Sadness and mild cognitive impairment as predictors for interferon-alpha-induced depression in patients with hepatitis C

Susanne Sarkar, Rahul Sarkar, Thomas Berg and Martin Schaefer

Background
Antiviral therapy with interferon-alpha (IFN-α) for hepatitis C virus (HCV) infection is associated with increased risk for depression. Poor baseline performance in the Trail Making Test A/B (TMT A/B) was observed during antiviral treatment in 91% of patients with chronic HCV infection without a history of psychiatric disorders. Cognitive function was evaluated using the Trail Making Test A/B (TMT A/B). (Trial registration at ClinicalTrials.gov: NCT00136318.)

Method
Depression (defined with the Montgomery–Åsberg Depression Rating Scale) was evaluated before and during antiviral treatment in 91 people with chronic HCV infection who participated in a prospective controlled trial investigating the prevention of depressive symptoms with escitalopram during antiviral treatment with PEG-IFN-α and ribavirin (for detailed information see Schaefer et al1). The reported prevalence of IFN-associated major depressive episodes is 20–40%, and up to 70% of patients may develop mild to moderate depressive episodes with a significant impact on their quality of life. This is likely to negatively affect adherence to treatment and may represent a major cause of treatment discontinuation.1–4 Recently, pre-emptive treatment with antidepressants, notably selective serotonin reuptake inhibitors (SSRIs), demonstrated a reduction of the incidence of mild to severe major depressive episodes in HCV-infected patients during antiviral treatment.5 The largest trial in patients without prior psychiatric disorders to date was conducted with escitalopram. When compared with controls without pre-emptive treatment, a lower incidence and severity of depression, defined by the presence of DSM-IV major depressive episodes criteria and Montgomery–Åsberg Depression Rating Scale (MADRS) scores, was observed during HCV treatment.5 Although antidepressant pre-treatment showed a high efficacy in reducing overall depression rates and the use of SSRIs was safe without negative effects on antiviral response, it also needs to be considered that a significant proportion of patients do not develop clinically relevant depressive episodes. Thus, a pharmacological prophylactic treatment might put them at risk of harm without apparent benefits. Concomitant treatment with antidepressants may cause additional side-effects including sexual dysfunction, insomnia, nausea and visual and cardiac symptoms.7 Hence, pre-emptive antidepressant treatment should be limited to patients who are at increased risk for depression, and reliable identification of high-risk patients for IFN-associated depressive syndromes is warranted. Screening criteria to identify those patients who might have the greatest benefits from pre-emptive antidepressant therapy are warranted for limiting potential disadvantages and maximising potential benefits of psychopharmacological pre-treatment in routine care. The aim of the present post hoc analysis was to identify neuropsychiatric risk factors that predict the development of depression during antiviral treatment with IFN-α.

Aims
To identify clinical predictors for IFN-α-induced depression during antiviral therapy for HCV infection.

Results
Depression during antiviral therapy was significantly associated with a baseline MADRS score of 3 or higher (P=0.006). In total, 89% (n=16) of patients who had a baseline score >0 for the single item sadness developed depression. Poor baseline performance in the TMT A (P=0.027) and TMT B (P=0.033) was predictive for severe depression.

Conclusions
Pre-treatment screening for subthreshold depressive and cognitive symptoms will help to identify those at risk for IFN-α-associated depression among patients with chronic hepatitis C.

Declaration of interest
M.S.: consultancy and member of speakers’ bureau at Roche Pharma (Germany/Switzerland), Janssen-Cilag, Servier and Shire. T.B.: consultancy or advisory board member, speakers’ bureau at Abbott, Bristol-Myers Squibb, Boehringer Ingelheim, Gilead Sciences, Janssen Pharmaceuticals, Merck & Co, Novartis, Roche and Vertex Pharmaceuticals Incorporated; research grants from Gilead Sciences, Janssen Pharmaceuticals, Novartis and Roche.

The combination of pegylated (PEG) interferon-alpha (IFN-α) plus ribavirin is established standard treatment of hepatitis C virus (HCV) infection. Antiviral therapy including IFN-α, however, is associated with significant psychiatric side-effects such as depressive symptoms, fatigue and insomnia.1–3 The reported prevalence of IFN-associated major depressive episodes is 20–40%, and up to 70% of patients may develop mild to moderate depressive episodes with a significant impact on their quality of life. This is likely to negatively affect adherence to treatment and may represent a major cause of treatment discontinuation.1–4 Recently, pre-emptive treatment with antidepressants, notably selective serotonin reuptake inhibitors (SSRIs), demonstrated a reduction of the incidence of mild to severe major depressive episodes in HCV-infected patients during antiviral treatment.5 The largest trial in patients without prior psychiatric disorders to date was conducted with escitalopram. When compared with controls without pre-emptive treatment, a lower incidence and severity of depression, defined by the presence of DSM-IV major depressive episodes criteria and Montgomery–Åsberg Depression Rating Scale (MADRS) scores, was observed during HCV treatment.5 Although antidepressant pre-treatment showed a high efficacy in reducing overall depression rates and the use of SSRIs was safe without negative effects on antiviral response, it also needs to be considered that a significant proportion of patients do not develop clinically relevant depressive episodes. Thus, a pharmacological prophylactic treatment might put them at risk of harm without apparent benefits. Concomitant treatment with antidepressants may cause additional side-effects including sexual dysfunction, insomnia, nausea and visual and cardiac symptoms.7 Hence, pre-emptive antidepressant treatment should be limited to patients who are at increased risk for depression, and reliable identification of high-risk patients for IFN-associated depressive syndromes is warranted. Screening criteria to identify those patients who might have the greatest benefits from pre-emptive antidepressant therapy are warranted for limiting potential disadvantages and maximising potential benefits of psychopharmacological pre-treatment in routine care. The aim of the present post hoc analysis was to identify neuropsychiatric risk factors that predict the development of depression during antiviral treatment with IFN-α.

Method

Settings and participants
This article reports on a total of 91 people with chronic HCV infection who participated in a prospective controlled trial investigating the prevention of depressive symptoms with escitalopram during antiviral treatment with PEG-IFN-α and ribavirin (for detailed information see Schaefer et al1) and who were randomised to placebo treatment.

All patients were treatment-naive to HCV treatment, older than 18 years and had a chronic HCV infection with serum HCV-RNA levels of 1000 IU/mL or higher. Psychiatric exclusion criteria were a lifetime diagnosis of an affective disorder, drug misuse in the past 12 months, treatment with antidepressants during the past 3 years or a history of any other Axis I disorder according to DSM-IV criteria.1 Medical exclusion criteria were pre-treatment with IFN or immunotherapy, other chronic infections or a severe somatic comorbid condition. The study protocol was approved by the ethics committee of the Charité – University Medicine Berlin and confirmed by local ethics
committees. The study was conducted in accordance with the principles of the Declaration of Helsinki and local laws and regulations. All participants provided written informed consent. The trial is registered at ClinicalTrials.gov: NCT00136318.

**Treatment**

Eligible patients participated in a multicentre, double-blind, prospective, randomised and placebo-controlled phase-III study consisting of three different study periods as described elsewhere. The original study compared pre-emptive treatment with escitalopram v. placebo in patients undergoing antiviral therapy with PEG-IFN-2α plus ribavirin; here we report on the 91 patients who had been randomised to placebo. During a pre-treatment period of 14 weeks before antiviral therapy was started, patients were monitored for spontaneously developing symptoms of depression. Next, all patients received antiviral therapy with PEG-IFN-2α plus ribavirin (Roche, Grenzach-Wyhlen, Germany). Patients with HCV genotype 1 or 4 were treated for 48 weeks with PEG-IFN-2α at a dose of 180 μg per week, and ribavirin at a dose of 1000 mg per day (body weight <75 kg) or 1200 mg per day (body weight ≥75 kg). Patients with HCV genotype 2 or 3 received PEG-IFN-2α (180 μg/week) and ribavirin (800 mg daily) for 24 weeks. Doses of ribavirin and PEG-IFN-2α could be adjusted in response to side-effects. Benzodiazepines were not allowed during the trial.

**Psychiatric assessments and end-points**

Psychiatric assessments were performed 14, 8, and 2 weeks before antiviral therapy was started as well as after 2, 4, 12 and 24 and, for patients with genotypes 1 and 4, 48 weeks of antiviral therapy and, finally, 24 weeks after antiviral treatment (follow-up assessment). The Mini-International Neuropsychiatric Interview was used at baseline as a screening tool to detect pre-existing psychiatric disorders. The Structured Clinical Interview for DSM-IV Disorders was used to verify the incidence of major depressive disorder before and during treatment. In addition we used the MADRS11 and the 21-item Beck Depression Inventory (BDI)12 to define and identify the incidence of all clinically relevant (mild to moderate) depressive symptoms. The MADRS scale consists of 10 items assessing symptoms associated with depression (apparent and reported sadness, inner tension, reduced sleep, reduced appetite, loss of concentration, lassitude, inability to feel, pessimistic thoughts and suicidal thoughts), each item scored from 0 to 6 according to severity. We applied three different definitions of depressive mood disorder: (a) those who fulfilled criteria of a current major depressive episode according to DSM-IV (independent of severity), and two groups defined on the basis of MADRS scores independent of fulfilling full DSM-IV major depressive episodes criteria; (b) MADRS scores ≥13 were used as cut-off for clinically relevant depression; (c) MADRS ≥25 for severe depression.5,11,13 In addition, a self-rating instrument, the BDI was administered to screen for potential risk factors for developing depressive episodes during antiviral therapy. To evaluate the relationship between specific subclinical depressive symptoms at baseline and the development of an IFN-associated depressive syndrome (MADRS scores ≥13, major depressive episodes, MADRS ≥25), items measured on the MADRS scale were used to perform different regression trees. Independent variables were the 10 single items of the MADRS scale (apparent and reported sadness, inner tension, inability to feel, loss of concentration, pessimistic thoughts, reduced sleep, reduced appetite, lassitude and suicidal thoughts). On the basis of decision trees, we defined cut-off points for each symptom of depression of the MADRS scale at baseline to discriminate between patients with or without IFN-associated depression. There is a need for cut-off points for subclinical depressive symptoms to construct rules for making predictions about individual cases and assist in identifying homogeneous groups with high or low risk for the development of IFN-associated depression. To evaluate the influence of subclinical cognitive dysfunction at baseline, multiple logistic regression analysis was used to calculate the odds ratios of the incidence of IFN-associated depressive symptomatology (MADRS ≥13, major depressive episodes, MADRS ≥25) adjusted for gender, age, BMI, genotype (1 and 4 v. 2 and 3), baseline MADRS ≥3 and baseline BDI >4.

Results

**Development of depressive symptoms during antiviral treatment**

None of the 91 patients exhibited any clinically relevant depressive syndromes (defined as a MADRS score ≥13) before the initiation of antiviral therapy.
of INF treatment. A total of 88% (80/91) of the patients finished antiviral therapy according to the protocol. The overall incidence of adverse events was 87%. Dose reduction of PEG-IFN-α2a was necessary for 19% (n = 17) of patients and ribavirin dose was decreased in 29% (n = 26) of patients. Of the 91 patients, 54% (n = 49, 95% CI 48–69%) developed MADRS scores ≥ 13 during antiviral therapy, 19% developed a major depressive episode according to DSM-IV criteria (n = 17, 95% CI 12–28%) and 12% had severe depressive syndromes (MADRS score ≥ 25) (n = 11, 95% CI 7–21%) (Table 1).

**Baseline factors and INF-associated depression**

Baseline factors were compared between patients with any kind of depressive syndrome during antiviral therapy with PEG-IFN-α2a and ribavirin (MADRS score ≥ 13), severe depressive syndrome (MADRS score ≥ 25), major depressive episode (defined by DSM-IV criteria) and patients without depression. No differences between patients with depression and those without were found for age, genotype and BMI. However, more women (n = 28, 65%) than males (n = 21, 44%) developed depressive syndromes (MADRS ≥ 13, χ² = 4.166, d.f. = 1, P = 0.041).

Patients with clinically relevant depression during IFN treatment had significant higher MADRS (t = −3.858, d.f. = 89, P < 0.001) and BDI scores (t = −2.012, d.f. = 88, P = 0.05) at baseline and exhibited significantly higher MADRS scores after 2 (t = −4.155, d.f. = 89, P < 0.001) and 4 weeks (t = −4.861, d.f. = 88, P < 0.001) of antiviral treatment. Patients who developed a major depressive episode during antiviral therapy demonstrated higher MADRS scores at baseline (t = −2.437, d.f. = 88, P = 0.017), and after 2 (t = −4.245, d.f. = 89, P < 0.001) and 4 weeks (t = −3.071, d.f. = 89, P = 0.013) of antiviral treatment. In contrast to this, no significant differences were found between individuals with depression and those without for the BDI score (P = 0.19). Patients who developed severe depressive syndromes (MADRS score ≥ 25) also showed significantly higher MADRS scores at baseline (t = −2.440, d.f. = 89, P = 0.017), as well as after 2 (t = 4.552, d.f. = 89, P < 0.001) and 4 weeks (t = 2.976, d.f. = 80, P = 0.004) of antiviral treatment. No influence on the development of major depressive episodes or a severe depressive syndrome was found for gender, age, genotype and BMI.

Table 2 shows the single-item analysis of the MADRS scale before antiviral treatment was initiated. Patients who developed any kind of an IFN-associated depressive syndrome (MADRS score ≥ 13) had significantly higher scores in the following single MADRS items at baseline: apparent sadness (P = 0.009), reported sadness (P = 0.002), inability to feel (P = 0.015), loss of concentration (P = 0.001), pessimistic thoughts (P = 0.005), reduced sleep (P = 0.050), reduced appetite (P = 0.009) and lassitude (P = 0.004). Patients with major depressive episodes had higher scores on the following five MADRS items at baseline: reduced appetite (P = 0.016), loss of concentration (P < 0.001), lassitude (P = 0.016), pessimistic thoughts (P = 0.001) and suicidal thoughts (P = 0.022). The development of severe depression (MADRS score ≥ 25) during antiviral therapy was associated with higher scores for reduced appetite (P = 0.001), loss of concentration (P = 0.001), lassitude (P = 0.045) and pessimistic thoughts (t = 2.636, d.f. = 90, P = 0.010).

We also examined the possible role of cognitive function for the development of depression with the following neuropsychological tests: TMT A, TMT B and the Regensburger word fluency test. Pre-treatment results were compared between patients with and without IFN-associated depressive syndromes. Patients with any clinically relevant depressive syndrome during therapy (MADRS ≥ 13) showed lower scores on TMT B (t = 2.580, d.f. = 83, P = 0.012) and semantic word fluency (t = 2.219, d.f. = 85, P = 0.03).

**Table 1. Comparison of pre-treatment demographic and clinical characteristics between patients without and with depressive syndromes during antiviral treatment**

<table>
<thead>
<tr>
<th>Baseline</th>
<th>All (n = 91)</th>
<th>No depression (n = 42)</th>
<th>Major depressive episode (n = 17)</th>
<th>MADRS score ≥ 13 (n = 49)</th>
<th>MADRS score ≥ 25 (n = 11)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Male, gender, n (%)</td>
<td>48 (53)</td>
<td>27 (64)</td>
<td>7 (41)</td>
<td>21 (43)</td>
<td>4 (36)</td>
</tr>
<tr>
<td>Age, years</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean (s.d.) range</td>
<td>48.5 (11.3) 21–71</td>
<td>47.9 (11.3) 21–66</td>
<td>50.8 (9.7) 31–71</td>
<td>49 (11.5) 23–71</td>
<td>50 (11.3) 31–71</td>
</tr>
<tr>
<td>&lt; 40 years, n (%)</td>
<td>22 (24)</td>
<td>11 (26)</td>
<td>1 (6)</td>
<td>11 (22)</td>
<td>10 (9)</td>
</tr>
<tr>
<td>Ethnicity, White, n (%)</td>
<td>80 (88)</td>
<td>35 (83)</td>
<td>16 (94)</td>
<td>45 (92)</td>
<td>11 (100)</td>
</tr>
<tr>
<td>Body mass index, kg/m²</td>
<td></td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Mean (s.d.) range</td>
<td>25.8 (8.0) 17–39</td>
<td>25.8 (4.2) 17–34</td>
<td>24.7 (3.4) 22–36</td>
<td>25.8 (10.1) 17–39</td>
<td>25.4 (4.0) 22–36</td>
</tr>
<tr>
<td>&gt; 25 kg/m², n (%)</td>
<td>40 (44)</td>
<td>23 (55)</td>
<td>5 (30)</td>
<td>17 (35)</td>
<td>5 (46)</td>
</tr>
<tr>
<td>Genotype, n (%)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>59 (65)</td>
<td>24 (57)</td>
<td>12 (71)</td>
<td>35 (71)</td>
<td>9 (82)</td>
</tr>
<tr>
<td>2</td>
<td>5 (5)</td>
<td>3 (7)</td>
<td>2 (12)</td>
<td>2 (12)</td>
<td>1 (10)</td>
</tr>
<tr>
<td>3</td>
<td>21 (23)</td>
<td>11 (26)</td>
<td>2 (12)</td>
<td>10 (20)</td>
<td>1 (9)</td>
</tr>
<tr>
<td>4</td>
<td>6 (7)</td>
<td>4 (10)</td>
<td>1 (6)</td>
<td>2 (4)</td>
<td>0 (0)</td>
</tr>
<tr>
<td>MADRS, mean (s.d.) range</td>
<td>2.7 (3.9) 0–16</td>
<td>1.4 (2.4) 0–10</td>
<td>5.4 (5.8) 0–16</td>
<td>4.6 (4.8) 0–16</td>
<td>5.9 (5.9) 0–16</td>
</tr>
<tr>
<td>Beck Depression Inventory, mean (s.d.) range</td>
<td>4.0 (4.1) 0–18</td>
<td>4.0 (3.8) 0–13</td>
<td>6.4 (3.9) 0–15</td>
<td>5.6 (4.2) 0–18</td>
<td>5.5 (3.7) 0–13</td>
</tr>
<tr>
<td>Trail Making Test, s: mean (s.d.) range</td>
<td>39.5 (16.4) 15–100</td>
<td>36.4 (11.6) 19–65</td>
<td>48.2 (19.6) 20–88</td>
<td>42.1 (19.3) 15–100</td>
<td>59.5 (14.4) 35–88</td>
</tr>
<tr>
<td>Test A</td>
<td>94.6 (36.1) 30–205</td>
<td>81.8 (27.2) 35–145</td>
<td>109.5 (35.0) 60–190</td>
<td>108.4 (40.7) 30–205</td>
<td>130.9 (28.1) 100–190</td>
</tr>
<tr>
<td>Test B</td>
<td>32.4 (10.4) 12–61</td>
<td>29.8 (9.8) 13–47</td>
<td>35.6 (11.9) 20–61</td>
<td>34.7 (10.9) 12–61</td>
<td>37.2 (12.5) 20–61</td>
</tr>
<tr>
<td>Phonematic word fluency test, mean (s.d.) range</td>
<td>22.1 (9.4) 7–77</td>
<td>21.5 (6.6) 7–35</td>
<td>24.4 (15.8) 11–77</td>
<td>22.9 (4.9) 18–36</td>
<td>22.4 (8.6) 11–34</td>
</tr>
<tr>
<td>Semantic word fluency test, mean (s.d.) range</td>
<td>20.6 (4.2) 18–38</td>
<td>19.2 (2.6) 18–31</td>
<td>22.6 (5.6) 18–33</td>
<td>21.7 (4.9) 18–38</td>
<td>22.6 (5.6) 18–33</td>
</tr>
</tbody>
</table>

MADRS, Montgomery-Åsberg Depression Rating Scale.
a. hepatitis C virus (HCV) subtypes/genotypes 1, 2, 3 or 4 that are important for different treatment strategies: patients with HCV genotype 1 or 4 received treatment for 48 weeks with pegylated PEG-interferon-alpha (IFN-α2a, 180 μg weekly and ribavirin 1000 mg daily (body weight < 75 kg) or 1200 mg daily (body weight ≥ 75 kg). Patients with genotype 2 or 3 received PEG-IFN-α2a, 180 μg weekly and ribavirin 800 mg daily, for 24 weeks.
P = 0.029) before IFN treatment was started. Patients with a major depressive episode during IFN therapy demonstrated significantly lower scores on TMT A (t = 2.104, d.f. = 83, P = 0.038), and semantic word fluency (t = 2.265, d.f. = 85, P = 0.026) compared with patients without depression at baseline. Finally, patients with a severe depressive syndrome (MADRS score ≥25) had significantly lower scores in the TMT A (t = 3.927, d.f. = 87, P < 0.001) and TMT B at baseline (t = 3.105, d.f. = 83, P = 0.003).

With respect to the SCAG, patients with IFN-associated depressive syndrome and major depressive episodes had significant higher scores before INF therapy was started (MADRS ≥13: t = 2.901, d.f. = 88, P = 0.005; major depressive episodes: t = 2.256, d.f. = 88, P = 0.027).

**Prediction of INF-associated depression**

A multivariable regression analysis was performed to identify those risk factors that maximally discriminated between patients who developed any kind of a clinically relevant depressive syndrome (MADRS scores ≥13) during IFN treatment and those who remained without clinically relevant depressive mood changes (Fig. 1). A baseline MADRS score of 3 or higher was associated with a 6.4-fold increased higher risk for the development of an IFN-associated clinically relevant depressive syndrome (95% CI 1.7–24.0, P = 0.006). A BMI less than 25 at baseline resulted in a significantly higher risk for the development of a clinically relevant depressive syndrome (OR = 4.63, 95% CI 1.5–14.8, P = 0.010). In contrast, logistic regression analysis revealed that neither major depressive episodes according to DSM-IV criteria nor severe depressive syndromes (MADRS score ≥25) were associated with any baseline factors (data not shown).

A multiple logistic regression analysis was used again to calculate odds ratios of the incidence of INF-associated depressive syndromes (MADRS ≥13, major depressive episodes, MADRS ≥25) adjusted for TMT A, TMT B, Regensburger verbal fluency test semantic and phonematic word fluency and the SCAG score. Reduced performance on the TMT A (OR = 1.1, 95% CI 1.0–1.1, P = 0.027) and TMT B (OR = 1.0, 95% CI 1.0–1.1, P = 0.033) predicted the development of severe depression (MADRS ≥25) during antiviral treatment. Higher scores on the SCAG (OR = 1.23, 95% CI 1.0–1.4, P = 0.014) was found to be the only

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**Table 2  Pre-treatment depression symptoms on the Montgomery–Åsberg Depression Rating Scale (MADRS) items**

<table>
<thead>
<tr>
<th>MADRS single items at baseline</th>
<th>Mean (s.d.)</th>
<th>All (n = 91)</th>
<th>No depression (n = 42)</th>
<th>Major depression (DSM-IV) (n = 17)</th>
<th>Depression, MADRS score &gt; 13 (n = 49)</th>
<th>Severe depression, MADRS score &gt; 25 (n = 11)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Apparent sadness</td>
<td>0.3 (0.6)</td>
<td>0.1 (0.4)</td>
<td>0.5 (0.7)</td>
<td>0.5 (0.8)**</td>
<td>0.4 (0.7)</td>
<td></td>
</tr>
<tr>
<td>Reported sadness</td>
<td>0.3 (0.7)</td>
<td>0.1 (0.3)</td>
<td>0.5 (0.9)</td>
<td>0.5 (0.8)**</td>
<td>0.4 (0.7)</td>
<td></td>
</tr>
<tr>
<td>Inner tension</td>
<td>0.4 (0.8)</td>
<td>0.3 (0.7)</td>
<td>0.5 (1.0)</td>
<td>0.5 (0.9)</td>
<td>0.5 (0.9)</td>
<td></td>
</tr>
<tr>
<td>Reduced sleep</td>
<td>0.7 (1.0)</td>
<td>0.5 (0.9)</td>
<td>0.8 (1.0)</td>
<td>0.9 (1.1)*</td>
<td>1.1 (1.1)</td>
<td></td>
</tr>
<tr>
<td>Reduced appetite</td>
<td>0.3 (0.7)</td>
<td>0.1 (0.3)</td>
<td>0.6 (1.1)*</td>
<td>0.4 (0.9)**</td>
<td>0.9 (1.3)*</td>
<td></td>
</tr>
<tr>
<td>Loss of concentration</td>
<td>0.3 (0.7)</td>
<td>0.1 (0.5)</td>
<td>0.6 (0.9)**</td>
<td>0.5 (0.7)**</td>
<td>0.9 (0.9)**</td>
<td></td>
</tr>
<tr>
<td>Lassitude</td>
<td>0.3 (0.6)</td>
<td>0.1 (0.4)</td>
<td>0.6 (0.7)*</td>
<td>0.5 (0.7)**</td>
<td>0.6 (0.7)*</td>
<td></td>
</tr>
<tr>
<td>Inability to feel</td>
<td>0.1 (0.4)</td>
<td>0.0 (0.2)</td>
<td>0.2 (0.6)</td>
<td>0.2 (0.5)**</td>
<td>0.3 (0.6)</td>
<td></td>
</tr>
<tr>
<td>Pessimistic thoughts</td>
<td>0.3 (0.7)</td>
<td>0.1 (0.4)</td>
<td>0.8 (1.1)**</td>
<td>0.5 (0.8)*</td>
<td>0.8 (1.2)**</td>
<td></td>
</tr>
<tr>
<td>Suicidal thoughts</td>
<td>0.0 (0.2)</td>
<td>0.0 (0.0)</td>
<td>0.1 (0.3)*</td>
<td>0.06 (0.2)</td>
<td>0.09 (0.3)</td>
<td></td>
</tr>
</tbody>
</table>

*P < 0.05, **P < 0.01.

**Fig. 1  Odds ratios for developing any clinically relevant depressive syndrome (Montgomery–Åsberg Depression Rating Scale (MADRS) scores ≥13) during interferon treatment.**

BMI, body mass index; BDI, Beck Depression Inventory.
relevant risk factor for the development of an INF-associated depressive syndrome (MADRS ≥ 13) during antiviral therapy.

**Decision tree for the prediction of INF-associated depression**

As a MADRS score of 3 or higher was determined as the only significant predictor for an INF-associated depressive syndrome, we analysed the individual items of the MADRS. Items were included in the decision tree regression analyses in order to define cut-off points for each symptom of depression on the MADRS scale at baseline to discriminate between patients with or without an INF-induced depressive syndrome (Fig. 2). Patients were classified according to their values on the MADRS items as independent variables, and according to the emergence of a depressive syndrome (MADRS ≥ 13). The final decision tree analysis distinguished between patients with clinical depression and those without with a sensitivity of 73.6%. For developing INF-induced depression (MADRS ≥ 13), statistically significant predictors at baseline evaluation were reported sadness, loss of concentration, pessimistic thoughts and reduced sleep.

In total, 89% \( (n = 16) \) of patients with a baseline score for ‘reported sadness’ greater than 0 developed a clinically relevant depressive syndrome during INF-therapy. Further analyses revealed that 75% \( (n = 9) \) of patients without reported sadness, but with loss in concentration (score > 0) developed an INF-associated depressive syndrome (MADRS ≥ 13). The prediction rate for absence of depression with INF-treatment (specificity) for this decision tree was 73.8%.

**Discussion**

**Main findings**

Clinically relevant depressive symptoms emerged in up to 70% of patients with chronic hepatitis C who received antiviral therapy including IFN-α. Two recently published meta-analyses demonstrated that pre-emptive antidepressant treatment with SSRIs significantly reduces the incidence of INF-associated depression. However, strategies are demanded that help to discriminate between patients at risk of INF-induced depression and those who are not. Our results show significant differences at baseline between patients with and without INF-associated depressive syndromes. Pre-existing subclinical depressive mood (MADRS ≥ 3) resulted in a 6.4-fold increased risk of clinically relevant INF-associated depressive syndromes in patients without...
a history of psychiatric disorders. By using decision tree analysis, four single items of the MADRS scale – reported sadness, loss of concentration, pessimistic thoughts and reduced sleep – stood out in predicting the development of a clinically relevant depressive syndrome (MADRS ≥ 13) with a sensitivity of 73.8%. A total of 89% (n = 16) of patients with reported sadness at baseline (MADRS single item, score ≥ 0) developed a clinically relevant depressive syndrome during INF-therapy.

Our results are in line with previous studies demonstrating that patients with higher MADRS scores at baseline have a higher risk for the development of depressive symptoms during cytokine therapy. Capuron and colleagues identified similar mood (sadness, pessimistic thoughts) and behavioural symptoms (sleep disturbance) as significant predictors of depressive symptoms at the end of the first month of cytokine treatment. These findings support the notion that even subclinical deterioration of the affective state of patients before the initiation of INF therapy may indicate an increased individual vulnerability for the development of clinically significant INF-associated depressive symptoms. Interestingly, the number or severity of these symptoms seem not to be decisive factors. Rather, our results indicate an ‘all-or-nothing’ effect: the risk of developing clinically relevant depressive symptoms during INF treatment was increased whenever patients reported any kind of subclinical depressive symptoms, especially sadness and in addition sleep disturbance, pessimistic thoughts and lack of concentration. Thus, screening of patients for any degree of these pre-existing symptoms is advisable.

The predictive value of subclinical depressive symptoms before the beginning of interferon therapy shows similarities with the concept of subsyndromal symptomatic depression. This has been defined as minimal depressive symptoms below the diagnostic threshold for minor, dysthymic or major depressive disorders. Subsyndromal symptomatic depression is frequently observed in patients with unipolar major depressive disorder and is considered as an integral component of the symptomatic course of illness. In addition, the prevalence of subsyndromal symptomatic depression shows significant associations with a higher prevalence of future and past major depressive episodes. A similar relationship seems to exist for INF-associated depression and subclinical depressive symptoms before the beginning of antiviral therapy. However, it should be emphasised that despite major depression being a serious and highly prevalent side-effect of INF therapy, most affected patients will respond to acute treatment with SSRIs, or depression might be prevented by pre-emptive antidepressant treatment. Thus, even with the increased risk for depression, it is not indicated to exclude patients with minimal depressive symptomatology from antiviral treatment with INF-α.

Another important and novel finding is that severe depression can be predicted by neuropsychological tests measuring cognitive flexibility (TMT A and B). It was evident that inferior cognitive performance predicts especially severe depression before the start of INF treatment. This increased vulnerability to severe depression was even found in patients without a history of any affective disorders. With respect to subjective cognitive problems as measured by the SCAG, patients with INF-associated depressive syndromes (MADRS ≥ 13) had significantly higher scores before INF treatment was started.

**Cognitive impairment and HCV infection**

Cognitive difficulties are associated with impaired work productivity. Of note, treatment-naive patients with chronic hepatitis C infection have an already increased risk of mild cognitive impairment. Chronic infection with HCV is suspected to cause lasting changes to the immune system and to neurotransmission, leading to mild cognitive impairment. Different hypotheses have been proposed to explain the occurrence of cognitive impairment in HCV infection, such as a direct action of the virus on the central nervous system (CNS), a delayed action of the virus on the CNS mediated by viral replication in neurons and cognitive deterioration as a side-effect of the inflammatory process. In contrast, a recently published study demonstrated no evidence of an association between HCV infection and cognitive impairment. The authors argue that a lack of rigorous selection criteria in previous studies may have resulted in an overestimation of the prevalence of cognitive impairment associated with HCV by including patients with a history of alcohol and/or illegal drug misuse, patients using psychotropic drugs or patients with depression. However, our trial, which excluded patients with psychiatric disorders, now confirms the presence of at least mild cognitive impairment in a subgroup of treatment-naive patients with chronic hepatitis C independent from depressive symptoms.

On the other hand, and independent from HCV infection, two recent meta-analyses support the presence of cognitive impairment in patients with major depressive disorder. These studies demonstrate that executive function mediated by the prefrontal cortex and measured with the TMT B is affected by depression, which may have an impact on cognitive performance. The finding that a reduced performance on TMT A and TMT B predicts the development of severe depression (MADRS ≥ 25) lends further support to a depression–executive dysfunction syndrome. Executive dysfunction is also one of the clinical expressions of abnormalities in the frontostriatal–limbic circuitry that might predispose to late-life depression.

**BMI**

In our trial a BMI ≥ 25 at baseline resulted in a significantly reduced risk of developing clinically relevant depression with INF treatment. Obesity is one of the core characteristics that has been associated with lower sustained viral response rates in patients with chronic hepatitis and might be associated with a higher relapse rate. Therapy with IFN-α in these patients might have less clinical antiviral effects, resulting in lower rates of neuropsychiatric side-effects.

**Limitations**

Although the present study generally supports the utility of the TMT as a screening tool for HCV patients before the beginning of INF therapy, further investigations are clearly required. For example, additional cognitive tests should be conducted in future studies to evaluate whether other cognitive functions also predict severe INF-associated depression (for example, attention or memory). A more extensive evaluation would provide a stronger and more rigorous basis for assessing the potential impact of cognitive functions at the beginning of INF therapy in relation to INF-associated depression. Although a reasonable number of patients was included, numbers did not allow for a more precise identification of risk factors for developing DSM-IV major depressive episodes or severe depression (MADRS ≥ 25). Finally, multiple testing might limit the results.

**Implications**

In summary, our data demonstrate that pre-existing subclinical depressive mood significantly increases the risk of clinically relevant IFN-associated depressive syndromes in patients without a history of psychiatric disorders. Mild cognitive disturbances are an independent risk factor for the development of severe depression. Scoring on the MADRS single items reported sadness,
loss of concentration, pessimistic thoughts and reduced sleep prior to treatment, or a MADRS total score > 3 at baseline may identify patients at risk of treatment-emergent depression. Pre-emptive treatment with antidepressants is recommended in these high-risk patients.

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First received 18 Nov 2013, final revision 17 Jun 2014, accepted 19 Jun 2014

Acknowledgements

We thank Professor Heinz Grunze for a critical review of the paper and helpful advice.

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BJP published online October 30, 2014 Access the most recent version at DOI: 10.1192/bjp.bp.113.141770