Systematic Review of Depression in Patients With Multiple Sclerosis And Its Relationship to Interferonβ treatment.

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ABSTRACT

Background: Multiple sclerosis is a chronic disease considered the major cause of neurological disability in young adults worldwide. While depression is considered a determinant factor of impaired quality of life and poorer prognosis among patients with multiple sclerosis, it is very often dismissed and undertreated by physicians. Depression has been related to treatment with some immunomodulatory drugs, such as IFNβ. Data from patients who committed suicide during the pivotal study of interferon used as a disease modifying treatment in multiple sclerosis support this association. Moreover, there is plenty of evidence of neuropsychiatric toxicity caused by the use of IFNα as a treatment for other medical conditions. Although this link still remains relatively unknown, the presence of warnings regarding the possible relationship between depression and IFNβ led to restriction in medical indications in these patients. The purpose of this paper is to try to understand the reasons for an increased prevalence in depression in multiple sclerosis and to examine the impact that IFNβ treatment has on their mood.

Methods: we performed a literature search on MEDLINE and Google Scholar databases applying PRISMA guidelines for systematic reviews. Studies were included if the participants were diagnosed with MS and prescribed IFNβ as the main treatment. We excluded non-english and full-text non available papers, as well as the articles where mental health was assessed exclusively as a feature of quality of life. The sample includes articles from 1980 to 2014, although filtration by year of publication was not applied and contains data from IFNβ-1a and IFNβ-1b. The Cochrane Collaboration Tool assessing risk of bias was used to determine the quality of the studies.

Results: ten studies met full criteria for inclusion and final data extraction. The articles have heterogeneity regarding the samples, the methodology used and the expression of the results. Only three studies support the evidence of a relationship between depression and interferon, which is statistically significant in some patients at the beginning of the treatment. They suggest that only patients on IFNβ treatment with a past history of depression may develop a major depression episode during the first six months. The remaining articles reviewed (including BENEFIT, BEYOND, and LTF trials) suggest the absence of an association.

Conclusion: The reviewed studies conclude that there is not a clear relationship between IFNβ and depression. A history of depression is a risk factor for developing depression during the first 6 months of treatment, nevertheless, it is not sufficient to contraindicate it. The development of new strategies is crucial for early detection of depressive symptoms. An adequate treatment can both improve the mood and deal with the neurological disease by increasing treatment adherence and interfering with inflammation. Chronic destructive brain changes and serotonergic depletion due to inflammatory factors have been proposed as the underlying cause of depression in these patients. It is suggested that these patients would have fewer functional reserve remaining to deal with stressful life events, which could precipitate a depressive disorder.

Keywords: systematic review, multiple sclerosis, depression, depressive symptoms, interferonβ, relationship.

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1. INTRODUCTION
Multiple sclerosis (MS) is a chronic autoimmune inflammatory disease characterised by demyelination of the central nervous system (CNS). It is considered the most common cause of neurological disability in young and middle-aged adults. MS is thought to affect more than 2.1 million people worldwide(1)(2).

Depression is the strongest determinant of impaired quality of life in MS patients. Lifetime prevalence of major depression has been estimated at around 50%(3). This represents almost three times the rate reported in the general population, and it is specially high in young women with less than ten years of disease duration. Major depression is also associated with non-adherence to treatment and with an increased suicide risk. Depression in patients with MS is under-recognised by physicians and therefore, under-treated. Almost 67% of patients were not receiving antidepressant medications and a 33% of suicidal patients had not received psychological assistance (4).

Several immunomodulatory and immunosuppressive drugs have been approved for treating MS. IFNβ (a glycoprotein naturally secreted in response to viral infections) has proven to show beneficial effects in reducing relapse rates and delaying physical disability in relapsing-remitting multiple sclerosis (RRMS) and secondary-progressive multiple sclerosis (SPMS)(5).

A link between depression and IFNβ has been suggested based on data from the pivotal study during which five patients, all on active treatment, attempted suicide (6). Since then, there have been cases describing patients suffering from depression after starting IFNβ treatment. This association is supported by the evidence of IFNα's neuropsychiatric toxicity reported in cancer chemotherapy and chronic hepatitis (7)(8).

As a result of these facts, manufacturers include warnings regarding the possible relationship between depression and IFNβ which leads to a restriction in medical indications. Thus, MS patients cannot fully enjoy the right to benefit from medical treatment. However, when this has been evaluated in clinical trials using validated rating scales, no association has been found. These discrepancies have been little explored and controversy on this topic is still present nowadays.

The aim of this article is to review the existing evidence available for IFNβ therapy causing or exacerbating depression, as well as assessing the reasons for an increased major depression prevalence amongst MS patients in order to improve knowledge on neuropsychiatric manifestations and its possible triggers.
2. METHODS
2.1. Search strategy
We performed a literature search on MEDLINE database using the search terms: "Interferon-beta"[MeSH] OR ("Interferon beta-1b" OR "Interferon beta-1a") OR ("Interferon Type I"[MeSH] NOT "Interferon-alpha") AND Depress* AND “Multiple Sclerosis”. We used the PRISMA (Preferred Reporting Items for Systematic reviews and Meta-Analyses) guidelines as a framework for our review. In addition, we examined the first 10 pages in Google Scholar using the terms Interferon-beta AND Cause AND Depression AND Multiple Sclerosis. We did not apply filtration by year of publication and studies contained data from IFNβ-1a as well as data from IFNβ-1b treatment.

2.2. Inclusion and exclusion criteria
Studies were included if participants were primarily diagnosed with MS and prescribed IFNβ as a disease modifying treatment. We also included studies that compared other immunomodulatory drugs (i.e. IFNα, glatiramer acetate) to IFNβ as an aetiological factor for depression, but excluded those where INFβ was not the main treatment. Depression symptoms were mainly evaluated with specific affective scales such as The Beck Depression Inventory (BDI) or The Hamilton Rating Scale for Depression (HRSD). However, studies regarding depression related to fatigue or using other non validated scales for depression were also considered. We excluded the articles where mental health was exclusively assessed as a feature of quality of life. Non-english papers and full-text not available articles were also excluded. If there were doubts on whether an abstract should be included for full text retrieval, it was decided that they would be included.

2.3. Quality assessment
The Cochrane Collaboration Tool assessing risk of bias was used to determine the quality of the studies (9). This is a two part tool addressing six specific domains of bias (sequence generation, allocation concealment, blinding of participants, personnel and outcome assessors, incomplete outcome data, selective outcome reporting and “other sources of bias”). Each domain includes one or more specific entries in a “risk of bias” table. Within each entry, the first part of the tool describes what was reported to have happened in the study, whereas the second part assigns a judgement to the risk of bias. This is achieved by assigning a “low risk”, “high risk” or “unclear risk” of bias to each study.

3. RESULTS AND DISCUSSION
3.1 Description of studies
Initially, 119 studies were identified. After duplicates were excluded, 114 abstracts remained. All abstracts were read and 33 were selected for full text retrieval. Overall, 10 studies met full criteria for inclusion and final data extraction. A PRISMA flowchart describing the results of the search is shown in figure 1.

A description of the included studies and the most conclusive results regarding the main aim (i.e.relationship between interferon and depression) that are summarized in table 1, are discussed.
3.2. Quality assessment of studies
After assessing the risk of bias of all the studies using the Cochrane guidelines, we found that the majority of studies had a high risk of bias in at least two of the six domains. (see table 2). All of the studies reported inadequate randomisation and concealment of allocation. Ziemssen et al.(4) presented insufficient information to allow for any judgement. In Feinstein et al.(10) there was uncertain outcome data and was deemed high risk of bias in blinding and selective outcome reporting. In Pattern et al.(11) the blinding and ‘other’ sources of bias were also high. In Siegert et al.(12) the outcome measurement is likely to be influenced by a lack of blinding. There was also a high risk of bias in selective outcome and unclear outcome data. Insufficient information is reported in Gold et al.(13) so it cannot be entered in a meta-analysis.
<table>
<thead>
<tr>
<th>Authors</th>
<th>Study design</th>
<th>IFN type</th>
<th>Definition of depression</th>
<th>Diagnostic criteria/instrument</th>
<th>Association between IFN and depression</th>
</tr>
</thead>
<tbody>
<tr>
<td>Patten S (2005)</td>
<td>Pooled data from 6 controlled studies and 17 non controlled clinical trials 1995-2002</td>
<td>IFN beta-1a-sc</td>
<td>Aggravated depression: either new onset, newly recognized or worsened</td>
<td>Physician examination (DSM-IV criteria were not necessarily applied)</td>
<td>Statistically significant association between depression and IFN use during the first six months of treatment</td>
</tr>
<tr>
<td>Siegert R (2005)</td>
<td>Review 1879-2004</td>
<td>IFN beta</td>
<td>Unipolar or major depression</td>
<td>Structured psychiatric interview</td>
<td>Concerns about interferon treatment and depression may have been undue</td>
</tr>
<tr>
<td>Ziemssen T (2009)</td>
<td>Review 1984-2008</td>
<td>IFN beta</td>
<td>Major depressive disorder</td>
<td>Beck Depression Inventory (BDI)</td>
<td>IFN beta may aggravate depression</td>
</tr>
<tr>
<td>Gold S (2009)</td>
<td>Review 1980-2005</td>
<td>IFN beta-1a and other cytokines</td>
<td>Depressive symptoms</td>
<td>Sickness behaviour model</td>
<td>Subtypes of MS depression may be linked to inflammatory markers</td>
</tr>
<tr>
<td>Nikfar S (2010)</td>
<td>Meta-analysis 9 controlled trials reviewed in PubMed, Scopus, Cochrane 1966–May 2010</td>
<td>IFN beta</td>
<td>Depression</td>
<td>Not mentioned</td>
<td>There were no significant differences in depression between IFN beta and placebo.</td>
</tr>
<tr>
<td>Feinstein A (2011)</td>
<td>Review 1984-2011</td>
<td>IFN beta-1b</td>
<td>Syndrome of major depression</td>
<td>Clinical interview</td>
<td>Poor adherence with a disease modifying drug is associated with depression</td>
</tr>
<tr>
<td>Plosker G (2011)</td>
<td>Review in MEDLINE, EMBASE and AdisBase 1980-Nov 2010</td>
<td>IFN beta-1b</td>
<td>Depression</td>
<td>Not mentioned</td>
<td>Depression occurs with increased frequency in association with interferon use</td>
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<tr>
<td>Reder A (2014)</td>
<td>Review of adverse events in 3 clinical trials (BENEFIT, BEYOND, and the 16-year Long-Term Follow-up (LTF))</td>
<td>IFN beta-1b</td>
<td>Depression</td>
<td>Mandated questions and patients’ self-reports</td>
<td>There was not an increase in rates of depression due to interferon β-1b treatment</td>
</tr>
</tbody>
</table>
Table 2. Quality assessment of the included studies based on risk of bias

<table>
<thead>
<tr>
<th>Study</th>
<th>Sequence generation</th>
<th>Allocation concealment</th>
<th>Blinding of participants, personnel and outcome assessors</th>
<th>Incomplete outcome data</th>
<th>Selective outcome reporting</th>
<th>Other sources of bias</th>
</tr>
</thead>
<tbody>
<tr>
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<td>High</td>
<td>Low</td>
<td>Unclear</td>
<td>Low</td>
<td>Low</td>
</tr>
<tr>
<td>Patten S (2005) 11</td>
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<td>High</td>
<td>High</td>
<td>Low</td>
<td>Low</td>
<td>High</td>
</tr>
<tr>
<td>Siegert R (2005) 12</td>
<td>High</td>
<td>High</td>
<td>High</td>
<td>Unclear</td>
<td>High</td>
<td>High</td>
</tr>
<tr>
<td>Goeb J (2006) 15</td>
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<td>High</td>
<td>High</td>
<td>Unclear</td>
<td>Low</td>
<td>Low</td>
</tr>
<tr>
<td>Ziemssen T (2009) 4</td>
<td>Unclear</td>
<td>Unclear</td>
<td>Unclear</td>
<td>Unclear</td>
<td>High</td>
<td>High</td>
</tr>
<tr>
<td>Gold S (2009) 13</td>
<td>High</td>
<td>High</td>
<td>Unclear</td>
<td>Low</td>
<td>High</td>
<td>Low</td>
</tr>
<tr>
<td>Nikfar S (2010) 16</td>
<td>High</td>
<td>High</td>
<td>Low</td>
<td>Low</td>
<td>Low</td>
<td>Low</td>
</tr>
<tr>
<td>Feinstein A (2011) 10</td>
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<td>High</td>
<td>High</td>
<td>Unclear</td>
<td>High</td>
<td>High</td>
</tr>
<tr>
<td>Plosker G (2011) 17</td>
<td>High</td>
<td>High</td>
<td>Low</td>
<td>Low</td>
<td>Low</td>
<td>Low</td>
</tr>
<tr>
<td>Reder A (2014) 18</td>
<td>High</td>
<td>High</td>
<td>Low</td>
<td>Low</td>
<td>Low</td>
<td>Low</td>
</tr>
</tbody>
</table>

3.3. Interferon and depression

One of the articles that approached the question of whether interferon induced depression for the first time was Feinstein et al.(14). They evaluated all the publications available on the topic since 1993. The authors considered the possibility of the IFNβ-1b having an inherent mood destabilizing effect, but highlighted the methodological weaknesses of the studies. They exposed that the use of rating scales such BDI could influence the results, as it contains numerous questions linked to somatic complaints.

Five years later, Siegert et al.(12) reviewed similar data and concluded that, although depression should be a feature of medical management in MS patients, the relationship between interferon beta and depression had been unwarranted. They also emphasised on the importance of treating depression in these patients.

Other studies reported comparable results. For example, Goeb et al.(15) reviewed sixteen articles about psychiatric side effects under IFNβ. Fourteen articles discarded an association and one found that prevalence rates for major depression declined during treatment.

On the other hand, several authors presented data supporting the hypothesis of a relationship.
Pattern et al. (11) evaluated 2819 patients including patients with RRMS and SPMS. Of these, 824 received placebo and IFNβ-1a was administered to 1995. Depression was defined based on mood clinical changes detected by the treating physician. They recorded data from a six-month assessment period. Two-year and six-year data were recorded from subsets of the controlled population, comprised of 1178 patients of whom 392 were receiving placebo from 1-2 or 1-3 years and 786 were receiving interferon. During the first six months, between 5% and 18% of patients included in the treatment group reported depression compared to the 8% of patients taking placebo. Depression was a reason for abandoning therapy in 1.3% of patients receiving IFN and for only 0.6% on the placebo group. No significant differences were found among groups after two and six years of treatment. Ziemssen et al. (4) assumed that interferon increases the risk of depression in susceptible individuals and even proposed an algorithm for its management in MS patients. They suggested an initial screening for depression using the BDI at baseline. Patients who scored ≥13 on the test should undergo a deep diagnostic interview. For those with a previous history of depression or a current confirmed depressive episode, other non-interferon treatments should be chosen.

Gold et al. (13) speak of an ‘inflammatory’ kind of depression. Depressive symptoms including anxiety, dysphoria, anhedonia, fatigue, anorexia, cognitive and slowness can occur after 1 to 3 months of therapy and persist unless treatment is terminated or supplemented with antidepressants. In healthy human volunteers, endogenous cytokines have been proven to develop transient neuropsychiatric symptoms. It is then conceivable that the depression observed in MS patients reflects the cytokine network activation partly triggered by interferon treatment.

Plosker et al. (17) published an updated review of the pharmacology, clinical efficacy and tolerability of interferon in the management of patients with MS. The authors extracted data from 9 placebo controlled and active-comparator trials in RRMS and SPMS patients. Depression was reported as one of the most serious adverse effects and pointed out that it occurred with major frequency among patients using IFNβ-1b. However, they do not provide further information regarding this issue.

Recent publications indicate that IFNβ is not directly linked to depression in these patients. In 2011 Feinstein et al. (10) established that developing depression during interferon treatment was mainly determined by a prior history of depression. They suggested that in case of depression, interferon treatment should not be discontinued.

Nikfar et al. (16) analysed 9 randomized, double-blind trials assessing the efficacy and tolerability of interferon beta in patients with MS. Specific adverse events of interest such as depression were analyzed individually and compared between IFNβ and placebo. The accumulative relative risk for depression in three trials was 0.99. There were no significant differences between groups.

Reder et al. (18) reviewed results from three trials. The BENEFIT trial assessed the potential effects of early vs. delayed treatment of 292 MS patients receiving IFNβ-1 and 176 receiving placebo. In the first 2 years, 10.3% of patients on treatment experienced depression compared to
11.4% in the placebo group. The BEYOND study compared efficacy, safety and tolerability of doses of 250μg of IFNβ-1b in 887 patients, 500μg in 888 patients and in 445 patients receiving glatiramer acetate. Authors reported incidence rates of depression of 17% in both groups receiving IFNβ-1b and 14% in those with glatiramer. The 16-year LTF study provided additional long-term follow-up for 328 patients from the original pivotal trial of IFNβ-1b in RRMS. After two years, 42% of patients on treatment reported depression compared to 46.3% of those who were not receiving interferon.

The results from detailed and methodologically sound clinical trials assessing this association were collected and a Forest Plot was produced. Figure 2, Figure 3.

**Figure 2.** Forest Plot including clinical trials reporting data of incidence of depression among patients receiving treatment with interferon beta-1a treatment compared to placebo

<table>
<thead>
<tr>
<th>Study</th>
<th>Treatment</th>
<th>Events (Treat)</th>
<th>Events (Cont)</th>
<th>OR</th>
<th>IC(95%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>The Nordic SPMS Study Group</td>
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<td>25</td>
<td>1.52</td>
<td>0.9 2.6</td>
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<tr>
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<td>52</td>
<td>0.55</td>
<td>0.4 0.8</td>
</tr>
<tr>
<td>CHAMPS STUDY GROUP</td>
<td>yes</td>
<td>39</td>
<td>25</td>
<td>1.67</td>
<td>1.2 2.9</td>
</tr>
<tr>
<td>SPECTRIMS Study Group</td>
<td>yes</td>
<td>67</td>
<td>29</td>
<td>1.18</td>
<td>0.7 1.9</td>
</tr>
<tr>
<td><strong>Summary</strong></td>
<td></td>
<td>1</td>
<td>0.8</td>
<td>1.3</td>
<td></td>
</tr>
</tbody>
</table>

**Figure 3.** Forest Plot including clinical trials reporting data of incidence of depression among patients receiving treatment with interferon beta-1b compared to placebo

<table>
<thead>
<tr>
<th>Study</th>
<th>Treatment</th>
<th>Events (Treat)</th>
<th>Events (Cont)</th>
<th>OR</th>
<th>IC(95%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>BENEFIT Study Group</td>
<td>yes</td>
<td>30</td>
<td>20</td>
<td>0.89</td>
<td>0.5 1.6</td>
</tr>
<tr>
<td>IFNB Multiple Sclerosis Study Group</td>
<td>yes</td>
<td>23</td>
<td>15</td>
<td>1.34</td>
<td>0.6 2.8</td>
</tr>
<tr>
<td><strong>Summary</strong></td>
<td></td>
<td>1.1</td>
<td>0.7</td>
<td>1.7</td>
<td></td>
</tr>
</tbody>
</table>

3.4. Depression in multiple sclerosis

Depression is a common and important occurrence in people with MS. Despite the afterthought that depression in these patients is primarily reactive to the diagnosis and prognosis, the heterogenous nature of the syndrome suggests that many factors are at play. No correlation has been found between disease severity and depression; and interventions to provide social support have failed to show any improvement (19)(20). In addition, major depression is seen more often in MS compared to other chronic progressive diseases. (21) Apparently, it does not seem to be a clear genetic predisposition to major depression in MS (22). With the development of neuroimaging tests a link between depression and structural changes has been established. MS
patients with brain lesions were more depressed than those patients with only spinal cord lesions, specially if the lesions were located in the left anterior temporal regions. Although certain relationships/links were observed, authors suggest that depression would more likely be a consequence of chronic destructive brain changes than a result of the acute hyperintense lesions. They suggest that MS patients would have fewer functional reserve remains to deal with stressful life events which could precipitate a depressive disorder. (23)(24) On the other hand, the nervous system inflammation by itself could be the underlying cause of depression in MS. Meta-analyses indicate that depression is associated with an increase in circulating levels of the pro-inflammatory cytokine, interleukin-6, and elevated levels of this cytokine have been found in adults with major depression (25)(26)(27). In the case of MS, the breakdown of the blood-brain barrier may allow the entry of inflammatory cells into the CNS as well as an increased local production of cytokines as a result of the inner brain inflammation. In the last few years, it has been proposed that cytokines could produce a serotonergic depletion in CNS, and thus lead to depression.(28)(29)(30) One of the studied mechanisms is the induction of indolamine-2,3-dioxygenasa (IDO), an enzyme that metabolizes tryptophan, the precursor of serotonin. Some studies have found an increased secretion of interleukine-6 and IDO stimulation due to IFNβ administration. (31)(32)(33). In addition, elevated levels of corticotropin releasing hormone due to a hyperactivation of the hypothalamus pituitary adrenal axis seen in stressed and depressive patients can aggravate the inflammatory response and perpetuate the brain lesion. 

The articles included in this review have heterogeneity regarding the samples, methodology used and expression of results. The samples of the studies are in many cases small, and vary according to the MS type and the IFNβ formulation. Few studies used a structured, validated clinical interview. In addition, a valid rating scale specific for mood in these patients should be developed. Alcohol and substance abuse, which is known to be closely related to major depression should be taken into account. The distinction between mild symptoms of depression and the diagnoses of depression as a whole syndrome can also influence the results.

4. DISCUSSION

To our knowledge, this is the first systematic review conducted to evaluate the relationship between IFNβ treatment and depression in MS patients. Although clinical trials suggest a link between IFNβ and depression, data from reviewed studies do not support this relationship. Several narrative reviews have been published, showing contradictory results. In the first review of the topic, Feinstein (14) highlights that no conclusions can be achieved due to the weakness of the methodology of the studies. Siegert et al (12) pointed out that the relationship between IFNβ and depression had been unwarranted established. In the review conducted by Goeb et al (15) no evidence of increase in depression in MS patients treated with IFNβ was found. Similar results were found by Reder et al (18), who concluded that there was no increase in the depression rate due to IFNβ-1b after reviewing data from three clinical trials. Ziemssen (4) assumed that IFNβ increased the risk of depression in susceptible individuals, but this conclusion is not based on an exhaustive review of the subject and reported that the impact of IFNβ on depression is controversial. Plosker et al (17) reviewed data from nine clinical trials and mentioned that depression was one of the most serious adverse events reported with IFNβ-1b and that depression
occurs with increased frequency with IFN use, but no incidence rate or increased risk rate was described.

In the analysis of pooled data from 6 controlled studies and 17 non controlled studies, Patten et al (11) concluded a statistically significant association between depression and IFN use within the 6 initials months of treatment, however, no significant differences were found among groups after two nor six years of treatment.

The best methodological study available is the meta-analysis of 9 controlled trials conducted by Nifkar et al (16), and no significant difference in depression rate was found between the IFNβ and placebo group.

The conducted forest plot supports data showing a lack of association between interferon beta and depression in the articles reviewed. The results are more robust for interferon beta-1a than for interferon beta-1b, as more homogeneous interferon beta-1a clinical trials assessing this issue are available in the literature.

5. CONCLUSIONS
As mentioned above, data from our systematic review does not support a clear link between treatment with IFNβ and depression. Only a history of depression has been found to be a risk factor for developing depression in MS patients. According to the available data, it seems controversial to contraindicate IFNβ treatment in MS patients. A close follow-up of patients with a history of depression is recommended when they undergo treatment with IFNβ. Cooperation between a neurologist and a psychiatrist is required in order to detect major depression. Designing new specific structured interviews and counseling patients that depression may develop can help to recognise depressive symptoms early on in the course of the disease. While untreated depression in these patients is likely to worsen, it is particularly responsive to the use of antidepressants. Antidepressant treatment is believed to decrease inflammatory markers. Detecting and treating depression not only may reduce the inflammatory process, but could also increase the patient’s compliance with a disease modifying drug, which has proved efficacy in slowing down the progression of the disease. Nevertheless, we are aware that methodological limitations prevent from drawing definite conclusions on these issues and we encourage authors to carry out further investigation.

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