Chapter 4

Bipolar disorder and complementary medicine: current evidence, safety issues, and clinical considerations

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Abstract

Background:
Bipolar Disorder (BD) is a debilitating syndrome that is often undiagnosed and under-treated. Population surveys show that persons with BD often self-medicate with complementary and alternative medicine (CAM) or integrative therapies in spite of limited research evidence supporting their use. To date no review has focused specifically on non-conventional treatments of BD.

Objectives:
To present a comprehensive review of non-conventional (complementary and integrative) interventions examined in clinical trials on BD, and to offer provisional guidelines for the judicious integrative use of CAM in the management of BD.

Methods:
PubMed, CINAHL, Web of Science and Cochrane Library databases were searched for human clinical trials in English during mid-2010 using Bipolar Disorder and CAM therapy and CAM medicine search terms. Effect sizes (Cohen's d) were also calculated where data were available.

Results:
Several positive high-quality studies on nutrients in combination with conventional mood stabilizers and antipsychotic medications in BD depression were identified, while branched-chain amino acids and magnesium were effective (small studies) in attenuating mania in BD. In the treatment of bipolar depression evidence was mixed regarding omega-3, while isolated studies provide provisional support for a multi-nutrient formula, n-acetylcysteine, and L-tryptophan. In one study acupuncture was found to have favorable, but non-significant effects on mania and depression outcomes.

Conclusions:
Current evidence supports the integrative treatment of BD using combinations of mood stabilizers and select nutrients. Other CAM or integrative modalities used to treat BD have not been adequately explored to date; however, some early findings are promising. Select CAM and integrative interventions add to established conventional treatment of BD and may be considered when formulating a treatment plan. It is hoped that the safety issues and clinical considerations addressed in this article may encourage the practice of safety-conscious and evidence-based integrative treatment of BD.
Introduction

Bipolar disorder (BD) is a debilitating heritable mental illness that has profound personal and socioeconomic effects. Emerging research findings reviewed in this paper suggest that non-conventional therapies may have a potentially significant role in improving quality of life, reducing side-effects, improving adherence with conventional medications, and reducing the severity of bipolar symptoms. Complementary and Alternative Medicine (CAM) consists of therapies such as acupuncture or naturopathy, and medicinal interventions such as herbal medicines or nutrients (Sarris & Wardle, 2010). CAM is commonly used by people diagnosed with mood disorders with up to 50% of psychiatric outpatients using one or more CAM therapies in a given year (Knaudt et al., 1999; Hoenders et al., 2006). Studies reveal especially high CAM use rates in women, the highly educated, the elderly, and those with chronic diseases (Keaton et al., 2009; Kessler et al., 2001; Wu et al., 2007). Aside from specific CAM interventions used in conjunction with pharmacotherapies (reviewed in this paper), according to users, CAM may provide general beneficial effects on physical and mental health and quality of life (Sirois, 2008). Furthermore, research has shown that administration of CAM within an holistic model can be beneficial for mental health (Cooley et al., 2009). Thus, adjuvant use of CAM within an integrated whole-person framework (combined with pharmacological and psychosocial interventions) holds the potential to reduce the severity of bipolar symptoms and risk of relapse.

Despite this, few studies have been done on non-conventional treatments of BD. Other than a review by Andreescu and colleagues (Andreescu et al., 2008), there is a paucity of peer-reviewed publications reporting clinical trial outcomes and safety issues associated with non-conventional treatments of BD. While their article is significant for providing the first review in this area, its focus is primarily on unipolar depression and it does not review nutritional interventions in detail, many of which are effective adjunctive interventions when combined with conventional pharmacotherapies (Sarris et al., 2009). The Andreescu et al. review is also limited by excluding studies published over 10 years ago. In the present paper we critically review evidence for CAM and integrative treatments of bipolar depression, mania and hypomania, and make recommendations for provisional clinical guidelines.

Methods

PubMed, CINAHL, Web of Science and Cochrane Library databases were searched during mid 2010 firstly for research in the overarching area of bipolar disorder; then for human clinical trials using the terms “Bipolar Disorder”, “Bipolar Depression”, “Bipolar Mania”, “Mania”, “Hypomania”, “Cyclothymia” together with terms for CAM therapies and products (e.g. herbal and nutritional medicine), and dietary and lifestyle factors. A forward search of identified papers was subsequently performed using Web of Science cited reference search. Reasons for exclusion included: a higher level of evidence being available or inadequate methodological rigor. Included clinical trials were open or controlled human studies that recruited people diagnosed with BD, examined CAM as a monotherapy or as an adjuvant with a conventional medication, and measured outcomes using established psychiatric rating scales. The term “significant” was used in studies with \( p \) values of <0.05. Effect sizes were calculated in all RCTs where data were available. From the results of the clinical
trials we calculated an effect size as Cohen’s \( d \) (Cohen, 1988) by firstly subtracting the differences between the intervention and placebo scores on the scale used, then dividing this by the pooled standard deviation at baseline (clinical effect: 0.2 = small, 0.5 medium, > 0.8 large). Studies involving exercise and psychological therapies were omitted from analysis as, for purposes of this review, they are considered mainstream interventions (however they are briefly discussed in the Clinical Considerations section). Results are grouped under omega-3 fatty acids, amino acids, vitamins and minerals, herbal medicines, and acupuncture.

**Bipolar Disorder**

**Etiology**

Bipolar disorder (BD) is a debilitating heritable mental illness that affects approximately 1-2% of adults in their lifetime. When milder subclinical presentations are included the prevalence rate increases to approximately 4% (Merikangas et al., 2007). First-degree relatives of bipolar individuals are significantly more likely to develop the disorder than the population at large and twins have a 70% risk of sharing the disorder (Gurling et al., 1995). Although not yet fully elucidated, BD symptoms are probably caused by dysregulation of serotonergic and dopaminergic pathways, and diminished activity in the hippocampus and pre-frontal cortex (Konradi et al., 2004; Miklowitz & Johnson, 2006). It has been suggested that abnormal activity in hypothalamic circuits involved in maintaining normal circadian rhythms manifest as the affective and behavioral symptoms of BD (Miklowitz & Johnson, 2006).

**Diagnosis**

According to current conventional Western psychiatric nosology, BD diagnosis is divided into Bipolar Disorder Type I (BD I) and Bipolar Disorder Type II (BD II: see table 1), and can be differentiated from unipolar depression (major depressive disorder: MDD) by the presence of manic or hypomanic (lesser) episodes (American Psychiatric Association, 2000). A manic episode is a complex symptom pattern that may encompass disparate affective, behavioral and cognitive symptoms, including pressured speech, racing thoughts, euphoric or irritable mood, agitation, inflated self-esteem, distractibility, excessive or inappropriate involvement in pleasurable activities, increased goal-directed activity, diminished need for sleep, and in severe cases, psychosis (American Psychiatric Association, 2000).
Table 1. Unipolar depression and bipolar depression: differential diagnoses*

<table>
<thead>
<tr>
<th>Major Depressive Disorder</th>
<th>Bipolar Disorder I (mania)</th>
<th>Bipolar Disorder II (hypomania)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Two weeks or more of persistent low mood and/or anhedonia (loss of interest in pleasurable activities)</td>
<td>Episodes of mania or a mixed episode lasting 5 or more days. Often presents with cycling between mania and depression</td>
<td>Hypomanic episodes that do not meet criteria for full mania, in addition to cycling to one or more episodes of MDD</td>
</tr>
<tr>
<td>Low mood impairs work and/or social functioning, may require hospitalization for severe depression</td>
<td>Manic episodes impairs work and/or social functioning, and often requires hospitalization</td>
<td>Hypomanic episodes not as severe as BD I mania, and do not significantly impair work/social functioning</td>
</tr>
<tr>
<td>Changes in body weight, digestion, and sleep patterns</td>
<td>Manic phase (euphoric or irritable mood, grandiosity or psychosis, decreased need for sleep)</td>
<td>Hypomanic phase (euphoric or irritable mood, decreased sleep, increased talkativeness, goal planning and sexual focus)</td>
</tr>
<tr>
<td>Psychological changes (e.g. suicidal ideation, guilt, self-worthlessness, agitation)</td>
<td>Depressive phase can present with similar psychological and somatic changes to MDD</td>
<td>Depressive phase can present with similar psychological and somatic changes to MDD (more common in BD II than BD I)</td>
</tr>
<tr>
<td>Not due to bereavement, medical comorbidity, drugs or alcohol, or medication</td>
<td>Not due to medical comorbidity, drugs or alcohol, or medication</td>
<td>Not due to medical comorbidity, drugs or alcohol, or medication</td>
</tr>
</tbody>
</table>

* Adapted from DSM-IV TR (APA, 2000)

A history of depressive episodes is not required for a formal diagnosis of BD I according to the Diagnostic and Statistical Manual of Mental Disorders version IV (DSM-IV-TR) (APA, 2000). In contrast, BD II can be diagnosed only in cases when at least one hypomanic episode and at least one depressive episode have been documented. In both disorders moderate or severe depressive episodes typically alternate with manic symptoms, however in “mixed mania” symptoms of mania and depressed mood overlap. Another variant called rapid cycling is diagnosed when at least four complete cycles of depressed mood and mania occur during any 12-month period. A mild variant of BD, cyclothymic disorder, is diagnosed when several hypomanic and depressive episodes take place over a two-year period in the absence of severe manic, mixed or depressive episodes (APA, 2000). The emerging concept of a “mood spectrum” hypothesis suggests that patterns of depressive and manic symptoms occur along a continuum, and that “mood disorders” do not exist as discrete diagnostic entities (Benazzi, 2007; Cassano et al., 2002). Evidence supports that unipolar depression and bipolar II depression occur across a spectrum, with 30% of patients diagnosed with major depressive disorder (MDD) experiencing various bipolar symptoms (e.g. agitation, racing thoughts, decreased sleep) (Benazzi, 2007). While BD I presents as a distinct psychiatric disorder that fulfills specific diagnostic criteria, BD II and cyclothymic disorder are more variable in presentation.
It is estimated that two thirds of individuals diagnosed with BD experience moderate or severe symptoms in any given year (Suppes et al., 2005). Bipolar patients experience depressive symptoms three times more often than mania, and five times more often than rapid cycling or mixed episodes (Judd et al., 2002). A diagnosis of BD is one of the highest risk factors for suicide (Pompili et al., 2009). BD usually presents in a rhythmic manner, oscillating between episodes of mania and depression, with several risk factors and triggers contributing to the rate of relapse and failed response to treatment (see figure 1).

**Conventional treatment**

Conventional pharmacotherapies are an important and often necessary treatment of both the depressive and manic phases of BD. First-line treatments of BD include mood stabilizers (e.g. lithium carbonate, carbamazepine, and valproate), antidepressants, antipsychotics and sedative-hypnotics (Miklowitz & Johnson, 2006; Suppes et al., 2005). Antipsychotics are used to treat agitation and psychosis, which occur frequently in acute mania, and select antipsychotics have been found to be effective mood stabilizers. Sedative-hypnotics are sometimes prescribed for the severe insomnia that accompanies mania, as well as for daytime management of agitation and anxiety (Cousin & Young, 2007).

Side effects often occur with these medications, having a mixed record of success due to their limited efficacy and high rates of treatment discontinuation (Pompili et al., 2009). Fewer than half of patients who take a conventional mood stabilizer or other psychotropic medications following an initial manic episode report sustained control of their symptoms (Culver et al., 2007). As many as one half of all bipolar patients who take mood stabilizers do not experience good control of their symptoms or refuse to take medications, and approximately 50% discontinue their medications because of serious adverse effects including tremor, weight gain, thyroid dysfunction elevated liver enzymes, and many others (Fleck et al., 2005). People diagnosed with bipolar disorders should be maintained on a consistent long-term pharmacotherapeutic regimen to reduce the rate of re-hospitalization and increase chances of full remission (Perlick et al., 1999). In patients diagnosed with BD, stressors, seasonal change, reduced sleep and stimulants or recreational drug use may provoke an episode of hypomania (although sometimes the trigger may have no apparent cause (Miklowitz & Johnson, 2006). Regular exercise, good nutrition, a strong social support network, and a predictable low-stress environment help reduce relapse risk (Suppes et al., 2005; Benjamin, 2007; Miklowitz & Scott, 2009; Lakhan & Vieira, 2008).

Psychotherapy and psychosocial interventions in stable bipolar patients may potentially reduce relapse risk by providing psychological support, enhancing medication adherence, and helping patients address warning signs of recurrning depressive or manic episodes before more serious symptoms emerge (Miklowitz & Scott, 2009). Relapse prevention usually involves use of "lifecharts" and an effective stress management plan. An example is the novel BD relapse prevention program (called MAPS), which has evidence in reducing relapse of mania and BD depression. The Australian developed MAPS program was studied in a clinical trial involving 84 participants with BD, and was conducted over 12 weeks (Castle et al., 2010). Participants were randomized to either the program involving education on BD (symptoms, monitoring triggers, and symptom management skills), in addition to goal setting, medication management, and relapse prevention planning; or a control group consisting of treatment as usual plus telephone calls. Results revealed that
participants who received the group-based intervention were significantly ($p=0.04$) less likely to have a relapse of mania or depression, and spent less time unwell.

**Clinical Considerations**

When considering recommending CAM or integrative treatments for BD (in concert with traditional psychotropic medications), it is essential to first carefully examine the evidence for both conventional and nonconventional therapies. In this paper we propose an integrative model that incorporates select CAM therapies in combination with conventional psychotropic medications. We recommend against the use of CAM or integrative therapies whose safety and efficacy is not supported by strong research evidence. This caveat applies even more so to the management of severe symptoms of bipolar depression or mania. Mental healthcare providers should always recommend those therapies supported by the highest level of evidence (Hoenders et al., 2010), and in spite of their limited efficacy and unresolved safety issues, mood stabilizers and select antipsychotics should always be regarded as the first-line treatments of the severe form of bipolar illness (i.e. BD I) (Nivoli, 2011).

Prior to diagnosing bipolar mood disorder a period of mania or hypomania lasting several days must be established in the context of a broader pattern of impairment that may include decreased need for sleep, flight of ideas, grandiosity, excessive or unrealistic spending, or hyper-sexual activity. A life chart can help establish a pattern of cyclic mood changes and associated impairments in social activity, academic performance and work. An important initial consideration when prescribing CAM in persons diagnosed with BD, depends upon which phase are they in (i.e. mania, depression, or remission). A careful history is needed to establish a persisting pattern of mood changes fluctuating between depression and mania or hypomania. Conventional laboratory tests and functional brain imaging studies can be used to rule out medical disorders that can mimic symptoms of depressed mood or mania including, for example, thyroid disease, strokes (especially in the right frontal area of the brain), multiple sclerosis, seizure, or other neurologic disorders (Kumar & Clark, 2002). Irritability or euphoria alternating with periods of depressed mood is sometimes associated with chronic abuse of stimulants, marijuana or other drugs. Thus, screening for substance abuse should be done before a formal diagnosis of BD is made (Swann, 2010).

In persons in the manic phase of BD type I, hospitalization is usually required, at which time adjunctive use of magnesium, choline, branched-chain amino acids, or L-tryptophan may be appropriate adjuvants in combination with mood stabilizers or antipsychotics. When prominent symptoms of anxiety or agitation are present, effective integrative strategies should prioritize treatment of those symptoms. In addition to sedating pharmacotherapies, the use of botanical or nutritional anxiolytics (such as *Piper methysticum*, magnesium, L-theanine, or L-tryptophan) may be beneficial (Lakhan & Vieira, 2008; Sarris et al., 2009a).
For patients in the depressed phase the treatment approach is different. It is important to note that depressed symptoms are often misdiagnosed as major depressive disorder when in fact they are the depressive phase of BD. Thus careful screening is essential in all patients presenting with depressed mood including assessment of the length, frequency and severity of depressive episodes, identification of precipitating stressors, and evaluation of suicidality. Assessment should include a urine drug and alcohol screen and a review of their sleep pattern and level of stress. CAM therapies for bipolar depressed patients include select nutrients (S-adenosyl methionine, L-tryptophan, omega-3 fatty acids (mixed evidence)) (Sarris et al., 2009b), botanicals (Hypericum perforatum, Rhodiola rosea, Crocus sativus) (Sarris, 2007), select acupuncture protocols (Wang et al., 2008), regular exercise (Barbour et al., 2007), good nutrition (Jacka et al., 2010), and psychological interventions (Cuijpers et al., 2008).

All bipolar patients being treated for depressed mood are at risk of “switching” to mania and should be closely monitored. A personalized relapse prevention program including patient and family education about early warning signs of recurring depressed mood or mania, and a well thought-out plan is an essential component of care for all bipolar patients. Important safety considerations are raised.
by the use of CAM therapies that have efficacy in major depressive disorder but that have not been established as safe in bipolar depressed mood and found to not increase the risk of switching to mania. Case reports implicate select natural products used to treat major depressive disorder, such as *Hypericum perforatum* (Moses & Mallinger, 2000), *Ephedra sinica* (Boerth & Caley, 2003), or S-adenosyl methionine (Papakostas et al., 2003) as potentially inducing mania. Several case reports of serotonin syndrome have been associated with the combined use of *Hypericum perforatum* and SSRIs (Sarris et al., 2009c). In addition it should be noted that extracts high in the active constituent hyperforin may induce cytochrome P450 3A4 enzymes and the P-glycoprotein drug efflux pump resulting in reduced serum levels of many drugs including oral contraceptives, anticoagulants, protease inhibitors, and anti-seizure medications (Whitten et al., 2006).

A significant percentage of individuals diagnosed with BD use non-pharmacological modalities adjunctively with prescription medications. However, with the exception of the nutrient interventions reviewed in this article, there is relatively little evidence for the safety and efficacy of the majority of such integrative treatments. Finally, while exercise should be encouraged in all BD patients because of established benefits to general health and mood, those patients taking lithium should consult their physician before starting a rigorous exercise program. Lithium is excreted with perspiration and strenuous physical activity that involves significant sweating may lower lithium levels in the blood; at least one such case has been reported (Waring, 2006).

**Clinical Evidence of CAM in Bipolar Disorder**

*Omega-3 fatty acids*

Countries where there is high fish consumption have relatively lower prevalence rates of BD (Hibbeln, 1998). Several clinical trials have studied omega-3 fatty acids including fish oil and purified eicosapentaenoic acid (EPA) and docosahexaenoic acid (DHA), as a mono-therapy or as an adjuvant intervention in BD. An early RCT involving 44 participants using a combination of EPA and DHA (9.6 g/day) with conventional drug therapies revealed positive results on measures of depressed mood in terms of response and remission rates on The Hamilton Depression Rating Scale (HAMD) ($d = 1.40$) (Stoll et al., 1999; Hamilton, 1960). No significant effect on mania outcomes occurred. A 26-week open-label adjuvant study by Osher and colleagues (2005) in 12 participants with BD I, revealed that eight out of ten participants with one month of EPA were responders on HAMD ($d=1.23$). It should be noted that while these open-label studies are positive, confidence in these should be tempered as they are not controlled. A 12-week, 3-arm controlled study involving 75 participants using 1 g or 2 g of EPA combined with any class of psychotropic medication revealed a small but significantly greater reduction on the HAMD from either dose, compared with placebo (1 g: $d = 0.90$, 2 g: $d = 0.50$) (Frangou et al., 2006). However, no significant effect for mania was achieved on The Young Mania Rating Scale (YMRS) (Young et al., 1978). A later RCT conducted by the research group (Frangou et al., 2007) using 2 g of EPA versus placebo over 12 weeks in 14 female participants with BD I revealed positive, but not statistically significant effects on depression outcomes. A novel 4-week adjuvant study involving 21 participants with BD I by Hirashima and colleagues (2004), revealed no significant differences between EPA 5 g plus DHA 3 g and a non-treatment control. Interestingly however,
brain resonance imaging showed that T2 levels were reduced in the treatment group, denoting increased neuronal cell membrane fluidity.

A larger study \((n = 121)\) using 6 g of EPA in combination with at least one mood stabilizer in patients diagnosed with rapidly cycling bipolar disorder also found no benefit over placebo on reducing mania on YMRS (Keck et al., 2006). Further, a small RCT with 15 participants, using 4.4 g of EPA and 6.6 g of DHA/day adjuvantly with 20 mg/kg/day of valproate also revealed no benefit over placebo for reducing mania (Chiu et al., 2005). Two open-label studies have been published on omega-3s in pediatric bipolar disorder: an adjuvancy study and a monotherapy study. Clayton et al. (2009) conducted a 6-week study involving 18 adolescents with BD I or II using omega-3 (DHA 1560 mg and EPA 360 mg per day) and found significantly reduced clinician-rated mania and depression from baseline. A study by Wozniak and colleagues (2007) also revealed significant reductions on YMRS \((d = 0.90)\) and BPRS \((d = 0.83)\) compared to baseline using 1290 mg-4300mg of fish oil in 20 adolescents who met DSM-IV-TR criteria for BP and had a YMRS score of >15.

Current evidence weakly supports use of omega-3 preparations in combination with conventional psychotropic medications in the depressive phase of BD; however, omega-3s probably have little or no clinical effect in attenuating mania. Although considered a very safe intervention, rare cases of increased bleeding times, but not increased risk of bleeding, have been reported in patients taking aspirin or anti-coagulants together with omega-3s (Sarris et al., 2009b).

**Amino acids**

Bipolar patients may be genetically susceptible to mood swings when certain amino acids or other micronutrients are lacking in the diet (Kaplan et al., 2007). Findings of two small RCTs suggest that certain branched-chain amino acids (BCAA) may rapidly improve acute mania by interfering with synthesis of norepinephrine and dopamine (Scarna et al., 2003). In one study 25 bipolar patients randomized to a blend of the branched-chain amino acids leucine, isoleucine and valine (60 g/day) versus placebo experienced significant reductions in the severity of mania within six hours (Scarna et al., 2003). Improvements in mania were sustained with repeated administration of the amino acid drink. N-acetylcysteine (NAC) is an amino acid with strong anti-oxidant properties that has been used to treat a range of inflammatory disorders (Dodd et al., 2008). In novel research, Berk and colleagues (2008) conducted a 24-week RCT using 1 g of NAC versus placebo in a sample of 75 participants stable on medication or therapy with DSM-IV-TR diagnosed BD I or BD II. Results revealed that NAC significantly reduced bipolar depression on the Montgomery Depression Rating Scale (MADRS) (Montgomery & Asberg, 1979) and the Bipolar Depression Rating Scale with strong effect sizes respectively of 1.04 and 0.83. No significant effect was found on mania outcomes, although it should be noted that YMRS mania levels were very low, thus significant changes were unlikely to occur.

Restricting or excluding L-tryptophan from the diet may increase the susceptibility of bipolar patients to depressive mood swings; however, research findings to date are highly inconsistent (Bell et al., 2005). L-tryptophan has demonstrated beneficial effects in reducing depressed mood in people with unipolar depression (Sarris et al., 2009b). A 1985 2-week study used 12 g of the amino acid in 24 participants with mania (Chouinard et al., 1985). In this two-phase trial, measures of mania were significantly reduced with L-tryptophan on Clinical Global
Inventory \( (d = 1.47) \) in the initial open phase, and continued but lessened during the controlled phase.

**Vitamins and minerals**

Magnesium may be an effective adjuvant therapy for treatment of acute mania or rapidly cycling BD. In a small open trial oral magnesium supplementation had comparable efficacy to lithium in rapid cycling bipolar patients (Chouinard et al., 1990). In a small case series intravenous magnesium sulfate used as an adjuvant with lithium, haloperidol and a benzodiazepine in bipolar patients with severe treatment-resistant mania resulted in significant improvement in global functioning and reduction in the severity of mania (Heiden et al., 1999). Many patients treated with intravenous magnesium sulfate remained stable on lower doses of conventional medications. An RCT used 375 mg of magnesium oxide versus glucose placebo over 16 weeks in 20 participants with prior DSM-IV diagnosed mania and > 6 months on a stable mood stabilizer (verapamil). Results revealed a significant reduction on mania compared to control at week 16 (Giannini et al., 2000).

Two clinical trials using inositol (12 g and 5-20 g) adjuvantly with maintenance doses of mood stabilizers have been conducted. Both RCTs (Chengappa et al., 2000; Evins et al., 2006) had small samples \((n=24\text{ and } 26, \text{ respectively})\) and were conducted over six weeks. Results in both studies revealed no significant differences between inositol or placebo on depression or mania outcome scales. Clinical response however was noted in 12/21 participants taking inositol on pooled results of both studies on HAMD and MADRS.

Folic acid has been studied in one early trial as an adjuvant in BD patients stabilized on lithium. A 52-week RCT by Coppen and colleagues (1986) compared 200 mcg of folic acid versus placebo tablets in 102 participants taking lithium. Results revealed that the completers in the folic acid group \((n=41)\) had significantly lower BDI scores than the control group, with a strong effect size of 1.07.

A proprietary 36-ingredient formula of vitamins and minerals may significantly reduce symptoms of mania, depressed mood and psychosis in bipolar patients when taken alone or used adjunctively with conventional mood stabilizers. Six out of eleven completers had clinical response with strong effect sizes (HAMD: 1.70, YMRS: 0.83). In one case series 11 bipolar patients who completed a 6-month protocol were able to reduce their conventional mood stabilizers by half while improving clinically (Kaplan et al., 2001). In another case series, 13 out of 19 bipolar patients who continued on the nutrient formula remained stable after discontinuing conventional mood stabilizers (Simmons, 2003). Some patients stopped taking the formula because of nausea and diarrhea and three patients resumed conventional mood stabilizers because of recurring manic symptoms. Researchers believe the formula works by correcting metabolic errors that result in bipolar-like symptoms in genetically predisposed individuals when certain micronutrients are deficient in the diet (Popper et al., 2001; Kaplan et al., 2001).

Choline is necessary for the biosynthesis of acetylcholine, and abnormal low brain levels of acetylcholine may contribute to some cases of mania (Leiva, 1990). Findings of a small open study suggest that phosphatidylcholine (15 gm to 30 gm/day) may reduce the severity of mania and depressed mood in bipolar patients (Stoll et al., 1996). It should be noted that two non-responders were also taking hypermetabolic doses of thyroid medication. Clinical improvement correlated with increased intensity of the basal ganglia choline signal as measured on proton
magnetic resonance imaging (MRI). The effect of choline on depressive symptoms was variable (Stoll et al., 1996).

Findings of a small open study suggest that patients diagnosed with BD who exhibit mania or depressed mood may respond to low doses (50 micrograms with each meal) of a natural lithium preparation (Fierro, 1988). Post-treatment serum lithium levels were undetectable in patients who responded to trace lithium supplementation.

Findings from animal research and a small open study suggest that bipolar patients who take potassium 20 mEq twice daily with their conventional lithium therapy experience fewer side effects, including tremor, compared to patients who take lithium alone (Tripuraneni, 1990). No changes in serum lithium levels were reported in patients taking potassium. Pending confirmation of these findings by a larger double-blind trial, potassium supplementation may provide a safe, cost-effective integrative approach for the management of bipolar patients who are unable to tolerate therapeutic doses of lithium due to tremor and other adverse effects (patients who have cardiac arrhythmias or are taking anti-arrhythmic medications should consult their physicians before considering taking a potassium supplementation).

Herbal medicines
Findings of a large 12-week placebo-controlled trial involving a sample of 58 BD patients suggest that a proprietary Chinese compound herbal formula consisting of at least 11 herbs may enhance the effect of conventional mood stabilizers for treatment of the depressive phase of BD (Zhang et al., 2005). Bipolar depressed (but not manic) patients randomized to the herbal formula plus carbamazepine experienced significantly greater reductions in the severity of depressed mood compared to matched patients receiving a mood stabilizer only. These findings were replicated in a subsequent study, which confirmed that bipolar depressed patients treated with the herbal formula improved more than patients treated with a placebo, with a strong effect size ($d=0.98$) between treatments at week 12 (Zhang et al., 2007).
Table 2. Adjuvant nutrient interventions in Bipolar Disorder

<table>
<thead>
<tr>
<th>Intervention</th>
<th>BD (depression)</th>
<th>BD (mania)</th>
<th>Level of evidence</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Depression</td>
<td>Mania</td>
<td></td>
</tr>
<tr>
<td>Omega-3</td>
<td>✔</td>
<td>A</td>
<td>D</td>
</tr>
<tr>
<td>N-acetylcsteine</td>
<td>★</td>
<td>B</td>
<td>C</td>
</tr>
<tr>
<td>BCAA</td>
<td>✔</td>
<td>NK B</td>
<td>C</td>
</tr>
<tr>
<td>Inositol</td>
<td>✔</td>
<td>C</td>
<td>NK</td>
</tr>
<tr>
<td>Choline</td>
<td>✔</td>
<td></td>
<td>C</td>
</tr>
<tr>
<td>Follic acid</td>
<td>✔</td>
<td>B</td>
<td>C</td>
</tr>
<tr>
<td>Magnesium</td>
<td>✔</td>
<td></td>
<td>C A</td>
</tr>
<tr>
<td>Chelated minerals</td>
<td>✔</td>
<td></td>
<td>B</td>
</tr>
</tbody>
</table>

✔ = Potential use in bipolar mania or bipolar depression  
NK= Not known  
BCAA= Branched-chain amino acids  
A= Several repeated clinical trials (with some positive results)  
B= Unreplicated positive study  
C= Small study, or inconclusive results  
D= Several clinical trials reveal mainly negative results

Early studies suggested that the Ayurvedic plant medicine *Rauwolfia serpentina* and an alkaloid derivative, reserpine, was an effective treatment of BD by augmenting the anti-manic efficacy of lithium without risk of toxic interactions (Bacher & Lewis, 1979; Berlant, 1986). However, therapeutic use of this plant in various Western countries is restricted due to the presence of the alkaloid reserpine, which has potent effects on blood pressure and the CNS. A previous review published in *JACM* (Sarris et al., 2009d) of placebo-controlled trials comparing *Hypericum perforatum* to placebo or conventional antidepressants concluded that the herbal medicine might be beneficial for mild to moderate depressive symptoms. Although this herb may potentially be beneficial in the depressive phase of BD, no studies on this have been conducted to date. Several case reports of mania induction with St. John’s wort (Moses & Mallinger, 2000; Fahmi et al., 2002) and potential serious interactions with many drugs (Izzo, 2004) have resulted in limited use of this herbal for the treatment of BD.
Figure 2. Treatment decision tree

**Bipolar Disorder**

**Assess Risk and Establish Particulars**
- Current or history of manic episode with or without depressive episode? (BD I); or hypo-mania and depressive episodes? (BD II); or concurrent manic and depressive symptoms? (mixed mania)
- Dominant symptoms? (e.g., euphoric, irritability, depressed mood, insomnia, psychosis)

**Determine Causative Factors**
- Family history of bipolar disorder, other psychiatric disorders?
- Pre-existing psychiatric disorder?
- Underlying medical disorder?
- Effects of a prescription drug?
- Underlying substance abuse?
- Severity of functional impairment?

**Formulate Integrative Treatment Plan**
- Confirm diagnosis of bipolar disorder and rule out co-morbid psychiatric or medical disorders
- Document conventional, CAM and integrative therapies that have already been tried
- Identify core symptoms that will be the focus of clinical attention for treatment
- Stabilize patient’s (hypo) mania as rapidly as possible starting with conventional or integrative treatment protocols
- Start with most validated conventional or integrative treatment protocols
- When more substantiated modalities are not effective, consider less validated modalities with patient’s informed consent
- After hypo/mania has resolved, be aware of any relapse into a depressive phase and treat accordingly

**Referrals**
- Urgent referral of BP I pts to emergency room if acutely manic, suicidal or psychotic (e.g. delusional, paranoid, auditory hallucinations)
- Non-urgent referral of BP II pts to primary care physician to evaluate thyroid, rule out possible medication adverse effects, and rule out substance abuse
- Non-urgent referral to psychotherapist for long-term supportive therapy

**Diagnostic investigations**
- Thorough history to determine symptoms type and severity and evaluate need for psychiatric evaluation and/or hospitalization
- Check serum drug level (lithium and valproic acid) in patients taking these medications
- FT4 and TSH to rule out hyperthyroidism
- Urinary toxicology screen to rule out drug abuse

**Integrative CAM Treatment Options**
- Mood Stabilizers, Antipsychotics + e.g.
  - Omega-3 fatty acids
  - Branched-chain amino acids
  - N-acetylcysteine
  - Folic acid
  - Magnesium
- Conventional and natural antidepressants often helpful but may risk mania induction
- Encourage exercise and mind-body practice for relapse prevention

**Communication and Follow up Protocols**
- Frequency of follow-up appointments reflects severity and risk
- Continue to monitor for suicide risk and mania
- Encourage phone or e-mail contact for questions about plan or concerns about adverse effects
- On-going discussion of valid treatment choices and patient preferences
- Review appropriate therapies for core symptoms and severity
- Educate patient re relapse risk, warning signs of recurring manic or depressive episodes, and priority on routine stress management
- Review all pertinent safety concerns
- Consider patient preferences and cost constraints
Acupuncture
The art of acupuncture has been used for millennia in Eastern cultures for treating a range of illnesses including mental disorders. Research in the area of unipolar depression has revealed mixed but mainly positive results. A two-part clinical trial has been conducted examining the safety, effectiveness, and acceptability of adjunctive acupuncture in the treatment of hypomania and depression associated with bipolar disorder (Dennehy et al., 2009). In the first study 20 patients experiencing symptoms of mood elevation were given targeted acupuncture (points specific to symptoms) versus ‘sham’ acupuncture (non-acupoint needling) over 12 weeks, while for 26 patients experiencing symptoms of depression targeted acupuncture was compared to (off the meridian) acupuncture for nonpsychiatric health concerns over 8 weeks. While the results revealed that acupuncture treatment reduced symptoms (mood elevation in Study I, depression in Study II), a non-significant difference occurred between treatments, with all patients experiencing improvement over the course of study.

Conclusions
Bipolar disorder may potentially be effectively managed using an integrative approach. While use of select nutrients combined with medication in BD is supported by strong evidence, to date herbal medicines have not been adequately evaluated. Surprisingly, other CAM modalities potentially used by bipolar patients are currently not supported by clinical trial evidence. Interventions such as manual therapies (e.g. acupuncture, massage), dietary modification, meditation, or mind-body practices (e.g. tai chi or yoga) are supported by some evidence in psychiatric disorders; however, their beneficial effects in patients diagnosed with BD are currently unknown or inconclusive. However, while these interventions may not be ‘directly effective’ in the treatment of BD, they have general beneficial effects on physical and mental health and quality of life, are probably associated with reduced stress and improved functioning in general, and thus may potentially reduce the severity of bipolar symptoms and risk of relapse. We hope this article illuminates the current evidence of CAM for the integrative management of BD, and may encourage further discussion and research in this area.
Conflict of Interest

No conflicts of interest noted.

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