prevalence of low-grade inflammation to be about 25% in a sample of adult US women. Khera et al. (2005) reported the prevalence of CRP >3 mg/L to be >30% in US adult males and females. One reason for these high prevalence reports in general population samples could be that these studies include both healthy and diseased individuals including those with chronic inflammatory physical illness. Therefore, for a more accurate comparison of the prevalence of inflammation between depressed cases and healthy controls, we have used studies that included cases matched to healthy controls for the calculation of odds ratios. In our results, the stability of the odds ratios for elevated CRP in depressed patients compared with healthy controls across different CRP thresholds (ORs = 1.46 for CRP levels >3 mg/L; OR = 1.47 for CRP levels >1 mg/L; and OR = 1.52 for CRP levels >10 mg/L) provides confidence that patients are more likely to have evidence of inflammation than healthy controls. Furthermore, excluding patients with very high levels of inflammation did not significantly affect the odds ratio for low-grade inflammation (>3 mg/L) in depressed subjects (OR = 1.44).

Strengths of this work include the systematic nature of the literature search, which identified a large number of relevant studies comprising 13 541 patients and 155 728 controls from different countries and settings, and spanning diverse ethnic and age groups. The methods were laid out prospectively and published on PROSPERO (Osimo et al., 2018a). We assessed the studies for quality using the validated Newcastle–Ottawa Scale (Stang, 2010). We conducted multiple sensitivity analyses to examine the robustness of the findings. There was no evidence of publication bias, suggesting that we covered a range of results spanning the whole expected distribution of means.

Limitations of this work include sample heterogeneity: the studies included used different methods to assess depression (albeit a valid method was required for inclusion), and samples were recruited from different sources making it difficult to test the association between the prevalence of inflammation and depression severity. However, we have reported meta-analytic results separately by sample source (community, inpatient, etc.), which could be taken as an indirect indicator of depression severity. Inflammation prevalence did not differ much by sample source. However, due to the lack of comparable data on depression severity, we could not assess this directly. Study setting and
sample characteristics could account for some of the observed heterogeneity. We used random-effects meta-analyses in order to take care of inter-study variability. Another limitation is that we were not able to account for comorbidities, partly because for some studies this information was not reported. By design we have focused on dichotomous measure of inflammation, so we cannot comment on the distributions of continuous CRP values in patients/controls; these have been subject to previous meta-analyses reporting higher mean levels of CRP in depression compared with controls (Howren et al., 2009; Dowlati et al., 2010; Haapakoski et al., 2015; Goldsmith et al., 2016).

In summary, this systematic review and meta-analysis provides a robust estimate of the prevalence of low-grade inflammation in depressed patients, which is about one in four. We also report that depressed patients are about 50% more likely to have evidence of inflammation as compared to matched non-depressed controls. These findings are relevant for future treatment studies of anti-inflammatory drugs and for clinical practice, particularly for predicting response to antidepressants and for predicting co-morbid, immune-related physical illness, such as CVD.

Supplementary material. The supplementary material for this article can be found at https://doi.org/10.1017/S0033291719001454

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