The Bipolar Spectrum in Migraine, Cluster and Chronic Tension Headache

a report by
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There has been a great deal of research describing the co-morbidity of migraine/tension headache with anxiety and depression. Previous studies have documented the increased association with migraine and the bipolar spectrum. Those with the bipolar spectrum have also been shown to be more likely to suffer from migraine. For cluster headache and chronic tension headache (without migraine), there have been few studies examining the relationship with bipolar. The study described in this article was performed in order to assess the prevalence of bipolar in three distinct headache types: migraine, cluster and chronic tension (without migraine).

The bipolar spectrum tends to be underdiagnosed, with the full clinical spectrum being an evolving concept. The mild end of the bipolar spectrum (bipolar II, cyclothymia, bipolar not otherwise specified [NOS]) is often missed. It is likely that 4% (or more) of the general population suffers from the bipolar spectrum. As bipolar complicates treatment in a variety of ways, the clinical stakes for missing bipolar are enormous. Bipolar and migraine share common genetic links and both are multifactorial in origin.

Methods

One thousand, two hundred consecutive migraine patients, 275 cluster patients and 292 patients with chronic tension headache without migraine were evaluated. They were all patients at our headache centre. The diagnoses were based on criteria set down by the International Headache Society (IHS). The evaluation was based on a chart review, a mood disorder questionnaire (MDQ), a patient health questionnaire (PHQ)-9 and interviews with patients and families. The inclusion criteria stipulated that the patients should be >20 years of age and that a diagnosis of migraine, cluster or chronic tension headache (without migraine) should be made. The lifetime prevalence of bipolar was assessed, including the milder end of the spectrum.

Bipolar illness was defined according to the criteria established by the Diagnostic and Statistical Manual of Mental Disorders, Fourth Edition (DSM-IV). In addition, the modifications to DSM-IV by Akiskal were utilised in defining bipolar disorders. The following four conditions were defined according to the DSM-IV criteria.

Bipolar I disorder was defined when there had been at least one episode, currently or in the past, of true mania.

An assessment of bipolar II was made if there had been one or more major depressive episodes, plus at least one hypomanic episode, no mania or mixed episodes, and these episodes must have caused significant distress or impairment in patient functioning.

Cyclothymic disorder was defined as at least two years of numerous periods of hypomania and numerous episodes of depressive symptoms that do not meet criteria for major depressive episode. During the two-year period, the patient could not have been without the symptoms for more than two months at a time, and no major depressive episode, manic episode or mixed episode could have been present during the first two years of the disturbance. These symptoms had to cause clinically significant distress or impairment in functioning, and could not be due to substance abuse or a medical condition.

Bipolar disorder NOS was defined with additions according to Akiskal. Examples of patients included in this category are those with rapid alterations between manic and depressive symptoms that do not meet minimal criteria for a full manic episode or for a major depressive episode; recurrent hypomanic episodes without intercurrent depressive symptoms; the presence of a hypertymic temperament as the prevalent, long-term functioning of the person; and the presence of a persistently agitated, angry and moody personality (temperamental instability), particularly with a strong family history of bipolar disorder and/or a hypomanic reaction to an antidepressant (e.g. up all night, mind racing). Increased energy and lability of mood were also used as additional indicators of bipolarity. In addition, criteria that were considered in the diagnosis of the ‘softer’ end of the spectrum include: early onset of depression (prior to 25 years of age, and certainly prior to 17 years of age), atypical or psychotic depressive episodes, post-partum depression and lack of response to three or more antidepressant trials.

Results

Migraine

One thousand, two hundred consecutive migraine patients were evaluated according to DSM-IV guidelines. The results were as follows: bipolar I, 24; bipolar II, 28; bipolar NOS, 34; and cyclothymia, 17. Total bipolar spectrum for migraineurs was 103 (8.6% of the total).

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Headache

Cluster

Two hundred and eighty-seven cluster headache patients were seen over 18 years. The episodic cluster cohort totalled 141. The results showed: bipolar I, two; bipolar II, four; bipolar NOS, one; and cyclothymia, two; the total episodic cluster was nine (6.4%). The chronic cluster cohort totalled 146: bipolar I, two; bipolar II, two; bipolar NOS, two; and cyclothymia, four. Total chronic was 10 (6.8%). Total bipolar spectrum for cluster patients was 19 (6.6% of the total).

Chronic Tension (without Migraine)

Two hundred and ninety-two patients with chronic tension-type headache without migraine were evaluated. The results were: bipolar I, five; bipolar II, three; bipolar NOS, three; and cyclothymia, two. The total bipolar spectrum for chronic tension headache was 13 (4.5%).

Discussion

The bipolar spectrum is seen relatively often in headache patients, particularly among migraineurs. The clinical implications for not diagnosing bipolar is enormous, with the result that patients tend to move from antidepressant to antidepressant, with generally poor results, without being on adequate mood stabilisers. Accurate diagnosis of bipolar is crucial, because it is not an easy condition to treat or to deal with, for family members as well as the patient. Treatment with adequate mood stabilisers may help, but we are often left with less than desirable results. As is the case with headache preventatives, we need better medications for the bipolar spectrum.

The clinical spectrum of bipolar is evolving. Mania is better recognised as well as for treating the depression associated with acute mixed mania. Its efficacy in preventing bipolar depression is less clear. The atypical (second-generation) antipsychotics may be of benefit for some patients with bipolar depression, as well as for treating the mania/hypomania. Quetiapine has the best data at present, but does carry at least a mild to moderate liability for the metabolic syndrome. While this class may help certain aspects of the bipolarity, the medications are probably most useful as adjuncts to lithium or lamotrigine.

The recognition of the bipolar spectrum is crucial if we are to effectively treat the patient with headaches.