Risk of bipolar disorder and schizophrenia in relatives of people with attention-deficit hyperactivity disorder

Henrik Larsson, Eleonore Rydén, Marcus Boman, Niklas Långström, Paul Lichtenstein and Mikael Landén

Background
Attention-deficit hyperactivity disorder (ADHD) is associated with bipolar disorder and schizophrenia, and it has been suggested that combined bipolar disorder and ADHD is aetiologically distinct from the pure disorders.

Aims
To clarify whether ADHD shares genetic and environmental factors with bipolar disorder and schizophrenia.

Results
First-degree relatives of the ADHD proband group were at increased risk of both bipolar disorder (odds ratio (OR) = 1.84–2.54 for parents, offspring and full siblings) and schizophrenia (OR = 1.71–2.22 for parents, offspring and full siblings). The risks of bipolar disorder and schizophrenia among second-degree relatives were substantially lower than among full siblings.

Conclusions
These findings suggest that the co-occurrence of ADHD and bipolar disorder as well as ADHD and schizophrenia is due to shared genetic factors, rather than representing completely aetiologically distinct subsyndromes.

Declaration of interest
None.

Attention-deficit hyperactivity disorder (ADHD), bipolar disorder and schizophrenia are highly heritable psychiatric disorders that sometimes co-occur.1–4 The extent to which ADHD shares aetiological factors with bipolar disorder and/or schizophrenia has important implications for both clinical practice and research. Studies suggest shared genetic susceptibility loci as well as genetic deletions and duplications (i.e. copy number variants, CNVs) that overlap between these disorders.5,6 However, no adequately sized family study has explored the degree to which ADHD shares genetic and environmental risk factors with bipolar disorder and schizophrenia.

To clarify whether ADHD shares genetic and environmental factors with bipolar disorder and schizophrenia, and it has been suggested that combined bipolar disorder and ADHD is aetiologically distinct from the pure disorders.

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None.

Method
The study was based on data from Swedish longitudinal national registers held by the National Board of Health and Welfare and Statistics Sweden, which were linked through each individual’s unique personal identification number. The Patient Register has nationwide coverage for psychiatric in-patient care since 1973 and information on psychiatric out-patient care (not provided by a general practitioner) since 2001. Every record has a discharge date, a primary discharge diagnosis and secondary diagnoses assigned by the treating medical doctor according to the World Health Organization’s International Statistical Classification of Diseases and Related Health Problems: ICD-8 for 1969–1986, ICD-9 for 1987–1996 and ICD-10 from 1997.10–12

The Swedish Prescribed Drug Register is a national healthcare register with data on prescribed and dispensed pharmaceuticals. Information regarding drug identity according to the Anatomical Therapeutic Chemical classification system (ATC code), quantity and dosage of the prescribed drug, and date of prescription/dispensing has been registered since July 2005 along with specific patient information (gender, age and residential area). The register covers the entire population of Sweden, and the identity of the patients is available for over 99.7% of the population.13

From the Total Population Register we obtained information on gender, birth year and migrant status for the entire Swedish population. At the time of the construction of our database, this register covered all persons born up to 2009 and all migration events during 1969–2009. Information on vital status (a registered death date) was taken from the Cause of Death Register. At the time of the analyses this register covered essentially all deaths from 1952 to 2009. The Multi-Generation Register includes personal identification numbers of index persons and their biological and adoptive parents and was used to identify parents, offspring, full siblings and half-siblings of index persons. Index persons were individuals born 1932–2009 and registered as living in Sweden at any time during 1961–2009 or who migrated here together with one or both parents and obtained permanent citizenship before age 18 years.

Classifications of ADHD
Patients obtaining a diagnosis of ADHD between 1987 and 2009 were identified in the patient register (ICD-9 code 314; ICD-10 code F90). All discharges and physician appointments (other
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than with general practitioners) with a psychiatric diagnosis were included and no distinction was made between primary and secondary diagnoses. Patients treated with stimulant or non-stimulant medication for ADHD – methylphenidate (N06BA04), atomoxetine (N06BA09), amfetamine (N06BA01) or dexamfetamine (N06BA02) – at any time between July 2005 and December 2009 were identified via the Prescribed Drug Register and assigned to the ADHD proband group. In our study patients aged 3–65 years at the time of their first ADHD diagnosis (or first prescription of stimulant or non-stimulant medication for ADHD) were included as probands. National guidelines for medication of ADHD, issued by the Swedish National Board of Health and Welfare in 2002, state that pharmacological treatment should be reserved for cases in which other supportive interventions have failed. This indicates that individuals with ADHD drug prescriptions represent more severe cases of the disorder. The authority to prescribe ADHD drugs in Sweden is restricted to specialist physicians familiar with the treatment of this disorder.

We used psychiatric symptom data from 20 000 twins (born 1992–2001) from the Swedish Twin Register to explore the validity of the register-based ADHD diagnosis. Symptoms of ADHD in twins were assessed using the Autism – Tics, ADHD and Other Comorbidities Inventory (A-TAC), which covers 96 specific child neuropsychiatric symptoms. One study with extensive psychometric analyses found excellent validity for the A-TAC ADHD measures. The mean ADHD score of twins who also had at least one register-based ADHD diagnosis or medication prescription according to our definition (mean score 9.05, s.d. = 5.32) was substantially higher (Cohen’s d = 1.74) than in the total sample (mean 1.73, s.d. = 2.68). In addition, about 70% of twins with a register-based ADHD diagnosis were also screen-positive for parent-rated ADHD. Similar results were obtained when these validity checks were restricted either to ADHD cases identified through ICD diagnoses (obtained from the Patient Register) or to pharmacological ADHD treatment (obtained from the Prescribed Drug Register).

Classification of bipolar disorder and schizophrenia

Bipolar disorder was defined as at least one discharge diagnosis of the disorder (ICD-8 codes 296.1, 296.3, 296.8; ICD-9 codes 296A/B/C/D/E/W; ICD-10 codes F30–F31). Individuals with bipolar disorder who had ever been diagnosed with schizoaffective disorder (ICD-8 code 295.7, ICD-9 code 295H or ICD-10 code F25) or schizophrenia (see below) during 1973–2009 were excluded. Similarly, we defined schizophrenia as at least one schizophrenia diagnosis (ICD-8 codes 295.0–295.4, 295.6, 295.8–295.9; ICD-9 codes 295A–295E, 295G, 295W, 295X; ICD-10 code F20). Individuals with schizophrenia who also had a diagnosis of bipolar disorder or schizoaffective disorder (for criteria, see above) during 1973–2009 were excluded. All psychiatric discharges were included in this study and no distinction was made between primary and secondary diagnoses.

Statistical analysis

The statistical analyses were performed using nested case–cohort designs. First, we explored the associations between ADHD and bipolar disorder or schizophrenia, by comparing ADHD probands with controls. Second, we addressed the familial overlap between ADHD and bipolar disorder or schizophrenia by comparing relatives of the ADHD proband group with relatives of the control group (parents, offspring, full siblings, and maternal and paternal half-siblings).

Results

In total we identified 61 187 individuals with ADHD according to study criteria, of whom 41 603 (68%) were male.

Bipolar disorder and schizophrenia in proband group

Among members of the ADHD proband group without schizophrenia (n = 60 655), 4.9% had a co-occurring bipolar disorder diagnosis. In contrast, 0.2% of the control group had a bipolar disorder diagnosis. Hence, the proband group were 24 times more likely to be diagnosed with bipolar disorder compared with the control group (Table 1). Of those in the ADHD proband group without bipolar disorder (n = 58 133), 0.8% were also diagnosed with schizophrenia compared with 0.1% of the control group, corresponding to a substantially increased risk of schizophrenia (OR = 6.7; Table 1).

Relatives of probands v. relatives of controls

A family design was used to study the genetic and environmental sources of overlap between the disorders. We compared the risk of bipolar disorder and schizophrenia in relatives of probands with ADHD with risk in relatives of matched controls. To each proband–relative pair, ten randomly selected control–relative pairs were matched by birth year and gender of both proband and relative. This method avoids bias introduced by individuals in the population registries entering the study at different times (left truncation) and allows equal follow-up periods of the relatives to the probands and controls. Further, to study specifically the effect of shared genetic and environmental factors, we adjusted for the potential existence of aetiologically distinct subsyndromes (e.g. an ADHD plus bipolar disorder subtype) by excluding from the proband and control groups anyone with schizophrenia or bipolar disorder, and by excluding relatives of probands or controls with ADHD. Shared familial (genetic and environmental) risk factors are indicated when probands with the index disorder have relatives with the other two disorders but not the index disorder. We analysed first-degree and second-degree relatives separately to assess whether the observed familial association was due to genetic and/or shared environmental influences. This set of analyses were based on the following assumptions: first-degree relatives (who share 50% of their co-segregating genes) are more similar genetically than second-degree relatives (who share 25% of their co-segregating genes), and maternal half-siblings are more similar with regard to shared environmental exposures than paternal half-siblings because children continue to live predominantly with their mother following parental separation.

To describe associations, we used odds ratios (ORs) with 95% confidence intervals obtained from conditional logistic regression models in PROC PHREG in SAS version 9.3 on Unix. When studying associations within families, confidence intervals were obtained with a robust sandwich estimator function to adjust for non-independence (PROC PHREG, covsandwich option).

Bipolar disorder v. controls

We initially compared the risk of bipolar disorder and schizophrenia in the ADHD proband group with that among matched controls. For each case, we randomly selected 10 control group members matched by birth year and gender. In line with well-established procedures for nested case–cohort designs, control group participants were alive and living in Sweden and not diagnosed with ADHD at the time of the first ADHD diagnosis of the proband.
Bipolar disorder and schizophrenia in proband group relatives

First-degree relatives of those in the ADHD proband group were more likely to have been diagnosed with bipolar disorder than first-degree relatives of control group participants (OR = 1.84–2.54; Table 2), supporting familial influences for the overlap between ADHD and bipolar disorder. Second-degree relatives of those in the ADHD group were also more likely to have been diagnosed with bipolar disorder than the relatives of controls. The risk of bipolar disorder was statistically significant and similar in magnitude among maternal (OR = 1.17) and paternal (OR = 1.22) half-siblings, but substantially lower than for full siblings. Further, first-degree relatives of probands with ADHD were more likely to have schizophrenia than relatives of controls (OR = 1.71–2.22; Table 2). The risk of schizophrenia was similar among maternal (OR = 1.11) and paternal (OR = 1.06) half-siblings and substantially lower than for full siblings.

Table 1 Risk of bipolar disorder and schizophrenia in the attention-deficit hyperactivity disorder (proband) group (cases) and the control group matched for birth year and gender

<table>
<thead>
<tr>
<th></th>
<th>Cases n (%)</th>
<th>Controls n (%)</th>
<th>OR (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bipolar disorder</td>
<td>2989 (4.9)</td>
<td>1363 (0.2)</td>
<td>24.0 (22.5–25.7)</td>
</tr>
<tr>
<td>Schizophrenia</td>
<td>467 (0.8)</td>
<td>715 (0.1)</td>
<td>6.7 (5.9–7.5)</td>
</tr>
</tbody>
</table>

a. Proband sample n = 60,655 for analysis of bipolar disorder risk.  
b. Proband sample n = 58,133 for analysis of schizophrenia risk.

Discussion

We conducted a large-scale nationwide family study to elucidate whether ADHD shares genetic and environmental risk factors with bipolar disorder and schizophrenia. The results suggested increased risks of both bipolar disorder and schizophrenia in relatives of the proband group. Moreover, risks among half-siblings were considerably lower than in full siblings, but similar in maternal and paternal half-siblings. This pattern indicates that these disorders share genetic rather than environmental risk factors, consistent with prior twin study results for ADHD, bipolar disorder and schizophrenia, suggesting substantial heritability and a limited role of shared environmental effects.17–19

Prior family studies have indicated that relatives of people with ADHD and co-occurring bipolar disorder have an increased risk of bipolar disorder, whereas relatives of people with ADHD only do not.7,8 One interpretation of those results is that the two disorders are transmitted together in families, representing a familial distinct syndrome. At odds with this notion, however, is our finding that pure ADHD in probands actually predicted pure bipolar disorder in relatives. A potential explanation of these conflicting results is that the previous studies were insufficiently powered to detect a familial association between pure ADHD and bipolar disorder. The comorbidity of ADHD and schizophrenia is biased, as the nested case–cohort method allows equal follow-up

<table>
<thead>
<tr>
<th></th>
<th>n (%)</th>
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<th>OR (95% CI)</th>
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<tbody>
<tr>
<td>Bipolar disorder</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>First-degree relatives</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>Parents</td>
<td>986 (0.95)</td>
<td>4835 (0.51)</td>
<td>1.84 (1.72–1.97)</td>
</tr>
<tr>
<td>Offspring</td>
<td>52 (0.28)</td>
<td>187 (0.12)</td>
<td>2.54 (1.92–3.35)</td>
</tr>
<tr>
<td>Full siblings</td>
<td>319 (0.52)</td>
<td>1212 (0.23)</td>
<td>2.22 (1.98–2.50)</td>
</tr>
<tr>
<td>Second-degree relatives</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Maternal half-siblings</td>
<td>88 (0.44)</td>
<td>135 (0.32)</td>
<td>1.26 (1.01–1.58)</td>
</tr>
<tr>
<td>Paternal half-siblings</td>
<td>84 (0.38)</td>
<td>132 (0.29)</td>
<td>1.34 (1.06–1.71)</td>
</tr>
</tbody>
</table>

Strengths and limitations

The main strength of this study was our ability to address the risk of bipolar disorder and schizophrenia primarily in first-degree and second-degree relatives of a proband group with pure ADHD and a control group. Second, the study was powered to yield robust, unambiguous results. One potential limitation is that the outpatient register is relatively new (started in 2001). Also, as already mentioned, individuals enter and leave the registers at different ages. Both these features, inherent in using registers for research, mean that the percentage of individuals with a disorder does not equal the true lifetime prevalence. However, associations between ADHD and bipolar disorder or schizophrenia are not biased, as the nested case–cohort method allows equal follow-up
periods of the proband and control groups as well as of their relatives. Another limitation is the non-standardised register diagnoses. However, validity studies of bipolar disorder and schizophrenia, as well as our own validity checks of ADHD using data from the Swedish twin register, support high specificity for the register-based diagnosis. Specifically, bipolar disorder using data from the Swedish twin register, support high specificity. However, validity studies of bipolar disorder and schizophrenia, as well as our own validity checks of ADHD diagnoses indicated 94% agreement when compared with re-assessed diagnostic status based on patients’ medical records, whereas in-patient schizophrenia diagnoses indicated 94% agreement when compared with research diagnoses based on semi-structured interviews and medical records.

References


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