Comparison of depressive episodes in bipolar disorder and in major depressive disorder within bipolar disorder pedigrees

Philip B. Mitchell, Andrew Frankland, Dusan Hadzi-Pavlovic, Gloria Roberts, Justine Corry, Adam Wright, Colleen K. Loo and Michael Breakspear

Background
Although genetic epidemiological studies have confirmed increased rates of major depressive disorder among the relatives of people with bipolar affective disorder, no report has compared the clinical characteristics of depression between these two groups.

Aims
To compare clinical features of depressive episodes across participants with major depressive disorder and bipolar disorder from within bipolar disorder pedigrees, and assess the utility of a recently proposed probabilistic approach to distinguishing bipolar from unipolar depression. A secondary aim was to identify subgroups within the relatives with major depression potentially indicative of ‘genetic’ and ‘sporadic’ subgroups.

Method
Patients with bipolar disorder types 1 and 2 (n = 246) and patients with major depressive disorder from bipolar pedigrees (n = 120) were assessed using the Diagnostic Interview for Genetic Studies. Logistic regression was used to identify distinguishing clinical features and assess the utility of the probabilistic approach. Hierarchical cluster analysis was used to identify subgroups within the major depressive disorder sample.

Results
Bipolar depression was characterised by significantly higher rates of psychomotor retardation, difficulty thinking, early morning awakening, morning worsening and psychotic features. Depending on the threshold employed, the probabilistic approach yielded a positive predictive value ranging from 74% to 82%. Two clusters within the major depressive disorder sample were found, one of which demonstrated features characteristic of bipolar depression, suggesting a possible ‘genetic’ subgroup.

Conclusions
A number of previously identified clinical differences between unipolar and bipolar depression were confirmed among participants from within bipolar disorder pedigrees. Preliminary validation of the probabilistic approach in differentiating between unipolar and bipolar depression is consistent with dimensional distinctions between the two disorders and offers clinical utility in identifying patients who may warrant further assessment for bipolarity. The major depressive disorder clusters potentially reflect genetic and sporadic subgroups which, if replicated independently, might enable an improved phenotypic definition of underlying bipolarity in genetic analyses.

Declaration of interest
C.K.L has received lecture honoraria from Eli Lilly and Wyeth, and grant support from Neuronetics.

There is growing interest in the recognition and treatment of bipolar depression.1–7 Although genetic epidemiological studies have confirmed increased rates of major depressive disorder among the relatives of people with bipolar affective disorder,8–5 no report has compared the clinical characteristics of depression between these individuals and their relatives with bipolar disorder within the same pedigrees. Confirming differences in these subgroups could be highly informative, first in the debate over phenomenological differences between bipolar disorder and major depression,6–8 and second regarding which cases of major depressive disorder in bipolar disorder pedigrees have more ‘genetic’ or ‘sporadic’ aetiologies.5,8 Given the overlapping presentations of the two forms of depression, the predominance of depressive features among people with bipolar disorder10 and the implications of misdiagnosis,11,12 the usefulness of a method for distinguishing between the two conditions based on clinically observable depressive features is clear. Recently published guidelines from the International Society for Bipolar Disorders (ISBD) Diagnostic Task Force have argued for a dimensional rather than categorical distinction between bipolar depression and major depressive disorder,8,13 leading to the development of a ‘probabilistic’ approach to the diagnosis of bipolar depression.8 A range of depressive features and symptoms more likely to be associated with a diagnosis of bipolar disorder type 1 were identified from the literature in the development of those guidelines (see Appendix), with the presence of five or more specific features being proposed as indicative of bipolar depression, although this cut-off has yet to be empirically validated. In a similar vein, focusing on the finding of a greater prevalence of major depressive disorder among relatives of bipolar disorder probands and the implications of this for linkage studies, Blacker et al estimated that 65–74% of these relatives with major depressive disorder had genetically determined illness.3,14 They proposed developing an index of ‘genetic bipolarity’ for these individuals, based on a combination of potentially discriminating phenomenological and longitudinal characteristics (not dissimilar to that proposed in the ISBD guidelines). To assess the nature of depressive episodes in both bipolar disorder and major depressive disorder within bipolar disorder pedigrees, we examined a data-set of participants in a molecular genetics study of bipolar disorder. We also tested the utility of the ISBD probabilistic approach in identifying bipolar depression in this sample,8 and sought to identify subgroups within the major depressive disorder cases which could represent sporadic v. genetically driven forms of illness, i.e. a potential phenotype for underlying bipolarity.
Method

Participants were recruited through the Australian Bipolar Disorder Molecular Genetics Study,15 undertaken at the University of New South Wales in collaboration with the Prince of Wales Medical Research Institute, the Garvan Institute of Medical Research and Macquarie University in Sydney, Australia. A total of 1128 individuals were recruited in medium-to-large unilineal multigenerational bipolar pedigrees. Initial data on family and illness history were obtained from the proband using the Family Interview for Genetic Studies,18 and each participating family member was subsequently assessed using the Diagnostic Interview for Genetic Studies (DIGS) version 2.0.17 Interviews were conducted by experienced medical practitioners, psychologists and psychiatric nurses who received training in the use of each instrument. Best-estimate Research Diagnostic Criteria and DSM-IV diagnoses were made by senior research psychiatrists and psychiatric nurses who received training in the use of each instrument. Best-estimate Research Diagnostic Criteria and DSM-IV diagnoses were made by senior research psychiatrists after independent evaluation of DIGS interviews, family informant data and medical records.18 The study was approved by the human research ethics committee of the University of New South Wales, and complies with the guidelines of the Australian National Health and Medical Research Council. The sample used did not overlap with the bipolar disorder and major depressive disorder samples described in previous studies by our group, which investigated phenomenological differences between the two forms of depression.19,20

Inclusion criteria

Inclusion in the analysis was restricted to participants with a best-estimate DSM-IV diagnosis of bipolar type 1 disorder, bipolar type 2 disorder or major depressive disorder, the last diagnosis requiring a history of at least two major depressive episodes. For the original molecular genetics study, written informed consent was obtained after complete description to the participants.

Statistical analysis

For this study we combined participants with bipolar disorder types 1 and 2 into a single category as there were relatively few with type 2 disorder. Continuous variables were non-normally distributed, and comparisons were made using the Mann–Whitney U-test. Categorical data were compared using Pearson’s χ² test. All symptoms significantly associated with diagnosis at the bivariate level were included in binary logistic regression models, to identify whether weighted combinations of symptoms could differentiate bipolar disorder from major depression. Each symptom was entered as a single predictor, and then in a multivariate model to assess independent associations with diagnosis after accounting for the effects of other symptoms. Results are reported as age- and gender-adjusted odds ratios for a diagnosis of bipolar disorder. All analyses were carried out using Stata version 10 on Windows XP.

To assess the utility of the probabilistic approach in identifying bipolar depression, we included nine clinical features and symptoms from the current data-set that had previously been included in the operational criteria for the probabilistic approach (see Appendix).8 The DIGS featured items that related to hypersomnia, hyperphagia, weight gain, psychomotor retardation, delusions and hallucinations, pathological guilt, mixed features, early onset and multiple episodes. As all participants had a family history of bipolar disorder, we were unable to include this as a potential distinguishing feature. The number of positive ‘bipolar’ features was summed for each participant, ranging from 0 to 9. For each possible score we compared sensitivity and specificity against a diagnostic standard (best-estimate diagnosis), and plotted receiver operating characteristics curves.

To identify subgroups among the major depressive disorder cases that might correspond to either ‘genetic’ or ‘sporadic’ depression, propensity scores were calculated for each participant, which were the estimated probabilities of bipolar v. unipolar group membership from the previously described bivariate logistic regression. The propensity score, ranging from 0 to 1, is an index of the probability of ‘caseness,’ or a diagnosis of bipolar disorder, with a higher score indicating a greater likelihood that the individual has the disorder. We predicted that major depressive disorder cases with an underlying bipolar genetic liability would more closely resemble the ‘true’ bipolar disorder cases than the major depressive disorder cases with a ‘sporadic’ depression, and therefore have higher propensity scores. Consistent with the probabilistic approach, these cases would be more likely to show clinical and phenomenological features typically associated with bipolar depression, based on the stronger underlying genetic component to their illness. After calculating propensity scores, we carried out a hierarchical cluster analysis, using between-groups linkage, to identify any groupings within the major depressive disorder cases, and compared clinical feature and symptom profiles between clusters.

Results

Of the 1128 participants, 423 (37.5%) met initial diagnostic criteria. Of these, 57 had never had a major depressive episode (i.e. they reported only manic episodes) or provided incomplete data on depressive episodes for the DIGS interview. Of the final 366 participants included in the analysis, 202 (55.2%) had been diagnosed with DSM-IV bipolar type 1 disorder, 44 (12.0%) with bipolar type 2 disorder and 120 (32.7%) with major depressive disorder. All participants with major depressive disorder had at least one first-degree relative with type 1 bipolar disorder.

Sociodemographic characteristics

The sociodemographic characteristics of the sample are reported in Table 1; no significant difference was observed between the two diagnostic groups in age or gender ratio.

Longitudinal characteristics

Data on the illness course and clinical characteristics of bipolar disorder and major depressive disorder participants are shown in Table 1. Bipolar disorder was significantly associated with a greater number of lifetime depressive episodes, with a higher proportion of participants with bipolar disorder (48.4%) reporting at least five lifetime depressive episodes compared with the major depressive disorder group (26.4%). No difference was found in either age at onset or duration of the most severe depressive episode.

Treatment and suicidal behaviour

Patterns of help-seeking and treatment did not vary significantly across the groups, with the majority of participants seeking help from a mental health professional or being prescribed medication during their most severe depressive episode (Table 2). Rates of hospital admission during the most severe depressive episode were significantly higher among those with bipolar disorder (46.3%) compared with major depressive disorder (26.7%). Rates of any lifetime suicide attempt were comparable among those with bipolar disorder (32.5%) and major depressive disorders (28.3%). There was no significant difference across the diagnostic...
groups in the number of reported attempts or age at first suicide attempt.

**Symptom profile during most severe depressive episode**

Table 3 shows the prevalence of each depressive symptom for the major depressive disorder and bipolar disorder groups. In the bivariate analysis the bipolar disorder group was characterised by a significantly greater prevalence (compared with major depressive disorder) of psychomotor retardation, difficulty in thinking or concentrating, early morning wakening, morning worsening, delusions, hallucinations and the presence of three or more concurrent mixed features.

Where a significant bivariate association was found, each symptom was entered into a logistic regression model with diagnosis as the outcome variable. Age- and gender-adjusted odds ratios are reported in Table 4. Compared with the major depressive disorder group, bipolar depression was characterised by significantly higher rates of psychomotor retardation (OR = 2.14), difficulty thinking (OR = 2.44), early morning wakening (OR = 1.68), morning worsening (OR = 1.75), delusions (OR = 2.23) and hallucinations (OR = 3.85). Mixed features were not significantly associated with bipolar disorder. Each item was then included in a multivariate model to assess the independent association between specific symptoms and diagnostic category after controlling for the effect of other symptoms (Table 4). After adjusting for age, gender and the other items, only psychomotor retardation (OR = 1.63) remained a significant predictor of a bipolar disorder diagnosis compared with major depressive disorder.

**Probabilistic approach to the diagnosis of bipolar depression**

To assess the usefulness of the probabilistic approach, we focused on nine clinical features and symptoms that have previously been associated with a bipolar diagnosis, for which relevant data were available in the version of the DIGS used for this data-set (see Appendix). The number of positive features endorsed by each

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### Table 1  Sociodemographic and clinical characteristics of participants in the major depressive disorder and bipolar type 1 and 2 disorder samples

<table>
<thead>
<tr>
<th></th>
<th>Unipolar (MDD) (n = 120)</th>
<th>Bipolar type 1/2 (n = 246)</th>
<th>Test statistic</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Gender</strong>, n (%)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Male</td>
<td>37 (30.8)</td>
<td>88 (36.1)</td>
<td>0.98&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>Female</td>
<td>83 (69.2)</td>
<td>156 (63.9)</td>
<td></td>
</tr>
<tr>
<td><strong>Age, years: median (IQR)</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>At interview</td>
<td>50 (30.5–63)</td>
<td>48 (36–57)</td>
<td>0.35&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
<tr>
<td>At first mood episode</td>
<td>20.5 (16–28)</td>
<td>20 (16–29)</td>
<td>−0.26&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
<tr>
<td>At first depressive episode</td>
<td>20.5 (16–28)</td>
<td>22 (17–31)</td>
<td>−1.05&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
<tr>
<td>At most severe episode</td>
<td>28 (21–41)</td>
<td>33 (25–42)</td>
<td>−1.64&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
<tr>
<td><strong>Number of lifetime depressive episodes, n (%)</strong></td>
<td>23 (26.4)</td>
<td>77 (48.4)</td>
<td>11.27&lt;sup&gt;***&lt;/sup&gt;</td>
</tr>
<tr>
<td>Five or more episodes</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Duration of most severe depressive episode, n (%)</strong></td>
<td>70 (64.8)</td>
<td>127 (55.2)</td>
<td>3.15&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>Less than 3 months</td>
<td>18 (16.7)</td>
<td>55 (23.9)</td>
<td></td>
</tr>
<tr>
<td>3 to 6 months</td>
<td>20 (18.5)</td>
<td>48 (20.9)</td>
<td></td>
</tr>
</tbody>
</table>

IQR, interquartile range; MDD, major depressive disorder.

*Total does not equal 366 because of missing data.

**Chi-squared test.

**Mann–Whitney test.

* P < 0.05, ** P < 0.01, *** P < 0.001.

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### Table 2  Treatment characteristics and suicide history

<table>
<thead>
<tr>
<th></th>
<th>Unipolar (MDD) (n = 120)</th>
<th>Bipolar type 1/2 (n = 246)</th>
<th>Test statistic</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Treatment</strong>, n (%)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sought help from professional</td>
<td>94 (78.3)</td>
<td>208 (84.5)</td>
<td>2.16&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>Prescribed medication</td>
<td>84 (70.0)</td>
<td>188 (76.4)</td>
<td>1.74&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>Received ECT</td>
<td>15 (12.5)</td>
<td>49 (19.9)</td>
<td>3.08&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>Any treatment</td>
<td>96 (80.0)</td>
<td>211 (85.8)</td>
<td>1.99&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>Hospitalised for depression</td>
<td>32 (26.7)</td>
<td>114 (46.3)</td>
<td>13.02&lt;sup&gt;****&lt;/sup&gt;</td>
</tr>
<tr>
<td><strong>Suicide history</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ever attempted suicide, n (%)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>34 (28.3)</td>
<td>80 (32.5)</td>
<td>0.66&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td><strong>Number of attempts</strong>, n (%)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>One</td>
<td>23 (67.6)</td>
<td>38 (48.7)</td>
<td>3.53</td>
</tr>
<tr>
<td>Two</td>
<td>6 (11.8)</td>
<td>17 (21.8)</td>
<td></td>
</tr>
<tr>
<td>Three or more</td>
<td>8 (20.6)</td>
<td>23 (29.5)</td>
<td></td>
</tr>
<tr>
<td>Age at first suicide attempt, years: median (IQR)</td>
<td>17 (16–23)</td>
<td>24.5 (17–33)</td>
<td>−1.74&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
</tbody>
</table>

ECT, electroconvulsive therapy; IQR, interquartile range; MDD, major depressive disorder.

*Total does not equal 80 because of missing data.

**Chi-squared test.

**Mann–Whitney test.

* P < 0.05, ** P < 0.01, *** P < 0.001.
A number of robust differences were identified in the phenomenological and longitudinal characteristics of depressive episodes between cases of bipolar disorder and major depressive disorder within these bipolar disorder pedigrees. This is the first study to identify subgroups within major depressive disorder and bipolar depression.

**Identifying subgroups within major depressive disorder**

The propensity scores (probabilities from the logistic regression) for each of the 120 major depressive disorder cases were entered into a hierarchical cluster analysis, with a two-cluster solution providing the best fit. Cluster 1 consisted of 90 cases, with the remaining 30 cases assigned to cluster 2. We compared depressive symptoms across the two clusters, with several key differences emerging (Table 5). Cluster 1 was characterised by significantly higher rates of anhedonia, hypomania, psychomotor retardation, feelings of worthlessness, difficulty thinking/concentrating, morning worsening and delusions. In the Cluster 2 group, patients were more likely to have reported five or more major depressive episodes and to have been admitted to hospital during their most severe depressive episode. There was no significant difference in other clinical features or symptoms between the two clusters.

**Discussion**

A number of robust differences were identified in the phenomenological and longitudinal characteristics of depressive episodes between cases of bipolar disorder and major depressive disorder within these bipolar disorder pedigrees. This is the first study to identify subgroups within major depressive disorder and bipolar depression.
report comparing depressive features within such families, and supports growing evidence for substantive differences between bipolar and unipolar depression. First, there were a number of differences in the phenomenology of the most severe lifetime depressive episode across these disorders. In cases of bipolar disorder, patients reported a significantly higher prevalence of early morning waking, morning worsening, psychomotor retardation, difficulty thinking/concentrating, delusions and hallucinations (after controlling for age and gender). Psychomotor retardation was the most robust feature distinguishing between the two groups, supporting previous reports of psychomotor disturbance as a cardinal feature of bipolar depression,\textsuperscript{16–21} as well as historical clinical descriptions.\textsuperscript{5} Psychomotor retardation may have more relevance to bipolar type 1 depression, with Benazzi, for example, finding no difference in rates of retardation between bipolar type 2 disorder and major depressive disorder samples, but rather significantly higher rates of agitation among the group with type 2 bipolar disorder.\textsuperscript{22} In addition, we found a greater prevalence of psychotic features during depressive episodes in the bipolar disorder group, again consistent with numerous prior reports.\textsuperscript{23} The increased frequency of early morning waking in the bipolar group has previously been reported both in patients with a diagnosis of major depressive disorder who were found to have bipolar disorder on longitudinal follow-up,\textsuperscript{24} and in a number of cross-sectional comparisons with major depressive disorder,\textsuperscript{25} although there have been negative reports.\textsuperscript{26} Overall, the findings are consistent with bipolar disorder being characterised by both melancholic\textsuperscript{19} and psychotic features.

Second, in terms of longitudinal course and treatment, bipolar disorder cases had a greater number of lifetime depressive episodes (a finding which has been widely reported),\textsuperscript{3,26,27} and more hospital admissions, again consistent with previous studies.\textsuperscript{28,29} There was, however, no difference in rates of help-seeking or in usage of antidepressants or electroconvulsive therapy. Despite consistent prior reports of a younger age at onset for patients with bipolar disorder compared with non-related patients with major depressive disorder,\textsuperscript{30,31} no significant difference was found in the age at onset of the first depressive episode in this study. As age at onset of depression has been shown to be highly familial,\textsuperscript{32,33} the lack of a significant difference in this study may be due to the shared family history among cases. In a study comparing probands with bipolar or schizoaffective-bipolar disorder and unwell relatives, there was no difference in the age at onset for the first mood episode between the groups.\textsuperscript{34} Although higher rates of mixed features were observed among cases of bipolar disorder in the initial bivariate comparisons, this difference did not remain significant when logistic regression was used. This finding is perhaps surprising, given reports of the high rates of mixed symptoms in bipolar depression,\textsuperscript{35} although this may further suggest that some clinical differences traditionally seen between major depressive disorder and bipolar disorder were attenuated in the current sample owing to the shared family history.

Third, this study has been the first to empirically test the utility of a proposed probabilistic approach for distinguishing between bipolar and unipolar depression.\textsuperscript{2} Depending on the cut-off employed, the positive predictive value ranged from 74% to 82%. These values are consistent with or higher than those reported in numerous studies employing the Mood Disorders Questionnaire to differentiate between bipolar and unipolar cases, which have ranged from 36% to 79%.\textsuperscript{36,37} In terms of sensitivity and specificity, the optimal cut-off would appear to lie in the range of three to five features; final determination will require validation in independent samples.

Other approaches

Other methods have been proposed for distinguishing between bipolar disorder and major depressive disorder. Solomon \textit{et al} developed the Screening Assessment of Depression – Polarity (SAD-P), using three clinical features showing the greatest disparity between patients with the two disorders (number of episodes, family psychiatric history and psychotic features).\textsuperscript{38} This screen achieved a sensitivity of 82%, a specificity of 61% and a positive predictive value of 36%. In a study of non-psychotic bipolar depression and major depressive disorder involving outpatients recruited to three large multicentre trials (one of bipolar disorder type 1 and two of major depressive disorder), Perlis \textit{et al} found that family history, age at onset, number of episodes and a number of items on the Montgomery–Åsberg Depression Rating Scale and Hamilton Rating Scale for Depression differed between these groups.\textsuperscript{27} A stepwise logistic regression was undertaken which correctly classified 87% of patients, with a sensitivity of 69% and specificity of 95%. Excluding family history resulted in considerable loss of sensitivity (43%), highlighting the critical predictive utility of family history in distinguishing between bipolar and major depression. Furthermore, that model has not been tested in independent data-sets.

Identification of subgroups

The third aim of this study was to test if any differentiating features between bipolar depression and major depressive disorder could be used to identify subgroups within the major depressive disorder sample that might delineate ‘genetic’ and ‘sporadic’ populations, i.e. to determine an improved phenotype or case index of underlying bipolarity. Rice has argued that such a ‘caseness’ index would represent – in the absence of a diagnostic gold standard – a reasonable balance between the strict binary categories of diagnostic systems (such as DSM) and the clinical reality of more equivocal diagnoses.\textsuperscript{39} He suggested that this could be used to weight cases in genetic linkage studies based on the probability that they were truly genetically affected, thereby improving validity of classification and increasing the capacity to identify susceptibility genes.\textsuperscript{40} To our knowledge there has only been one report of a study looking for potential subgroups among major depressive disorder cases in bipolar disorder families. McMahon \textit{et al} examined age at onset in bipolar disorder pedigrees as an index of genetic liability, finding similarities between those with type 1 and type 2 disorders.\textsuperscript{9} However, there was no similarity in age at onset with the major depressive disorder cases, for which the authors reported a wide variance in age at onset. McMahon \textit{et al} interpreted this as indicating that the individuals with major depressive disorder represented a heterogeneous population with both genetic and sporadic cases, but it was not possible to statistically identify distinct subgroups. In two case–control studies, Blacker \textit{et al} compared illness features between unipolar depression in people with a family history of bipolar disorder and that in relatives of those with major depressive illness,\textsuperscript{14} or those in a control group with non-affected families,\textsuperscript{41} but these features showed little discriminatory power in either report. Gershon \textit{et al}, reporting on depression in relatives of patients with major depressive disorder and bipolar disorder, suggested that greater impairment and multiple lifetime episodes were possible clinical markers of a familial or genetically driven illness, consistent with the higher rates of hospital admission and greater number of lifetime episodes reported here in cluster 1.\textsuperscript{42}

We predicted that people with major depression with a postulated ‘genetic’ illness would be more likely to report features associated with bipolar depression. Using cluster analysis we
identified two groups, one of which included two-thirds of all major depressive disorder cases and was characterised by higher rates of symptoms found to be associated with bipolar depression in both this study and prior reports. Conversely, the other group demonstrated significantly lower rates of these ‘bipolar’ depressive features. Clearly this finding suggests some genetic subgroup must be regarded cautiously, requiring replication in independent samples before it can be confidently included in phenotypic definitions for linkage or other genetic analyses.

Limitations of the study
Several limitations should be considered when interpreting the findings of this paper. First, as the study recruited individuals from within bipolar disorder pedigrees, the results may not be generalisable to other populations; the major depressive disorder cases included here are not representative of major depressive disorder in general. Despite this, many findings that have previously been reported in the broader clinical literature on bipolar and unipolar depression were replicated in this genetically defined sample, wherein the shared genetic origins would be expected to diminish the likelihood of demonstrating such differences. Second, the sample size, although consistent with many other comparative studies, was not large enough to enable analysis of the bipolar type 1 and type 2 groups separately. The few studies that have separately examined the two types suggest that there may be important differences between them compared with major depressive disorder. Third, this validation of the probabilistic approach was limited, as not all variables included in the original list operational criteria could be included because of the limitations of the version of the DIGS used in this dataset. Finally, the assessment of depressive features was retrospective, as it was premised on the most severe lifetime episode.

Implications
In conclusion, a number of clinical features previously associated with bipolar depression were replicated in this sample, which was recruited through bipolar disorder pedigrees. This first empirical validation of the probabilistic approach in differentiating between unipolar and bipolar depression is consistent with dimensional distinctions between the two disorders and appears to offer clinical utility in identifying patients with depression who may warrant further assessment for bipolarity. Finally, cluster analysis of the major depressive disorder sample provided evidence suggestive of subgroups that might represent ‘genetic’ and ‘sporadic’ cases – a provocative finding that requires replication in other bipolar disorder pedigree samples.

Appendix
Criteria for a ‘probabilistic’ approach to the diagnosis of bipolar depression

A greater likelihood of diagnosis of bipolar type 2 disorder should be considered if five or more of the following features are present:

Symptoms and mental state signs
- Hypersomnia and/or increased daytime napping
- Hyperphagia and/or increased weight
- Other atypical depressive symptoms such as ‘leaden paralysis’
- Psychomotor retardation
- Psychotic features and/or pathological guilt
- Lability of mood and manic symptoms

Course of illness
- Early onset of first depression (<25 years of age)
- Multiple (five or more) prior episodes of depression

Family history
- Family history of bipolar disorder
  (Adapted from the criteria published by Mitchell et al)
  a. Data were available from the Diagnostic Interview for Genetic Studies (DIGS) to include this symptom in the probabilistic model.
  b. No relevant data were available in the DIGS, therefore this symptom was not included in the probabilistic model.

References
Depression in bipolar disorder pedigrees


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