Cardiometabolic disease and features of depression and bipolar disorder: population-based, cross-sectional study

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Background
The relative contribution of demographic, lifestyle and medication factors to the association between affective disorders and cardiometabolic diseases is poorly understood.

Aims
To assess the relationship between cardiometabolic disease and features of depression and bipolar disorder within a large population sample.

Method
Cross-sectional study of 145 991 UK Biobank participants: multivariate analyses of associations between features of depression or bipolar disorder and five cardiometabolic outcomes, adjusting for confounding factors.

Results
There were significant associations between mood disorder features and ‘any cardiovascular disease’ (depression odds ratio (OR) = 1.15, 95% CI 1.12–1.19; bipolar OR = 1.28, 95% CI 1.14–1.43) and with hypertension (depression OR = 1.15, 95% CI 1.13–1.18; bipolar OR = 1.26, 95% CI 1.12–1.42). Individuals with features of mood disorder taking psychotropic medication were significantly more likely than controls not on psychotropics to report myocardial infarction (depression OR = 1.47, 95% CI 1.24–1.73; bipolar OR = 2.23, 95% CI 1.53–3.57) and stroke (depression OR = 2.46, 95% CI 2.10–2.80; bipolar OR = 2.31, 95% CI 1.39–3.85).

Conclusions
Associations between features of depression or bipolar disorder and cardiovascular disease outcomes were statistically independent of demographic, lifestyle and medication confounders. Psychotropic medication may also be a risk factor for cardiometabolic disease in individuals without a clear history of mood disorder.

Declaration of interest
None.

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Bipolar disorder and major depression are common affective disorders, affecting approximately 2%1 and 16%2 of the population respectively. In addition to significant psychiatric morbidity, they also have an adverse impact on social and occupational functioning, quality of life and physical health,3–8 and often coexist with cardiovascular and metabolic diseases.9 The association of both major depressive disorder and bipolar disorder with cardiometabolic risk factors, vascular disease and poor cardiovascular outcomes is well recognised.10–12 Although psychotropic medications alter cardiovascular risk profiles (through weight gain, hypertension, dyslipidaemia and glucose dysregulation), their use is also associated with a range of demographic and lifestyle factors, including social deprivation, smoking and alcohol use.13 To date, there has been no large-scale population-based study in the UK that has assessed associations between a lifetime history of depressive or bipolar features and adverse cardiometabolic outcomes while also taking account of a wide range of potential confounding factors, including psychotropic medications.

Beyond the UK, there have been a number of large-scale reports assessing the association between mood disorders and cardiometabolic disease, including the National Comorbidity Study Replication2 and the World Mental Health Survey.7 Although very informative, these studies did not assess the wide range of covariates we have been able to include in our analyses. The landmark UK Biobank cohort,14 comprising over half a million adults in middle age, represents a unique opportunity to explore these associations at a population level (both cross-sectionally and prospectively) and has the potential to inform future mechanistic studies and the development of population-level interventions. Here we assess relationships between five cardiometabolic diseases (myocardial infarction, angina, hypertension, diabetes and stroke) and lifetime features of affective disorders (major depression and bipolar disorder) within UK Biobank. In additional subanalyses we also assess the effect of psychotropic medication on cardiometabolic disease.

Method

Data source
UK Biobank is a large prospective cohort of more than 502 000 residents of the UK, aged between 40 and 70 years.15 It is one of the largest resources worldwide for studying the genetic, environmental, medication and lifestyle factors that cause or prevent disease in middle and older age. Recruitment was undertaken over a 4-year period, from 2006 to 2010. In the final 2 years of UK Biobank recruitment, 172 751 participants were assessed in detail with respect to lifetime features of bipolar disorder and major depression. Eligibility for inclusion in our study was restricted to the 145 991 participants who provided complete data on lifetime features of mood disorder and complete data on self-reported cardiometabolic disease status.15

Data collection and ethical approval
Participants attended one of 22 assessment centres located across Great Britain. They completed a touchscreen questionnaire that collected information on demographics (including age, gender,
ethnicty and postcode), health-related behaviours (including smoking status and alcohol consumption) and a self-report of physician-diagnosed medical conditions, including diabetes, hypertension, myocardial infarction, angina and stroke. Current medication was recorded with assistance from a member of trained clinic staff, who also used standard operating procedures to measure height and weight for body mass index (BMI) calculation. This study was conducted under generic approval from the NHS National Research Ethics Service (approval letter dated 17 June 2011, Ref 11/NW/0382) and full written informed consent was gained from participants at the point of data collection.

Definitions
Criteria for a lifetime history of clinically significant features of bipolar disorder or depression were based on responses to questions within the baseline touchscreen questionnaire (see online supplements DS1 and DS2). Although not diagnostic of bipolar disorder or major depressive disorder, these questions were similar to questions assessing mood disorders within structured diagnostic assessment instruments. To some extent the validity of these questions has been supported internally within this data-set by comparisons of gender distribution, socioeconomic status, self-reported health rating, current depressive symptoms and smoking status.15

For the purposes of this study, participants were categorised into three groups: those with bipolar disorder features, those with features of major depression and a control group (Appendix).15,16 'Any cardiovascular disease' was defined as the self-report of a previous physician diagnosis of hypertension, myocardial infarction, angina and/or stroke. Diabetes was also defined by self-report. BMI was determined though anthropometric measurements carried out at the assessment centre and categorised into underweight (< 18.5 kg/m²), normal weight (18.5–24.9 kg/m²), overweight (25.0–29.9 kg/m²), class I obese (30.0–34.9 kg/m²), class II obese (35.0–39.9 kg/m²) and class III obese (≥ 40 kg/m²). Smoking status, frequency of alcohol consumption and ethnic group were self-reported on the touchscreen questionnaire. Smoking status was classified as 'current smoker', 'previous smoker' or 'never smoked' and alcohol use was classified as 'daily/almost daily', '3–4 times per week', '1–2 times per week', '1–3 times/month', 'special occasions only' or 'never'. Ethnicity was categorised as: 'White', 'mixed', 'Asian/Asian British', 'Black/Black British', 'Chinese', and 'other'. The Townsend deprivation index – an area-based measure of socioeconomic status derived from information collected in the census on car ownership, overcrowding, owner-occupation and unemployment – includes both positive and negative values, with positive values indicating higher levels of deprivation.17 Townsend scores were divided into quintiles (within the study population) to facilitate comparisons. The use of current medication (including psychotropics) was self-reported by participants and a comprehensive list of commonly used psychotropics was identified (online supplement DS3). Participants were classified as using psychotropic medication if they reported currently taking any of these medications.

Statistical analyses
Differences in baseline characteristics between the three groups (control, depressive features and bipolar features) were analysed using the chi-squared test for categorical data and the chi-squared test for trend for ordinal data. We used separate logistic regression models to examine the associations between mood group and cardiometabolic disease categories ('any cardiovascular disease' (not including diabetes), diabetes, hypertension, myocardial infarction, angina or stroke), with controls as the referent group. These primary outcomes were used to provide a more comprehensive analysis of common cardiometabolic comorbidities in the depressive and bipolar features groups. Models were initially adjusted for age, gender, socioeconomic deprivation and ethnicity (a partially adjusted model) and then repeated including additional covariates: smoking status, frequency of alcohol use, BMI and the use of psychotropic medication (fully adjusted model). Interaction tests for gender were undertaken and subgroup analyses carried out as appropriate.

To explore the relative contribution of psychotropic medication to cardiometabolic risk across the mood spectrum, we carried out a subanalysis to assess the effect of current psychotropic medication and mood group status on cardiometabolic disease across the mood spectrum. Six groups were created: individuals with no mood disorder features and not currently taking psychotropic medication (our referent group); individuals with no mood disorder features but currently taking one or more psychotropic medications; individuals with depressive features not currently taking psychotropic medication; individuals with depressive features currently taking one or more psychotropic medications; individuals with features of bipolar disorder not currently taking any psychotropic medication; and individuals with features of bipolar disorder currently taking psychotropic medication.

Differences in cardiometabolic outcomes between different mood and medication status groups were reported as prevalences and then analysed using the chi-squared test for categorical data. We again used separate logistic regression models to examine associations between medication and mood disorder group with cardiometabolic disease (diabetes, myocardial infarction, angina, hypertension or stroke). As noted above, controls who were not currently taking psychotropic medication were chosen as the most appropriate referent group. These models were adjusted for age, gender, socioeconomic deprivation and ethnicity, smoking, alcohol and BMI. All statistical analyses were performed using Stata version 12.1. Statistical significance was defined conservatively as P < 0.001.

Results

Group characteristics: bipolar features, depression features and controls
A total of 172 751 UK Biobank participants were assessed at baseline with respect to lifetime history of depressive and bipolar features. Of these, complete data on mood disorder features and cardiometabolic disease status were available for 145 991 (84.5%). From this sample, according to our criteria in the Appendix, 1557 (1.07%) had features of bipolar disorder, 30 990 (21.23%) had features of major depression and 113 444 (77.71%) had no significant features of mood disorder (controls) (Table 1).

Participants with features of depression were younger and more likely to be women (Table 1). Current smoking was most common in the bipolar features group (21.3%), followed by the depressive features group (12.7%) and then the non-mood-disordered control group (8.8%) (Table 2). The proportions of different levels of socioeconomic deprivation were mixed between the groups, but of note was that 34.6% of the bipolar features group were in the most deprived quintile, compared with 22.4% for the depressive features group, and 19.2% for the control group. White ethnicity was most common in the depressive features group and least common in the bipolar features group (Table 1).

Patterns of alcohol use were similar between groups (Table 2) and obesity was more common in both the depressive features and
bipolar features groups compared with the control group (Table 1). Rates of current psychotropic medication use were highest in the groups with bipolar (32.1%) and depressive (20.6%) features and lowest within the control group (3.4%) (Table 2).

Prevalence of cardiometabolic disease
The prevalence of ‘any cardiovascular disease’ was highest in the bipolar features group (34.5%), followed by the depressive features group (30.6%) and lowest in the control group (28.7% (Table 2). Cardiometabolic disease frequencies were also calculated for men and women. Men had consistently higher rates of all cardiometabolic diseases and, in general, rates of cardiometabolic disease were highest in the bipolar features group, followed by the depressive features group and the control group (Table 2).

Partially adjusted model
On multivariate logistic regression (adjusting for age, gender, socioeconomic deprivation and ethnicity), odds ratios (ORs) for all cardiometabolic outcomes were significantly increased in the depressive features and the bipolar features groups relative to controls (all \( P < 0.001 \), Table 3, see also online Table DS1 for a more detailed version of this table that includes an analysis by gender). In general, the odds of each cardiometabolic outcome, as well as the odds of having ‘any cardiovascular illness’, increased across the two mood groups, in the depressive features group and the bipolar features group (Table 3). It should be noted, however, that findings for the bipolar features group with respect to diabetes and stroke were not statistically significant. The highest odds ratios in the partially adjusted model for the combined male and female group were for myocardial infarction (OR = 1.90, 95% CI 1.44–2.51) and angina (OR = 1.69, 95% CI 1.30–2.19) within the bipolar features group relative to controls, and for diabetes (OR = 1.29, 95% CI 1.22–1.37) in the depressive features group relative to controls (Table 3).

There was a significant interaction between mood disorder and gender in predicting risk of having ‘any cardiovascular illness’ (\( P < 0.001 \)) and subgroup analyses by gender were subsequently carried out (Table DS1). Men with bipolar features had elevated odds ratios for myocardial infarction (2.02, 95% CI 1.50–2.72). The odds ratio for myocardial infarction in women was not significantly elevated in the partially adjusted group (1.37, 95% CI 0.65–2.92). Women with bipolar features had slightly higher odds ratios for any cardiovascular disease (1.55, 95% CI 1.32–1.83 in women v. 1.46, 95% CI 1.25–1.69 in men) and for hypertension (1.46, 95% CI 1.24–1.73 in women v. 1.42, 95% CI 1.22–1.66 in men) (Table DS1).

The odds ratios for specific conditions within the depressive features group were broadly similar for men and women. For angina this was OR = 1.54 (95% CI 1.40–1.69) for men and OR = 1.43 (95% CI 1.27–1.61) for women. Similarly, for diabetes the OR for men was 1.32 (95% CI 1.23–1.44) and for women the OR was 1.25 (95% CI 1.15–1.36). In women with depressive features, the odds ratio for stroke was 1.70 (95% CI 1.47–1.97) compared with 1.53 (95% CI 1.33–1.76) for men (Table DS1).

Fully adjusted model
On additional adjustment for smoking status, frequency of alcohol consumption, BMI and the use of psychotropic medication, the risk of having ‘any cardiovascular illness’ remained significantly elevated in both the depressive (OR = 1.15, 95% CI 1.12–1.19) and the bipolar features groups (OR = 1.28, 95% CI 1.14–1.43)
This pattern was also largely present in a subgroup analysis by gender (Table DS1). In the fully adjusted model, odds ratios for hypertension and myocardial infarction were also significantly elevated in the combined male and female groups (hypertension: depression features group OR = 1.15, 95% CI 1.13–1.18, bipolar features group OR = 1.26, 95% CI 1.12–1.42; myocardial infarction: depression features group OR = 1.18, 95% CI 1.08–1.30) (Table 3).

Differences in odds ratios for angina and stroke within the fully adjusted, combined group analyses were not statistically significant. There were, however, elevations in odds of cardiovascular conditions within the depressive features group, most notably for stroke (OR = 1.26, 95% CI 1.13–1.30) (Table 3). Associations with diabetes were not significant in the fully adjusted model in either the depressive features or bipolar features groups. Odds ratios were reduced by approximately a quarter for many of the outcome measures on the additional adjustment for BMI, smoking, alcohol consumption and psychotropic medication.

**Current use of any psychotropic medication**

We carried out a subanalysis to assess the contribution of current psychotropic medication use to cardiometabolic risk in the control group, the depressive features group and the bipolar features group. In total, 109,557 individuals without features of mood disorder who were not taking psychotropic medication were identified as a control group (Table 4). There were 3867 individuals who did not meet the criteria for depressive or bipolar features but who reported taking one or more psychotropics (Table 4). Although the indication for these medications was not known, individuals who did not meet the criteria for depressive or bipolar disorder but reported taking one or more psychotropics displayed higher prevalences of all cardiometabolic diseases compared with controls. Of particular note were elevated rates of diabetes (8.97%), angina (5.87%), hypertension (36.07%) and stroke (3.98%) (Table 4).

As illustrated in Table 4, compared with controls not on psychotropic medication, individuals with depressive features plus no psychotropic medication had similar rates of diabetes, myocardial infarction, angina, hypertension and stroke. Participants with depressive features taking psychotropic medication had higher rates of all cardiometabolic diseases compared with controls not on psychotropic medication. Individuals with bipolar features not taking psychotropic medication had higher rates of cardiometabolic disease relative to controls not on psychotropic medication. As expected, cardiometabolic diseases were also more prevalent for individuals with bipolar features who reported taking psychotropic medication.

**Fully adjusted medication analyses**

To assess the relative contribution of current psychotropic medication use in cardiometabolic disease across the mood...
Cardiometabolic disease and features of depression and bipolar disorder spectrum we carried out a multivariate logistic regression adjusting for age, gender, socioeconomic deprivation, ethnicity, smoking status, frequency of alcohol consumption and BNI (Fig. 1(a)–(f)). In general, odds ratios for most cardiometabolic diseases remained significantly elevated in each of the mood and medication groups relative to controls (Fig. 1(a)–(f)). Exceptions were for diabetes in the depressive features group not taking psychotropic medication, the bipolar features group not taking psychotropic medication, and the bipolar group currently taking psychotropic medication. Increases in odds ratios in the bipolar features group not on psychotropic medication were also not significantly elevated for angina (OR = 1.58, 95% CI 1.22–2.06) and stroke (OR = 1.70, 95% CI 1.20–2.41).

Furthermore, odds ratios for angina and stroke were not significantly elevated for both the bipolar features groups (those currently not taking psychotropic medication, as well as those taking psychotropic medication) (Fig. 1(d) and (f)). In general, however, the odds of reporting a history of an adverse cardiometabolic outcome were associated with both psychotropic medication use and with mood disorder (Fig. 1(a)–(f)) and the size of these associations increased with mood disorder severity (from depression to bipolar).

Discussion

We found that in a very large population sample of adults with lifetime features of depression and bipolar disorder there was an increased risk of comorbid cardiovascular disorders, even after adjusting for a wide range of confounding factors. In general, these associations were more pronounced for individuals with features of bipolar disorder than features of depression. Perhaps unsurprisingly, we also identified an association between current use of psychotropic medication and risk of cardiometabolic disease in individuals with a history of depressive and bipolar features. It is, however, notable that this association also occurred in individuals with no definite history of mood disorder who were currently taking psychotropic medication.

Both the bipolar features and depressive features groups had significantly higher odds ratios for ‘any cardiovascular disease’

### Table 3

<table>
<thead>
<tr>
<th>Cardiovascular disease any</th>
<th>Partially adjusted&lt;sup&gt;b&lt;/sup&gt;</th>
<th>Fully adjusted&lt;sup&gt;c&lt;/sup&gt;</th>
</tr>
</thead>
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<tr>
<td><strong>OR (95% CI)</strong></td>
<td><strong>P</strong></td>
<td><strong>OR (95% CI)</strong></td>
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<td>Control (Reference)</td>
<td>1 (Reference)</td>
<td>1 (Reference)</td>
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<tr>
<td>Depression</td>
<td>1.29 (1.25–1.33)</td>
<td>&lt;0.001</td>
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<tr>
<td>Bipolar</td>
<td>1.50 (1.34–1.68)</td>
<td>&lt;0.001</td>
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<tr>
<td>Diabetes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Control (Reference)</td>
<td>1 (Reference)</td>
<td>1 (Reference)</td>
</tr>
<tr>
<td>Depression</td>
<td>1.29 (1.22–1.37)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Bipolar</td>
<td>1.37 (1.15–1.67)</td>
<td>0.002</td>
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<tr>
<td>Hypertension</td>
<td></td>
<td></td>
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<tr>
<td>Control (Reference)</td>
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<td>1 (Reference)</td>
</tr>
<tr>
<td>Depression</td>
<td>1.27 (1.23–1.31)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Bipolar</td>
<td>1.44 (1.29–1.61)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Myocardial infarction</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Control (Reference)</td>
<td>1 (Reference)</td>
<td>1 (Reference)</td>
</tr>
<tr>
<td>Depression</td>
<td>1.38 (1.26–1.51)</td>
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<tr>
<td>Bipolar</td>
<td>1.90 (1.44–2.51)</td>
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<tr>
<td>Angina</td>
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<td></td>
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<tr>
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<td>1 (Reference)</td>
</tr>
<tr>
<td>Depression</td>
<td>1.49 (1.39–1.61)</td>
<td>&lt;0.001</td>
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<tr>
<td>Bipolar</td>
<td>1.69 (1.30–2.19)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Stroke</td>
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<td>1 (Reference)</td>
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<tr>
<td>Bipolar</td>
<td>1.80 (1.27–2.54)</td>
<td>0.001</td>
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</tbody>
</table>

<sup>a</sup> See online Table DS1 for a more detailed version of this table that includes data for men and women separately.

<sup>b</sup> Partially adjusted for age, gender, deprivation and ethnicity.

<sup>c</sup> Fully adjusted for age, gender, deprivation, ethnicity, body mass index, smoking status, alcohol consumption and psychotropic medication.

### Table 4

<table>
<thead>
<tr>
<th>Mood group</th>
<th>Total, n</th>
<th>Diabetes</th>
<th>Myocardial infarction</th>
<th>Angina</th>
<th>Hypertension</th>
<th>Stroke</th>
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</thead>
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<tr>
<td>Controls not on psychotropic medication</td>
<td>109,577</td>
<td>5640 (51.5)</td>
<td>2262 (20.6)</td>
<td>2921 (26.7)</td>
<td>28,287 (25.81)</td>
<td>1316 (12.2)</td>
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<td>Controls on psychotropic medication</td>
<td>3867</td>
<td>347 (8.97)</td>
<td>136 (3.52)</td>
<td>227 (5.87)</td>
<td>1395 (36.07)</td>
<td>154 (3.98)</td>
</tr>
<tr>
<td>Depressive features not on psychotropic medication</td>
<td>24,603</td>
<td>1199 (4.87)</td>
<td>508 (2.06)</td>
<td>690 (2.80)</td>
<td>609 (2.86)</td>
<td>370 (1.50)</td>
</tr>
<tr>
<td>Depressive features on psychotropic medication</td>
<td>6387</td>
<td>531 (8.31)</td>
<td>168 (2.63)</td>
<td>321 (5.03)</td>
<td>2065 (32.33)</td>
<td>192 (3.01)</td>
</tr>
<tr>
<td>Bipolar features not on psychotropic medication</td>
<td>1057</td>
<td>108 (10.22)</td>
<td>55 (5.20)</td>
<td>63 (5.96)</td>
<td>484 (45.79)</td>
<td>34 (3.22)</td>
</tr>
<tr>
<td>Bipolar features on psychotropic medication</td>
<td>500</td>
<td>48 (9.60)</td>
<td>25 (5.00)</td>
<td>24 (4.80)</td>
<td>165 (33.00)</td>
<td>16 (3.20)</td>
</tr>
</tbody>
</table>
relative to controls, within partially and fully adjusted models. Given the broad range of common confounding variables that were adjusted for, this suggests an independent association between mood disorder and cardiometabolic diseases. Within the partially adjusted (for age, gender, deprivation, ethnicity) versus fully adjusted models (additionally adjusted for BMI, smoking status, alcohol consumption and psychotropic medication), odds ratios for ‘any cardiovascular disease’ fell from 1.29 (partial) to 1.15 (full) within the depressive features group and from 1.50 (partial) to 1.28 (full) in the bipolar features group. Although depression has been considered a risk factor for cardiovascular disease for some time,18 there has been debate about the relative contribution of lifestyle factors for this group of patients. We have been able to control for some of these lifestyle factors.

**Strengths and limitations**

The strengths of this study include the general population design, the breadth of coverage of confounding factors and the very large sample size. However, several limitations are acknowledged. Cardiometabolic disorders previously diagnosed by a physician were self-reported by the participants. Similarly, mood disorder features were self-reported, rather than assessed using a formal diagnostic interview, although a structured approach was used (Appendix). Psychiatric diagnoses based on formal interviews...
were not practical within UK Biobank, so we took a pragmatic approach to mood disorder groupings. It is possible that our criteria for mood disorder are less stringent than formal diagnostic criteria. It should be noted, however, that we are reporting features of mood disorders at a population level, rather than formal clinical diagnoses, and we were unable to validate diagnoses because of the lack of availability of definitive diagnostic information. Similarly, because of a lack of definitive cardiometabolic diagnostic information we were unable to validate outcome measures. This lack of validation for both cardiometabolic and psychiatric diagnoses has implications in terms of the likely sensitivity and specificity of our groupings, which are based on self-report. Linkage to routine hospital and general practice health records will be available in the future and will allow some validation of these groups.

There is, however, evidence for the validity of these groupings using internal variables, for example, the gender distributions of approximately 1:1 for bipolar disorder and approximately 2:1 (women:men) for depression are consistent with a large body of epidemiological research on lifetime rates of mood disorder in men and women. Furthermore, the lifetime prevalence rates for bipolar disorder (1.1%) and recurrent major depressive disorder (21.2%) are consistent with other population-based lifetime estimates.

Although we are reasonably confident that members of the control groups did not have significant features of depression or bipolar disorder, it should be noted that a proportion may have fulfilled criteria for other mental illnesses, such as anxiety disorder or, less likely, schizophrenia, both of which have been associated with poor cardiometabolic health. There are also limitations with our broad definition of current psychotropic medication use that groups together different classes of medication. In a subanalysis of individuals on psychotropic monotherapy, we found that selective serotonin reuptake inhibitors (SSRIs), other antidepressants and sedatives/hypnotics were all associated with greater risk of myocardial infarction relative to psychotropic-free controls (SSRIs OR = 1.82, 95% CI 1.51–2.21; other antidepressants OR = 1.50, 95% CI 1.20–1.87; and sedatives/hypnotics OR = 2.53, 95% CI 1.76–3.66). It is therefore possible that adverse cardiovascular outcomes in depressive disorders are not limited to the use of antipsychotics but may also be a consequence of other classes of psychotropic medications.

A relatively low proportion of the mood disorder features groups reported taking psychotropic medication (bipolar features 32.1%, depressive features 20.6% and controls 3.4%) but, as noted above, the focus in this study was not clinically diagnosed mood disordered groups but rather lifetime features of mood disorder at a population level. Poor adherence with medication may be an issue. According to some estimates, up to 50% of patients prescribed psychotropic medication do not take their medication regularly. Although adherence for physical health medications may be slightly better than this, it is likely that the factors that cause poor adherence with psychotropic medication may also lead to poor adherence with cardiovascular medications in individuals with mood disorders. This might explain the proportionately worse cardiovascular outcomes in a less adherent mood disordered group compared with a more adherent non-mood disordered group. As a result of the level of information on medication status that was collected at baseline, we were unable to assess the impact of duration of exposure to psychotropic medication on adverse cardiometabolic outcomes.

We included a range of possible confounding variables in the regression models but were unable to control for physical activity. The available data on physical activity were not collected in terms of standardised measures. Unfortunately, it was not possible to create a standardised measure, such as the metabolic equivalent of task (MET). Similarly, we were unable to adjust for the potential confounding effects of diet. When considering appropriate variables to include as confounding variables, it is important to note that there is uncertainty as to the extent that these variables might represent confounders or mediators. For example, obesity may represent an important component of the pathophysiology for depression, bipolar disorder and coronary heart disease. A prospective cohort study is required to address these concerns and is planned as part of future work in this cohort.

Our analyses did not assess the additive effects of sociodemographic and lifestyle factors. Rather, given the large number of confounding factors involved, we took a broad approach, focusing on cardiovascular outcomes after partial and full adjustment for confounders.

Severity of mood disorder may have contributed to the associations we observed. We therefore conducted an additional analysis of the impact of illness severity on cardiovascular disease outcomes. The depressive features group were divided into those who had a history of a probable single episode of major depression, versus those with a history of probable moderate recurrent depression and those with probable severe recurrent depression (Appendix). The bipolar group was also split into those with features of probable bipolar type I illness and those with features of probable bipolar type II illness (Appendix). In general, individuals in the more severe mood disorder groups had higher risk of adverse cardiovascular outcomes. For the bipolar type I group the odds ratio of ‘any cardiovascular illness’ was 1.39 (95% CI 1.23–1.58), for the recurrent severe depression group the OR was 1.33 (95% CI 1.18–1.50) and for the bipolar type II group the OR was 1.21 (95% CI 1.09–1.32).

Findings in the context of previous work

Many studies have identified an association between major depression and cardiovascular disease but relatively few studies have examined this with respect to bipolar disorder. Although some reports in the literature support the association between bipolar disorder and cardiovascular disease, these studies have not been able to adjust for the same range of confounding factors as our study.

Relative to controls, women with features of bipolar disorder had a greater risk of ‘any cardiovascular disease’ than men with bipolar features (OR 1.36 v. 1.19). This finding may be particularly noteworthy given that men are known to have increased rates of cardiovascular disease within the general population. In their 2013 population study, Crump and colleagues reported increased all-cause mortality for women with bipolar disorder compared with men with bipolar disorder (adjusted hazard ratio of 2.13 compared with 1.74). Furthermore, the same study reported higher hazard ratios of death from ischaemic heart disease for women with bipolar disorder compared with men with bipolar disorder (2.14 compared with 1.73). Our findings add to the evidence that women with bipolar disorder may be disproportionately affected by cardiovascular disease.

In our fully adjusted model for individuals with depressive or bipolar features, men (but not women) had an increased risk of myocardial infarction, and women (but not men) were at higher risk of stroke (Table DS1). There are several possible explanations for this, including studies that have found that men are more likely to receive a diagnosis of myocardial infarction than women and that stroke is more commonly diagnosed in middle-aged women than in men. We also found that, compared with both controls and women, men with depressive features had a higher risk of ‘any cardiovascular disease’ and hypertension,
supporting previous findings in this area. Furthermore, we found that in the combined male and female results, depressive features were associated with an increased risk of ‘any cardiovascular disease’. Although odds ratios were not as large as previous studies, this may underscore the importance of adjusting for a wide variety of confounding factors where possible.

Possible mechanisms

There are several possible underlying mechanisms for the associations we have observed, including shared behavioural factors, inequalities in treatment, shared genetic vulnerabilities and shared pathophysiology. Individuals with mood disorder may have difficulties in accessing preventative medical interventions, as well as higher rates of risky lifestyle factors. Such factors include smoking, physical inactivity and reduced adherence with cardiovascular medications. In keeping with other work, smoking was more common in the bipolar features mood disorder and cardiometabolic disease. Our results also show a significant attenuation in odds ratios when moving from partially adjusted to fully adjusted analyses, underlining the impact of smoking, alcohol, medication and BMI on cardiovascular risk in individuals with depressive and bipolar features.

Although our study adds to knowledge on the possible contribution of lifestyle and demographic factors to the association between cardiometabolic disease and mood disorder, it cannot contribute substantially to understanding pathophysiological mechanisms. Several biological systems may contribute to the comorbidity between mood disorder and cardiometabolic disease, including (but not limited to) abnormalities of the hypothalamic–pituitary–adrenal (HPA) axis, oxidative stress processes, shared genetic vulnerability, abnormal inflammatory responses, endothelial dysfunction and haemodynamic reactivity.

We did, however, find that the use of psychotropic medication was associated with an increased risk of cardiovascular disease in all groups, even those without features of mood disorder. The metabolic effects of psychotropic medications are known to increase cardiometabolic risk by various mechanisms, including weight gain and impaired glucose tolerance. Given that our analyses were conducted on the basis of current use of any psychotropic medication, it is perhaps noteworthy that adverse cardiometabolic effects may not be limited to antipsychotics. There are reports in the literature of other psychotropic medications used in the treatment of mood disorder, such as antidepressants, lithium and valproic acid, having adverse metabolic effects.

Future work

There remain several unanswered questions and opportunities for further research. There is a need for longitudinal and mechanistic studies in this area to better understand causal pathways. To date, very few studies have tested specific genetic associations between lifestyle-related risk factors for cardiometabolic disease across the broad clinical spectrum of mood disorders. In addition, further mechanistic studies examining epigenetic factors, as well as oxidative stress, endothelial dysfunction and HPA axis abnormalities will be important to better understand comorbidity between mood disorder and cardiometabolic disease.

Funding

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Appendix

Criteria for lifetime features of bipolar disorder and depression

Bipolar disorder features

(a) Features of bipolar disorder, type I: ever ‘manic or hyper’ for at least 2 days OR ever ‘irritable/argumentative’ for 2 days; plus at least 3 features from ‘more active’, ‘more talkative’, ‘needed less sleep’ and ‘more creative/more ideas’; plus duration of a week or more; plus ‘needed treatment or caused problems at work’.

(b) Features of bipolar disorder, type II: ever ‘manic or hyper’ for at least 2 days OR ever ‘irritable/argumentative’ for 2 days; plus at least 3 features from ‘more active’, ‘more talkative’, ‘needed less sleep’ and ‘more creative/more ideas’; plus duration of a week or more.

Major depression features

(a) Features of single-episode major depression: ever depressed/down for a whole week; plus at least 2 weeks duration; plus only one episode; plus ever seen a general practitioner (GP) or a psychiatrist for ‘nerves, anxiety, depression’ OR ever anhedonic (unenthusiasm/uninterest) for a whole week; plus at least 2 weeks duration; plus only one episode; plus ever seen a GP or a psychiatrist for ‘nerves, anxiety, depression’.

(b) Features of recurrent major depression (moderate): ever depressed/down for a whole week; plus at least 2 weeks duration; plus at least two episodes; plus ever seen a GP (but not a psychiatrist) for ‘nerves, anxiety, depression’ OR ever anhedonic (unenthusiasm/uninterest) for a whole week; plus at least 2 weeks duration; plus at least two episodes; plus ever seen a GP (but not a psychiatrist) for ‘nerves, anxiety, depression’.

(c) Features of recurrent major depression (severe): ever depressed/down for a whole week; plus at least 2 weeks duration; plus at least 2 episodes; plus ever seen a psychiatrist for ‘nerves, anxiety, depression’ OR ever anhedonic (unenthusiasm/uninterest) for a whole week; plus at least 2 weeks duration; plus at least two episodes; plus ever seen a psychiatrist for ‘nerves, anxiety, depression’.

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Unique data identifier (UDI) codes

Features of bipolar disorder (type I)
Either: 4642 Ever manic/hyper for at least 2 days or 4653 Ever irritable/argumentative for at least 2 days, plus
At least 3 from 6156.01 (more active), 6156.02 (more talkative), 6156.03 (needed less sleep), and 6156.04 (more creative/more ideas), plus
5663 Duration of a week or more, plus
5674 Needed treatment or caused problems at work

Features of bipolar disorder (type II)
Either: 4642 Ever manic/hyper for at least 2 days or 4653 Ever irritable/argumentative for at least 2 days, plus
At least 3 from 6156.01 (more active), 6156.02 (more talkative), 6156.03 (needed less sleep), and 6156.04 (more creative/more ideas), plus
5663 Duration of a week or more

Single (probable) episode of major depression
EITHER:
4598 Ever depressed/down for a whole week, plus
4609 At least 2 weeks duration, plus
4620 Only one episode, plus
2090 Ever seen a GP or 2100 a psychiatrist for nerves, anxiety, depression

OR:
4631 Ever anhedonic (unenthusiasm/uninterest) for a whole week, plus
5375 At least 2 weeks, plus
5386 At least one episode, plus
2090 Ever seen a GP (but not a psychiatrist) for nerves, anxiety, depression

Probable recurrent major depression (moderate)
EITHER:
4598 Ever depressed/down for a whole week, plus
4609 At least 2 weeks duration, plus
4620 At least two episodes, plus
2090 Ever seen a GP (but not a psychiatrist) for nerves, anxiety, depression

OR:
4631 Ever anhedonic (unenthusiasm/uninterest) for a whole week, plus
5375 At least 2 weeks, plus
5386 At least two episodes, plus
2090 Ever seen a GP (but not a psychiatrist) for nerves, anxiety, depression

Probable recurrent major depression (severe)
EITHER:
4598 Ever depressed/down for a whole week, plus
4609 At least 2 weeks duration, plus
4620 At least two episodes, plus
2100 Ever seen a psychiatrist for nerves, anxiety, depression

OR:
4631 Ever anhedonic (unenthusiasm/uninterest) for a whole week, plus
5375 At least 2 weeks, plus
5386 At least two episodes, plus
2100 Ever seen a psychiatrist for nerves, anxiety, depression
Online supplement DS2
Questions and answers used to assess mood disorder features

Bipolar features questions

(a) Have you ever had a period of time lasting at least 2 days when you were feeling so good, ‘high’, excited or ‘hyper’ that other people thought you were not your normal self or you were so ‘hyper that you got into trouble?’ (4642) or ‘Have you ever had a period of time lasting at least 2 days when you were so irritable that you found yourself shouting at people or starting fights or arguments? (4653)’
Select: Yes, No, Don’t know, Prefer not to answer

(b) Please try to remember a period when you were in a ‘high’ or ‘irritable’ state and select which of the following apply
Select:
I was more active than usual (6156.01)
I was more talkative than usual (6156.02)
I needed less sleep than usual (6156.03)
I was more creative or had more ideas than usual (6156.04)
All of the above
None of the above

(c) What is the longest time period that these ‘high’ or ‘irritable’ periods have lasted? (5663)
Select:
At least 2 days, but less than a week
Less than a week
A week or more
Do not know
Prefer not to answer

(d) How much of a problem have these ‘high’ or ‘irritable’ periods caused you? (5674)
Select:
Needed treatment or caused problems with work, relationships, finances, the law or other aspects of life
Do not know
Prefer not to answer

Depressive features questions

(a) Looking back over your life, have you ever had a time when you were feeling depressed or down for at least a whole week? (4598)
Select: Yes, No, Don’t know, Prefer not to answer

(b) How many weeks was the longest period when you were feeling depressed or down? (4609)
Participants asked to enter the number of weeks on the touchscreen.

(c) How many periods have you had when you were feeling depressed or down for at least a whole week?
Participants asked to enter the number of periods on the touchscreen.

(d) Have you ever seen a general practitioner (GP) for nerves, anxiety, tension or depression? (2090) or Have you ever seen a psychiatrist for nerves, anxiety, tension or depression? (2100)
Select: Yes, No, Don’t know, Prefer not to answer

Anhedonia

(a) Have you ever had a time when you were uninterested in things or unable to enjoy the things you used to for at least a whole week? (4631)
Select: Yes, No, Don’t know, Prefer not to answer

(b) How many weeks was the longest period when you were uninterested in things or unable to enjoy the things you used to?’
Participants asked to enter the number of weeks on the touchscreen.

(c) How many periods have you had when you were uninterested in things or unable to enjoy the things you used to for at least a whole week?’
Participants asked to enter the number of periods on the touchscreen.

(d) Have you ever seen a general practitioner (GP) for nerves, anxiety, tension or depression? (2090) or Have you ever seen a psychiatrist for nerves, anxiety, tension or depression? (2100)
Select: Yes, No, Don’t know, Prefer not to answer
## List of psychotropic medications

### 1 Mood stabilisers
- Lithium product
  - Priadel (lithium)
  - Camcolit (lithium)
- Sodium valproate
  - Epilim (sodium valproate)
- Valproic acid
  - Carbamazepine product
  - Tegretol (carbamazepine)
  - Teril (carbamazepine)
  - Teril retard (carbamazepine)
  - Timonil retard (carbamazepine)
  - Epimav (carbamazepine)
- Sodium valproate
  - Epilim (sodium valproate)
- Carbamazepine
  - Cipramil (carbamazepine)
  - Tegretol (carbamazepine)
  - Teril (carbamazepine)
  - Teril retard (carbamazepine)
  - Timonil retard (carbamazepine)
  - Epimav (carbamazepine)

### 2 Antidepressant – selective serotonin reuptake inhibitors
- Paroxetine
- Seroxat (paroxetine)
- Fluoxetine
- Prozac (fluoxetine)
- Citalopram
- Cipramil (citalopram)
- Esctalopram
- Cipralex (escitalopram)
- Sertraline
- Lustral (sertraline)
- Fluvoxamine

### 3 Antidepressant – other
- Mirtazapine
- Zispin (mirtazapine)
- Duloxetine
- Cymbalta (duloxetine)
- Venlafaxine
- Effexor (venlafaxine)
- Amitriptyline
- Elavil (amitriptyline)
- Tryptizol (amitriptyline)
- Lentizol (amitriptyline)
- Amitriptyline+perphenazine
- Tripafon (amitriptyline+perphenazine)
- Amitriptyline+chlodiazepoxide
- Limbitorl 10 (amitriptyline+chlodiazepoxide)
- Limbifrol 5 (amitriptyline+chlodiazepoxide)
- Phenelzine
- Monoamine oxidase inhibitor – phenelzine
  - Nardil (phenelzine)
  - Moclobemide
  - Manerix (moclobemide)
  - Imipramine
  - Tofranil (imipramine)
  - Trimipramine
  - Surmontil (trimipramine)
  - Dothepin
  - Dosulepin
  - Prothiaden (dosulepin)
  - Thaden (dosulepin)
  - Clomipramine
  - Anafranil (clomipramine)
  - Lofepramine
  - Gamani (lofepramine)
  - Lomont (lofepramine)
  - Mianserin
  - BoVIdon (mianserin)
  - Norval (mianserin)

### 4 Antipsychotic – traditional
- Chlorpromazine
  - cpz-chlorpromazine
  - Largactil (chlorpromazine)
  - Haloperidol
  - Haldol (haloperidol)
  - Serenace (haloperidol)
  - Fluphenazine decanoate
  - Fluphenazine
  - Mocdecate (fluphenazine)
  - Moditen tablet (fluphenazine)
  - Moditen enanthate (fluphenazine)
  - Flupenthixol
  - Flupenthixol (flupenthixol)
  - Depxol (flupenthixol)
  - Fluoxal (flupenthixol)
  - Zuclopenthixol
  - Cloxipixi (Zuclopenthixol)
  - Tixipixi (cloxipixi)
  - Loxapine
  - Loxapac (loxapine)
  - droperidol
  - Dropleptan (droperidol)
  - trifluoperazine
  - Stelazine (trifluoperazine)
  - thioridazine
  - Mellaril (thioridazine)

### 5 Antipsychotic – second generation
- Quetiapine
- Seroquel (quetiapine)
- Risperdol
- Risperdal (risperdolone)
- olanzapine
  - Zyprexa (olanzapine)
  - aripiprazole
  - Abilify (aripiprazole)
  - amisulpride
  - Solian (amisulpride)
  - clozapine
  - Clozaril (clozapine)

### 6 Sedatives and hypnotics
- Diazepam
  - Diazepam product
  - Valium tablet (diazepam)
  - Valium syrup (diazepam)
  - Valium suppl (diazepam)
  - temazepam
  - Normison (temazepam)
  - Euhypnos (temazepam)
  - Zopiclone
  - Zimovane (zopiclone)
  - zaleplon
  - Sonata (zaleplon)
  - zolpidem
  - Stilnox (zolpidem)
  - nitrazepam
  - Mogadon (nitrazepam)
  - Nitrados (nitrazepam)
  - Remnos (nitrazepam)
  - Sonnite (nitrazepam)
  - Nocasted (nitrazepam)
  - Suren (nitrazepam)
  - Unisomna (nitrazepam)
  - flunitrazepam
  - Rohypnol (flunitrazepam)
  - triazolam
  - Halcion (triazolam)
Table DS1  Logistic regression analysis of cardiometabolic disease associated with mood disorder

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<th>Women</th>
<th>Men</th>
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<td>Fully adjusted\textsuperscript{b}</td>
<td>Partially adjusted\textsuperscript{a}</td>
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<td>P</td>
<td>OR (95% CI)</td>
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<td>1 (Reference)</td>
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<td>1.29 (1.25-1.33)</td>
<td>&lt;0.001</td>
<td>1.21 (1.17-1.26)</td>
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<tr>
<td>Bipolar</td>
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<td>&lt;0.001</td>
<td>1.55 (1.32-1.83)</td>
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<td></td>
<td>1 (Reference)</td>
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<td>0.002</td>
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<td>1 (Reference)</td>
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<tr>
<td>Depression</td>
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<td>1.19 (1.14-1.24)</td>
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<tr>
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<tr>
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<tr>
<td>Bipolar</td>
<td>1.69 (1.30-2.20)</td>
<td>&lt;0.001</td>
<td>1.17 (0.93-1.58)</td>
</tr>
</tbody>
</table>

\textsuperscript{a}Partially adjusted for age, gender, deprivation and ethnicity.  
\textsuperscript{b}Fully adjusted for age, gender, deprivation, ethnicity, body mass index, smoking status, alcohol consumption and psychotropic medication.