The prevalence of pain in bipolar disorder:

A systematic review and large scale meta-analysis

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**Running title:** Clinical Pain in bipolar disorder
Abstract

Objective

To conduct a meta-analysis investigating the prevalence of pain in people with bipolar disorder (BD).

Method

A systematic review and random effects meta-analysis searching major electronic databases from inception till 01/2014 in accordance with the PRISMA statement. We included articles reporting quantitative data on the prevalence of pain in people with BD with or without a healthy control group. Two independent authors conducted searches, extracted data and completed methodological quality assessment.

Results

Twenty two cross-sectional studies were included, representing 12,375,644 individuals (BD n = 171,352, n controls = 12,204,292). The prevalence of pain in people with BD was 28.9% (95% CI = 16.4 - 43.4%, BD n = 171,352). The relative risk (RR) of pain in BD compared to controls was 2.14 (95% CI = 1.67 - 2.75, n = 12,342,577). The prevalence of migraine was 14.2% (95% CI = 10.6 - 18.3%, BD n = 127,905) and the RR was 3.30 (95% CI = 2.27 - 4.80, n = 6,732,220). 23.7% (95% CI = 13.1 - 36.3%, n = 106,214) of people with BD experienced chronic pain. Age, percentage of males, methodological quality and method of BD classification did not explain the observed heterogeneity.

Conclusion

People with BD experience significantly increased levels of pain (particularly chronic pain and migraine). The assessment and treatment of pain should form an integral part of the management of BD.

Key words: Pain, chronic pain. Migraine, bipolar disorder, severe mental illness
Summations

- The pooled prevalence of clinical pain in people with bipolar disorder is approximately 28.9%, whilst 23.7% and 14.2% are affected by chronic pain and migraines respectively.
- Compared to the general population, people with bipolar disorder are at significantly increased risk of reported clinically relevant pain (Relative Risk= 2.14) and migraine (RR = 3.30).
- Since pain has a range of deleterious impacts on an individual’s health and quality of life and may worsen psychiatric symptoms, we recommend that pain assessment and treatment should form part of the routine care of people with bipolar disorder.

Considerations

- There was inconsistency in the assessment methods used to measure pain across the studies.
- There was considerable heterogeneity in each of the pooled analysis that could not be explained by mean age, percentage of males, method of diagnosing bipolar disorder and methodological quality of the included studies.
- There was insufficient information to determine the influence of the phase of illness and polarity as well as acuity of bipolar symptoms on the observed results.
Introduction

Pain has a deleterious impact on an individual’s health and wellbeing (1) and common painful conditions, such as chronic musculoskeletal disorders, contribute to a significant number of years lived with disability across the globe (2). Chronic pain in particular is associated with greatly reduced quality of life, difficulties with activities of daily living (ADL), and often has a negative impact on an individual’s emotional and mental health (3). A substantial body of literature suggests that those with chronic pain have higher rates of depressive and anxiety symptoms than those without chronic pain (4-6).

Despite this, the prevalence of chronic pain in persons with severe mental illness (SMI) has received little attention (7, 8). This is surprising as persons with SMI such as schizophrenia and bipolar disorder have a highly increased risk for a plethora of painful physical illnesses including cardiopulmonary diseases, metabolic diseases, bone disorders, viral infections, and cancer (9-12). In addition, pain in people with SMI is also associated with a worsening of psychiatric symptoms (7). Despite this increased risk of severe co-morbid physical illnesses, most persons with SMI do not receive adequate physical healthcare provision and treatment (13-15). Mental health specialists report barriers limiting their ability to treat physical co-morbidity and people with SMI are less likely to recognize or monitor co-occurring medical conditions than the general population (16, 17). Additionally many healthcare professionals fail to take people with severe mental illness seriously when they report physical health problems (18). When compared to those without SMI, persons with SMI appear to have an increased likelihood of experiencing conditions that cause pain whilst at the same time having a lower likelihood of receiving adequate care to manage it (9, 10).

A recent systematic review established that people with schizophrenia, who have been known to have a higher pain threshold for pain than the general population, have a lower prevalence of pain than people with other psychiatric disorders, particularly compared to those with bipolar disorder (19). However, to date, no systematic review or meta-analysis of pain in individuals
with bipolar disorder exists, despite the fact this group appears to be particularly more likely to experience chronic pain and less likely to seek medical help (8). In fact, people with bipolar disorder reported almost 4 pain complaints at any one time (20). Moreover, people with bipolar disorder who are treatment adherent report statistically lower levels of pain than their non-treatment adherent counterparts (21). Clearly, a better understanding of the risk and burden of pain is an important step toward improving clinical outcomes for individuals with bipolar disorder.

Aims of the study

In recognition of the potential for pain to be problematic for people with bipolar disorder, the paper had the following two aims: (1) to establish the prevalence of pain and its moderators in people with bipolar disorder, and (2) to compare the prevalence of pain in bipolar disorder with general population controls.
Method

This systematic review was conducted according to the PRISMA statement (22) following a predetermined, but unpublished protocol.

Inclusion and exclusion criteria

Studies were eligible that fulfilled the following criteria: (1) inclusion of participants with bipolar disorder, diagnosed according to diagnostic criteria (e.g. DSM IV (23) or ICD 10 (24)), a valid screening measure (e.g., Alcohol Use Disorder and Associated Disabilities Interview Schedule—DSM-IV Version) or through medical record review. When we encountered studies containing groups of mixed participants (e.g., with major depressive disorder), we contacted the authors up to two times over a month period to ascertain the variables of interest in bipolar disorder subjects. If these data were not available, we excluded the study. (2) Reporting of the prevalence of pain (of any type) or assessment of pain with a continuous measure with or without comparison to a control group that did not have a mental illness. When a study measured pain with a continuous measure, but did not specify prevalence rates with a cut-off point, we contacted the authors up to two times to obtain this information.

We did not place a language restriction upon our searches. If we came across studies that reported data from the same sample at different time points, we used the most recent data and/or the largest data set. We excluded studies that (1) reported pain as an adverse event of a drug trial (e.g., for headache), (2) reported the prevalence of bipolar disorder in a sample of patients who all had pain (no other comorbidities were excluded), or (3) in which the pain was experimentally induced. When we encountered studies without a control group that assessed pain in a sample with a continuous measure (e.g., SF 36 bodily pain scale, (25)), but did not have a cut-off to determine the prevalence of pain, we excluded the study if the authors did not respond to requests for additional data.

Information sources
Two reviewers (BS, DV) independently conducted searches on Academic Search Premier, MEDLINE, EMBASE, Psychology and Behavioral Sciences Collection, PsycINFO, SPORTDiscus, CINAHL Plus and Pubmed. In addition, the reference lists of all eligible articles and recent systematic reviews of the literature were scanned to assess eligibility of additional studies.

**Searches**

Two independent reviewers (BS, DV) employed the predetermined search strategy using the key words ‘bipolar disorder’ and ‘pain’ or ‘pain perception’ or ‘pain management’ or ‘pain measurement’ or ‘musculoskeletal pain’ or ‘pain intensity’ or ‘chronic pain’ or ‘neuropathic pain’ or ‘pain*’.

**Study Selection**

After the removal of duplicates, two independent reviewers (BS, DV) screened the titles and abstracts of all potentially eligible articles. Both authors applied the eligibility criteria, and a list of full text articles was developed through consensus. Two reviewers (BS, DV) then considered the full texts of these articles and the final list of included articles was reached through consensus.

**Data Extraction**

Two authors (BS, DV) independently conducted data extraction using a predetermined form. The data collected from each article included: study design, geographical location, bipolar sample and control sample characteristics (number, % male, mean age), bipolar diagnosis method, method of pain assessment (including site, severity, and interference of pain where available) and the prevalence of pain in people with bipolar disorder and controls as defined by the authors.

**Methodological quality assessment**

Two independent authors (BS, DV) completed methodological quality assessment of included articles using the Newcastle Ottawa Scale (NOS; (26)). Due to the anticipated paucity of data, we also
included studies without a control group. These studies were considered as case control studies for the purposes of methodological assessment in accordance with a previous review (27). The NOS is utilised to assess the methodological quality of non-randomised trials and has acceptable validity and reliability (26). The assessment tool focuses on three main methodological features: (1) the selection of the groups, (2) the comparability of the groups and (3) the ascertainment of the outcome of interest. The NOS can be modified and we adapted the NOS to take into account age and gender as comparability measures and considered pain assessment in the exposure category. Studies are given a score from 0-9, with a score of 5 or greater being indicative of satisfactory methodological quality. We anticipated studies without a control group would score below this and present their results with due consideration.

**Meta-analysis**

We pooled individual study data using DerSimonian- Laird proportion method (28). Our predetermined protocol stipulated that heterogeneity would be assessed with the Cochran Q statistic (29). Since we found significant heterogeneity (Cochran Q = 66988.29 (df = 24) P < 0.0001) a random effects meta-analysis was employed using StatsDirect. We calculated the relative risk (RR) to investigate the differences in pain between those with bipolar disorder and members of the general population when there were three or more studies (Aim 2). When possible, we conducted subgroup analyses to investigate the prevalence of migraine and chronic pain since the literature has suggested these are prevalent in people with bipolar disorder (8). In order to investigate sources of heterogeneity, we conducted moderator analysis with mean age, percentage of males, NOS score and the method of bipolar disorder classification (comparing DSIM, ICD or any other screening measure). We assessed publication bias with a visual inspection of funnel plots, yet gave priority to quantitative testing through the Begg-Mazumdar Kendall's tau (30) and Egger bias tests (31).
Results

Study selection

The original search yielded 2,713 potential hits which were reduced to 2,319 after the removal of duplicates. At the eligibility screening stage, a total of 72 articles were deemed potentially eligible and full texts were obtained and reviewed by two authors. In total, 50 articles were excluded with reasons and 22 articles met the eligibility criteria and were included in the review (8, 32-52). The full search strategy including reasons for exclusion is presented in figure 1.

Inset figure 1 about here

Study characteristics

In total 171,352 people with bipolar disorder and 12,204,292 general population controls (total sample size = 12,375,644) were included in the 22 meta-analysed studies. Details of the included studies are presented in table 1. All of the studies adopted a cross-sectional measurement of pain and 7 of these (n with bipolar disorder = 138,285; (8, 33, 38, 41, 44, 49, 52)) had a control group without a mental illness. The sample size of persons with bipolar disorder across the studies ranged from 10 (40) to 96,186 (8) and the control populations ranged from 32,333 (44) to 4,247,684 (8). The mean age of participants with bipolar disorder ranged from 39 (34) to over 65 years (51).

Methodological quality

The NOS summary score for each article is presented in table 1. All seven studies that had a control group scored high (mean NOS score 7.2±0.48) and were considered good quality. The 15 studies that did not have a control group all scored lower than 5 on the NOS, which was attributable to the absence of a control group; these studies scored zero (out of a possible 5 points) in the areas that compare the bipolar and control groups on selection, comparability, and exposure.

Measurement and location of pain in the bipolar populations
A range of different types of pain were considered. The most commonly investigated pain was headache/migraine (8, 32, 33, 35, 39, 40, 42-44, 47, 52) whilst six studies investigated chronic pain (8, 34, 45, 48-50). A wide range of methods were employed to ascertain pain in people with bipolar disorder and are presented in table 1.

*Prevalence of pain in persons with bipolar disease*

In total, 25 types of pain were investigated and the pooled prevalence of pain was 28.9% (95% CI: 16.4-43.4%, n=171,352, Cochran Q = 66988.29 (df = 24) P < 0.0001, figure 2a). The funnel plot was asymmetrical (figure 2b), however, both the Begg-Mazumdar (Kendall's tau = -0.013; P = 0.908) and Egger bias (Kendall's tau = 11.51; P = 0.4897) tests did not demonstrate any evidence of publication bias. Next, we pooled the prevalence of pain using only one pain measurement from each of the 22 studies, thus including only the highest prevalence of pain from 3 studies that contained data on pain at two sites (33, 35, 43). The prevalence of clinical pain across 22 studies was 28.4% (95% CI = 15.0-44.1%, Cochran Q =66477.17 (df = 21) P < 0.0001). Within this analysis, there was also no evidence of publication bias (Egger: bias = 12.44, P = 0.5176, Begg-Mazumdar: Kendall's tau = 0.021, P = 0.9113).

*Moderators of the prevalence of pain in people with bipolar disorder*

Ten studies (32-36, 38, 41, 42, 45, 49) had sufficient data on mean age, percentage of males and bipolar diagnosis method to enable moderator analyses. The moderator analyses demonstrated that mean age (b1 = -0.038, z = -0.311, P = 0.75), % male (b2 = -0.074, z = -1.013, P = 0.311) and method of diagnosing bipolar disorder (b3 = -0.0935, z = -0.092, P = 0.92) did not explain the heterogeneity in the prevalence of pain. We investigated the effect methodological quality (NOS score) on the prevalence of pain across the 22 studies and this suggested that a low NOS score was associated with a high prevalence of pain but this did not reach statistical significance (b1=0.532,
Lastly, we investigated the influence of the method of bipolar disorder diagnosis on the prevalence across all studies and this demonstrated that the classification used to diagnose bipolar disorder had no significant effect on the prevalence of pain (b1=0.310, z = 0.524, P = 0.59).

Comparing the prevalence of pain in people with bipolar disorder versus control groups
In each of the 7 studies with a control group, persons with bipolar disease consistently reported a higher prevalence of pain than the comparison group. One study (33) provided pain data for two different types of pain and was corrected for multiple comparisons in the pooled analysis. In total, data from 12,342,577 unique individuals (n with bipolar disorder =138,285 and control n= 12,204,292) indicated that the relative risk of pain in people with bipolar disorder was 2.14 (95% CI = 1.67 - 2.75 , Chi-square = 36.623 (df = 1) P < 0.0001; Cochran Q = 1078.49 (df = 7) P < 0.0001). The results from the meta-analysis are presented in figure 3. The funnel plot of the 7 included studies was not symmetrical indicating possible publication bias. However, the Eggers test (10.931 P = 0.013), but not the Begg Mazumdar: test (Kendall's tau = 0.14; P = 0.7195) showed evidence of publication bias.

Insert figure 3 about here

Pooled prevalence of Migraine in people with bipolar disorder
We also calculated the pooled prevalence of migraine in 127,905 individuals across 9 studies (8, 32, 39, 40, 42-44, 47, 52) and this yielded a prevalence of 14.2% (95% CI = 10.6% - 18.3%; Cochran Q = 1080.29 (df = 8) P < 0.0001).

Comparing the prevalence of migraine in people with bipolar disorder versus control groups
It was possible to pool the data from 3 comparative studies (8, 44, 52) involving 6,732,220 unique individuals (n with bipolar disorder=126,956, n controls = 6,605,264). The RR was 3.30 (95% CI=2.27-4.80, Chi-square test = 39.408 (df = 1) P < 0.0001).
Pooled prevalence of chronic pain in people with bipolar disorder

It was possible to calculate the pooled prevalence of chronic pain in 106,214 individuals with bipolar disorder across 6 studies (8, 34, 45, 48-50). The pooled prevalence of chronic pain was 23.7% (95% CI = 13.1-36.3, Cochran Q= 2200.77 (df = 5) P < 0.0001). Only two comparative studies (8, 49) contained data on chronic pain and it was therefore not possible to meta-analyse these data.
Discussion

To our knowledge this is the first systematic review and meta-analysis investigating the prevalence of pain in people with bipolar disorder. In this large review involving 171,352 persons with bipolar disorder and 12,204,292 controls, we found that a substantial proportion of patients with bipolar disorder reported clinically relevant levels of pain. The overall pooled analysis of pain in people with bipolar disorder was 28.9% and the relative risk was over double for people with bipolar disorder compared to members of the general population. In terms of specific types of pain, the pooled prevalence of chronic pain was high with almost one in four (23.7%) being affected. In addition, migraine affected one in seven (14.2%) persons with bipolar disorder and the comparative analysis demonstrated that people with bipolar disorder are over three times more likely to experience migraines than members of the general population.

Increased levels of pain in persons with bipolar disorder may be explained by several mechanisms. For instance, bipolar disorder and migraine appear to share some specific polymorphisms, with the KIAA0564 gene being particularly implicated, thus suggesting a close association (53, 54). Also, people with bipolar disorder have an increased prevalence of depression (8, 36), and depression has been associated with increased physical complaints, and, possibly, greater pain sensitivity (55), opposite to findings in schizophrenia (19). For example, neuroimaging studies in major depressive disorder indicate that heightened amygdala activity, in part, explains the high comorbidity of pain and depression when these conditions become chronic (56). However, due to limitations in the available data, we could not investigate the influence of depressive symptoms on the observed results. Other studies have suggested serotonergic and noradrenergic pathway involvement (7, 57). In addition, specific neuroinflammatory mechanisms responsible for an elevated risk of painful physical comorbidity in people with bipolar disorder may contribute to the higher levels of observed pain (58). Previous research (59) has found that migraine and bipolar disorder symptoms are closely related and the presence of migraine can influence pain perception.
Since we found that 14.2% of people with bipolar disorder experienced migraine, this could have influenced the variance in the prevalence of pain. Lastly, recent findings (60) also suggest that limited cognitive flexibility and memory capacities may be linked to the mechanisms of pain chronicity and probably also to its neuropathic quality. This may imply that people with bipolar disorder who are known to have deficits in executive functioning or memory have a greater risk of pain chronicity after a painful event. This seems particularly pertinent given the fact that we found across 106,214 individuals with bipolar disorder that almost one in four is affected by chronic pain.

**Clinical implications**

The results of this review are concerning since pain and in particular chronic pain in people with bipolar disorder is associated with impaired recovery (45), greater functional incapacitation (44, 61), lower quality of life (8), and increased risk of suicide compared to people without pain (62). Since bipolar disorder is already associated with a greatly increased risk of suicide (63), it is imperative that this population receives adequate pain assessment and management (36). A central component to this is the training and education of psychiatrists who are in a critical place to oversee the pharmacological management of pain (7). We advocate that systematic assessment of pain should be undertaken as part of the management of bipolar disorders, and that pain should be monitored during the course of treatment. Equally, healthcare professionals dealing with pain should consider mental health complications. Previous work suggests clinicians are more likely to attend to pain than mental distress (64). The potential benefits of early identification and treatment of pain may not only include a reduction in pain and of its impact on the individual, but may also extend to a reduction of health-care costs and improvement of mental health outcomes.

Of great concern are the high levels of chronic pain experienced by people with bipolar disorder. A better understanding of the association of bipolar disorder and chronic pain could help limit harmful/adverse pharmacological side effects. For instance, in the general population chronic
pain is often managed with tri-cyclical antidepressants (65), yet prescription of such medication to a person with bipolar disorder may inadvertently trigger a manic phase of illness if prescribed in the absence of a mood stabilizer (66). Commonly used analgesic medications also need careful consideration. For instance, there is sound evidence that non-steroidal anti-inflammatory medications can increase serum lithium levels, impairing renal lithium excretion and possibly eliciting lithium toxicity (67). Similarly, some stronger analgesic medications such as opioids may have mood altering qualities increasing the risk of eliciting a manic episode (68).

Limitations of the review

Several limitations, especially of the included literature need to be considered when interpreting the results of our review. First, bipolar disorder is a complex and heterogeneous disorder, and reporting of pain likely varies according to different phases, polarity and acuity of the disease. The paucity of information regarding these illness characteristics made it impossible to systematically evaluate their effects on pain prevalence in patients with bipolar disorder. In addition, the perception and therefore prevalence of pain is known to vary according to the type of bipolar disorder (I or II; (59)) but due to limitations in the data we were not able to disentangle this relationship. In addition, gender may also cause some variance, but our moderator analysis did not elucidate any evidence of a gender effect. Second, all of the included studies utilised a cross-sectional measurement of pain and did not correlate pain with mood state or severity of symptoms. Therefore, prospective longitudinal studies that assess pain prevalence and severity over time and in relationship to mood symptoms and treatments are essential. Third, our results may have been suspect to Berkson’s bias, which states that clinical samples are more impaired and experience more pain than non-clinical samples due to self-referral to a clinical setting. Berkson’s bias has been observed in the mood dimensions of bipolar disorder (69) and may account for an underreporting within the pooling of epidemiological data. Fourth, none of the included studies used a validated pain assessment scale and subsequently information about the severity, location, variability, and
interference of pain during activities is lacking. Fifth, all of the meta-analytic results were heterogeneous and some demonstrated a degree of publication bias. In our moderator analysis, we were not able to explain the heterogeneity with mean age, % males, or the methodological quality of method of classification of bipolar disorder. This finding demonstrates that unknown/unmeasured factors contribute to the observed heterogeneity. Regarding publication bias, the funnel plot for the main analysis (figure 2b) appeared asymmetrical, yet the quantitative investigation of bias did not demonstrate any evidence to support this. This discrepancy may be due to the fact that there is a trend for publication bias, but its magnitude is insufficient to reach statistical significance according to the Eggers test or Begg-Mazumdar test. In addition, the comparative analysis (figure 3) demonstrated some publication bias with the Eggers test, but this finding should be interpreted with caution due to the low number of studies (<10 (70)). Sixth, there was insufficient information about psychotropic and analgesic medication within the bipolar disorder cohorts to enable statistical investigation of these variables on the observed results. Future research should seek to investigate the influence of psychotropic and analgesic medications on pain and particular attention should be paid to the prevalence of pain in people with bipolar disorder who are drug naïve. In the same way, future research should investigate the role of psychiatric co-morbidities including anxiety and substance use disorders on the prevalence of clinical pain in these patients. Finally, we included 15 studies that received low methodological quality ratings. However, the low methodological quality ratings were due to the absence of a control group, and the moderator analysis demonstrated that these studies had no significant effect on the observed results. Despite the aforementioned, higher levels of pain were reported consistently among people with bipolar disorder than in the comparison groups.

**Future research**

It is essential that future research seeks to clearly assess pain characteristics including noting the site, severity, variability and chronicity. There were insufficient data to analyse these pain
characteristics in our meta-analysis. In addition, only one study (8) measured psychogenic pain and it would be important to investigate if this differs from physiological pain in people with bipolar disorder. An important question, also unaddressed, is what is the impact of comorbid pain, particularly chronic pain, on daily activities? It is likely that chronic pain amplifies the effect of bipolar disorder on disability and reduced quality of life. Future prospective studies should be conducted in order to truly capture the prevalence of pain and disentangle its impact and contributing factors. Such research should establish how pain impacts on a person’s mental health and wellbeing, with longitudinal studies being most important. Future research should also explore the extent to which those with bipolar disorder are more or less responsive to behavioral, pharmacological, and non-pharmacological treatments for pain. For example, studies have not yet examined the impact on pain of anti-epileptic medications such as lamotrigine, valproate and topiramate among persons with bipolar disorder. In addition, in the general population the promotion of physical activity is a key factor preventing the onset of chronic pain but is also encouraged to treat it (71) and many people with chronic pain are inactive (72). However, research (73) has established that most people with bipolar disorder are sedentary. Therefore, strategies to encourage people with bipolar disorder to become active that do not exacerbate their pain are likely to be key in the prevention and management of pain and physical therapists can lead this process (74). In addition, the barriers and facilitators to pain management should be explored in people with bipolar disorder with emphasis on the perspective of the patient and the treating multidisciplinary team. Lastly, within our review, there were limited studies assessing pain in patients with bipolar disorder and those with other psychiatric conditions, making it impossible to directly compare the prevalence of clinical pain in people with bipolar disorder and other psychiatric diagnosis. More research is required to directly compare clinical pain across different psychiatric disorders.

Conclusion
Almost 30% of persons with bipolar disorder experience clinically relevant pain, which was twice as common compared to general population controls. Chronic pain was prevalent affecting almost one in four people, and migraine was over 3 times as common than in the general population. Pain has a range of adverse and deleterious impact upon the individual and may impede recovery, reduce quality of life and have adverse effects on psychiatric symptoms. Therefore, it is essential that treating psychiatrists and the wider multidisciplinary team seek to provide adequate assessment and treatment of pain in people with bipolar disorders.
Conflict of Interest

BS, AJM, AS, SR and LE have no conflicts of interest to declare.

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MDH declares that he has been a consultant for, received grant and/or research support and honoraria from, and been on the speakers’ bureaus and/or advisory boards of the following companies: Astra Zeneca, Bristol-Myers Squibb, Eli Lilly, Janssen-Cilag, Lundbeck, Pfizer, Sanofi Aventis and Takeda.

CC has been a consultant and/or advisor to or has received honoraria from: Actelion, Alexza; Bristol-Myers Squibb, Cephalon, Eli Lilly, Genentech, Gerson Lehrman Group, IntraCellular Therapies, Janssen/J&J, Lundbeck, Medavante, Medscape, Merck, Otsuka, Pfizer, ProPhase, Roche, Sunovion, Takeda, Teva, and Vanda. He has received grant support from BMS, Janssen/J&J, Novo Nordisk A/S and Otsuka.

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<table>
<thead>
<tr>
<th>Study No</th>
<th>Location and design</th>
<th>Bipolar diagnosis</th>
<th>Participant with BD characteristics</th>
<th>Control participant characteristics</th>
<th>NOS score</th>
</tr>
</thead>
<tbody>
<tr>
<td>8</td>
<td>USA</td>
<td>ICD-9-CM</td>
<td>N=96,186</td>
<td>N=4,247,684</td>
<td>7</td>
</tr>
<tr>
<td></td>
<td>Cross sectional design collecting data over 1 year</td>
<td>Derived from patient electronic medical records</td>
<td>Age &lt;35-&gt;80 years Males 81,757 (85.0%) No data on BD severity or medication</td>
<td>Age &lt;35-&gt;80 years Males 3,882,806 (91.4%)</td>
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<tr>
<td>32</td>
<td>Brazil</td>
<td>DSM IV</td>
<td>N= 339 split in two groups Migraine (n=115) 41.6 ±11.20 years Males 16 (17.4%) None migraine (n=224) 41.5 ±12.32 years Male 60 (26.7%).</td>
<td>No control group</td>
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<tr>
<td>33</td>
<td>USA</td>
<td>ICD-9</td>
<td>N= 3,557</td>
<td>N=726,262</td>
<td>7</td>
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<tr>
<td></td>
<td>Cross sectional study collecting data over 5 year period</td>
<td></td>
<td>39.3 ±11.8 years Males 1395 (39.2%). BD more likely have substance use disorder OR 2.92; (95% CI, 2.59 –3.29) &amp; alcohol use disorder AOR 19.63; (95% CI, 17.59-21.90)</td>
<td>37.7±12.8 years Male 345,146 (47.5%)</td>
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<tr>
<td>34</td>
<td>USA</td>
<td>Composite International Diagnostic Interview Version 3.0</td>
<td>N=740 39 (±10.6) years Males 414 (56%) No data on BD severity or medication</td>
<td>No control group</td>
<td>3</td>
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</table>
Table 1: Included study characteristics and methodological quality

<table>
<thead>
<tr>
<th>Study</th>
<th>Location and design</th>
<th>Bipolar diagnosis</th>
<th>Participant with BD characteristics</th>
<th>Control participant characteristics</th>
<th>NOS score</th>
</tr>
</thead>
</table>
| 35    | Australia           | DSM IV            | N=67  
Males 35.8% (n=24)  
40.4 (±13.5) years  
**BDRS= 11.5 ± 9.3**| No control group | 3         |
| 36    | Spain               | DSM-IV-TR         | N=121  
50.7 years (± 12.3)  
Males 45 (37.8%)  
50.7% had suicidal ideation| No control group | 3         |
| 37    | Italy               | DSM-IV-TR         | N= 248  
Demographic information not available| No control group | 3         |
| 38    | USA                 | AUDADIS-IV        | N=883  
36.9±0.3 years  
Males 380 (43%)  
N= 42,210  
45.4±0.1 years  
Males 20,261 (48%)| 7         |
| 39    | UK                  | Not stated        | N=169  
Demographic information not available| No control group | 3         |
| 40    | Taiwan              | DSM-IV-TR         | N=10  
Demographic data not available| No control group | 3         |
| 41    | USA                 | ICD-9 medical records | N=4,310  
Males 3879 (90%)  
53±13 years  
**BD more likely have SUD (p<.0001)**| N=3,408,760  
Males 3,067,884 (90%)  
58 years| 8         |
Table 1 Included study characteristics and methodological quality

<table>
<thead>
<tr>
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<th>Location and design</th>
<th>Bipolar diagnosis</th>
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<th>NOS score</th>
</tr>
</thead>
<tbody>
<tr>
<td>42</td>
<td>USA</td>
<td>DSM IV</td>
<td>N=111 44.8±13.2 years Males 35 (32.4%)</td>
<td>No control group</td>
<td>3</td>
</tr>
<tr>
<td>43</td>
<td>Italy</td>
<td>DSM III</td>
<td>N= 30 Demographic data not available</td>
<td>No control group</td>
<td>3</td>
</tr>
<tr>
<td>44</td>
<td>Canada</td>
<td>CIDI</td>
<td>N= 938 Age 25-64 years Males 436 (46.4%)</td>
<td>N=32,333 Demographic information not available</td>
<td>7</td>
</tr>
<tr>
<td>45</td>
<td>USA</td>
<td>ICD-9 criteria</td>
<td>N= 384 42.07±11.3 years Males 128 (33.3%)</td>
<td>No control group</td>
<td>3</td>
</tr>
<tr>
<td>46</td>
<td>Australia</td>
<td>ICD 9</td>
<td>N= 27 10 males (37.0%)</td>
<td>No control group</td>
<td>3</td>
</tr>
<tr>
<td>47</td>
<td>Canada</td>
<td>DSM IV</td>
<td>N=296 with BD 1 and BD 2 49.8 ± 12.7 years % males not available</td>
<td>No control group</td>
<td>3</td>
</tr>
<tr>
<td>48</td>
<td>South Korea</td>
<td>DSM-IV</td>
<td>N=190 Demographic data not available</td>
<td>No control group</td>
<td>3</td>
</tr>
</tbody>
</table>
### Table 1 Included study characteristics and methodological quality

<table>
<thead>
<tr>
<th>Study</th>
<th>Location and design</th>
<th>Bipolar diagnosis</th>
<th>Participant with BD characteristics</th>
<th>Control participant characteristics</th>
<th>NOS score</th>
</tr>
</thead>
<tbody>
<tr>
<td>49</td>
<td>Scotland</td>
<td>GP databases</td>
<td>N=2,582</td>
<td>N=1,421,796</td>
<td>8</td>
</tr>
<tr>
<td></td>
<td>Cross sectional</td>
<td></td>
<td>54.5 years</td>
<td>47.9 years (p&lt;0.001)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>retrospective analysis</td>
<td></td>
<td>Males 1,021 (39.5%)</td>
<td>Males 698,408 (49.1%)</td>
<td></td>
</tr>
<tr>
<td>50</td>
<td>Singapore</td>
<td>CIDI 3.0</td>
<td>N=93</td>
<td>Not reported</td>
<td>3</td>
</tr>
<tr>
<td></td>
<td>Cross sectional</td>
<td></td>
<td>Age 18-65&gt; years</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Males 47 (50.5%)</td>
<td></td>
<td></td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>66% BD I -had severe or moderate manic/hypomanic &amp; 100% respondents with BP-II reported mild clinical severity on the YMRS</td>
<td></td>
<td></td>
</tr>
<tr>
<td>51</td>
<td>United States</td>
<td>ICD 9</td>
<td>N= 24206</td>
<td>No control group</td>
<td>3</td>
</tr>
<tr>
<td></td>
<td>Cross sectional</td>
<td></td>
<td>All &gt;65 years nursing home residents</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>No specific data on demographics</td>
<td></td>
<td></td>
</tr>
<tr>
<td>52</td>
<td>United States</td>
<td>ICD 9</td>
<td>N=27,054</td>
<td>N=2,325,247</td>
<td>8</td>
</tr>
<tr>
<td></td>
<td>Cross sectional</td>
<td></td>
<td>Demographics not available</td>
<td>Demographics not available</td>
<td></td>
</tr>
<tr>
<td></td>
<td>retrospective analysis</td>
<td></td>
<td></td>
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</tr>
</tbody>
</table>

**Key:** BD = bipolar disease, GP=General practitioner, ICD-9-CM=International Classification of Diseases, 9th Revision, Clinical Modification, DSM-IV-TR=Diagnostic and Statistical Manual of Mental Disorders Fourth Edition Text Revision, CIDI 3.0 = World Mental Health Composite International Diagnostic Interview version 3.0, AUDADIS-IV=NIAAA Alcohol Use Disorder and Associated Disabilities Interview Schedule-DSM-IV Version, NOS = Newcastle Ottawa Scale, BDRS = Bipolar Depression Rating Scale, YMRS= Young Mania Rating Scale
Table 2 Results of Pain in included studies

<table>
<thead>
<tr>
<th>Study</th>
<th>Type of pain</th>
<th>Method of pain assessment/ascertainment</th>
<th>Pain results bipolar disorder</th>
<th>Pain results in control</th>
<th>Other results</th>
</tr>
</thead>
<tbody>
<tr>
<td>8</td>
<td>Arthritis</td>
<td>ICD-9-CM based on electronic patient records</td>
<td>Total n=96,186 in BD sample</td>
<td>Total n= 4,247,684 in control sample</td>
<td>OR comparing BD and controls:</td>
</tr>
<tr>
<td></td>
<td>Back pain</td>
<td></td>
<td>Any pain 61.3% (n=58,983)</td>
<td>Any pain 42.3% (n=1,795,600)</td>
<td>Any pain OR 2.17 (CI 2.14-2.19)*</td>
</tr>
<tr>
<td></td>
<td>Chronic Pain</td>
<td></td>
<td>Arthritis 45.3% (n=43,595)</td>
<td>Arthritis 32.2% (n=1,365,901)</td>
<td>Arthritis OR 1.75 (CI 1.73-1.77)*</td>
</tr>
<tr>
<td></td>
<td>Migraine</td>
<td></td>
<td>Back pain 33.5% (n=32,264)</td>
<td>Back pain 17.0% (n=721,372)</td>
<td>Back pain OR 2.47 (CI 2.43-2.50)*</td>
</tr>
<tr>
<td></td>
<td>Other Headache</td>
<td></td>
<td>Chronic Pain 3.4% (n=3,316)</td>
<td>Chronic Pain 0.7% (n=27,758)</td>
<td>Chronic Pain OR 5.43 (CI 5.23-5.63)*</td>
</tr>
<tr>
<td></td>
<td>Psychogenic</td>
<td></td>
<td>Migraine 4.9% (n=4,677)</td>
<td>Migraine 1.1% (n=46,015)</td>
<td>Migraine OR 4.67 (CI 4.53-4.82)*</td>
</tr>
<tr>
<td></td>
<td>Neuropathic</td>
<td></td>
<td>Other Headache 6.7% (n=6,419)</td>
<td>Other headache 2.0% (n=86,126)</td>
<td>Other headache OR 3.46 (CI 3.37-3.55)*</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Psychogenic pain 0.9% (n=833)</td>
<td>Psychogenic pain 0.1% (n=3,646)</td>
<td>Psychogenic pain OR 10.17 (CI 9.43-10.96)*</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Neuropathic pain 5.4% (n=5,180)</td>
<td>Neuropathic pain 3.7% (n=156,393)</td>
<td>Neuropathic pain OR 1.49 (CI 1.45, 1.53)*</td>
</tr>
<tr>
<td>32</td>
<td>Migraine</td>
<td>Physician diagnosis</td>
<td>33.9% (n = 115) had migraines</td>
<td>No Control Group</td>
<td>Migraine group higher nr of psychiatric comorbidity (72.6%) vs. non migraine group (47.4%) p&lt;0.001</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Migraine group more likely anxiety disorder (p&lt;0.001) and depressive polarity</td>
</tr>
<tr>
<td>Study</td>
<td>Type of pain</td>
<td>Method of pain assessment/ascertainment</td>
<td>Pain results bipolar disorder</td>
<td>Pain results in control</td>
<td>Other results</td>
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</tr>
<tr>
<td>33</td>
<td>LBP</td>
<td>Elixhauser Comorbidity Index</td>
<td>25.8% (n=919/3557) had LBP</td>
<td>13.3% (n=96,201/726,262) had LBP</td>
<td>BD more likely to have LBP (p&lt;0.0001) and headaches (p&lt;0.0001).</td>
</tr>
<tr>
<td></td>
<td>Headache</td>
<td></td>
<td>19.3% (n=685/3557) headaches</td>
<td>5.7% (n=41,234/726,262) headaches</td>
<td></td>
</tr>
<tr>
<td>34</td>
<td>Chronic pain interfering with ADL</td>
<td>Single item question</td>
<td>46% (n=338/641) had chronic pain interfering with ADL</td>
<td>No control group</td>
<td></td>
</tr>
<tr>
<td>35</td>
<td>1) migraines and 2) body aches</td>
<td>Self-report 4 point Likert scale questions rating pain during depressive episode</td>
<td>68.5% sample (46/67) had headaches; 29.9% slight, 20.9% moderate and 17.9% major problem</td>
<td>No control group</td>
<td>Current BDRS score predicted headaches (p = 0.012) BDRS severity not related to body aches p=0.3</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>62.6% sample had body aches; 22.4% slight, 19.4% moderate, and 20.9 major problems</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Study</td>
<td>Type of pain</td>
<td>Method of pain assessment/ascertainment</td>
<td>Pain results bipolar disorder</td>
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<td>Other results</td>
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</tr>
<tr>
<td>36</td>
<td>General pain</td>
<td>VAS interviewer administered to assess pain over last 6 weeks (&gt;40 on VAS). Noted severity, duration and interference with ADL</td>
<td>51.2% (n= 62/121) had pain</td>
<td>No control group</td>
<td>Older age associated with pain (OR 1.03 (CI 1.00-1.07))</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Duration of pain 62.5 (±90.9) months</td>
<td></td>
<td></td>
<td>Sex, education, marital status, diagnostic group, depressed mood, sleep disorders and depression not related to pain.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Severity 67.5 (±14.9)</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td></td>
<td></td>
<td>Interference with ADL 67.7 (±21.2)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Location of pain: Head 66.1%; Neck 66.1%; Back 74.2%; Limbs 67.7%; Joints 64.5%</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Nr. of pain locations: 3.44 (±1.46)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>75% of pain musculoskeletal pathology</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>37</td>
<td>Painful somatic symptoms</td>
<td>Medical records</td>
<td>22.6% (n=56/248)</td>
<td>No control group</td>
<td></td>
</tr>
<tr>
<td>38</td>
<td>General pain interfering with activities</td>
<td>Single item question about pain interfering with ADL over past 4 weeks (0-not at all to 5 extremely)</td>
<td>24.8% (n=219/883) had moderate or worse pain interfering with ADL</td>
<td>11.9% (n=5023/ 42,210) had moderate or worse pain interfering with ADL</td>
<td>Comorbid anxiety (OR 1.72, 95% CI 1.41–2.10), being married (OR 1.33, 95% CI 1.08–1.64) and SUD (OR 1.91, 95% CI 1.56–2.34) associated with interfering pain. Age, lower income associated with pain (p&lt;0.001).</td>
</tr>
<tr>
<td>Study</td>
<td>Type of pain</td>
<td>Method of pain assessment/ascertainment</td>
<td>Pain results bipolar disorder</td>
<td>Pain results in control</td>
<td>Other results</td>
</tr>
<tr>
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<td>--------------</td>
</tr>
<tr>
<td>39</td>
<td>Migraine</td>
<td>Unclear, searched patient records</td>
<td>4.7% (n=8/169)</td>
<td>No control group</td>
<td></td>
</tr>
<tr>
<td>40</td>
<td>Migraine/ headache</td>
<td>ICHD-2</td>
<td>70% (n=7/10) had migraine</td>
<td>No control group</td>
<td></td>
</tr>
<tr>
<td>41</td>
<td>LBP</td>
<td>LBP classified as present yes/no from national patient electronic records database</td>
<td>15.4% (n=663/4310) had LBP</td>
<td>10.6% (n=361,868/3,408,760) P&lt;0.0001</td>
<td>BD more likely to have LBP (p&lt;0.0001)</td>
</tr>
<tr>
<td>42</td>
<td>Migraine</td>
<td>Question on lifetime prevalence of migraine</td>
<td>39.8% (n=43/108) had lifetime prevalence of migraine</td>
<td>No control group</td>
<td>Number of psychiatric admission higher in BD without migraine (p=.046), no difference in suicide attempts or SUD</td>
</tr>
<tr>
<td>43</td>
<td>Migraine/ headache</td>
<td>Physician diagnosed and classified</td>
<td>20% (n=6/30) migraine</td>
<td>No control group</td>
<td></td>
</tr>
<tr>
<td>44</td>
<td>Migraine</td>
<td>Survey on previous diagnosis by physician</td>
<td>14.9% for males and 34.7% for females had migranes</td>
<td>5.8% for men and 14.7% females had migraines</td>
<td>BD males with migraine more likely to report earlier BD onset (p&lt;.05), and anxiety (p&lt;.05).</td>
</tr>
<tr>
<td>Study</td>
<td>Type of pain</td>
<td>Method of pain assessment/ ascertainment</td>
<td>Pain results bipolar disorder</td>
<td>Pain results in control</td>
<td>Other results</td>
</tr>
<tr>
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</tr>
<tr>
<td>45</td>
<td>Arthritis/ chronic pain</td>
<td>Single item self-report question</td>
<td>48.9% (n=188/384) had chronic pain</td>
<td>No control group</td>
<td>Chronic pain associated with worse physical HRQOL (p&lt;0.001) but better mental HRQOL scores (p=0.01)</td>
</tr>
<tr>
<td>46</td>
<td>Joint pain</td>
<td>Check list</td>
<td>18.5% (n=5/27) had joint pain</td>
<td>No control group</td>
<td>Migraine associated with BD diagnostic subtype (p&lt;0.001), History suicidal behaviour (p=.03), social phobia 7 panic disorder (p&lt;.001), OCD and anxiety (p&lt;.001)</td>
</tr>
<tr>
<td>47</td>
<td>Migraine</td>
<td>ID migraine questionnaire according to International Headache Society</td>
<td>23.9% (n=71/296) had migraine</td>
<td>No control group</td>
<td></td>
</tr>
<tr>
<td>48</td>
<td>Medically unexplained pain (MUS)</td>
<td>Asked if has pain lasting &gt;6 months in past year that was severe/interfered with ADL and could not be explained.</td>
<td>0.3% had severe chronic pain interfering with ADL (n=190/6328) OR 5.93 (1.71–20.60)</td>
<td>No control group data</td>
<td></td>
</tr>
<tr>
<td>49</td>
<td>Chronic pain</td>
<td>GP database ≥4 analgesic prescriptions in</td>
<td>17.5% (n=451/2582) Had chronic pain</td>
<td>8.8% (n=125,680/ 1,421,796) had chronic pain</td>
<td>OR 1.88 P&lt;0.001 Chronic pain in BD vs. control</td>
</tr>
<tr>
<td>Study</td>
<td>Type of pain</td>
<td>Method of pain assessment/ascertainment</td>
<td>Pain results bipolar disorder</td>
<td>Pain results in control</td>
<td>Other results</td>
</tr>
<tr>
<td>-------</td>
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<td>---------------------------------------------------</td>
</tr>
<tr>
<td>50</td>
<td>Chronic pain</td>
<td>Modified CIDI checklist for medical disorders</td>
<td>40.4% (n=38/93) had chronic pain</td>
<td>Not reported</td>
<td>BD associated with chronic pain OR 3.0 (CI 1.5-5.8) p&lt;0.001*</td>
</tr>
<tr>
<td>51</td>
<td>General pain</td>
<td>Classified from medical records present =yes/no</td>
<td>18.1% had pain (n=4381/24206)</td>
<td>No control group</td>
<td></td>
</tr>
<tr>
<td>52</td>
<td>Migraine</td>
<td>Defined from medical records</td>
<td>2.0% migraine (n=530/27,054)</td>
<td>0.7% (n=16,383/2,325,247)</td>
<td></td>
</tr>
</tbody>
</table>

**Key:** BD= bipolar disorder, VAS = visual analogue scale, ADL=activities of daily living, ICHD-2=International Classification of Headache Disorders, 2nd edition, LBP=low back pain, SF 36= short form 36, nr=number, SUD= substance use disorder, **BDRS = Bipolar Depression Rating Scale**, OCD=obsessive compulsive disorder
Records identified through database searching (N=2,713)

Additional records identified through other sources (N=4)

Records after duplicates removed (N = 2,319)

Records excluded on title abstract level (N = 1859)

Records screened (N = 460)

Records excluded (N = 388)

Full-text articles assessed for eligibility (N = 72)

Studies included in narrative synthesis (N=22: BPD n= 171,352 and control n= 12,204,292)

Full-text articles excluded (n=50), with reasons:
N=22 did not report pain prevalence/measure pain
N=10 not persons with bipolar disorder
N=6 not relevant
N=5 contacted authors to request data for inclusion but no response
N=2 selection bias/not representative
N=2 overlap
N=2 case studies
N=1 contacted authors and exclude as meet exclusion criteria
Figure 2: Random effects pooled prevalence of pain in bipolar samples (N=22, n=171,352)

Pooled proportion = 28.9% (95% CI = 16.4 - 43.4%)

Cochran Q = 66988.29 (df = 24) P < 0.0001
Figure 2: Random effects pooled prevalence of pain in bipolar samples (N=22, n= 171,352)

Figure 2b Funnel plot

Begg-Mazumdar: Kendall's tau = -0.013, P = 0.90

Egger: bias = 11.510, P = 0.48
Figure 3 Relative risk of pain in people with bipolar disorder compared to controls (N=7, n=12,342,577)

Pooled relative risk = 2.14 (95% CI = 1.676 - 2.75), Chi squared= 36.623  (df = 1) P < 0.0001
Cochran Q = 1078.49 (df = 7)  P < 0.0001