Inflammatory insults and mental health consequences: does timing matter when it comes to depression?

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It has become widely accepted that the immune system, and specifically increased levels of inflammation, play a role in the development of depression. However, not everyone with increased inflammation develops depression, and as with all other diseases, there are risk factors that may contribute to an increased vulnerability in certain individuals. One such risk factor could be the timing of an inflammatory exposure. Here, using a combination of PubMed, EMBASE, Ovid Medline and PsycINFO, we systematically reviewed whether exposure to medically related inflammation in utero, in childhood, and in adolescence, increases the risk for depression in adulthood. Moreover, we tried to determine whether there was sufficient evidence to identify a particular time point during the developmental trajectory in which an immune insult could be more damaging. While animal research shows that early life exposure to inflammation increases susceptibility to anxiety- and depressive-like behaviour, human studies surprisingly find little evidence to support the notion that medically related inflammation in utero and in adolescence contributes to an increased risk of developing depression in later life. However, we did find an association between childhood inflammation and later life depression, with most studies reporting a significantly increased risk of depression in adults who were exposed to inflammation as children. More robust clinical research, measuring direct markers of inflammation throughout the life course, is greatly needed to expand on, and definitively address, the important research questions raised in this review.

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Key words: Depressive disorder, early life, inflammation.

Introduction

One of the most important developments in translational mental health is the observation that the inflammatory system is involved in the pathogenesis of major depressive disorder (MDD) (Bufalino et al. 2013; Valkanova et al. 2013; Zunszain et al. 2013; Kiecolt-Glaser et al. 2015; Miller & Raison, 2015). Activation of the immune system in subsets of depressed patients plays a role not only in disease progression, but also in determining the success of antidepressant therapy (Zunszain et al. 2011, 2013; Haroon et al. 2012; Strawbridge et al. 2015). However, the temporal relationship remains largely unclear: is increased inflammation a cause or an effect of MDD?

Interestingly, exposure to increased inflammation may indeed play a causal role in the pathogenesis of depression, as shown from research detailing how a high proportion of patients undergoing treatment with interferon (IFN)-α, a cytokine used for cancer or viral hepatitis C, go on to develop depression (Capuron et al. 2001; Bonacorso et al. 2002; Horikawa et al. 2003; Loftis & Hauser, 2004; Raison et al. 2005). Furthermore, various longitudinal studies support that diabetes mellitus, obesity, and cardiovascular disease, all characterized as low-grade chronic inflammatory states, are significant predictors of depression in later life (Mezuk et al. 2008; Nabi et al. 2010; Hare et al. 2014; Luppino et al. 2015).

However, not everyone exposed to increased inflammation develops depression. As with all other diseases, there are risk factors that contribute to an increased vulnerability in certain individuals. Such risk factors could include the type, severity, frequency and/or the timing of an inflammatory challenge. Indeed, research has already shown that the timing and/or age at exposure may be an important predictor of future psychopathology, with exposure to adverse experiences particularly in early life being consistently associated with increased susceptibility to a variety of neuropsychiatric disorders (Turecki et al. 2014; Visser et al. 2014; Kalnakis & Chandler, 2015; Trotta et al. 2015).

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In two prospective studies, although depression in adulthood was associated with an accumulation of stressors across the life course, most originated in the first years of life (Clark et al. 2010; Colman et al. 2014). Furthermore, childhood adversity was either directly associated with adolescent, early adulthood, and mid-life affective disorder psychopathology (Clark et al. 2010), or was associated with intermediate risk factors that subsequently increased the risk of future depression (Colman et al. 2014). Thus, it would seem that adversity in early life has important effects on the life course of depression, and that timing of such adversity may be an important factor in the etiology of the disorder.

Admittedly, there may even be a time point in early life when individuals are most vulnerable to adverse exposure. In 2012, Bosch and colleagues demonstrated how the sequelae of early life adversity depended on the age at the time of exposure, showing how timing of an adverse event could differentially alter the functioning of the hypothalamic-pituitary-adrenal (HPA) axis in later life. Specifically, they showed how hypercortisolism was a potential consequence of adverse exposure between 6 and 11 years, while adverse exposure between 12 and 15 years contributed to hypocortisolism. Interestingly, individuals exposed before the age of five had no alterations in stress responsivity (Bosch et al. 2012). This suggests that the first 5 years of life could represent a stress-hypo-responsive period (Sapolsky & Meaney, 1986) – a temporary and well-documented developmental period in life characterized by attenuated stress responsivity (Gunnar et al. 1996; Larson et al. 1998; Gunnar & Donzella, 2002) – the evolutionary purpose of which is thought to promote maternal-infant attachment, with low levels of cortisol attributed to maintaining and reinforcing this attachment (Moriceau & Sullivan, 2007). Poignantly, impaired maternal-infant attachment has been associated with poor emotional outcomes in offspring (Leckman-Westin et al. 2009), so further understanding of this physiological phenomenon could help extend our knowledge on the etiology of depression. Indeed, the idea that the stress system (and perhaps even the immune system) may be biologically more or less responsive depending on developmental age could explain why certain types of exposures at certain times in life may exert differential outcomes, and this study emphasizes not only the importance of the type of adversity and the frequency of its occurrence, but also the timing of exposure.

To date, no clinical study has established whether there exists a critical period in life when adversity in the form of a medically-related immune insult may increase one’s susceptibility to mental illness. However, one of the hallmarks of the developing immune system is that it exhibits an increased sensitivity for environmentally induced toxicity compared to the fully matured immune system of an adult (Dietert et al. 2000; Holladay & Smialowicz, 2000; Dietert, 2013). Moreover, an early life inflammatory insult can result in the impairment of a variety of biological systems involved in the etiology of MDD, including the neuroendocrine system (Rivest, 2010). Indeed, it seems highly plausible that a ‘critical window’ of vulnerability exists for immune system activation on mental health susceptibility, and that this window of vulnerability could exist when the immune system is still maturing in early life, which starts in utero, and continues until the age of 15 years (Hannet et al. 1992; Osugi et al. 1995; Holt & Jones, 2000).

**Review objectives**

The primary objective of this article is to review the current clinical literature in order to elucidate whether exposure to an inflammatory insult early in life, driven by medical or infective causes, increases the risk for depression later in life. Given that neural development extends from the embryonic period through to adolescence (Rice & Barone, 2000; Johnson, 2001), and that this coincides with the development of the immune system (Hannet et al. 1992; Osugi et al. 1995; Holt & Jones, 2000) we focus on studies reporting exposure to increased inflammation in three developmental life stages: antenatal, childhood (birth to age 12), and adolescence (age 13–18). We specifically exclude studies where inflammation was driven by exposure to psychosocial stress or adversity. Being able to identify when during the developmental trajectory an immune insult is more detrimental could have significant implications in terms of developing successful prevention strategies, raising awareness, and targeting more vulnerable individuals, and as such, it is important to establish whether timing does indeed matter.

**Method**

**Search strategy and limits**

A combination of PubMed, EMBASE, Ovid Medline and PsycINFO databases were used to systematically select studies for discussion, and reference lists of selected papers were manually searched to check for any additional studies. An independent systematic search of all aforementioned databases was carried for each life stage under study.

We included only clinical studies that were longitudinal in nature, and where there was a minimum of 2 years between exposure and mental health assessment, thereby minimizing any potential contamination of measures. No publication date restrictions were imposed,
but our searches were limited to English-language studies only.

**Inflammatory exposure: definition and limits**

We defined an inflammatory challenge as any illness pertaining to increased inflammation, which we classified as either a direct, or indirect, immune challenge.

A direct immune challenge was characterized as any bacterial, fungal, parasitic, allergic and/or viral infection/illness, either chronic or acute, in which the primary host’s response to infection/illness was inflammation. Chronic illnesses resulting in the sustained use of immunosuppressant medication, or any autoimmune disease/condition resulting in organ transplantation were excluded.

An indirect immune challenge was defined as any exposure to an illness/condition characterized as a systemic inflammatory state. Substantial evidence chronicles the activation of the immune system in diabetes mellitus (gestational, types I and II) (Donath & Shoelson, 2011; Calle & Fernandez, 2012), obesity (Kredel & Siegmund, 2014; de Jong et al. 2014), congenital heart disease (Sharma et al. 2003; Allan et al. 2010), and cardiovascular disease (Hansson, 2005; Mangge, 2014). Reciprocity between inflammation and these conditions have been consistently demonstrated, such that they are typically referred to as chronic low-grade inflammatory states. As such, we included all studies reporting (a) exposure to maternal obesity and diabetes antenatally (in utero), and (b) living with diabetes, cardiovascular disease, and/or obesity postnatally, in relation to the development of depression in later life. Although cancer is a well-known inflammatory state (Payne, 2014; Roxburgh & McMillan, 2014), due to the vast heterogeneity of this disease, studies on cancer were excluded.

**Depressive psychopathology: definition and limits**

Depressive psychopathology was conceptualized by the use of affective/mood symptoms and diagnoses. Most studies assessed depression and/or depressive symptomology in adult participants, i.e. aged >18 years, but we also included papers reporting symptoms or diagnoses of depression in adolescent participants, as the effect(s) of early-life adversity on mental health may have emerged by this time point (Pawlby et al. 2011, Plant et al. 2015). For studies assessing mental health in adolescence, we also accepted papers reporting symptoms of internalizing and externalizing behaviours, which have been shown to predict adulthood affective disorders (Roza et al. 2003; Clark et al. 2010).

For a full list of key words used in our searches see Supplementary online Appendix.

**Results**

Fig. 1 highlights the number of studies identified at each stage of our search strategy for all stages combined. Of the 10 087 papers initially flagged up, only 22 clinical studies were eligible and ultimately included in our review.

**Exposure to increased inflammation antenatally**

Table 1 displays all clinical studies that directly and indirectly investigated the effect of an antenatal inflammatory challenge on depression susceptibility in later life.

**Exposure to maternal infection in utero**

Although several studies have associated maternal infections with adult psychiatric disorders, predominantly in schizophrenia, autism spectrum disorder, epilepsy and cerebral palsy (Knuesel et al. 2014), the impact of maternal infection on depression in offspring is less well-known. To date, only six clinical papers examined whether exposure to infection during pregnancy increased risk of offspring depression (Brown et al. 1995; Machón et al. 1997; Mino et al. 2000; Mellins et al. 2003; Gaughan et al. 2004; Pang et al. 2009). In one such study, a significant increase in depression was reported for individuals exposed during their second trimester to a Finnish influenza epidemic compared to control subjects born 6 years prior to the outbreak (13% v. 2%) (Machón et al. 1997). In contrast, two studies examining patients with mood disorders born during an influenza epidemic in Holland and Japan found no significant differences in the risk for depression in patients born during the epidemics (Brown et al. 1995; Mino et al. 2000). However, the contradictory findings between these studies could partially be explained by other environmental factors such as diet, and indeed, low incidence rates of inflammatory disease in Greenland Inuit and Japanese people, thought to be attributed to the large consumption of fish containing omega-3 fatty acids, has previously been reported (Simopoulos, 2002). Thus, it seems plausible that women in the Japanese cohort may have had diets rich in omega-3 fatty acids, which consequently may have bestowed some protection against depression (Lin & Su, 2007; Sublette et al. 2011; Martins et al. 2012).

Recently, however, a larger UK study examining the effect of a variety of prenatal viral infections and offspring depression found no overall increased risk for depression associated with viral exposure (Pang et al. 2009). Similarly, two large studies investigating the impact of antenatal exposure to human immunodeficiency virus (HIV) on the subsequent development of depression in later life found that HIV exposure did
not predict the development of poor emotional and/or behavioural outcomes. Although a high prevalence of behavioural problems did exist among HIV-infected children, these studies found that neither HIV infection nor prenatal drug exposure was the underlying cause (Mellins et al. 2003; Gaughan et al. 2004).

**Exposure to gestational diabetes and obesity in utero**

Interestingly, no studies on the effect of gestational diabetes or obesity on increased risk of offspring depression were found. Although gestational diabetes has been linked to altered brain development and behaviour in offspring, specifically in autism spectrum disorder (Xu et al. 2014) and attention-deficit hyperactivity disorder (Nomura et al. 2012), no study was found in relation to depression.

**Exposure to increased inflammation in childhood**

Clinical studies examining whether direct or indirect exposure to inflammation in childhood predicts depression in later life are given in Table 2.

**Chronic illness in childhood and depression in later life**

Eight studies pertaining to chronic or acute illness in childhood were found (Pless et al. 1989; Kokkonen & Kokkonen, 1993; Cohen et al. 1998; Packham et al., 2002; Gaughan et al. 2004; Goodwin, 2011; Ferro & Boyle, 2015; Khandaker et al. 2014). Of these, five used a broad definition of infection/illness, and as such had heterogeneous diagnostic groups, while the other two referred to a specific infection/illness.

*Exposure to heterogeneously defined physical illness in childhood*

One British birth cohort investigated the effect of chronic illness in childhood on the mental health well-being of participants at ages 26 and 36 years (Pless et al. 1989). The authors found no overall significantly increased reports of psychiatric disorders in chronically ill children compared to healthy controls (men: 6.1% v. 3.8%; women: 14.1% v. 9%). However, they did find that participants who were ill in childhood and again after 21 years of age were significantly more likely to have a psychiatric disorder compared to other members of the cohort. Additionally, when cohort members were re-interviewed at 36 years of age, specifically to assess current affective state, women who experienced childhood chronic illness had significantly higher depressive scores compared to controls. This study highlights that (a) repeated exposure to medically-related inflammation, across two life stages, may be necessary for eliciting psychopathology in some individuals, and (b) the effect of early life illness may operate only for a subset of individuals, with women, in this instance, being more vulnerable.

In a second study, the prevalence rates of mental disorders in adults who suffered from a variety of chronic
<table>
<thead>
<tr>
<th>Study</th>
<th>Objectives</th>
<th>Sample/design</th>
<th>Inflammatory exposure</th>
<th>Timing of exposure</th>
<th>Psychopathological outcome</th>
<th>Time of outcome</th>
<th>Findings</th>
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<tr>
<td><strong>Direct inflammatory insult</strong></td>
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<td>Brown <em>et al</em>. 1995</td>
<td>To examine whether exposure to an influenza epidemic in Holland would increase the risk for adult affective disorders</td>
<td>980697 participants (193,701 exposed) Ecological Study</td>
<td>Potential exposure to A2 influenza virus Those that were <em>in utero</em> from Sept. 1957 to July 1958</td>
<td>Antenatally</td>
<td>Major depressive disorder</td>
<td>23–31 years</td>
<td>a</td>
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<tr>
<td>Machón <em>et al</em>. 1997</td>
<td>To determine whether exposure to an influenza epidemic in Finland would increase the risk for adult major affective disorder</td>
<td>1378 participants (163 exposed) Ecological study</td>
<td>Potential exposure to A2/Singapore influenza virus All adults born from Nov. 1957, to Aug. 1958 with diagnosis of depression</td>
<td>Antenatally</td>
<td>Major depressive disorder</td>
<td>&gt; 30 years</td>
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<tr>
<td>Mino <em>et al</em>. 2000</td>
<td>To examine the relationship between the birth of patients with mood disorders and influenza epidemics in Japan</td>
<td>361 participants (61 exposed) Ecological study</td>
<td>Potential exposure to 2 strains of influenza virus: A2 Asian, and AB mixed type influenza All adults born 5 months after each wave of influenza: June-July 1957, Nov.-Dec. 1957, Apr.-May 1958, July-Sept. 1962 and 1965</td>
<td>Antenatally</td>
<td>Depressive disorder Diagnosed using ICD-10</td>
<td>&gt; 30 years</td>
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<tr>
<td>Mellins <em>et al</em>. 2003</td>
<td>To examine the long-term effects of <em>in utero</em> exposure to human immunodeficiency virus (HIV)</td>
<td>307 participants (96 HIV-infected) Prospective study</td>
<td>HIV infection Confirmed by routine clinical and laboratory/virology evaluations</td>
<td>Antenatally</td>
<td>Emotional and behavioural problems Conner’s Parent Rating Scale (CPRS), parental report</td>
<td>3–17 years</td>
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<tr>
<td>Gaugan <em>et al</em>. 2004</td>
<td>To examine the long-term effects of <em>in utero</em> exposure to human immunodeficiency virus (HIV)</td>
<td>2298 HIV-infected and 1021 HIV-exposed, uninfected Prospective study</td>
<td>HIV infection Confirmed by routine clinical and laboratory/virology evaluations</td>
<td>Antenatally</td>
<td>Depression</td>
<td>&gt;15 years</td>
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physical illnesses in childhood were compared to age-matched controls (Kokkonen & Kokkonen, 1993). Similar to findings from Pless and colleagues, this study found no significant difference in the prevalence rates of all types of depression in young adults with childhood illness compared to healthy controls (13% v. 12%). However, severe depression was significantly more common in patients exposed to childhood illness than in controls (6% v. 2%). Again, we find only a subset of vulnerable individuals, and, as in this case, unless studies examine diagnosis by severity, participants with milder forms of depression may mask the association between early life illness and more severe forms of depression.

Contrary to the findings of the two aforementioned studies, which all demonstrate an increased vulnerability to depression in only a subset of participants suffering from illness as children, three more recent studies find an overall significant association between childhood infection and the mental health of their cohort (Cohen et al. 1998; Goodwin, 2011; Ferro & Boyle, 2015). A large US cohort found that chronic physical illness in childhood predicted an increased risk of future depression in both adolescence [odds ratio (OR) 3.81, 95% confidence interval (CI) 1.55–9.39] and young adulthood (OR 4.04, 95% CI 1.54–10.62) independent of prior depressive episodes and other demographic covariates. Moreover, the authors also showed how immunologically mediated disorders, specifically atopic illness, hay fever and mononucleosis, exhibited strong associations with subsequent onset of depression in both adolescence and young adulthood (Cohen et al. 1998). Similarly, another large, but retrospective study, examined the association between infection in the first year of life and mental disorders among youth in a community sample, and reported that early life infection was associated with significantly increased odds of depression (OR 3.7, 95% CI 1.0–13.4) (Goodwin, 2011). However, despite the claim of significantly increased odds of depression, we should be mindful that the confidence intervals included one. Finally, in congruence with both Cohen and colleagues, and Goodwin, another large study found that those chronically ill in childhood reported significantly more symptoms of depression in adolescence compared to healthy controls (OR 2.71 v. 2.36) (Ferro & Boyle, 2015).

Exposure to specific physical illness in childhood: taking inflammatory markers into account

Thus far there appears to be fairly strong evidence to suggest that exposure to illness in childhood increases the risk for depression in later life, particularly for a subset of individuals. However, none of the
Table 2. Studies examining the association between childhood exposure to inflammation and risk for depression in later life

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<tr>
<th>Study</th>
<th>Objectives</th>
<th>Sample/design</th>
<th>Inflammatory exposure</th>
<th>Timing of exposure</th>
<th>Psychopathological outcome</th>
<th>Time of outcome</th>
<th>Findings</th>
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<tr>
<td>Direct inflammatory insult</td>
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<tr>
<td>Gaughan et al. 2004</td>
<td>To examine the long-term effects of postnatal exposure to human immunodeficiency virus (HIV)</td>
<td>2298 HIV-infected and 1021 HIV-exposed, uninfected</td>
<td>HIV infection</td>
<td>Median age 10 years</td>
<td>Depression</td>
<td>&gt;15 years</td>
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<td></td>
<td></td>
<td></td>
<td>Confirmed by routine clinical and laboratory/ virology evaluations</td>
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<td>Assessment for Children. Incidence of psychiatric hospitalizations obtained from the National Hospital Discharge Survey (NHDS)</td>
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<tr>
<td>Cohen et al. 1998</td>
<td>To assess the association between early life illness and mental health in later life</td>
<td>774 participants (233 chronically ill)</td>
<td>Used a checklist of chronic conditions (e.g. heart problems, chronic respiratory conditions, chronic pain, orthopaedic problems)</td>
<td>1–10 years</td>
<td>Depressive disorder</td>
<td>At 13, 16 and 22 years</td>
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<td></td>
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<td>Measured by both self, and mothers, report</td>
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<td>Packham et al. 2002</td>
<td>To investigate the effect of juvenile idiopathic arthritis (JIA) on mental health in later life</td>
<td>246 participants with JIA (no control group)</td>
<td>Meeting standard criteria for JIA</td>
<td>0–16 years</td>
<td>Depression</td>
<td>18–71 years</td>
<td>-b</td>
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<td>Measured by interview, clinical examination and notes review by the same rheumatologist</td>
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<td>Measured with the Hospital Anxiety and Depression (HAD) scale</td>
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<tr>
<td>Goodwin, 2011</td>
<td>To determine the association between bacterial infection in early life and mental health in a community sample</td>
<td>1285 participants (14 with infection) Retrospective design</td>
<td>A severe infection needing antibiotics Measured by parental report</td>
<td>0–1 year</td>
<td>Depressive disorder</td>
<td>9–17 years</td>
<td>-b</td>
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<td>Diagnosis determined using the DISC</td>
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<tr>
<td>Khandaker et al. 2014</td>
<td>To test whether higher serum levels of IL-6 and CRP would increase future risk for depression</td>
<td>4415 participants</td>
<td>Serum levels of IL-6 and CRP Measured in non-fasting blood samples</td>
<td>9 years</td>
<td>Depression</td>
<td>18 years</td>
<td>-b</td>
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<td>Diagnosis determined by Clinical Interview Schedule-Revised (CIS-R) and Mood and Feelings Questionnaire (MFQ)</td>
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<td>Study</td>
<td>Objectives</td>
<td>Sample/design</td>
<td>Inflammatory exposure</td>
<td>Timing of exposure</td>
<td>Psychopathological outcome</td>
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<tr>
<td>Ferro &amp; Boyle, 2015</td>
<td>To evaluate the impact of chronic physical illness on depression and anxiety</td>
<td>10,646 participants (1932 with illness)</td>
<td>Asthma, cerebral palsy, epilepsy, heart condition, kidney condition, any other long-term condition</td>
<td>0–11 years</td>
<td>Anxiety and Depression</td>
<td>14–15 years</td>
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<td></td>
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<td>Prospective study</td>
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<td>Ontario Child Health Study Checklist</td>
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<tr>
<td>Pless et al., 1989</td>
<td>To investigate the effect of chronic physical illness on mental health in later life.</td>
<td>5,362 participants (467 chronically ill)</td>
<td>Any physical, non-fatal condition lasting less than 3 months in a given year. Repeated episodes of acute physical illness excluded Measured by parental report and cross-referenced with hospital records</td>
<td>&lt;15 years</td>
<td>Emotional disturbance, using self-report</td>
<td>26 years</td>
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<td></td>
<td></td>
<td>Prospective study</td>
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<td>Affective state, using Present State Examination</td>
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<tr>
<td>Kokkonen &amp; Kokkonen, 1993</td>
<td>To determine whether chronic physical illness in childhood increases the risk for later life depression</td>
<td>530 participants (407 chronically ill, age-matched to 123 controls)</td>
<td>Chronic disorders: asthma, diabetes, epilepsy, growth hormone deficiency, motor handicaps, rheumatoid arthritis, congenital heart disease</td>
<td>Childhood, age range not defined</td>
<td>Depression</td>
<td>18–25 years</td>
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<tr>
<td></td>
<td></td>
<td>Prospective study</td>
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<td></td>
<td>Assessed by Present State Examination</td>
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<td>Indirect inflammatory insult</td>
<td>Kovacs et al., 1997</td>
<td>To determine prevalence rates and risk factors for psychiatric disorders associated with type 1 diabetes mellitus</td>
<td>Diagnosis of classic, acute-onset type 1 diabetes mellitus</td>
<td>8–13 years</td>
<td>Depressive disorder</td>
<td>Median 20 years</td>
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<td></td>
<td></td>
<td>92 participants with diabetes mellitus (no control group) Longitudinal, naturalistic design</td>
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<td></td>
<td>Assessed using standardized, semi-structured, symptom-based Interview Schedule for Children and Adolescents (ISCA)</td>
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<tr>
<td>Areias et al., 2013</td>
<td>To test for the effects of different demographic, clinical and psychosocial variables on psychiatric morbidity of participants with congenital heart disease (CHD).</td>
<td>150 CHD patients (no appropriate control group)</td>
<td>CHD. Identified through the paediatric cardiology or adult cardiology outpatient clinic</td>
<td>0–18 years</td>
<td>Major depressive disorder</td>
<td>26 years</td>
<td>c</td>
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<td></td>
<td></td>
<td>Retrospective design</td>
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<td>Assessed using schedule for affective disorders and schizophrenia (SADS-L) interview, YSR (Youth Self-Report) and ASR (Adult Self-Report) to assess recent behavioural and emotional problems</td>
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</table>
aforementioned studies have directly measured levels of inflammation, a concept we should be mindful of in the context of this review. However, there exist three clinical studies that provide some insight into the direct effect of inflammation on depression susceptibility.

One such study, assessing adults with juvenile idiopathic arthritis (JIA), found that patients with systemic-onset JIA had significantly higher levels of depression (10.7%) compared to other JIA subsets. Interestingly, the study also found that depression was most commonly seen when the age of onset was between 6 and 12 years (11.1%) compared to early (2.7%) or late (0%) onset, and that the first episode of depression tended to be between ages 15 and 25 years (38.5%). However, although these findings are highly novel in that they are the first to demonstrate that JIA onset specifically in childhood may exert a greater influence over the subsequent development of depression, the study additionally highlighted how joint inflammation based on the Thompson–Kirwan scale, likely representing the magnitude of systemic inflammation, was not a significant predictor of depression in this cohort (Packham et al., 2002).

Similarly, another study examining the long-term effects of postnatal exposure to HIV on mental health in later life found that although HIV infected children were at increased risk for depression compared to healthy controls, immunological and virological markers were not responsible for predicting first admission hospitalizations for depression (Gaughan et al. 2004).

Interestingly, these studies emphasize how psychological variables, rather than the acute effects of specific inflammatory factors, may explain the majority of variance seen in later life depression. However, one population-based study showed how participants with increased levels of systemic inflammation in childhood, indicated by higher serum levels of interleukin-6, were at increased risk for depression compared to healthy controls and behavioural problems (Khandaker et al. 2014). Unlike many of the already discussed papers, this study’s strength was that it controlled for adversity-related causes of inflammation, as well as children with medical conditions. Therefore, these findings are unique in that they show how raised levels of inflammation, not likely accounted for by adversity, infection or illness, can predict future psychopathology.

To compare mood state profiles in adult patients with childhood-onset and adulthood-onset type 1 diabetes mellitus, 214 participants with diabetes mellitus (no control group) were assessed by psychologists. Diagnosis of classic, acute-onset type 1 diabetes mellitus was identified through the Lithuanian Diabetes Registry. Evaluations were performed using the Profile of Mood States (Lašaitė et al. 2015). The results are as follows:

- No differences in risk of depression.
- Significantly increased risk of depression.
- Significantly increased risk of depression for particular subset of cohort.
Living with obesity, diabetes mellitus and congenital heart disease in childhood and later life depression

Widening our search to incorporate studies investigating the long-term impact of childhood obesity, congenital heart disease, and type I diabetes mellitus yielded an additional three papers.

Childhood obesity and depression in later life

Although previous studies have posited a link between adiposity and depression, with inflammation playing a key role in the disorder’s pathogenesis (Shelton & Miller, 2011), no published clinical studies investigating the impact of childhood obesity on future depression were found. However, this is unsurprising since only recently has the impact of obesity in relation to mental health been under full investigation. Given the time and resources involved in conducting longitudinal studies, insights into the impact of childhood obesity on mental health in later life is likely to emerge in subsequent years.

Type I diabetes mellitus and depression in later life

Two clinical studies investigating the effect of diabetes in childhood on affective disorder psychopathology were found, with contradictory conclusions as to when the sensitive period for disease onset may lie. One study reported an increased risk for depressive disorders in early adulthood for childhood-onset diabetic patients (Kovacs et al. 1997), while the other found that adulthood onset type I diabetic women reported higher levels of depression than childhood-onset diabetic patients (Lašaitė et al. 2015).

Congenital heart disease and depression in later life

The influence of congenital heart disease on later life affective disorder psychopathology has been investigated in only one longitudinal study, which found no overall difference in the mental health outcomes of these patients and healthy controls. However, similar to previous findings from other studies, the study did find that for a subset of the cohort – female patients, and those with more complex forms of the disorder – significantly higher levels of anxiety and depression were reported. Moreover, the authors found that age at assessment was important for evaluating the impact of these disorders on later mental health, finding that those aged 19–26 years had more symptoms of anxiety/depression than those aged 12–18 years (Areias et al. 2013).

Exposure to increased inflammation in adolescence

Table 3 displays all clinical studies that have assessed whether direct or indirect exposure to increased inflammation in adolescence predicts future depression.

Chronic illness in adolescence and depression in later life

A comprehensive search of the literature yielded one study exploring a direct link between adolescent infection/illness and the development of future depression. However, whilst the data for infection during adolescence was lacking, we found one large study evaluating the relationship of allergic rhinitis (AR) to the development of any depressive disorder in later life (Chen et al. 2013). In this study, adolescents with AR had a significantly higher prevalence of major depression (2.5% v. 1.2%) and any depressive disorder (4.9% v. 2.8%) in later life compared to control subjects.

Living with obesity, diabetes and cardiovascular disease in adolescence and depression in later life

Thus far only one study pertaining to adolescent direct exposure has been identified, which has limited our ability to identify any patterns. Broadening our search to incorporate studies looking at the effect of indirect inflammatory conditions yielded an additional five papers.

Adolescent obesity and depression in later life

Three studies exploring the relationship between obesity in adolescence and the subsequent development of future depression were identified. One study reported an overall positive association between obesity and depressive symptoms in adulthood, finding that a higher body mass index (BMI) at age 14 correlated with higher BMI, leptin, C-reactive protein, and depressive symptoms at age 17. Moreover, the study found that females who were obese in both adolescence and adulthood more frequently reported symptoms of depression (Herva et al. 2006). Interestingly, the remaining two studies found no overall association between adolescent obesity and depression, but did find that the increased risk for the development of future depression was gender specific. Anderson and colleagues found that adolescent obesity in females, but not in males, predicted an increased risk for the subsequent development of depression and anxiety disorder (Anderson et al. 2007). Similarly, Marmorstein and colleagues reported how only obesity in female adolescents predicted the onset of depression in early adulthood: specifically, it was an onset of obesity after the age of 14 that predicted the development of...
Table 3. Studies examining the association between adolescent exposure to inflammation and risk for depression in later life

<table>
<thead>
<tr>
<th>Study</th>
<th>Objectives</th>
<th>Sample/design</th>
<th>Inflammatory exposure</th>
<th>Timing of exposure</th>
<th>Psychopathological outcome</th>
<th>Time of outcome</th>
<th>Findings</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Direct inflammatory insult</strong></td>
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<tr>
<td>Chen et al. 2013</td>
<td>To assess whether allergic rhinitis (AR) increases the risk of depression in later life</td>
<td>8365 participants (1673 with AR)</td>
<td>Diagnosis of AR</td>
<td>12–15 years</td>
<td>Depressive disorder</td>
<td>&gt;22 years</td>
<td>No differences in risk of depression.</td>
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<tr>
<td></td>
<td></td>
<td>Prospective study</td>
<td>Based on World Health Organization classification system, using the International Classification of Disease (ICD-10)</td>
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<tr>
<td>Jacobson et al. 1997</td>
<td>To evaluate the psychological adjustment of young adults with diabetes mellitus</td>
<td>111 participants (57 with diabetes mellitus)</td>
<td>Diagnosis of type 1 diabetes mellitus</td>
<td>9–16 years</td>
<td>Depression</td>
<td>19–26 years</td>
<td>Significantly increased risk of depression.</td>
</tr>
<tr>
<td>Herva et al. 2006</td>
<td>To examine the association between obesity and depression</td>
<td>10,096 participants (377 with obesity)</td>
<td>Obesity classified as a BMI &gt;23.43 kg/m² (males), 23.81 kg/m² (females)</td>
<td>14 years</td>
<td>Depressive disorder</td>
<td>31 years</td>
<td>Significantly increased risk of depression for particular subset of cohort.</td>
</tr>
<tr>
<td>Anderson et al. 2007</td>
<td>To assess whether adolescent obesity is associated with risk for major depressive disorder (MDD) or anxiety disorder in later life</td>
<td>701 participants (45 with obesity)</td>
<td>Obesity classified as a BMI &gt;95th percentile</td>
<td>9–18 years</td>
<td>Depressive disorders</td>
<td>28–39 years</td>
<td>Significantly increased risk of depression for particular subset of cohort.</td>
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<tr>
<td></td>
<td></td>
<td>Prospective study</td>
<td>Baseline based on parental report; follow up based on self-report</td>
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<tr>
<td>Marmorstein et al. 2014</td>
<td>To examine prospective associations between obesity from early adolescence and early adulthood and depression</td>
<td>1512 participants</td>
<td>Obesity classified as a BMI &gt;95th percentile</td>
<td>11–24 years</td>
<td>Depressive disorder</td>
<td>14–24 years</td>
<td>Significantly increased risk of depression for particular subset of cohort.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Prospective twin study</td>
<td>Height and weight measured using a Detecto mechanical physician scale</td>
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<tr>
<td>Lašaitė et al. 2015</td>
<td>To compare mood state profiles in adult patients with childhood-onset and adulthood-onset type 1 diabetes mellitus</td>
<td>214 participants with diabetes mellitus (no control group)</td>
<td>Diagnosis of classic, acute-onset type 1 diabetes mellitus</td>
<td>0–18 years</td>
<td>Tension-anxiety, depression-dejection</td>
<td>&gt;18 years</td>
<td>Evaluated using the Profile of Mood States</td>
</tr>
</tbody>
</table>

*a* No differences in risk of depression.  
*b* Significantly increased risk of depression.  
*c* Significantly increased risk of depression for particular subset of cohort.
depression in early adulthood among females in the cohort (Marmorstein et al. 2014).

**Type 1 diabetes mellitus and depression in later life**

With respect to the impact of diabetes on depression susceptibility in later life only two clinical studies were identified. Both studies found no difference in the psychological outcome of adults diagnosed with diabetes in adolescence compared to healthy controls (Jacobson et al. 1997; Lašaitė et al. 2015). Interestingly, however, Jacobson and colleagues did find that in early adulthood individuals with diabetes had lower self-esteem (Jacobson et al. 1997), a considered predisposing factor for depression (Ferro & Boyle, 2015). Therefore, it is possible that these individuals may develop depression in later adulthood.

**Cardiovascular disease in adolescence and depression in later life**

Regarding the effect of cardiovascular disease in adolescence on mental health, no studies were found. However, this was anticipated given that the onset of cardiovascular disease is typically in adulthood.

**Conclusion**

**Main findings**

We have reviewed all available literature examining the effect of a medically related inflammatory challenge in early life, i.e. antenatally, in childhood and in adolescence, on depression susceptibility in later life. We found no clinical evidence to support that inflammation in utero contributes to an increased risk of developing future depression. This was somewhat surprising given that (a) animal research consistently supports that increased inflammation neonatally increases depressive-like behaviour in later life (Walker et al. 2004, 2006, 2008, 2009; Bilbo et al. 2005; Spencer et al. 2006, 2011; Galic et al. 2008; Roque et al. 2014), and (b) stress, particularly during pregnancy, is a potential predictor for later life psychopathology (Betts et al. 2015; Slykerman et al. 2015; Biaggi et al. 2016), which is pertinent given the bi-directional relationship between stress and inflammation (Chovatiya & Medzhitov, 2014; Slavich & Irwin, 2014). However, given that cytokines do not cross the placenta barrier in normal term foetuses, this may contribute for our findings (Zaretzky et al. 2004; Aaltonen et al. 2005).

However, we did find some converging evidence to support that exposure to increased inflammation in childhood increases the risk for adult depression. Furthermore, the evidence suggests that persistent physical health problems in childhood may relate to the presence of greater or more severe psychiatric disorders, and that psychiatric outcome may be gender specific, with females being more vulnerable to exposure. Interestingly, the evidence pertaining to studies assessing exposure to severe conditions in childhood was mixed, but did seem to suggest that for lifelong conditions, such as HIV, diabetes and congenital heart diseases, increased inflammation does not increase susceptibility to depression. However, it is important to note that, for such conditions, treatment strategies may be reducing the overall increased inflammation associated with disease state. Finally, when looking at the direct effect of inflammation in childhood on the susceptibility to depression, the evidence was both limited in quantity and inconsistent in findings, emphasizing the need for further clinical research directly measuring inflammatory markers and evaluating their role in the aetiology of depression.

Looking at adolescent exposure to increased inflammation on future depression risk, a limited quantity of evidence was found. However, we did find some evidence to support that an indirect immune challenge in the form of obesity may increase the risk for future depression, particularly for female adolescents.

**Limitations of the existing literature**

It is noteworthy that fewer clinical studies were identified for the antenatal and adolescent life stages, and this made it extremely difficult to establish consistent patterns pertaining to these developmental phases from the available evidence. Although we did find some evidence to support that increased inflammation in childhood, and to some extent in adolescence, increases depression risk in later life, many of these studies did not look specifically at levels of inflammation in their cohorts, and as such we cannot conclude that it was increased inflammation per se that predicted depression. Indeed, inflammatory markers were unavailable for the majority of the studies reviewed, predominately because their research question was not addressing the effect of increased inflammation on depression, and as such, this represents a major empirical limitation of conclusions in this review. Other psychological or biological factors associated with these disease states may be accounting for the observed associations.

Furthermore, there are several other limitations that weaken the evidence overall. First, most studies relied on self-reports and/or parental reports of physical illness and affective disorder psychopathology, and it is uncertain whether similar findings would be found with physician-diagnosed medical illnesses. Second, in some studies neither the definition nor timing of physical illness or psychopathology was clearly
described. Indeed, several studies assessed exposure to infection across multiple developmental stages, i.e. exposure any time from childhood to early adulthood, and given the large biological and psychosocial differences between the developmental stages, results should be interpreted with caution. Third, important confounders, such as perinatal complications, parental mental health, stress (biological and psychosocial), and childhood maltreatment were not measured and controlled for in analyses in most of the studies. We should be particularly cautious in attributing the observed associations found, especially given the evidence substantiating the involvement of the HPA axis (Pariante & Lightman, 2008), childhood maltreatment (Pawlyby et al. 2011; Lindert et al. 2014; Plant et al. 2015), parental psychopathology (McLaughlin et al. 2012), and obstetric complications (Rääkkönen et al. 2008; Tuovinen et al. 2010) in the pathogenesis of depression. Fourth, none of the included studies controlled for infections/illnesses across the life course, and it is possible that the increased risk of depression, found in several studies, may be a consequence of an inflammatory insult at another, more vulnerable, later life-stage, or due to the accumulation of inflammatory insults throughout the life course. Therefore, it is difficult to determine whether the immune system may be ‘primed’ to give an enhanced response after repeated inflammatory episodes throughout life. Indeed, elucidating whether an early life infection can ‘prime’ the developing organism’s sensitivity to subsequent environmental challenges is one research question that requires investigation in a clinical setting. Finally, several studies lacked an appropriate control group, making drawing firm conclusions tentative, and/or had a small sample size, potentially lacking the required power to thoroughly investigate the research question.

Despite these limitations, there was considerable agreement with regard to many of the findings pertaining to childhood exposure, and consistency was maintained for studies with both large and small sample sizes, and across individuals with different conditions. Moreover, the lack of an overall association between the prevalence rates of depression in the previously ill v. healthy participants in some of the other aforementioned studies may have been masked by the fact that participants were no older than 25 years at the time of assessment. Empirical research suggests that the average age of onset for mood disorders is 30 years (Kessler et al. 2005), and these studies could have potentially failed to capture the expression of depression in such young cohorts. Furthermore, for all prospective studies, there is a degree of confounding by severity, insofar as the participants with the most severe physical and/or mental disorders may not have been able to meet the demands of the studies in question, and a truly representative outcome may not have been achieved.

**Future work and implications**

Investigating the effect of medically-related inflammation at different life stages will ultimately help identify whether the timing of an immune response is relevant to the pathogenesis of depression, and more prospective longitudinal studies that measure depressive outcomes against number and severity of immune activation throughout life is necessary to confirm this link. Inflammatory markers must be measured in order to investigate the association between severity of immune response and the risk of depression developing. Moreover, controlling for important confounders pertaining to stress (e.g. measuring cortisol stress response), as well as parental psychopathology, and adversity throughout the life course is necessary to confidently establish whether there is a particular time point in life where susceptibility to inflammation predisposes individuals to depression.

In conclusion, this review is the first to evaluate the effects of early life inflammation on the development of future depression in a clinical setting. We show that increased inflammation in childhood may increase depression risk in later life, and although more robust clinical research is needed to definitively address the research questions raised by this article, we bring to light that timing may indeed matter. However, we should heed caution before classifying childhood as a potential ‘window of vulnerability’ owing to the vast limitations of the reviewed studies, and the insufficient research pertaining to the other two life stages. Being able to definitively pinpoint when in life individuals are at increased risk from an inflammatory challenge could (a) help practitioners and individuals better monitor and report these exposures/risk factors, (b) aid early intervention practices, through minimizing and efficiently treating inflammatory conditions during vulnerable stages of development, and (c) ultimately tailor treatment plans – all of which are much needed advancements in care practices.

**Supplementary material**

For supplementary material accompanying this paper visit http://dx.doi.org/10.1017/S0033291716000672.

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Declaration of Interest

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