Prediction of transition from common adolescent bipolar experiences to bipolar disorder: 10-year study

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Background
Although (hypo)manic symptoms are common in adolescence, transition to adult bipolar disorder is infrequent.

Aims
To examine whether the risk of transition to bipolar disorder is conditional on the extent of persistence of subthreshold affective phenotypes.

Method
In a 10-year prospective community cohort study of 3021 adolescents and young adults, the association between persistence of affective symptoms over 3 years and the 10-year clinical outcomes of incident DSM-IV (hypo)manic episodes and incident use of mental healthcare was assessed.

Results
Transition to clinical outcome was associated with persistence of symptoms in a dose-dependent manner. Around 30–40% of clinical outcomes could be traced to prior persistence of affective symptoms.

Conclusions
In a substantial proportion of individuals, onset of clinical bipolar disorder may be seen as the poor outcome of a developmentally common and usually transitory non-clinical bipolar phenotype.

Declaration of interest
None.

Recent research in adolescents and young adults has indicated that subthreshold phenotypes consisting of (hypo)manic symptoms are a common phenomenon in the general population. Although it has been established that these common developmental expressions of (hypo)manic symptoms are associated with an increased risk for adult-onset bipolar disorder, much less is known about which characteristics determine the poor outcome of bipolar disorder in only a small minority of all those with (hypo)manic symptoms. One of the missing pieces of information regarding the prediction–onset cycle in bipolar disorder is that because of the lack of prospective longitudinal data, little is known about the dynamics of the course of subthreshold phenotypes in relation to later onset of the disorder. In the current paper, the hypothesis was tested that differential course of (hypo)manic symptoms in adolescence would be associated with differential risk for transition to full-blown bipolar disorder, greater levels of persistence over time predicting greater likelihood of transition to a diagnosable disorder. Second, as bipolar disorder is often preceded by depressive symptoms, it was hypothesised that the course and level of persistence of depressive symptoms would be equally relevant in predicting transition. Both hypotheses were tested in a large representative cohort of adolescents followed over a period of up to 10 years. Given previous evidence of the effect of number of symptoms (symptom loading) on risk of transition, the effect of persistence of (hypo)manic and depressive symptoms was analysed in relation to symptom loading as well.

Study design
The study consists of a baseline survey (T₀, n = 3021) and three follow-up investigations (T₁, T₂, T₃), covering a time period of approximately 1.6 years (T₀–T₁, s.d. = 0.2), 3.4 years (T₀–T₂, s.d. = 0.3) and 8.3 years (T₀–T₃, range 7.4–10.6 years, s.d. = 0.7) respectively. Since the older cohort of adolescents, aged 18–24 years at baseline, was not interviewed at T₁, the current results are based on the time periods T₀–T₂ and T₀–T₃. Response rates were 84% at T₂ (n = 2548) and 73% at T₃ (n = 2210). For the younger cohort (n = 1228), the time periods T₀–T₁ and T₁–T₂ were aggregated to represent the interval T₀–T₂. For the current report, the risk set was defined as the set of individuals at risk of developing, for the first time, the clinical outcome at T₃. The risk set consisted of all individuals who: had post-baseline DIA-X/M–CIDI interviews with complete data at both T₂ and T₃ (n = 2029); and had never been diagnosed before T₀ and/or T₁ (n = 381), yielding a risk set of 1648 (i.e. 2029–381). After exclusion of both DSM (hypo)manic episodes prior to T₁ and mental healthcare use prior to T₃, a risk set of 1565 (i.e. 2029–464) participants remained.

Sample
This study is part of the Early Developmental Stages of Psychopathology (EDSP) study, a prospective longitudinal cohort community study. Detailed information about the design, sample, instruments, procedures and statistical methods of the EDSP is presented elsewhere. Data were collected in a random representative population sample of adolescents and young adults living in the Munich area (Germany), aged 14–24 years at baseline. The study sample was randomly drawn from the 1994 government population registers and comprised residents in Munich and its surrounding area.

Method

See pp. 87–88, this issue.
Assessment of affective-symptom groups

Affective symptoms were assessed at \( T_0 \) and \( T_3 \) using the 28 symptom items (DSM–IV and ICD–10) of the DIA–X/M–CIDI depression and dysthymia section (items regarding feeling depressed, loss of interest, loss of energy, hopelessness, decreased concentration, loss of appetite, weight loss, sleep disturbances, feelings of worthlessness or guilt, decreased self-esteem and suicidal ideation) and the 11 symptom items of the DIA–X/M–CIDI mania section (items regarding increase in goal-directed activity, psychomotor agitation, spending sprees, sexual indiscretions, increased talkativeness, flight of ideas, increased self-esteem or grandiosity, decreased need for sleep and distractibility). Symptom items were rated either yes or no. Depressive symptoms were only rated if present for at least 2 weeks; (hypo)manic symptoms if present for at least 4 successive days. If the symptom was the direct result of alcohol or drug use or somatic conditions, the item was not counted towards the diagnosis of a primary mood disorder. Furthermore, symptoms were only assessed and rated if at least one of the DIA–X/M–CIDI core depressive or core (hypo)manic symptoms was present. Only participants having core (hypo)manic symptoms that were either noticed by others or because of which participants experienced problems were included.

Two sum scores of symptom ratings were formed: a sum score of depressive symptoms with a minimum of 0 and a maximum score of 28 endorsements; and a sum score of (hypo)manic symptoms with a minimum of 0 and a maximum score of 11 endorsements. Subsequently, in both symptom groups, progressively stricter and overlapping subcategories of these sum scores, indicating the degree of symptom loading, were created (level 0, level 1, level 2, level 3; occurrence of symptoms twice, both at at least two, four and six symptoms respectively). Odds ratios for all phenotypes were adjusted for age.

Assessment of persistence

For each of the symptom groups, a persistence variable was created. ‘Persistence’ was defined as the number of times at the \( T_0 \) and \( T_3 \) interviews that participants scored positive on having depressive and/or manic symptoms, irrespective of which particular depressive and/or manic symptoms were present. The persistence variable thus had three levels: level 0, no symptoms at \( T_0 \) and \( T_3 \); level 1, occurrence of symptoms only once at \( T_0 \) or \( T_3 \); and level 2, occurrence of symptoms twice, both at \( T_0 \) and \( T_3 \).

Assessment of clinical outcome

In order to predict transition to clinical disorder, two clinical outcomes were used in the analyses. The first was defined as suffering from either a DSM–IV manic or a DSM–IV hypomanic episode (hereafter: DSM (hypo)manic episode) and the second as need for mental healthcare because of affective symptoms (mental healthcare use). Participants suffering from DSM (hypo)manic episodes were defined using the DIA–X/M–CIDI/DSM–IV diagnostic algorithm, as follows: participants suffering from either hypomanic nor manic episodes; or participants suffering from either hypomanic or manic episodes.

In order to assess need for mental healthcare use, data from two DIA–X/M–CIDI sections were used. First, participants were asked whether they were ever treated in a hospital or spoke to a professional because of (hypo)manic symptoms. Second, participants were shown a list of several types of out-patient, in-patient or day-patient institutions for mental health problems, ranging from a general practitioner or a school psychologist to psychiatric sheltered housing, after which they were asked if they had ever sought help at any of these institutions because of any mental health problems. All participants who responded positively to one or both of these questions were considered to have the mental healthcare use outcome.

Statistical analysis

The association (expressed as odds ratio) between persistence as the independent variable and clinical outcome (DSM (hypo)manic episodes and mental healthcare use) as the dependent variable was analysed for each symptom group ((hypo)manic, depressive and bipolar symptoms respectively) and each symptom loading (two, four and six symptoms respectively) using logistic regression in STATA, version 9.2 on Windows XP. First, in order to test for a monotonic trend in the association between level of persistence and transition to the clinical outcome, an ordinal variable was created that represented the number of symptoms present in each symptom group (values ranging from 0 to 2 for level 0, level 1 and level 2 of persistence respectively). Second, in order to test for a monotonic trend in the association between number of symptoms and transition to the clinical outcome, an ordinal variable was created that represented the number of symptoms present in each symptom group (values ranging from 0 to 3 for zero, two, four and six symptoms respectively). Odds ratios for all phenotypes were adjusted for age.

Results

Analyses regarding the development of DSM (hypo)manic episodes were conducted in a sample of 1902 adolescents. Gender distribution was approximately equal (52.3% males). Mean age at baseline was 18.3 years (s.d. = 3.3; range 14–24). In this risk set, 1.1% \((n = 21)\) developed an incident DSM (hypo)manic episode at \( T_3 \). Analyses regarding mental healthcare use were conducted in a sample of 1648 adolescents. Gender distribution was approximately equal (53.9% males). Mean age at baseline was 18.2 years (s.d. = 3.3; range 14–24). In this risk set, 10.4% \((n = 172)\) had incident mental healthcare use at \( T_3 \). Drop-out rates at \( T_3 \) (after excluding all participants without complete data at both \( T_2 \) and \( T_3 \), yielding a data-set of 2029
participants) were almost equal for the different levels of persistence for: (hypo)manic symptoms (18.8% persistence level 0 v. 21.1% level 1 v. 23.1% level 2); depressive symptoms (19.1% level 0 v. 19.2% level 1 v. 22.2% level 2); and bipolar symptoms (14.9% level 0 v. 22.6% level 1 v. 24.3% level 2).

**Presence of (hypo)manic symptoms and transition to clinical outcome**

More than a quarter (25.1%, n = 392) of 1565 participants displayed (hypo)manic symptoms once at T0 or T2, whereas 2.6% (n = 41) experienced symptoms twice (Table 1). The number of affected participants decreased with increasing level of symptom loading (Table 2).

Participants who never experienced two or more (hypo)manic symptoms (n = 1160) had an 0.7% risk of developing DSM (hypo)manic episodes and a 9.4% risk for mental healthcare use in the final follow-up. With greater levels of persistence, the risk of developing DSM (hypo)manic episodes increased from 0.7% to 2.0–3.2%, and the risk of mental healthcare use from 9.4% to 11.5–12.8% (Table 2).

Within the different categories of symptom loading (Table 2), an association was found with persistence level for both transition to DSM (hypo)manic episodes (summary increase in risk per unit increase in persistence level (hereafter: summary increase in risk) ranging from 2.10 to 3.13 depending on category of symptom loading) and transition to mental healthcare use (summary increase in risk ranging from 1.22 to 1.36), which was significant for all comparisons related to DSM (hypo)manic episodes (Table 2, columns 5 and 9). Likewise, within each level of persistence, a dose–response relationship was seen between the level of symptom loading and the risk of transition (Table 1).

Depicting level of persistence and level of symptom loading together revealed that level of persistence became increasingly relevant as the number of symptoms persisting increased (Figs 1 and 2).

**Presence of depressive symptoms and transition to clinical outcome**

Nearly half (45.1%, n = 706) of 1565 participants displayed depressive symptoms once at T0 or T2, whereas 14.5% (n = 227) experienced symptoms twice (Table 1). The number of affected participants decreased with increasing level of symptom loading (Table 2).

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**Table 1** Odds ratios (ORs) monotonic trend for impairment associated with symptom loading* by level of persistence and symptom group

| Persistence levelb | Total, % (n)c | (Hypo)manic episodesd,e | Mental healthcare usef | | | |
|--------------------|--------------|-------------------------|-----------------------| | | |
|                    |              | % (n) | ORa | 95% CI | P | % (n) | ORa | 95% CI | P |
| (Hypo)manic        |              |       |     |       |   |       |     |       |   |
| 1                  | 25.1 (392)   | 2.0 (10) | 1.62 | 1.13–2.33 | 0.009 | 12.3 (56) | 1.06 | 0.91–1.25 | 0.448 |
| 2                  | 2.6 (41)     | 3.2 (2) | 2.37 | 1.11–5.05 | 0.026 | 14.1 (9) | 1.42 | 0.92–2.06 | 0.063 |
| Depressive         |              |       |     |       |   |       |     |       |   |
| 1                  | 45.1 (706)   | 0.9 (8) | 1.00 | 0.72–1.40 | 0.996 | 11.9 (89) | 1.20 | 1.06–1.35 | 0.003 |
| 2                  | 14.5 (227)   | 2.4 (8) | 1.58 | 1.12–2.24 | 0.010 | 20.2 (53) | 1.50 | 1.29–1.74 | <0.001 |
| Bipolar            |              |       |     |       |   |       |     |       |   |
| 1                  | 39.7 (621)   | 1.2 (9) | 1.13 | 0.81–1.57 | 0.460 | 11.4 (76) | 1.14 | 1.01–1.29 | 0.039 |
| 2                  | 16.6 (259)   | 2.4 (9) | 1.71 | 1.22–2.39 | 0.002 | 17.0 (52) | 1.42 | 1.22–1.64 | <0.001 |

a. The ORs express summary increase in risk with 1 unit change in symptom loading (0: no symptoms; 1: at least 2 symptoms; 2: at least 4 symptoms; 3: at least 6 symptoms).
b. Level 1: symptoms at one time (T0 or T2), level 2: symptoms twice (T0 and T2).c. Risk set: all participants with data at T0, T1, T2, and T3 and no kind of impairment at both T2 and T3 (n = 1565).d. (Hypo)manic episodes: either hypomanic or manic episodes.e. Risk set: all participants with data at T0, T2, T3, and no mental healthcare use at both T2 and T3 (n = 1648).f. Risk set: all participants with data at T0, T1, T2, T3, and no kind of impairment at both T0 and T1 (n = 1902).

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**Table 2** Odds ratios (ORs) monotonic trend for impairment associated with persistence* by symptom loading and symptom group

| Symptom loading | Total % (n)c | (Hypo)manic episodesd,e | Mental healthcare usef | | | |
|-----------------|--------------|-------------------------|-----------------------| | | |
|                   |              | % (n) | ORa | 95% CI | P | % (n) | ORa | 95% CI | P |
| (Hypo)manic      |              |       |     |       |   |       |     |       |   |
| 2                | 25.8 (404)   | 1.9 (10) | 2.10 | 1.10–4.01 | 0.024 | 12.8 (61) | 1.30 | 1.00–1.70 | 0.053 |
| 4                | 16.5 (258)   | 2.3 (8) | 2.92 | 1.44–5.93 | 0.003 | 11.5 (38) | 1.22 | 0.88–1.69 | 0.224 |
| 6                | 5.2 (81)     | 3.3 (4) | 3.13 | 1.12–8.76 | 0.000 | 11.9 (15) | 1.36 | 0.83–2.23 | 0.222 |
| Depressive       |              |       |     |       |   |       |     |       |   |
| 2                | 51.5 (806)   | 1.2 (13) | 1.58 | 0.90–2.76 | 0.108 | 13.1 (114) | 1.77 | 1.44–2.18 | <0.001 |
| 4                | 40.3 (630)   | 1.5 (13) | 1.77 | 1.02–3.08 | 0.041 | 14.6 (101) | 1.79 | 1.45–2.21 | <0.001 |
| 6                | 27.7 (434)   | 1.9 (12) | 2.29 | 1.31–4.01 | 0.004 | 16.3 (78) | 1.90 | 1.51–2.39 | <0.001 |
| Bipolar          |              |       |     |       |   |       |     |       |   |
| 2                | 46.6 (729)   | 1.6 (15) | 2.03 | 1.19–3.46 | 0.009 | 12.8 (103) | 1.59 | 1.30–1.93 | <0.001 |
| 4                | 38.4 (601)   | 1.7 (14) | 2.24 | 1.31–3.81 | 0.003 | 13.9 (94) | 1.67 | 1.36–2.05 | <0.001 |
| 6                | 27.8 (435)   | 2.3 (14) | 2.85 | 1.65–4.91 | <0.001 | 14.8 (74) | 1.76 | 1.40–2.22 | <0.001 |

a. The ORs express summary increase in risk with 1 unit change in level of persistence (variable has 3 levels: level 0, no symptoms at T0 and T2, level 1, occurrence of symptoms only once at T0 or T2, level 2, occurrence of symptoms twice both at T0 and T2).b. Risk set: all participants with data at T0, T2, T3, and no kind of impairment at both T0 and T2 (n = 1565).c. (Hypo)manic episodes: either hypomanic or manic episodes.d. Risk set: all participants with data at T0, T2, T3, and no mental healthcare use at both T0 and T2 (n = 1648).e. Risk set: all participants with data at T0, T1, T2, T3, and no kind of impairment at both T0 and T1 (n = 1902).
Participants who never experienced two or more depressive symptoms (n=759) had an 0.9% risk of developing DSM (hypo)manic episodes and a 7.6% risk for mental healthcare use.

The risk of developing DSM (hypo)manic episodes was similar for persistence level 0 and 1 (0.9%), but increased to 2.4% for persistence level 2, whereas the risk of mental healthcare use increased with increasing persistence level from 7.6% to 11.9–20.2% (Table 1). Similarly, with increasing level of symptom loading, the risk of developing DSM (hypo)manic episodes increased to 1.2–1.9% and the risk of mental healthcare use to 13.1–16.3% (Table 2). Within the different categories of symptom loading (Table 2), an association was found with persistence level for both transition to DSM (hypo)manic episodes (summary increase in risk ranging from 1.58 to 2.29 depending on symptom category) and transition to mental healthcare use (summary increase in risk ranging from 1.77 to 1.90), which was significant for all but one comparison (Table 2).

Within the different categories of symptom loading, the risk of developing DSM (hypo)manic episodes increased with increasing persistence level from 7.6% to 11.9–20.2% (Table 1). Similarly, with increasing level of symptom loading, the risk of mental healthcare use to 13.1–16.3% (Table 2). Within the different categories of symptom loading (Table 2), an association was found with persistence level for both transition to DSM (hypo)manic episodes (summary increase in risk ranging from 1.58 to 2.29 depending on symptom category) and transition to mental healthcare use (summary increase in risk ranging from 1.77 to 1.90), which was significant for all but one comparison (Table 2). Similarly, within each level of persistence, a dose–response relationship was seen between the level of symptom loading and the risk of transition (Table 1). Depicting level of persistence and level of symptom loading together revealed that level of persistence became increasingly relevant as the number of symptoms persisting increased, but only for the outcome of DSM (hypo)manic episodes (Fig. 3) and not for the mental healthcare use outcome (Fig. 4).
The risk of mental healthcare use increased from 8.3 to 11.4–17.0% (Table 1). Similarly, with increasing level of symptom loading, the risk of developing DSM (hypo)manic episodes increased to 1.6–2.3% and the risk of mental healthcare use to 12.8–14.8% (Table 2). Again, within the different categories of symptom loading, an association was found with persistence level for both transition to DSM (hypo)manic episodes (summary increase in risk ranging from 2.03 to 2.85 depending on symptom category) and transition to mental healthcare use (summary increase in risk ranging from 1.59 to 1.76), which was significant for all comparisons (Table 2). Likewise, within each level of persistence, a dose–response relationship was seen between the level of symptom loading and the risk of transition (Table 1).

Depicting level of persistence and symptom loading together revealed that level of persistence became increasingly relevant as the number of symptoms persisting increased (Figs 5 and 6).

Proportion of clinical outcome with prior affective symptoms
Of the participants who developed DSM (hypo)manic episodes at T3 (n = 21), 47.6% (n = 10) had experienced two or more (hypo)manic symptoms prior to T3; of these 10, 2 (9.5% of total of 21) had experienced two or more (hypo)manic symptoms more than once. This compares to 61.9% (n = 13) and 38.1% (n = 8) of those who had experienced two or more depressive symptoms at least once or twice respectively, and 71.4% (n = 15) and 42.9% (n = 9) of those who had experienced two or more bipolar symptoms at least once or twice respectively.

Of the participants who developed a need for mental healthcare use (n = 172), 35.5% (n = 61) had experienced two or more (hypo)manic symptoms; of these 61, 9 (5.2% of total of 172) had experienced two or more (hypo)manic symptoms more than once. This compares to 66.3% (n = 114) and 30.8% (n = 53) of those who had experienced two or more depressive symptoms at least once or twice respectively, and 59.9% (n = 103) and 30.2% (n = 52) of those who had experienced two or more bipolar symptoms at least once or twice respectively.

The results suggest persistence of symptoms, relative to having symptoms per se, is predictive for transition to clinically relevant outcomes. The dose–response association between persistence and clinical outcomes became stronger as the number of symptoms persisting increased. The current results confirm the hypothesis that (hypo)manic symptoms are frequent in adolescence, most disappearing over time. However, the results also demonstrate that in some adolescents, (hypo)manic symptoms become persistent, representing a risk state that may progress to full-blown, clinically relevant bipolar disorder.

Explaining the role of persistence
The role of persistence of symptoms, in terms of clinical relevance, may be viewed in the light of the kindling-sensitisation model put forward by Post. According to this model, neurotransmitter pathways are activated by events and produce not only intermediate short-term effects, but also a series of events (i.e. intracellular changes at the level of gene transcription) that have long-lasting consequences for the organism. It is postulated that the type, magnitude and frequency of repetition of the event may be critical to these long-term effects. Thus, every time a person experiences an affective episode, the associated neurotransmitter and peptide alterations may leave behind memory traces that predispose to further episodes, a process referred to as ‘sensitisation’.

### Figure 5
Risk of incident (hypo)manic episodes following persistence of bipolar symptoms (odds ratios in figure quantified in table below figure).

<table>
<thead>
<tr>
<th>Persistence level</th>
<th>Symptom loading, OR (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>1.00</td>
</tr>
<tr>
<td>1</td>
<td>1.79 (0.55–5.86)</td>
</tr>
<tr>
<td>2</td>
<td>4.04 (1.39–11.72)</td>
</tr>
</tbody>
</table>

### Figure 6
Risk of incident mental healthcare (MHC) use following persistence of bipolar symptoms (odds ratios in figure quantified in table below figure).

<table>
<thead>
<tr>
<th>Persistence level</th>
<th>Symptom loading, OR (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>1.00</td>
</tr>
<tr>
<td>1</td>
<td>1.50 (1.00–2.24)</td>
</tr>
<tr>
<td>2</td>
<td>2.52 (1.70–3.74)</td>
</tr>
</tbody>
</table>

### Discussion
The role of persistence of symptoms, in terms of clinical relevance, may be viewed in the light of the kindling-sensitisation model put forward by Post. According to this model, neurotransmitter pathways are activated by events and produce not only intermediate short-term effects, but also a series of events (i.e. intracellular changes at the level of gene transcription) that have long-lasting consequences for the organism. It is postulated that the type, magnitude and frequency of repetition of the event may be critical to these long-term effects. Thus, every time a person experiences an affective episode, the associated neurotransmitter and peptide alterations may leave behind memory traces that predispose to further episodes, a process referred to as ‘sensitisation’.

### Presence of bipolar symptoms and transition to clinical outcome
Almost 40% (39.7%, n = 621) of 1565 participants displayed bipolar symptoms once at T1 or T2, whereas 16.6% (n = 259) experienced symptoms twice (Table 1). The number of affected participants decreased with increasing level of symptom loading (Table 2).

Participants who never experienced two or more bipolar symptoms (n = 836) had an 0.6% risk of developing DSM (hypo)manic episodes and an 8.3% risk for mental healthcare use.

With increasing levels of persistence, the risk of developing DSM (hypo)manic episodes increased from 0.6 to 1.2–2.4%, and
An interactive developmental model

In the literature, several explanations are given as to why (hypo)manic symptoms might develop during adolescence, with a focus on neurodevelopmental and environmental changes interacting with genetic risk.17–19 According to an interactive developmental model, the course of developmental subclinical expression of psychopathology is affected by interactions between the individual and the environment; exposure to additional environmental risk factors may thus explain why a minority of individuals deviate from a trajectory of good outcome of transient subclinical expressions to progression to the full-blown disorder.20,21

Clinical implications

Given the fact that risk factors for bipolar disorder may act by causing persistence of symptoms and subsequent transition from subthreshold expression to a clinical disorder, a window for intervention may exist. Intervention early in life may be particularly relevant, as adolescence represents a period in which the most critical stages of educational, occupational and social development are completed, disruption of which by psychiatric illness may lead to lifelong disability.22

Implications for classification

The results should be viewed from a public health perspective of risk, associated with distributed psychometric subthreshold states in the general population, which is different from the need of making a categorical diagnosis of a rare disease in clinical practice. In order to bridge the apparent divide, it has been proposed that the next revisions of DSM and ICD be open to spectrum interpretations of bipolar disorders, and that the same nosological material may be interpreted dimensionally (risk) or categorically (treatment) depending on the purposes of one's interpretation.23

Indeed, some investigators have suggested broadening current diagnostic concepts to include subthreshold states. Thus, Angst and colleagues24 suggest the inclusion of a broader concept of soft bipolarity, and Akiskal and colleagues25 reported empirical support for the inclusion of bipolar II 1/2 (cyclothymic temperament), bipolar III (antidepressant-induced hypomania) and bipolar IV (hyperthymic temperament) as distinct categories. Furthermore, several studies point to the existence of paediatric bipolar disorder, in which an early onset or a longer duration of bipolar IV (hyperthymic temperament) as distinct categories. Thus, Angst and colleagues24 suggest the inclusion of a broader concept of soft bipolarity, and Akiskal and colleagues25 reported empirical support for the inclusion of bipolar II 1/2 (cyclothymic temperament), bipolar III (antidepressant-induced hypomania) and bipolar IV (hyperthymic temperament) as distinct categories. Furthermore, several studies point to the existence of paediatric bipolar disorder, in which an early onset or a longer duration of symptoms predicts a worse outcome.26

Limitations

Several limitations need to be considered when interpreting the results. First, although a prospective design was used, the study necessarily became partly retrospective by implementing questions providing scientific advice on the design of the EDSP Study. The EDSP project and its family genetic supplement has been approved by the Ethics Committee of the Faculty of the Technische Universiteit Dresden (No. EK-1381).

References

1 Tijssen MJ, van Os J, Wittchen HU, Lieb R, Beesdo K, Mengelers R, et al. Transition from adolescent bipolar experiences to bipolar disorder similar across the different levels of persistence. Furthermore, previous analyses showed that mood disorders were not affected by selective attrition (details available from the author on request). Fourth, exclusion of individuals with bipolar impairment at T0 and T2, necessary to ensure that associations between persistence and impairment were truly predictive, resulted in a small number of individuals with a T3 clinical outcome and a decrease in statistical power. Therefore, it is possible that as a result of loss power the statistical significance of some associations was affected. The fact that effect sizes, albeit some non-significant, are in the expected direction and show dose–response relationships as expected, supports the validity of the results.

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