Multiple sclerosis and psychiatric disorders: comorbidity and sibling risk in a nationwide Swedish cohort

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Abstract

Background: Psychiatric disorders are known to be prevalent in Multiple Sclerosis (MS).

Objective: To study comorbidity between MS and bipolar disorder, schizophrenia and depression in a nation-wide cohort and to determine whether shared genetic liability underlies the putative association.

Methods: We identified ICD-diagnosed patients with MS (n=16,467), bipolar disorder (n=30,761), schizophrenia (n=22,781) and depression (n=172,479) in the Swedish National Patient Register and identified their siblings in the Multi-Generation Register. The risk of MS was compared in psychiatric patients and in matched unexposed individuals. Shared familial risk between MS and psychiatric disorders was estimated by sibling
Results: The risk of MS was increased in patients with bipolar disorder (hazard ratio [HR] 1.8, 95% confidence interval [CI] 1.6-2.2, p<0.0001) and depression (HR 1.9, 95% CI 1.7-2.0, p<0.0001). MS risk in schizophrenia was decreased (HR 0.6, 95% CI 0.4-0.9, p=0.005). The association between having a sibling with a psychiatric disorder and developing MS was not significant.

Conclusion: We found a strong positive association between MS and bipolar disorder and depression that could not be explained by genetic liability. The unexpected negative association between MS and schizophrenia might be spurious or indicate possible protective mechanisms that warrant further exploration.
Multiple sclerosis and psychiatric disorders: comorbidity and sibling risk in a nationwide Swedish cohort

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Author Contributions:

Viktoria Johansson, Cecilia Lundholm, Jan Hillert, Paul Lichtenstein, Mikael Landén and Christina M. Hultman were responsible for the study design and interpretation of the results. Paul Lichtenstein was responsible of the preparation of the data. Viktoria Johansson, Cecilia Lundholm and Christina M. Hultman worked with the data analysis and Cecilia Lundholm provided biostatistical expertise. Viktoria Johansson was responsible for literature search. Viktoria Johansson and Christina M. Hultman worked with the preparation of figures and tables. Viktoria Johansson, Cecilia Lundholm, Jan Hillert, Thomas Masterman, Paul Lichtenstein, Mikael Landén and Christina M. Hultman worked with the writing of the manuscript.
Key words:
Multiple Sclerosis, Bipolar disorder, Depression, Schizophrenia, Genetic liability, Cohort study

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Introduction

Multiple sclerosis (MS) is an inflammatory autoimmune disease resulting in demyelination and axonal loss of the central nervous system (CNS), which lead to disability with detrimental effects on quality of life (1). Emerging evidence suggests that inflammation also plays an important role in psychiatric disorders such as bipolar disorder, schizophrenia and depression (2). Several studies have shown increased rates of depression and bipolar disorder in patients with MS (3-14). In some studies, MS has also been associated with an increased risk of schizophrenia and non-affective psychosis (11, 15-17), while in other studies no association was demonstrated (14, 18).

The association between MS and these psychiatric disorders might be explained by shared familial liability. Family, twin and adoption studies in MS, as well as in bipolar disorder, depression and schizophrenia, indicate moderate to high degrees of genetic influence; and moderate to high genetic correlations have been observed between the psychiatric diagnoses (19-21). In a population-based cohort study of parents and siblings to MS patients, an increased risk of schizophrenia, but not bipolar disorder, was found (7, 17), and in another study depression was not more common in first-degree relatives of MS patients (n=221) than in unexposed individuals (10).

Some of these previous studies have had limited statistical power due to modestly sized data sets (10), while others have arrived at contradictory results with regard to non-affective psychosis and schizophrenia (11, 14-18) or failed to report risks for matched population controls. Here, with a view to studying the relationship between MS and bipolar disorder, schizophrenia and depression, we sought to overcome these limitations by using a Swedish national database, which constitutes, to our knowledge, the largest cohort ever employed for this purpose. In addition, we used a
matched cohort sibling design to investigate whether shared genetic liability might underlie the putative comorbidity between MS and the psychiatric disorders.

**Materials and Methods**

**Data sources**

The study was approved by the ethics committee of Karolinska Institutet. The Swedish National Patient Register (NPR, National Board of Health and Welfare, http://www.socialstyrelsen.se) is a nationwide register containing information on inpatient care (psychiatric diagnoses from 1973 and MS from 1987) and specialist care (from 2001). It has a high level of agreement with medical and psychiatric diagnoses (22). The Swedish MS Register (SMSreg; http://www.msreg.net) was started in 1996 and includes information on 13,558 unique individuals registered prior to January 1, 2010; MS diagnosis are decided by consultant neurologists (23). The Swedish Multi-Generation Register (Statistics Sweden, http://www.scb.se) is a national database with coverage of individuals born in Sweden since 1932 and living in Sweden after 1960, and of immigrants who became Swedish citizens before the age of 18. The Cause of Death Register (National Board of Health and Welfare) provides data regarding all deaths, and the Total Population Register (Statistics Sweden) contains information on immigration and emigration from the country.

**Disease classification**

Using codes from the International Classification of Diseases (ICD 8-10), we identified in the NPR patients diagnosed with bipolar disorder, schizoaffective disorder, schizophrenia, depression and MS (Table 1). We identified a subgroup of patients with bipolar disorder who had experienced one or more episodes of mania (designated bipolar disorder type I), and a subgroup of subjects with
depression who had experienced at least one episode of severe depression (designated severe depression). A validation study of the bipolar diagnosis in the NPR found that two independent inpatient admissions generated a positive predictive value of 0.81 (24). A requirement of two diagnoses to define schizophrenia resulted in frequencies comparable to internationally reported lifetime prevalence percentages (25). To secure diagnostic specificity, we required two or more diagnoses from the NPR for all diagnostic categories. To differentiate between bipolar disorder, schizophrenia and schizoaffective disorder, we did not allow any overlap between the three diagnoses, and subjects with more than one of these diagnoses were classified in a separate mixed category (designated mixed bipolar/psychosis).

We used the SMSreg to verify the quality of the MS diagnoses from the NPR where individuals are coded as having MS according to the McDonald criteria (26). 96% of the patients with a diagnosis of MS in the NPR were registered as having MS in the SMSreg.

**Study population and design**

To study associations between psychiatric disorders and MS, we used a matched cohort design based on individuals and pairs of siblings living in Sweden, whom we had identified from, register linkages with the help of the unique personal identification number assigned to each Swedish resident.

In a “comorbidity cohort” we assessed the association between a psychiatric diagnosis (exposure directly to the particular disorder) and risk of MS (outcome), and in a “sibling cohort” (familial analysis) we assessed the association between having a sibling with a psychiatric disorder (exposure to an inherited or environmentally mediated liability to the particular disorder) and risk
of MS (outcome; Figure 1). In the comorbidity cohort, we identified the exposed patients and 10 randomly selected unexposed individuals per patient, matched by birth year and sex. Unexposed individuals had not received an exposure diagnosis prior to the date of the patient’s first diagnosis or start of follow-up, if it occurred later. In the sibling cohort all pairwise combinations of sibling pairs were considered. For each sibling pair exposed to the disorder of interest, we randomly selected 10 unexposed sibling pairs, matched for birth year and sex (for both individuals in the pair). The index person of the selected unexposed pairs was required to be alive, living in Sweden and free of the diagnosis of the exposed index person at the start of follow-up.

The analyses were also performed in reverse order, to assess the association between MS or having a sibling with MS (exposure) and the risk of a psychiatric disorder (outcome; Figure 2).

**Analysis by time period**

To address the influence of ascertainment bias in the comorbidity analysis, we analyzed the risk of MS in three different time windows, as in Figure 3 and an alternative analysis, as in Figure 4.

**Sensitivity analysis**

To test the sensitivity of our results, we performed several additional analyses; patients with schizophrenia were analyzed without exclusion of cases with additional diagnoses of schizoaffective or bipolar disorder; individuals born between 1969 and 1991 were analyzed separately; the data was re-analyzed, with MS as exposure and the start of follow-up as 2001; the depression cohort was reanalyzed including patients with single diagnosis of depression; to assure the accuracy of the MS diagnoses, we analyzed the data using diagnostic information first from the SMSreg alone and then in combination with diagnostic information from the NPR.
**Statistical analysis**

Descriptive statistics are presented as means and standard deviations for continuous data and frequencies and percentages for categorical data. Estimates for the risk of being diagnosed with an event among exposed individuals were obtained by calculating hazard ratios (HR) with 95% confidence intervals (CI) by a Cox regression model using SAS version 9.3. To compensate for the clustering effect in the sibling analysis, we used robust sandwich estimates. Follow-up started in 1987 (or age 18) when MS was the event and in 1973 (or age 18) when psychiatric disorder was the event. Subjects were followed until an event occurred or until the subject was censored due to death, emigration or end of follow-up on December 31, 2009. Two-tailed probability values were calculated, with values less than 0.05 regarded as significant. In both the comorbidity and the sibling analyses, we adjusted for immigration status (born in or born outside of Sweden); effect modification was tested for by including interaction terms between sex and exposure.

**Results**

We identified 8,193,579 individuals born between 1932 and 1991 who were alive and residing in Sweden at the start of follow-up, Table 2.

**Comorbidity cohort—analysis of psychiatric disorders and MS**

We found a statistically significantly increased risk of MS in the patients with bipolar disorder as well as in the subcategory classified as bipolar disorder type I, Figure 5. In patients with schizophrenia, the risk of MS was significantly decreased. In schizoaffective disorder, MS risk was non-significantly decreased, whereas the risk in the mixed category did not differ from that of
unexposed subjects. In individuals with depression, MS risk was increased and the results remained significant in the subgroup of severely depressed patients. Adjustment for immigration status did not change the overall results. There was no evidence of sex-specific effects except for in bipolar disorder type I where the risk of MS was increased in males compared to females (interaction effect p=0.0008), Table 3. The results were similar when the risk of psychiatric disorders in the cohort of MS patients was analyzed, Table 4.

**Comorbidity cohort—analysis by time period**

The risk of MS was significantly higher in all three time windows (explained in Figure 3), and was not found to be associated with the date of onset of bipolar disorder, bipolar disorder type I and depression (explained in Figure 4), Table 5. The results for schizophrenia were not significant. The results remained unchanged when analyzed in the reverse direction.

**Sibling cohort—analysis of psychiatric disorders and MS**

The risk of MS in siblings to subjects with bipolar disorder, schizophrenia or depression did not differ from the risk in siblings to unexposed individuals, Figure 5. Sibling analyses were performed in the reverse direction, with similar results, Table 4.

**Sensitivity analysis**

Patients with schizophrenia, defined more widely, with inclusion of cases with additional diagnoses of schizoaffective or bipolar disorder, showed no significantly decreased risk of MS (HR 0.8, 95% CI: 0.7-1.0, p=0.1). To ensure that individuals would be no older than 18 years of age when entering the study in 1987 subjects born between 1969 and 1991 were analyzed. The results were in agreement with the main results but, in some cases, failed to reach statistical significance,
presumably due to lack of power. The outpatient register was started in 2001, which means that milder cases of depression might have been included from 2001 and onwards. To avoid selection bias, the comorbidity cohort was analyzed with MS as exposure, and the start of follow-up as 2001. The increased risk of depression in MS patients remained significant. We also analyzed the effect of depression on MS risk, including individuals with a single depression. In total, 303,763 individuals with depression were identified, and the increased risk of MS remained significant (HR 2.0 95% CI: 1.9-2.1, p<0.0001). Finally, using diagnostic information from the SMSreg alone and in combination with information from the NPR, we obtained similar results as in the main analysis.

Discussion

Comorbidity and familial liability between MS and bipolar disorder, schizophrenia and depression were studied in a nationwide Swedish cohort—to our knowledge, the largest study to date of MS and psychiatric conditions. A statistically significantly increased risk of comorbid MS was found in bipolar disorder and depression, but a significantly decreased risk of MS was found in schizophrenia. Compared to risks in unexposed siblings, the risk of MS was not increased in siblings to subjects with bipolar disorder, schizophrenia or depression, and the risk of these psychiatric disorders was not increased in siblings to MS patients. Our main conclusion is, therefore, that the associations between MS and the psychiatric disorders are not confounded by familial factors.

There are several strengths in this study: The large study base allowed us to perform analyzes by time period and to analyze subcategories such as bipolar disorder type I and severe depression. Potential confounders were accounted for by matching for age and sex and the data was analyzed
bidirectional. The quality of the MS diagnoses was assured by sensitivity analyses, using MS diagnosis from both the NPR and SMSreg. Limitations in our study include the circumstance that depression diagnoses in the NPR are not validated. Our lack of access to primary-care register data regarding depression may have resulted in a selective bias, in that we may have missed less severe cases of depression and therefore underestimated the magnitude of increased risks. However, inclusion of patients with a single depression diagnosis did not substantially affect estimates of MS risk. We were not able to adjust for socioeconomic status in childhood, but recent data indicate that socioeconomic status is not associated with MS (27) and it is not very likely that our results would be affected.

The comorbidity between MS and bipolar or depressive disorder were in agreement with previous studies in the field (3-14) and may be explained by several hypothetical mechanisms. First, the pathogenic events in MS, including the inflammatory process in the CNS, might give rise to depressive or even manic symptoms. Second, affective symptoms might be triggered by the psychological trauma of having been diagnosed with chronic, disabling disease such as MS. However, the analysis of the MS risk by time period revealed no temporal association between the occurrence of bipolar disorder or depression and the onset of MS, which argues against the role of purely psychological factors, Table 5. Also the increased risk of MS in the severe subgroups (bipolar disorder type I and severe depression) suggests involvement of biological mechanisms such as inflammation rather than psychological factors. Thirdly, the increased comorbidity could be an effect of ascertainment bias; however, the results from the analysis by time period do not support this explanation. A fourth possibility is that comorbidity is triggered by MS medications, such as interferon beta and corticosteroids, both known to cause psychiatric side effects (28), but again the results from the analysis by time period do not support this explanation. According to a
fifth possible mechanism, changes in behavior resulting from psychiatric disorders, such as
increased smoking or reduced exposure to ultraviolet light, might lead to reduced synthesis of
vitamin D, thus increasing the risk of MS (29). However, the notion that increased MS risk in
psychiatric conditions is mediated by smoking is not supported by our finding of decreased MS risk
in schizophrenia, given the high rate of tobacco smoking in patients with schizophrenia, compared
to rates in the general population (29). We also found gender differences as males with bipolar
disorder type I were at higher risk of MS compared to females, Table 3. The finding might deserve
further investigation.

The decreased risk of MS in patients with schizophrenia deviates from the findings of previous
studies (6, 11, 15-18) and there might be several reasons for the disparate results. The large
advantage with our data is the population-based approach, the large sample size and the limitation
to schizophrenia patients who did not exhibit overlap with schizoaffective disorder and bipolar
disorder. The significant negative association was only seen when no diagnostic overlap was
allowed, indicating that the results are specific to a particular schizophrenia phenotype lacking
bipolar affective symptoms. A negative association with schizophrenia has been observed in other
disorders mediated by inflammation, such as rheumatoid arthritis (30, 31). Like rheumatoid
arthritis, schizophrenia and MS have been shown to be associated with polymorphisms at human
leucocyte antigen (32, 33). Although no decreased familial risk was found between MS and
schizophrenia, it is tempting to speculate that risks conferred by alleles at HLA loci might explain
part of the negative association between MS and schizophrenia—as has previously been
demonstrated for DRB1*1501, which is a risk allele in MS and a protective allele in diabetes
mellitus type 1 (34). Finally protective effects, with regard to MS risk, of treatment with

http://mc.manuscriptcentral.com/multiple-sclerosis
antipsychotic medications or the possibility that patients with severe mental illness have a reduced ability to recognize and report symptoms to the healthcare-system, might also explain the findings.

Previous studies found no familial relationship between MS and bipolar disorder or depression—which is in accordance with our findings—but an increased risk of schizophrenia has been previously reported in patients with a family history of MS (7, 10, 17). Using a method with matched sibling controls, we found no significant familial link between MS and schizophrenia.

**Conclusion**

There is an increased risk of MS in bipolar disorder and depression, including bipolar disorder type I and severe depression, and, conversely, an increased risk of these psychiatric conditions in MS patients. In contrast to previous findings, in the present study, the risk of MS was found to be decreased in patients with schizophrenia, and the risk of schizophrenia decreased in patients with MS. However, MS was not found to be underrepresented in siblings to schizophrenic patients, nor was schizophrenia found to be underrepresented in siblings to MS patients—findings suggesting the involvement of other mechanisms than genetically determined liability. Clinicians should be aware of the increased risk of bipolar disorder and depression in patients with MS and pay particular attention to neurological symptoms in patients with schizophrenia.

**Funding**

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Conflict of interest statement

JH has received unrestricted research support from BiogenIdec, Merck-Serono, Bayer-Shering and Sanofi-Aventis and served on advisory boards or received compensation for lecturing for these companies as well as for TEVA and Novartis. VJ, CL, TM, PL, ML and CH declare no conflicts of interest.

References


Table 1 Diagnostic codes for disease classification according to International Classification of Diseases (ICD).

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Bipolar disorder</td>
<td>296.0-296.3, 296.8, 296.9</td>
<td>296.0-296.6, 296.8, 296.9</td>
<td>F30.0-F30.2, F30.8, F30.9, F31.1-F31.9</td>
</tr>
<tr>
<td>Schizophrenia</td>
<td>295.0-295.6, 295.9</td>
<td>295.0-295.6, 295.8, 295.9</td>
<td>F20</td>
</tr>
<tr>
<td>Schizoaffective disorder</td>
<td>2957</td>
<td>2957</td>
<td>F25</td>
</tr>
<tr>
<td>Depression</td>
<td>296.0, 300.4, 298</td>
<td>296.1, 300.4, 311, 298.0</td>
<td>F32, F33</td>
</tr>
<tr>
<td>Multiple sclerosis</td>
<td>340.99</td>
<td>340</td>
<td>G35</td>
</tr>
</tbody>
</table>

a Patients with only one code of bipolar disorder from ICD 8 and ICD 9 were classified as bipolar disorder if they had one or more of the following diagnosis (ICD8: 296.0, 296; ICD9: 2961)

b Patients who fulfilled the criteria of bipolar disorder and presented at least one of the following codes for several manic episodes (ICD8: 296.1; ICD9: 296.0, 296.2, 296.8, ICD10: F301, F302, F311, F312) were classified in the subgroup bipolar disorder type I

c Patients who fulfilled the criteria of depression and presented at least one of the following codes for severe depression (ICD8: 296.0, 298; ICD9: 296.1, 298.0; ICD10: F322, F323, F332, F333) were classified in the subgroup severe depression
Table 2 Characteristics for the patients with psychiatric disorders, multiple sclerosis and their siblings

### Psychiatric disorder as exposure and multiple sclerosis as the event

<table>
<thead>
<tr>
<th>Patients with a psychiatric disorder</th>
<th>Siblings to patients with a psychiatric disorder</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>No of patients</td>
</tr>
<tr>
<td>-------------------------------------</td>
<td>---</td>
</tr>
<tr>
<td>All psychotic disorders</td>
<td>71134</td>
</tr>
<tr>
<td>Bipolar disorder</td>
<td>30761</td>
</tr>
<tr>
<td>Bipolar type I</td>
<td>8695</td>
</tr>
<tr>
<td>Schizophrenia</td>
<td>22781</td>
</tr>
<tr>
<td>Schizo-affective disorder</td>
<td>1816</td>
</tr>
<tr>
<td>Mixed category</td>
<td>15776</td>
</tr>
<tr>
<td>Depression</td>
<td>172479</td>
</tr>
<tr>
<td>Severe depression</td>
<td>48329</td>
</tr>
</tbody>
</table>

### Multiple sclerosis as exposure and psychiatric disorder as the event

<table>
<thead>
<tr>
<th>Patients with multiple sclerosis</th>
<th>Sibling to patients with multiple sclerosis</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>No of patients</td>
</tr>
<tr>
<td>---------------------------------</td>
<td>---</td>
</tr>
<tr>
<td>All psychotic disorders</td>
<td>16437</td>
</tr>
<tr>
<td>Depression</td>
<td>16446</td>
</tr>
</tbody>
</table>

---

*Age of onset defined as age of the first healthcare contact due to the disorder.*
Table 3 The risk of multiple sclerosis (MS) in patients with psychiatric disorders stratified by sex.

<table>
<thead>
<tr>
<th>Diagnosis</th>
<th>Sex</th>
<th>Patients, n (MS affected, n)</th>
<th>HR</th>
<th>95% CI</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>All psychotic diagnosis</td>
<td>Male</td>
<td>34,628 (67)</td>
<td>1.24</td>
<td>0.97-1.60</td>
<td>0.0876</td>
</tr>
<tr>
<td></td>
<td>Female</td>
<td>36,506 (182)</td>
<td>1.29</td>
<td>1.11-1.50</td>
<td>0.0008</td>
</tr>
<tr>
<td>Bipolar disorder</td>
<td>Male</td>
<td>12,420 (42)</td>
<td>2.31</td>
<td>1.67-3.21</td>
<td>&lt;.0001</td>
</tr>
<tr>
<td></td>
<td>Female</td>
<td>18,341 (111)</td>
<td>1.70</td>
<td>1.40-2.06</td>
<td>&lt;.0001</td>
</tr>
<tr>
<td>Bipolar type I</td>
<td>Male</td>
<td>3631 (23)</td>
<td>4.76</td>
<td>2.94-7.70</td>
<td>&lt;.0001</td>
</tr>
<tr>
<td></td>
<td>Female</td>
<td>5064 (39)</td>
<td>1.76</td>
<td>1.27-2.44</td>
<td>0.0007</td>
</tr>
<tr>
<td>Schizophrenia</td>
<td>Male</td>
<td>14,525 (16)</td>
<td>0.71</td>
<td>0.43-1.18</td>
<td>0.1838</td>
</tr>
<tr>
<td></td>
<td>Female</td>
<td>8256 (20)</td>
<td>0.56</td>
<td>0.36-0.88</td>
<td>0.0122</td>
</tr>
<tr>
<td>Depression</td>
<td>Male</td>
<td>68,300 (202)</td>
<td>2.20</td>
<td>1.90-2.54</td>
<td>&lt;.0001</td>
</tr>
<tr>
<td></td>
<td>Female</td>
<td>104,179 (563)</td>
<td>1.77</td>
<td>1.63-1.92</td>
<td>&lt;.0001</td>
</tr>
<tr>
<td>Severe depression</td>
<td>Male</td>
<td>19,733 (48)</td>
<td>1.84</td>
<td>1.40-2.44</td>
<td>&lt;.0001</td>
</tr>
<tr>
<td></td>
<td>Female</td>
<td>28,596 (137)</td>
<td>1.36</td>
<td>1.15-1.60</td>
<td>0.0003</td>
</tr>
</tbody>
</table>

Hazard Ratios (HR) and 95% confidence limits (95% CI). The hazard ratios are not adjusted for immigration status.
Table 4 The risk of psychiatric disorders (PD) in patients with multiple sclerosis (MS) and siblings to patients with MS. (MS as exposure and PD as event)

<table>
<thead>
<tr>
<th>disorder</th>
<th>Patients, n (PD affected, n)</th>
<th>HR</th>
<th>95% CI</th>
<th>p-value</th>
<th>Siblings to patients, n (PD affected, n)</th>
<th>HR</th>
<th>95% CI</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>All psychotic disorders</td>
<td>16,467 (250)</td>
<td>1.37</td>
<td>1.20-1.56</td>
<td>&lt;.0001</td>
<td>26506 (306)</td>
<td>0.96</td>
<td>0.85-1.08</td>
<td>0.4665</td>
</tr>
<tr>
<td>Bipolar disorder</td>
<td>16,467 (154)</td>
<td>1.86</td>
<td>1.56-2.20</td>
<td>&lt;.0001</td>
<td>26506 (126)</td>
<td>0.93</td>
<td>0.77-1.11</td>
<td>0.4157</td>
</tr>
<tr>
<td>Bipolar type I</td>
<td>16,467 (62)</td>
<td>2.37</td>
<td>1.79-3.12</td>
<td>&lt;.0001</td>
<td>26506 (31)</td>
<td>0.77</td>
<td>0.53-1.10</td>
<td>0.1513</td>
</tr>
<tr>
<td>Schizophrenia</td>
<td>16,467 (36)</td>
<td>0.71</td>
<td>0.51-1.00</td>
<td>0.0478</td>
<td>26506 (93)</td>
<td>0.88</td>
<td>0.71-1.09</td>
<td>0.4424</td>
</tr>
<tr>
<td>Schizoaffective disorder</td>
<td>16,467 (3)</td>
<td>0.64</td>
<td>0.20-2.09</td>
<td>0.4629</td>
<td>26506 (4)</td>
<td>0.67</td>
<td>0.24-1.85</td>
<td>0.0411</td>
</tr>
<tr>
<td>Mixed category</td>
<td>16,467 (57)</td>
<td>1.28</td>
<td>0.97-1.68</td>
<td>0.0815</td>
<td>26506 (83)</td>
<td>1.15</td>
<td>0.92-1.45</td>
<td>0.2242</td>
</tr>
<tr>
<td>All depression</td>
<td>16,446 (763)</td>
<td>1.86</td>
<td>1.72-2.01</td>
<td>&lt;.0001</td>
<td>26474 (681)</td>
<td>1.02</td>
<td>0.95-1.11</td>
<td>0.5524</td>
</tr>
<tr>
<td>Severe depression</td>
<td>16,446 (180)</td>
<td>1.52</td>
<td>1.30-1.78</td>
<td>&lt;.0001</td>
<td>26474 (191)</td>
<td>0.99</td>
<td>0.85-1.15</td>
<td>0.8985</td>
</tr>
</tbody>
</table>

Hazard Ratios (HR) and 95% confidence limits (95% CI). The hazard ratios are not adjusted for immigration status.
**Table 5** Risk of multiple sclerosis (MS) in patients with psychiatric disorders (PD), and risk of PD in patients with MS, by time period.

<table>
<thead>
<tr>
<th>Risk of MS in patients with PD</th>
<th>Up to 2 years prior to date of PD onset (^a)</th>
<th>Date of onset of PD ±2 years (^a)</th>
<th>2 years after PD onset and onwards (^a)</th>
<th>From PD onset and onwards (^b)</th>
</tr>
</thead>
<tbody>
<tr>
<td>HR 95% CI</td>
<td>HR 95% CI</td>
<td>HR 95% CI</td>
<td>HR 95% CI</td>
<td>HR 95% CI</td>
</tr>
<tr>
<td>All psychotic disorders</td>
<td>1.63 1.27-2.09</td>
<td>1.73 1.28-2.34</td>
<td>1.16 0.97-1.39</td>
<td>1.10 0.93-1.30</td>
</tr>
<tr>
<td>Bipolar disorder</td>
<td>2.09 1.59-2.74</td>
<td>2.14 1.47-3.11</td>
<td>1.76 1.35-2.30</td>
<td>1.64 1.29-2.08</td>
</tr>
<tr>
<td>Bipolar type I</td>
<td>6.05 3.44-10.63</td>
<td>2.74 1.31-5.73</td>
<td>1.87 1.32-2.64</td>
<td>1.79 1.29-2.49</td>
</tr>
<tr>
<td>Schizophrenia</td>
<td>0.54 0.22-1.34</td>
<td>0.79 0.34-1.81</td>
<td>0.62 0.41-0.95</td>
<td>0.55 0.37-0.84</td>
</tr>
<tr>
<td>All depression</td>
<td>2.01 1.79-2.27</td>
<td>2.63 2.27-3.04</td>
<td>1.86 1.62-2.12</td>
<td>1.51 1.35-1.69</td>
</tr>
<tr>
<td>Severe depression</td>
<td>1.32 1.03-1.70</td>
<td>2.30 1.76-3.00</td>
<td>1.68 1.31-2.14</td>
<td>1.33 1.08-1.64</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Risk of PD in patients with MS</th>
<th>Up to 2 years prior to date of MS onset (^a)</th>
<th>Date of onset of MS ±2 years (^a)</th>
<th>2 years after MS onset and onwards (^a)</th>
<th>From MS onset and onwards (^b)</th>
</tr>
</thead>
<tbody>
<tr>
<td>HR 95% CI</td>
<td>HR 95% CI</td>
<td>HR 95% CI</td>
<td>HR 95% CI</td>
<td>HR 95% CI</td>
</tr>
<tr>
<td>All psychotic disorders</td>
<td>1.38 1.20-1.58</td>
<td>1.74 1.28-2.36</td>
<td>1.61 1.25-2.08</td>
<td>1.74 1.41-2.15</td>
</tr>
<tr>
<td>Bipolar disorder</td>
<td>1.82 1.52-2.18</td>
<td>1.98 1.35-2.89</td>
<td>1.93 1.45-2.57</td>
<td>2.08 1.63-2.65</td>
</tr>
<tr>
<td>Bipolar type I</td>
<td>2.25 1.69-3.01</td>
<td>2.76 1.25-6.12</td>
<td>4.93 2.69-9.05</td>
<td>4.76 2.85-7.97</td>
</tr>
<tr>
<td>Schizophrenia</td>
<td>0.73 0.51-1.02</td>
<td>0.91 0.39-2.11</td>
<td>0.74 0.32-1.70</td>
<td>0.89 0.48-1.65</td>
</tr>
<tr>
<td>All depression</td>
<td>1.53 1.33-1.76</td>
<td>2.78 2.39-3.22</td>
<td>1.80 1.58-2.04</td>
<td>2.13 1.93-2.36</td>
</tr>
<tr>
<td>Severe depression</td>
<td>1.44 1.11-1.88</td>
<td>2.72 2.07-3.59</td>
<td>1.08 0.81-1.44</td>
<td>1.55 1.25-1.92</td>
</tr>
</tbody>
</table>

Hazard Ratios (HR) and 95% confidence limits (95% CI). Significant HRs are indicated by bold text. The hazard ratios are not adjusted for immigration status.

\( ^a \) Analysis as described in Figure 3

\( ^b \) Analysis as described in Figure 4
Figure legends

**Figure 1** Flowchart of the study population in the comorbidity and the sibling cohorts (psychiatric disorders as exposure and multiple sclerosis as outcome).

**Figure 2** Flowchart of the study population in the comorbidity and the sibling cohorts (multiple sclerosis as exposure and psychiatric disorders as outcome).

**Figure 3** Analysis by time period of multiple sclerosis (MS) risk in the comorbidity cohort with psychiatric disorder (PD) as exposure:

MS risk was analyzed: A) from entry date until two years before date of PD onset; B) from two years before until 2 years after date of PD onset; C) from two years after date of PD onset until end of follow-up. The analyses were performed in a reverse manner, whereby PD risk was analyzed with MS as exposure.

**Figure 4** MS risk was analyzed after date of onset (entry date) of the psychiatric disorder (PD), until an event occurred or the subject was censored. Patients with an MS event before entry date were censored. The analyses were performed in a reverse manner, whereby PD risk was analyzed with MS as exposure.

**Figure 5** Boxplot showing the risk of multiple sclerosis (MS) in patients with a psychiatric disorder and in siblings to patients with a psychiatric disorder. HR = hazard ratio; CI = confidence interval. The hazard ratios are not adjusted for immigration status.
Figure 1 Flowchart of the study population in the comorbidity and the sibling cohorts (psychiatric disorders as exposure and multiple sclerosis as outcome).
190x142mm (300 x 300 DPI)
Figure 2 Flowchart of the study population in the comorbidity and the sibling cohorts (multiple sclerosis as exposure and psychiatric disorders as outcome).

190x142mm (300 x 300 DPI)
Figure 3 Analysis by time period of multiple sclerosis (MS) risk in the comorbidity cohort with psychiatric disorder (PD) as exposure:

MS risk was analyzed: A) from entry date until two years before date of PD onset; B) from two years before until 2 years after date of PD onset; C) from two years after date of PD onset until end of follow-up. The analyses were performed in a reverse manner, whereby PD risk was analyzed with MS as exposure.

69x18mm (300 x 300 DPI)
Figure 4 MS risk was analyzed after date of onset (entry date) of the psychiatric disorder (PD), until an event occurred or the subject was censored. Patients with an MS event before entry date were censored. The analyses were performed in a reverse manner, whereby PD risk was analyzed with MS as exposure.
Figure 5 Boxplot showing the risk of multiple sclerosis (MS) in patients with a psychiatric disorder and in siblings to patients with a psychiatric disorder. HR = hazard ratio; CI = confidence interval. The hazard ratios are not adjusted for immigration status.

208x136mm (300 x 300 DPI)