Pharmacological treatment of bipolar depression

Gustavo H. Vázquez, Leonardo Tondo, Juan Undurraga, Rodolfo Zaratiegui, Valerio Selle & Ross J. Baldessarini

SUMMARY

Bipolar depression remains a major challenge for psychiatric therapeutics. It is associated with disability and excess mortality, and accounts for three-quarters of the time spent in morbid states by treated patients with bipolar disorder. Major limitations of research on the treatment of depression in bipolar disorder include a paucity of short-term and lack of long-term trials, probably reflecting concern about inducing mania. In addition, polytherapy with multiple drugs appears to be widespread, but it is virtually untested for efficacy and safety. Here, we summarise the evidence concerning efficacy of treatment of bipolar depression with antidepressants, mood-stabilising anticonvulsants, lithium and second-generation antipsychotics.

LEARNING OBJECTIVES

• Gain critical appreciation of the paucity of research on the treatment of bipolar depression.
• Rationally balance the benefits and risks of using antidepressants in patients with bipolar disorder.
• Assess the evidence supporting a range of research-based treatment options for bipolar depression.

DECLARATION OF INTEREST
None.

Bipolar disorder is a disabling, largely episodic, recurrent illness associated with severe functional impairment, psychiatric and somatic comorbidity, and premature mortality from both suicide and medical illnesses (Baldessarini 2010a,b; Sanders 2010). Lifetime prevalence of bipolar disorder in the general population, taking into account both type I (with mania) and type II (with hypomania) disorder, is at least 1–2% (Merikangas 2011), and up to 10% if broad diagnostic criteria include major depression with subthreshold hypomania (Zimmermann 2009).

Depressive, dysthymic and mixed (dysphoric/agitated) states contribute to the total illness burden in bipolar disorder. These morbidity factors are strongly associated with, and predicted by, similar first lifetime episodes (Baldessarini 2010a, 2012). In several longitudinal studies of the treatment of bipolar disorder by community standards, the mean proportion of weeks in morbid states was 68% and three-quarters of that unresolved morbidity is accounted for by depressive illness (Baldessarini 2010a; Tondo 2013). Depressive components of bipolar disorder are associated not only with a high proportion of unresolved (treatment-resistant) morbidity, but also with psychiatric and medical comorbidity, disability and mortality from suicide in young patients and from co-occurring medical disorders in older patients – all resulting in very high levels of clinical and economic burden for patients, families and society (Osby 2001; Tondo 2007; Crump 2013).

Despite the high prevalence and major clinical, public health and economic significance of depression in bipolar disorder, few treatments have proved to be highly and consistently effective in acute episodes, and there is even less evidence of means of providing substantial long-term protection from recurrent episodes. In particular, there is considerable controversy about the value and risks of antidepressant drugs in treating bipolar depression (Pacchiarotti 2013; Vázquez 2013). In turn, lack of highly effective treatments encourages widespread empirical trials of combination therapies (polytherapy) that are largely untested for effectiveness and safety.

It is likely that the paucity of therapeutic studies for bipolar depression reflects a broadly accepted view that major depression is similar in its clinical characteristics as well as its responses to treatment in patients with bipolar as well as unipolar mood disorder (Baldessarini 2013a). Instead, there is considerable evidence that bipolar and unipolar mood disorders differ in many ways, including family history, gender distribution, age at onset, long-term diagnostic stability, episode duration, recurrence rates and response to treatment (Baldessarini 2013b).

These considerations indicate that bipolar depression remains a leading clinical problem and one of the most critical unsolved challenges.
for contemporary psychiatric therapeutics (Baldessarini 2010b, 2013a). We briefly discuss here the status of research evidence on drugs available to treat the depressive components of bipolar disorder, relying heavily on a series of recently reported meta-analyses (Undurraga 2012a; Tondo 2013; Vázquez 2013; Selle 2014). We focus on antidepressants as well as antimanic anticonvulsants and lithium (as putative mood-stabilising agents), and on emerging evidence for modern, second-generation antipsychotics.

Summaries are provided of efficacy estimates (response rate ratio (RR), usually based on ≥50% reduction in symptom ratings) after randomisation to the drugs discussed vs. placebo in controlled, randomised, monotherapy trials lasting an average of 8 weeks.

**Antidepressants**

**Bipolar depression**

The apparent ease and relative safety of treatment of major depressive episodes with antidepressants, combined with the strong wish of patients and their clinicians to minimise or avoid depression, has made antidepressants the leading treatment (Baldessarini 2008). As noted, it is likely that this circumstance reflects uncritical acceptance of the broad concept of major depression (Baldessarini 2013a). The tendency to view all forms of depression as similarly responsive to particular treatments has probably discouraged pharmaceutical manufacturers from conducting additional therapeutic trials specifically designed to test for treatment effects in depressive, dysthymic and mixed states of bipolar disorder, and to differentiate responses in bipolar type I and II disorder. An important contributing factor is that a known diagnosis of bipolar disorder has been an exclusion criterion from most controlled trials of antidepressants. This exclusion may be driven by a desire to avoid inducing mood switching or other potentially dangerous adverse behavioural effects, and associated potential legal liability, whether such concerns are warranted or not. The outcome is that antidepressants are both the most commonly employed treatment for bipolar disorder (Baldessarini 2008) and one of the most controversial (Pacchiarotti 2013; Vázquez 2013). Many experts call for caution in the use of antidepressants, discouraging their use in monotherapy and, if needed, recommend prescribing only in combination with mood-stabilising agents or second-generation antipsychotics (Pacchiarotti 2013).

Current expert opinions about the value and potential risks of antidepressants to treat bipolar depression are greatly constrained by the limited and inconsistent research-based information that is available despite more than half a century of research and clinical use (Gijssman 2004; Vázquez 2011, 2013; Sidor 2012; Baldessarini 2013a; Pacchiarotti 2013; Yatham 2013; Tondo 2014). Therapeutics research is very limited with regard to acute bipolar depression, and nearly lacking with respect to potential long-term benefits and risks (Ghaemi 2008; Pacchiarotti 2013). Moreover, treatments for important features of bipolar disorder that are depressive (dysthymia, dysphoric mixed-states) but that do not represent acute major depressive episodes are especially poorly studied. Working out optimal clinical procedures to manage depressive components of bipolar disorder requires major attention, with considerable urgency with respect to bipolar type II disorder, in which depression is the main clinical concern and in which suicide rates are about as high as in bipolar type I disorder (Tondo 2007).

**Efficacy in depressive episodes**

Well-designed, controlled monotherapy trials focusing on the efficacy of antidepressants for acute bipolar depression are surprisingly rare, variable in size and quality, and have yielded notably inconsistent findings (Vázquez 2011, 2013; Sidor 2012; Baldessarini 2013a; Pacchiarotti 2013; Yatham 2013; Tondo 2014).

Two large trials are often cited as providing compelling support for the lack of efficacy of antidepressant treatment in acute bipolar depression. They call for comment owing to designs that may limit interpretation of their findings. The problem is that third arms of trials often represent secondary interests and are often smaller than the main arms (usually test drug of commercial interest vs. placebo).

The first study, not a monotherapy trial, found no additional achievement of sustained remission of depressive symptoms by the addition of an antidepressant (paroxetine or bupropion) to a mood stabiliser (Sachs 2007). The second study randomised a small sample of patients with bipolar depression to paroxetine in an 8-week trial designed primarily to test the efficacy of quetiapine. Quetiapine (n = 492 patients) was statistically superior to placebo (n = 126 patients), but did not show a dose-dependent difference between 300 and 600 mg/day, whereas paroxetine (n = 122 patients) was not superior to placebo (McElroy 2010). In contrast, a recent meta-analysis including these and other relevant trials gives some support to the possible efficacy of antidepressants (Vázquez 2013) (Box 1).
There has been particular concern about pathological activation of mood and behaviour during treatment with antidepressants, stimulants or other mood-elevating drugs (e.g. corticosteroids) – either as a risk specific to having overt or potential bipolar disorder or as a psychotic effect. Risks specific to patients with bipolar type I disorder include potentially severe reactions involving mania, psychosis, aggression or irresponsible risk-taking, with associated liability concerns for clinicians and investigators. However, the frequency and severity of such reactions as well as the effects of measures that might limit such risks (e.g. co-treatment with a mood-stabilising or antimanic agent) remain unresolved matters requiring further research.

In a comprehensive review, we found that risk of spontaneous mania without antidepressants was high (averaging 13.8%), but that additional risk associated with antidepressant treatment increased risk by only 1.5% (Tondo 2010) (Box 2). In addition, evidence from randomised trials indicates that antidepressants vary significantly in their association with mood switching, which appears to be especially high with tricyclic antidepressants and the serotonin-noradrenaline reuptake inhibitor (SNRI) venlafaxine (Tondo 2010).

Available information about the epidemiology of mood-switching rates in patients is surprisingly limited. It does not include clear quantitative and qualitative distinctions between bipolar type I and type II disorder. In addition, there is a failure to report exposure times, estimates of risk-per-time or to define the time course of mood shifts with v. without antidepressant treatment.

**BOX 1** Efficacy of antidepressants for bipolar depression

A recent meta-analysis of ten randomised, placebo-controlled, monotherapy trials of antidepressants in acute bipolar depression (Vázquez 2013) found a highly significant pooled difference favouring antidepressant treatment (bipolar response rate ratio (RR) 1.43, 95% CI 1.11−1.84, z-score = 2.76, P = 0.006), with an estimated number needed to treat of 6.2 (95% CI 3.6−6.7); the addition of three recent trials to the meta-analysis supported the same conclusion (Selle 2014). Also notably, the pooled antidepressant/placebo RR in bipolar depression was not less than that found in a comprehensive meta-analysis of 122 randomised controlled trials in unipolar major depression (unipolar RR = 1.42 (95% CI 1.38−1.48), Undurraga 2012a) (Table 1). There also was no appreciable difference in responses between patients with bipolar and unipolar depression compared directly in the same trials (Vázquez 2011).

<table>
<thead>
<tr>
<th>BOX 2 Risk of mood switching</th>
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A comprehensive review of studies comparing spontaneous and antidepressant-associated mood switching into mania or hypomania in patients with bipolar type I or II depression found average rates of 13.8% v. 15.3%, without v. with an antidepressant, indicating a small effect of antidepressants but a high risk of spontaneous switching (Tondo 2010). Another review (Sidor 2012) found a pooled rate of switching of 8.0% in short-term, controlled treatment trials for bipolar depression, with a trivially greater risk with antidepressants (response rate ratio RR = 1.03, 95% CI 0.70−1.52). More recently, a review found an 8-week risk of 4.7% (95% CI 1.8−7.5) with placebo, and somewhat lower average risk of 3.7% (95% CI 2.1−5.3) after randomisation to a mood stabiliser or antipsychotic in 15 trials for bipolar depression (Selle 2014). These observations suggest that concerns about risks of mood switching specific to antidepressants in patients with bipolar disorder may be greater than is warranted, although risks of spontaneous mania and hypomania, especially without a mood-stabilising agent in place, are substantial in patients with bipolar depression.

**Mood switching**

There has been particular concern about pathological activation of mood and behaviour during treatment with antidepressants, stimulants
Lack of information about time-at-risk makes it difficult to distinguish between spontaneous and antidepressant-associated switch risk. Importantly, too, prospective, randomised comparisons of switching rates with v. without ongoing mood-stabilising or antipsychotic treatment are lacking. Meta-analysis even suggests, paradoxically, that risk of mood switching may be greater with a mood stabiliser included than with an antidepressant alone (Tondo 2010). However, such findings arise almost entirely from observations involving clinically selected treatments that surely involve confounding by indication, or use of mood stabilisers where risks are considered to be elevated based on history or current behaviour (Tondo 2010). Clinical and liability concerns about risk of mood switching (even if mainly spontaneous) contribute to the routine exclusion of patients with known bipolar disorder from treatment trials of agents with mood-elevating potential (Undurraga 2012b).

Despite this insufficient state of research, there is an evident clinical consensus that antidepressants should be used in patients with bipolar disorder only cautiously, briefly, in limited doses with short-acting agents, and in association with an effective mood-stabilising regimen, while monitoring closely for signs of emerging hypomania. We recommend that antidepressants not be used for bipolar depression if there is a history of mood switching during antidepressant treatment or if there is clinical evidence of current agitation or hypomanic symptoms (Pacchiarotti 2013). Tricyclic antidepressants and SNRIs should be used with extra caution. Such practices are plausible but require prospective testing to support sound clinical practice (Baldessarini 2013b).

**Long-term use of antidepressants**

The potential value and risks of long-term use of antidepressants in both patients with bipolar type I and II disorder, with the intent of limiting the risk of future depressive episodes, remains poorly studied. Again, the lack of research support appears to have little impact on empirical trials of such treatment in clinical practice (Baldessarini 2008, 2013a; Ghaemi 2008). Moreover, the value of long-term antidepressant treatment beyond the initial months of recovery from an index episode, even of unipolar major depression, remains uncertain, with a high risk of findings being confounded by clinically adverse effects of discontinuing treatment rather than due to lack of treatment (Baldessarini 2010c, 2013b). However, long-term use of antidepressants along with mood stabilisers may be appropriate in response to relapses after discontinuing an antidepressant, especially if discontinuation is gradual (Pacchiarotti 2013). It is not possible to recommend a specific duration for safe and effective continuation of antidepressant treatment. Generally, such treatment is managed in accordance with clinical response, and ideally with gradual removal of the antidepressant once the depression has remitted.

Two randomised controlled trials of continuation of antidepressants included patients who already had a favourable short-term response to the same treatments (Leverich 2006; Ghaemi 2010). Both studies suggest that a minority of patients may experience some delay or reduced frequency of depressive recurrences, with an even larger risk of mood switching. Moreover, rapid-cycling patients had an increased number of recurrent episodes with an antidepressant included in their treatment regimen, suggesting cycle acceleration (Ghaemi 2010; Pacchiarotti 2013).

**Anticonvulsants**

In recent decades, anticonvulsants have been widely used to treat bipolar disorder, based mainly on evidence of short-term antimanic effects (e.g. carbamazepine, valproate) and long-term protective effects of lamotrigine to limit risk of recurrent bipolar depressive episodes. However, the evidence of long-term, prophylactic effectiveness of anticonvulsants is less robust (Geddes 2013). Their use has also been encouraged to avoid the complexities of managing bipolar disorder with lithium (Baldessarini 2013a). Other anticonvulsants (e.g. gabapentin, levetiracetam, oxcarbazepine, pregabalin, topiramate) are either inadequately evaluated or have been found to be ineffective (Baldessarini 2013a; Geddes 2013; Reinares 2013). Despite the widespread use of anticonvulsants to treat mania and efforts to afford long-term protective effects in patients, evidence concerning the value and risks of this class of drugs to treat acute bipolar depression is limited, and evidence concerning long-term effects is even more limited (Reinares 2013).

Four small trials involving, in total, fewer than 100 patients suggest some value of divalproex as monotherapy for acute bipolar depression (Muzina 2011) (Table 1, Fig. 1). Lamotrigine may have some effect in acute bipolar depression, based on pooling inconsistent data across individual trials, some of which failed to show superiority over placebo (Table 1, Fig. 1). However, lamotrigine is approved by the US Food and Drug Administration (FDA) only for long-term prophylaxis in borderline personality disorder, with much greater effectiveness against recurrent bipolar depression than mania (Frye
2011). Moreover, the need for slow increments of doses to avoid dermatological reactions makes lamotrigine somewhat impractical for use in acute phases of bipolar disorder. Evidence concerning carbamazepine is very limited and controlled trials for other anticonvulsants are lacking (Reinares 2013; Selle 2014).

**Lithium**

There is remarkably little information concerning the effects of lithium in acute bipolar depression, despite its use as a fundamental treatment for bipolar disorder for more than six decades and being recommended as first-line treatment in some guidelines (Yatham 2013). Its explicit use for bipolar depression is based on a single modern controlled trial, in which it was included as the third arm of a study designed primarily to evaluate quetiapine (Young 2010; Table 1, Fig. 1). However, several small trials from the early 1970s, mostly based on crossover designs (usually lithium to placebo), suggest rates of response averaging about 73% in 100 patients with bipolar disorder and depression (Zornberg 1993).

Nevertheless, lithium may have some long-term benefits against recurrent bipolar depression as well as its more prominent effects against mania and hypomania (Baldessarini 2010a). Moreover, lithium appears to substantially reduce the risk of suicide in patients with bipolar disorder (Baldessarini 2006; Cipriani 2013). Some experts, based mainly on research in unipolar major depression, also recommend lithium to augment the effects of other treatments (Yatham 2013).

**Second-generation antipsychotics**

Antipsychotic drugs, including olanzapine combined with fluoxetine, as well as quetiapine and lurasidone, are currently the only medicines with FDA approval for the short-term treatment of acute major depressive episodes in bipolar disorder (Baldessarini 2013a; Selle 2014). Lurasidone received approval in June 2013, so the evidence base is as yet small (Loeble 2014). For the other drugs, responses are modest in adults, and quetiapine may not be effective in adolescent bipolar depression (DeBello 2009). Trials of quetiapine found no dose-dependent differences in efficacy (with 300 v. 600 mg/day), and only the lower dose is explicitly FDA-approved. The combination of olanzapine and fluoxetine produced superior benefits to those associated with lamotrigine in a rare head-to-head comparison (Brown 2006). However, olanzapine alone appears to be less effective than in combination with fluoxetine, although the combination has not been tested against placebo in sufficient trials to exclude the latter hypothesis (Zornberg 1993).

<table>
<thead>
<tr>
<th>Drug (polarity)</th>
<th>Trials, n</th>
<th>RR (95% CI)</th>
<th>z</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Valproate (bipolar)</td>
<td>4</td>
<td>2.08 (1.18–3.65)</td>
<td>2.54</td>
<td>0.01</td>
</tr>
<tr>
<td>Carbamazepine (bipolar)</td>
<td>1</td>
<td>1.84 (1.01–3.34)</td>
<td>1.98</td>
<td>0.05</td>
</tr>
<tr>
<td>Olanzapine + fluoxetine (bipolar)</td>
<td>1</td>
<td>1.84 (1.44–2.36)</td>
<td>4.84</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Lurasidone (bipolar)</td>
<td>1</td>
<td>1.72 (1.33–2.22)</td>
<td>4.15</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Antidepressants (bipolar)</td>
<td>10</td>
<td>1.43 (1.11–1.84)</td>
<td>2.76</td>
<td>0.006</td>
</tr>
<tr>
<td>Antidepressants (unipolar)</td>
<td>122</td>
<td>1.42 (1.38–1.48)</td>
<td>16.3</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Quetiapine (bipolar)</td>
<td>5</td>
<td>1.36 (1.24–1.49)</td>
<td>6.32</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Lamotrigine (bipolar)</td>
<td>5</td>
<td>1.25 (1.07–1.46)</td>
<td>2.81</td>
<td>0.005</td>
</tr>
<tr>
<td>Olanzapine (bipolar)</td>
<td>2</td>
<td>1.25 (1.08–1.44)</td>
<td>3.03</td>
<td>0.002</td>
</tr>
<tr>
<td>Lithium (bipolar)</td>
<td>1</td>
<td>1.12 (0.92–1.44)</td>
<td>1.10</td>
<td>0.27</td>
</tr>
<tr>
<td>Ziprasidone (bipolar)</td>
<td>2</td>
<td>1.02 (0.90–1.17)</td>
<td>0.34</td>
<td>0.73</td>
</tr>
<tr>
<td>Aripiprazole (bipolar)</td>
<td>1</td>
<td>0.88 (0.74–1.04)</td>
<td>0.69</td>
<td>0.49</td>
</tr>
</tbody>
</table>

Data adapted from recent meta-analyses of placebo-controlled monotherapy trials (Undurraga 2012a; Vázquez 2013; Selle 2014). Agents are ranked by drug/placebo response rate ratio (RR). There are additional data supporting the efficacy of lithium in several small, older trials not designed by currently standard methods (Zornberg 1993).

**FIG 1** Forest plot of results of random effects meta-analyses of placebo-controlled monotherapy trials of treatments for acute bipolar depression. Based on data summarised and references in Table 1. Numbers of reported trials for each treatment are shown in parentheses. Data are pooled drug–placebo response rate ratios (RRs, with 95% CIs) from 34 randomised placebo-controlled trials of antidepressants (10 trials), mood-stabilising anticonvulsants (10 trials), second-generation antipsychotics (13 trials) or lithium (1 trial) in acute bipolar depression. Horizontal bars are computed CIs; vertical dashed line is the null value of RR = 1.00. Aripiprazole, lithium and ziprasidone did not separate statistically from placebo, and effects of other, apparently effective, drugs are not clearly differentiated owing to overlapping CIs, thus precluding ranking by apparent efficacy.
Antidepressants, although widely used despite lack of explicit FDA approval for bipolar depression (even though ambiguously approved for major depressive episodes), have inconsistent support from controlled trials, as discussed earlier. In addition, lithium carbonate lacks explicit FDA approval for the treatment of acute depression in bipolar disorder, and it lacked efficacy in one trial not necessarily optimised to test lithium (Table 1, Fig. 1). Other agents with mood-elevating effects, notably stimulants such as methylphenidate and amphetamines and anti-narcolepsy agents such as modafinil and R-modafinil, as well as centrally active dopaminergic agonists such as pramipexole and ropinirole developed for the treatment of Parkinson’s disease, also remain experimental.

Anticonvulsants
In addition, several anticonvulsants lack specific evidence of efficacy in acute bipolar depression (e.g. gabapentin, levetiracetam) or have some research support but lack regulatory approval (e.g. carbamazepine, lamotrigine, pregabalin, valproate) (Table 1, Fig. 1).

Second-generation antipsychotics
Second-generation antipsychotics that appear to be ineffective or lack adequate testing for bipolar depression include aripiprazole, clozapine, iloperidone, paliperidone, risperidone and ziprasidone. Older antipsychotics are usually avoided in bipolar disorder owing to their adverse effects.

Other pharmacological treatments
The possible value of other pharmacological treatments has been considered, including calcium channel blockers, anticholinesterases, omega-3 fatty acids and other ‘nutriceuticals’, as well as exogenous thyroid hormones, but they require further testing in bipolar depression (Poon 2012). Promising findings have been recently reported for the glutamate N-methyl-D-aspartate receptor antagonist ketamine, which is inconvenient to administer (orally inactive) and short-acting. Similar agents, including S-ketamine, memantine and riluzole, remain to be tested and developed into clinically practical treatments with sustained benefits and, ideally, oral activity (Mathews 2013).

Non-pharmacological treatments
Non-pharmacological treatments may also be of value, or remain experimental. In particular, psychotherapies continue to be used widely even without adequate evidence of efficacy in bipolar depression, probably by hopeful analogy to the proven value of methods such as cognitive–behavioural and interpersonal psychotherapies in...
unipolar major depressive disorder (Schaub 2013) and evidence that psychosocial interventions augment mood stabilisation in bipolar disorder (Prasco 2013). Intense light therapy and sleep deprivation, too, are plausible candidate treatments but require adequate testing in bipolar depression.

Electroconvulsive therapy is probably effective in bipolar depression (Medda 2013). Vagal nerve stimulation is FDA-approved for otherwise treatment-resistant depression without specification of illness polarity (Baldessarini 2013a). Repeated transcranial magnetic stimulation and various forms of electrical stimulation of the brain remain experimental (Nierenberg 2008).

Conclusions

This overview was stimulated by the fact that depression in bipolar disorder is a clinically and economically burdensome, and sometimes dangerous, condition that presents major therapeutic challenges. Episodes of major depression in bipolar disorder, as well as dysthymia and dysphoric/agitated mixed states, are major contributors to residual morbidity, disability and excess mortality, even with treatment.

We summarised the limited research evidence of efficacy for several treatment modalities, finding encouraging results with some mood-stabilising anticonvulsants and second-generation antipsychotic drugs. However, a critical conclusion is that placebo-controlled trials of all plausible treatments in acute bipolar depression remain very scarce, notably including assessment of the oldest and best-established mood-stabilising agent, lithium. The value and risks of antidepressants in acute bipolar depression remain controversial, and research results are inconsistent. Nevertheless, trials that were identified yielded evidence of significant overall efficacy of antidepressants in bipolar depression that, remarkably, was not less than that found in unipolar depression (Table 1).

Moreover, adequately designed assessments of the value and risks of sustained, long-term treatment in bipolar disorder with antidepressants – either alone or more often in combination with mood-stabilising or antipsychotic treatments, with prophylactic intent – remain rare and insufficient to guide clinical practice. The paucity of compellingly effective alternative treatments encourages continued study of antidepressants in bipolar depression, and the value and potential added safety of their use in combination with mood-stabilising or antipsychotic agents require adequate testing. It is inevitable that lack of highly effective treatments for bipolar depression encourages widespread empirical applications of polytherapy, even though these remain largely untested for effectiveness and safety.

Further studies are required to advance therapeutic practices, despite difficulties that may be encountered, including concerns for potentially dangerous behavioural activation during antidepressant trials. There is also uncertain commercial interest in bipolar depression, as distinct from large and proven markets represented by unipolar major depressive and anxiety disorders, as well as concern for risk of inducing mania (which is unlikely with antipsychotic agents). Another important but unresolved question is whether particular aspects of psychopathology – including mild, subsyndromal hypomanic features or elements of mixed states that would not meet currently widely used but narrow diagnostic criteria – are predictive of poor responses to antidepressants in depression, as has been proposed recently. Moreover, it remains unclear whether such an effect would represent a characteristic of a syndrome type or a consequence of current agitation.

Also needed is further clarification of the efficacy and safety of antidepressants in varying doses for different forms of depressive morbidity in bipolar type I and II disorder, as well as for the emerging bipolar spectrum of disorders marked by recurrent depression and mild hypomanic features. Specifically, head-to-head comparisons are needed to compare the short- and long-term efficacy and safety of antidepressants and other medicines found to be effective in bipolar depression against each other and against lithium, administered both in monotherapy and in controlled combinations. Important additional concerns include whether some agents proposed for treating bipolar disorder may worsen aspects of mood, functional status or general health (e.g. excessive sedative effects or weight gain, metabolic syndrome).

In summary, we continue to be struck by the disparity between the seriousness of bipolar depression as an unresolved therapeutic challenge and the limited research effort that has been given to its experimental and clinical treatment over the past half-century of research into major mood disorders.

Funding

This work is supported by a Josep Font Research Grant from the Hospital Clinic of Barcelona to J.U.; by a grant from the Aretæus Association of Rome and the Lucio Bini Private Donors Research Fund to L.T.; and by a grant from the Bruce J. Anderson Foundation and the McLean Private Donors Research Fund to R.J.B.
References


Treatment of bipolar depression


MCQs

Select the single best option for each question stem.

1 Regarding treatment of bipolar depression with antidepressants:
   a most expert guidelines recommend them as first-choice monotherapy
   b they should always be avoided
   c they are more effective long-term than short-term, especially when added to lithium or an anticonvulsant
   d in practice, they are the most prevalent treatment provided to patients with bipolar disorder
   e they are about as well studied in bipolar as in unipolar major depression.

2 Which of the following has been associated with greatest risk of manic/hypomanic mood switching:
   a tricyclic antidepressants
   b selective serotonin reuptake inhibitors
   c monoamine oxidase inhibitors
   d venlafaxine
   e both a and d.

3 Which has yielded favourable number needed to treat estimates of <6:
   a olanzapine plus fluoxetine
   b olanzapine alone
   c lamotrigine
   d paroxetine
   e both a and c.

4 Which lacks evidence of efficacy in acute bipolar depression:
   a aripiprazole
   b paliperidone
   c gabapentin
   d only a and c
   e all of the above.

5 Which of the following statements is false:
   a therapeutics research is very limited regarding acute bipolar depression and nearly lacking for long-term, prophylactic benefits and risks
   b only 4 small trials (<100 patients) support the apparent value of divalproex as monotherapy for acute bipolar depression
   c there is little information concerning the effects of lithium in acute bipolar depression, despite its use as a fundamental treatment for bipolar disorder for more than six decades
   d quetiapine 600 mg/day was significantly more effective than 300 mg/day in at least 2 controlled trials
   e beneficial effects are not a universal or class effect of all modern antipsychotic agents.
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Access the most recent version at DOI: 10.1192/apt.bp.113.011460

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