Biological and psychological components of depression in patients receiving IFN-alpha therapy for hepatitis C

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A – research concept and design; B – collection and/or assembly of data; C – data analysis and interpretation; D – writing the article; E – critical revision of the article; F – final approval of the article

Abstract

Background. Depressive symptoms are frequent side effects of interferon α therapy (IFN-α). Both biological and psychological processes may occur concomitantly during hepatitis C treatment.

Objectives. This study was carried out to determine the impact of biological (immune response) and psychological factors on formation of depressive symptoms and major depressive disorder (MDD) during hepatitis C treatment.

Material and methods. A total of 99 patients receiving pegylated IFN-α and ribavirin for chronic C type hepatitis participated in the prospective cohort study. Symptoms of depression were assessed with the Montgomery-Åsberg Depression Rating Scale (MADRS) during treatment and 24 weeks after treatment. Neuroticism was measured with the Eysenck Personality Questionnaire – Revised (EPQ-R/N). Diagnosis of MDD was made using the Present State Examination (PSE-10) and DSM-IV-TR criteria. Factor analysis was used to detect factors adding up to total severity of depressive symptoms. Predictors of MDD were investigated using logistic regression analysis.

Results. Factor analysis returned 3 factors: 1st – MADRS scores at weeks 0–12, 2nd – MADRS and N scores before treatment, 3rd – MADRS at the 24th week of treatment and 24 weeks after treatment. The total severity of depressive symptoms consisted of 3 components: personality-related before treatment, IFN-α-related during treatment and dependent on the effect of treatment. Regression analysis showed that a history of psychiatric disorders (OR = 4.8) and MADRS scores before treatment (OR = 1.25) were predictors of MDD, as opposed to level of neuroticism.

Conclusions. The severity of depressive symptoms and MDD during the hepatitis C treatment was related to general depressive vulnerability, not to psychological factors related to neuroticism.

Key words: depression, hepatitis C, interferon α, neuroticism trait
Introduction

Interferon α (IFN-α) in combination with ribavirin used to be the main pharmacological treatment for the chronic hepatitis C virus (HCV) infection. Interferon α is known to induce several neuropsychiatric side effects including depression, anxiety, psychosis, hypomanic mood, and cognitive impairment. Depression is a particularly common side effect and in some rare cases it may be associated with suicidal ideation or suicide attempts. Previous psychiatric history was a strong risk factor for depression, anxiety and other psychiatric disorders during treatment, which might suggest that the overall susceptibility to depression and anxiety, including susceptibility of a psychological origin, underlay neuropsychiatric side effects induced by IFN-α.

Interferon-α-induced neuropsychiatric symptoms have been attributed to the release of pro-inflammatory cytokines modulating several neurophysiological and neuroendocrine systems involved in mood regulation. This is a multidirectional action involving neurotransmitter systems in the central nervous system (CNS), biochemical changes in the CNS and in the hypothalamic–pituitary–adrenal (HPA) axis. Interferon α is a potent inducer of pro-inflammatory cytokines including interleukins 1 and 6 (IL-1, IL-6) and tumor necrosis factor α (TNF-α). These cytokines play an important role in the development of sickness behavior – a set of psychological and behavioral changes that may lead to depressive symptoms in approx. half of the patients treated with IFN-α. The presumably depressogenic effect is not directly caused by cytokines but is a consequence of sickness behavior. This cytokine-induced behavioral syndrome is associated with alterations in the metabolism of neurotransmitters such as serotonin, norepinephrine and dopamine in brain regions essential to the regulation of emotions, including the limbic system, as well as the regulation of psychomotor and reward functions, including the basal ganglia. Cytokines activity is associated with: 1) significant alterations in diurnal HPA axis activity including the flattening of the adrenocorticotropic (ACTH) and cortisol diurnal fluctuations, and an increase in evening ACTH and cortisol concentrations; 2) increased activity of the metabolic enzyme indoleamine 2,3-dioxygenase responsible for degrading tryptophan to kynurenine which is then metabolized to quinolinic acid interfering with CNS activity; 3) decreased brain-derived neurotrophic factor (BDNF) levels, and 4) altered function of the glucocorticoid receptors.

The effect of IFN-α seems to be purely biological rather than psychological, but patients during an antiviral treatment are going through a difficult phase in their life caused by neuropsychiatric symptoms such as emotional lability, cognitive decline and insomnia, as well as somatic symptoms including fever, nausea, lack of appetite, and weakness. In vulnerable individuals, this stressful situation may cause depression. Significantly, both biological and psychological processes may occur concomitantly during hepatitis C treatment; in this light, the present paper offers an analysis of their mutual interdependence.

Material and methods

Participants and treatment

A total of 99 patients (50 men and 49 women) were included in the study. All of them had a chronic HCV infection with a detectable serum HCV-RNA concentration and a compensated liver disease. The patients were scheduled for treatment with pegylated IFN-α and ribavirin in the Ward of Infectious Diseases at the Department of Infectious Diseases, Liver Diseases and Acquired Immune Deficiencies at Wroclaw Medical University, Poland. All participants were over 18 years of age and were recruited from native Polish populations in the Lower Silesian region. The inclusion scheme was based on the order in which the patients had showed up for hepatitis C treatment, with sex stratification used to compensate for the number of men and women in the study group. The exclusion criteria entailed comorbidity with severe somatic disease, autoimmune diseases, neurological disorders (including dementia or brain injury), substance dependence (except for nicotine), active psychotic disorders, depression, and pregnancy. The study protocol was approved by the Wroclaw Medical University Ethics Committee. All patients provided their written informed consent.

Initially, the subjects were receiving weekly doses of 180 µg Pegintron® (PEG-IFN-a2a, Hoffmann-LaRoche, Basel, Switzerland) combined with either 1,000 mg/day or 1,200 mg/day of Rebetol (ribavirin, Schering-Plough Corporation, Kenilworth, USA), depending on their body weight (those weighing 75 kg and more were administered the higher dose). Except for patients infected with genotype 3 virus, who were treated for 24 weeks; in the case of all other subjects the treatment lasted 48 weeks.

Psychiatric assessments

The study followed a prospective longitudinal cohort design. The subjects were evaluated before the treatment (week 0), at weeks 2, 4, 8, 12, and 24, and 24 weeks after the conclusion of the treatment. Because patients with HCV genotype 3 were treated for 24 weeks and the remainder for 48 weeks, assessment point at the end of treatment was omitted in analysis. At each assessment point blood samples were collected and 2 psychometric scales were administered: Montgomery-Åsberg Depression Rating Scale (MADRS) and the Present State Examination (PSE-10). Neuroticism score was assessed once, at the beginning of the study, with the Eysenck Personality Questionnaire – Revised (EPQ-R/N).
The MADRS scale consists of 10 items assessing symptoms associated with depression, each item scored from 0 to 6 according to severity.\textsuperscript{10} We used the MADRS scale as adapted by Mazurek et al., which proved its good psychometric properties.\textsuperscript{11} The PSE-10 scale is a part of the Schedules for Clinical Assessment in Neuropsychiatry (SCAN 2.1) – a set of instruments and manuals aimed at assessing, measuring and classifying psychopathology and behavior associated with major psychiatric disorders of adult life.\textsuperscript{12} We utilized only sections 6 and 7 of the PSE-10 (i.e., depressed mood and ideation, dysthymia, recurrent brief depressive disorder and thinking, concentration energy, interests), adopting time criteria for depression. The Polish adaptation of SCAN was a collateral result of the international EDEN (European Day Hospital Evaluation) project and as such was successfully tested in several clinical applications.\textsuperscript{13–15} The PSE-10 served as a diagnostic tool for depression according to the DSM-IV criteria at all time points. Psychiatric disorders prior to treatment were assessed by a psychiatrist following a clinical interview. The scale of neuroticism was derived from the EPQ-R questionnaire. It consists of 24 items referring to a minor emotional distress. Neuroticism is an indicator of vulnerability to depression understood as a reaction to stressful life events.\textsuperscript{16–18} We used the Polish adaptation of the EPQ-R authored by Brzozowski and Drwal.\textsuperscript{19}

Depressive symptoms resulting from stressful life events are the effect of the interaction between the burden of stressful life events (SLE) and neuroticism level (N).\textsuperscript{17} Depressive symptoms due to biological processes are not dependent on personality structure, and thus are not related to neuroticism level. We assumed that the severity of depressive symptoms is the sum of 2 components: the biological (IF) and psychological (neuroticism, response to treatment effects) context.

The logistic regression/stepwise model method was used to determine predictors for major depressive disorder (MDD). Demographic, biological and psychiatric variables were included in the marginal results table using logistic regression forward stepwise procedure with sigma-restricted parameterization for categorical predictors. The dependent MDD variable was coded as MDD diagnosis for a good code and no MDD for a bad code. Variables which reached a statistically significant impact on the accuracy of a discriminative function were included in the analysis. All the statistical computations were performed with Dell STATISTICA v. 13.1 for Windows (Microsoft, Armonk, USA).

Results

Out of 99 patients enrolled, 85 (85.9\%) completed the treatment. The authors decided to perform analysis for MDD predictors in all patients except for 1 who had dropped out after 2 weeks of the treatment. The remainder stayed in treatment for at least 8 weeks which was considered sufficient for depressive symptoms to emerge. Table 1 shows pre-treatment demographic and clinical characteristics of the sample. None of the patients was diagnosed with MDD before the treatment but 14 patients (14\%) reached MADRS score of 13 and over, and 4 patients reached MADRS score of 18 and above, which meant that some of them were actually depressed although they did not meet the formal DSM-IV-TR criteria for a major depressive episode.\textsuperscript{20} The initial MADRS mean score in men (5.7) was slightly lower than in women (7.0), but the difference was not significant (t = –1.38, p = 0.17). Of the 98 patients 42 (42.9\%) met criteria for MDD at any time within the 24 weeks of treatment, women more often (49.0\%) than men (36.8\%), but this difference was not statistically significant (χ² = 1.5, p = 0.22); neither was the difference in mean maximal MADRS score between men (16.4) and women (18.5) significant (t = –1.44, p = 0.15).

The mean severity of depressive symptoms fluctuated in the course of treatment: it doubled during the first 4 weeks, then remained stable until week 24, and dropped slightly below the initial level 24 weeks after the treatment (Fig. 1).

In order to assess the structure of depressive symptoms and their association with the level of neuroticism (i.e., psychological factors), factor analysis was performed in 85 (85.9\%) patients that completed the treatment. Eigenvalues of the first 3 factors were 5.62, 0.88 and 0.62, respectively. The remaining eigenvalues were below 0.35, so analysis for a model with 3 factors was carried out. Factor analysis (Table 2) separates MADRS scores during treatment (weeks 2, 4, 8, 12, and 24 – factor 1) from those after treatment (week 24 of the treatment and the 24\textsuperscript{th} week after
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The association between the severity of depressive symptoms and neuroticism scores observed in this model points towards a psychogenic origin of these symptoms. This association is only significant before the treatment, so the rise of the severity of depressive symptoms during treatment is not related to initial neuroticism level, which is attributable to personality factors.

Stepwise logistic regression method returned 3 variables that were found to significantly affect the diagnosis of MDD, i.e., the initial MADRS score and individual history of psychiatric disorders. After these variables had been entered into the logistic regression model, only 2 proved to be statistically significant, i.e., the initial MADRS score and individual history of psychiatric disorders. The results of logistic regression calculations are presented in Table 3.

**Discussion**

Shifts in the severity of depressive symptoms during antiviral treatment may appear as a continuous process. Mild symptoms which occur prior to treatment increase during treatment and decrease after treatment. In our cohort study, we managed to capture the diversity of this process. Factor analysis revealed 3 different components of depression changing with time. The main component (factor 1 in the 3-factor model) relates to IFN action. Depressive symptoms due to IFN administration persisted throughout the treatment. The biological impact of IFN...
on mood did not depend on psychological factors related to neuroticism. On the other hand, the 2 remaining components, which appear to be much weaker, seemed to be related to psychological factors. The 3rd factor includes depressive symptoms at 2 timepoints, i.e., at the 24th week of the treatment and 24 weeks after the treatment. Therefore, it might be assumed that this component of depression is an effect of stress which in fact corresponds to worries over the outcome of the antiviral treatment – those patients who had recovered were less depressive. Towards the end of treatment, the patients might have already begun to suspect what the results of their treatment would be, and at the endpoint of this study (24 weeks after the treatment) some of them gained certainty – for example, after receiving information about the presence of HCV RNA in blood plasma at the end of treatment. Given that the second factor includes MADRS and neuroticism ratings before treatment, it reflects the psychological reaction to a stressful life situation that the liver disease certainly is. Patients suffering from different somatic diseases are known to have depressive symptoms correlating with the level of their neuroticism: the more severe illness, the stronger the correlation between neuroticism and depression.21 Somatic illness as a stressful life situation may cause depressive symptoms in vulnerable individuals, and the neuroticism score is a measure of this vulnerability.22 The results of this study did not confirm this relationship – the severity of depressive symptoms before treatment turned out to be a stronger predictor of MDD than the level of neuroticism. This relationship is quite common in other studies on depression in which the initial severity of depressive symptoms was a strong predictor of depressive symptomatology.23–26 Depressive symptoms in general seem to be the main predictor of depressive disorders and broad psychiatric and somatic symptomatology. A similar situation has been replicated in the present study – depressive symptoms, despite their origin, foreshadowed MDD in patients treated for hepatitis C. Moreover, the diagnosis of depression during treatment depended solely on the initial severity of depressive symptoms.

In the 3-factor model, the pre-treatment depressive symptoms are almost evenly dispersed between factors 1 (biological) and 2 (psychological). The initial severity of symptoms seems to be a superposition of 2 depressive processes, i.e., the biological one which is related to HCV infection and evolves considerably during treatment, and the psychological one which is bound to neuroticism ratings and gradually loses its significance. Factor 1 related to the IFN-induced depressive symptoms (46% of variance) proved much stronger than factor 2 related to neuroticism (22% of variance); the point of reference for such a distinction is the overall severity of symptoms among which IFN-induced depression predominates. Seemingly, there exists a direct link between pre-treatment depression and IFN-induced depression. The inflammation processes, endogenic cytokines activity and direct effects of administered INF-α constitute the base for this interconnection.

Hepatitis C infection causes a chronic mild inflammation affecting both the liver and the brain, especially its white matter.27,28 Hepatitis type C virus replicates in the mononuclear cells of the immune system and within the brain cells.29,30 The inflammation is usually very mild and does not cause noticeable signs of encephalitis, but may still trigger a mild cognitive dysfunction in the form of attention and memory decline observable in the results of neuropsychological cognitive tests. Moreover, the inflammation may also be detected with the use of sophisticated neuroimaging methods such as the computed tomography diffusion tensor.31 It is very likely that IFN-induced cytokines activity during the treatment is the continuation of inflammation-induced cytokines activity from before the treatment. Both may lead to MDD through illness behavior.

The results of the study showed that patients who previously suffered from mental disorders, as well as patients with a high level of depressive symptoms before treatment, are more susceptible to the development of a depressive disorder during treatment with IFN-α. Giving more attention to this group of patients will facilitate a rapid implementation of antidepressant treatment, which may prevent cases of treatment discontinuation.

The results of the study suggest that active assessment of the severity of depressive symptoms during control examinations (not only responding to patients’ complaints) is necessary to capture the depressive disorder at the sub-clinical stage, in order to start a more effective treatment.

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**Table 3. Results of logistic regression analysis**

<table>
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<th>Effect</th>
<th>Wald statistics</th>
<th>OR</th>
<th>OR lower CL (95%)</th>
<th>OR upper CL (95%)</th>
<th>p-value</th>
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<td>–</td>
<td>–</td>
<td>–</td>
<td>0.0035</td>
</tr>
<tr>
<td>MADRS 0</td>
<td>8.16</td>
<td>1.25</td>
<td>1.07</td>
<td>1.46</td>
<td>0.0043</td>
</tr>
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<td>Neuroticism</td>
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<td>1.04</td>
<td>0.93</td>
<td>1.17</td>
<td>0.49</td>
</tr>
<tr>
<td>History of psychiatric disorders</td>
<td>6.18</td>
<td>4.79</td>
<td>1.40</td>
<td>16.47</td>
<td>0.013</td>
</tr>
</tbody>
</table>

OR – odds ratio; CL – confidence limit.