Effect of Pharmacological and Neurostimulation Interventions for Cognitive Domains in Patients with Bipolar Disorder: A Systematic Review and Network Meta-Analysis of Randomized Controlled Trials

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Introduction: The priority of interventions to alleviate cognitive deficits in patients with bipolar disorder (BD) is inconclusive. We systematically evaluate the efficacy of pharmacological or neurostimulation interventions for cognitive function in BD through a network meta-analysis.

Methods: The PubMed, PsycINFO, Embase, and Cochrane Library databases were searched from database inception to September 30, 2021. Following PRISMA guidelines, all eligible studies were randomized controlled trials of adult bipolar patients that provided detailed cognitive outcomes. Studies were excluded if participants limited to comorbid substance use disorder or the intervention was a psychotherapy. Network meta-analysis comparing different interventions was conducted for 8 cognitive domains. Partially ordered set with Hasse diagram was used to resolve conflicting rankings between outcomes. The study was pre-registered on PROSPERO database (CRD42020152044).

Results: Total 21 RCTs including 42 tests for assessing intervention effects on cognition were retrieved. Adjunctive erythropoietin (SMD = 0.61, 95% CI = 0.00–1.23), Withania somnifera (SMD = 0.58, 95% CI = 0.03–1.13), and galantamine (SMD = 1.22, 95% CI = 0.10–2.35) was more beneficial for attention, working memory, and verbal learning in euthymic BD patients than treatment as usual, respectively. Hasse diagram suggested ranking of choice when multiple domains were combined.

Conclusion: Considerable variability in measurements of cognitive domains in BD was observed, and no intervention resulted in superior benefits across all domains. We suggested interventions priority can be tailored according to individual patients’ cognitive deficits. As current findings from relatively small and heterogeneous dataset, future trials with consensus should be applied for building further evidence.

Keywords: cognitive function, systematic review, network meta-analysis, Hasse diagram, partially ordered set

Introduction

According to the Global Burden of Disease Study 2013, bipolar disorder (BD), a major psychiatric condition characterized by early onset and chronicity, results in considerable years lost due to disability. The clinical course of BD consists of recurrent episodes of mania and depression interspaced by euthymia, and approximately 40% to 60% of cases are associated with cognitive deficits, even in remission status. Cognitive impairment in patients with BD is a susceptibility factor for...
BD recurrence and negatively affects both social functioning and inter-episode recovery. Therefore, cognitive dysfunction is increasingly recognized as a critical symptom of BD.

In a systematic review from Miskowiak et al., the findings regarding intervention efficacy on cognition in BD were overall disappointing or preliminary. Moreover, there were several major methodological limitations to overcome in future evaluation. Conventional pharmacotherapies as treatment as usual (TAU) for BD, which include lithium, anticonvulsants, antidepressants, and atypical antipsychotics, have varying effects on cognitive function. Little evidence supports lithium treatment improves cognitive performance in patients with BD, and some studies included in a literature review reported the opposite. The effects of anticonvulsants on cognition in BD have also not been extensively studied. Some evidence suggests that atypical antipsychotics and antidepressants improve cognitive performance in patients with schizophrenia and major depressive disorder by alleviating psychosis or mood-related symptoms; however, few studies have indicated similar benefits for patients with euthymic BD and more long-term studies are needed to better understand the impact of atypical antipsychotics on BD patients’ neurocognitive functioning. Given the stagnant state of research on targeting cognitive function in BD through first-line TAU, studies are beginning to examine the potential of cognitive enhancement agents adjunctive to TAU to prevent or alleviate cognitive deficits. Examples include cholinesterase inhibitors, antioxidants, and neurostimulations (eg, transcranial magnetic stimulation [TMS] and transcranial direct current stimulation [tDCS]), which are used in dementia and chronic inflammatory disease and for neuroplasticity enhancement, respectively. Despite the increasing number of studies investigating novel pharmacological or neurostimulation interventions over the past decade, no robust evidence for therapeutics targeting cognitive impairment in BD has emerged.

One of the complexities is the diversity of cognitive impairment in BD, for which the International Society of Bipolar Disorder (ISBD) has defined 8 relevant domains: processing speed, attention (or vigilance), working memory, verbal learning, visual learning, problem-solving, social cognition, and executive function. Patients with BD often exhibit varying degrees of impairment across various domains, with the domains of executive function and verbal learning reportedly accounting for the most impairments. Therefore, clinical trials on cognitive function in BD should evaluate intervention effects on multiple domains. Naturally, this would make assessing the efficacy of interventions considerably challenging. To the best of our knowledge, no such systematic comparisons have been conducted to overcome this problem. Furthermore, variation in the tools used to assess these domains in clinical trials creates heterogeneous outcomes, and drawing a generalized conclusion is difficult. In the present study, we evaluated the efficacy of pharmacological and neurostimulation interventions for all 8 cognitive domains outlined by the ISBD. We first conducted a network meta-analysis (NMA) for direct and indirect comparisons across all available treatments and rank the options by their efficacy in each domain. We then used the partially ordered set to sort out the ranking of treatments when considering potentially conflicting cross-domain results of multiple outcomes.

**Methods**

**Systematic Literature Review**

A systematic literature review was performed to look for randomized controlled trials (RCTs) on adult patients with BD that assessed the effect of pharmacological or neurostimulation interventions compared with placebo or TAU on cognitive function. The Embase, PubMed, Cochrane Library, and PsycINFO databases were searched comprehensively from database inception to September 30, 2021. The keywords used were medical subject headings of “Bipolar disorders” combined with “cognition” and filters with RCTs. In this searching process, “cognition” was also replaced with following cognitive domains: “processing speed”, “attention” or “vigilance”, “working memory”, “verbal learning”, “visual learning”, “problem solving”, “social cognition”, and “executive function.” To identify potential additional studies, reference citations were also examined. The review focused on evidence from RCTs because they are regarded as the gold standard trial for evaluating intervention effectiveness and carry a minimal risk of confounding factors. The inclusion criteria were as follows: RCTs of patients with BD aged >18 years that provided detailed descriptions and cognitive measurements of the outcomes. No restrictions on sex or ethnicity were imposed, but studies were excluded if participants limited to BD comorbid with certain substance use disorders or if psychotherapy was the intervention. Duplicate studies were also excluded. Full texts of all included studies were then assessed, and study authors were
contacted for additional information if needed. Conference abstracts were not included. The methods and analysis of the present study were followed the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) guidelines and prespecified and registered on the PROSPERO database (CRD42020152044).

Data Extraction and Bias Risk Assessment
During data extraction, publications describing the same trial were compiled into a single entry to prevent double counting patients. Extracted data included sample size, age, sex, disease diagnosis (bipolar type I or II), mood phase, intervention duration and protocols, and types and time points of the outcome assessments. Two independent reviewers (WY Chen and YC Cheng) performed both the screening and the data extraction, and any discrepancy between reviewers was resolved by a third independent reviewer. Bias risk assessment was conducted according to the recommendations in the Cochrane Handbook for Systematic Reviews of Interventions.16

Outcome Selection and Network Development
A feasibility assessment was performed to determine possible approaches for conducting NMA. Two approaches were considered: (1) Assessing studies evaluating the same cognitive test and (2) assessing studies evaluating the same cognitive domain, regardless of test. Although it is less prone to validity bias, we could not perform a meaningful NMA by using the first approach because of the large variety and seldom overlapping of cognitive tests used as endpoints among review studies. To overcome these problems, we used the second approach, in which the most commonly applied measurements together with other tests for each domain were evaluated. The priority of cognitive tests selected in each domain is summarized in Supplementary Table S1.

Studies have suggested that episodes of acute depression or mania negatively affect cognitive function in patients with BD.17,18 Considering studies of patients with BD in euthymia with those of patients experiencing mood episodes was not appropriate. To avoid introducing bias into the effects of adjunct interventions, we analyzed studies of patients in the euthymic, depressive, or manic phase separately and provided NMA results for each domain by mood status.

Statistical Analysis
We performed NMA for each domain to simultaneously assess the effects of more than 2 treatments. A NMA synthesizes both direct and indirect evidence over an entire network of treatments within one analysis. Given the expected clinical and methodological heterogeneity among the studies, we used a random effects model but switched to using a fixed effects model if the number of included studies was insufficient to estimate the random effects robustly. The standardized mean difference (SMD) of cognitive measurements in the same domain in changes from baseline to the end of follow-up was the outcome variable. The ranking of treatment formats was estimated according to the surface under the cumulative ranking area (SUCRA), which was based on the estimated random effects models. All analyses were performed using the Network package in Stata, version 14.19

We further used partially ordered set to resolve possible conflicting rankings between cognitive outcomes. The Hasse diagram was used to graphically illustrate the partially ordered set. This statistical method revealed incomparability between treatment rankings or insufficient data for some outcomes.20 We first combined all the outcomes and then deleted them one by one until structured Hasse diagrams, constructed using the hasseDiagram package (version 0.1.3) in R software (version 3.6.1), were formed.

Results
Systematic Literature Review
Figure 1 presents the PRISMA flowchart of studies through the systematic review process. Of the 1475 reference citations retrieved initially, 227 remained after duplicates and irrelevant citations were removed. First-stage screening of the citations identified 44 potentially relevant trials based on their abstracts. After the full texts were accessed for more detailed evaluation, 30 remained for extraction. However, detailed measurement data for 9 trials were inaccessible, leaving 21 RCTs for quantitative analysis. The 9 inaccessible trails were investigating adjunctive effect of N-acetylcysteine (NAC),21 melatonin,22 methylene blue,23 pramipexole,24 repetitive TMS (rTMS),25 docosahexaenoic acid (DHA),26 and atorvastatin27 among euthymic BD patients; and adjunctive tDCS,28 olanzapine,29 for depressed BD patients. Across these 21 trials for quantitative analysis, 42 cognitive tests were used to assess intervention effects on the 8 domains. A summary of the tests applied to each domain
and details on all 21 RCTs are presented in Supplementary Table S1 and Table S2, respectively.

Cognitive Outcomes for Network Development in Euthymic BD

Thirteen of the 21 RCTs reported results related to euthymic BD. Ten out of the 13 trials, comparing for adjunctive interventions, including memantine, rTMS, lurasidone, NAC, tDCS, quetiapine, erythropoietin (EPO), Withania somnifera, intranasal insulin, and galantamine to TAU. Three studies, which compared monotherapies of quetiapine or lithium with placebo, were excluded from the network plot, because of the absence of a common link with other interventions, leaving 10 studies with TAU as the control condition. Except trial for memantine, other 9 studies evaluated processing speed; 9 evaluated verbal learning (adjunctive rTMS, lurasidone, NAC, quetiapine, EPO, Withania somnifera, intranasal insulin, memantine and galantamine); 7 evaluated executive function (adjunctive lurasidone, NAC, tDCS, quetiapine, EPO, intranasal insulin, and galantamine); 6 evaluated working memory (adjunctive rTMS, lurasidone, NAC, quetiapine, EPO, Withania somnifera); 5 evaluated visual learning (adjunctive rTMS, lurasidone, tDCS, memantine and intranasal insulin) and attention (adjunctive rTMS, lurasidone, memantine, quetiapine, EPO); 2 evaluated social cognition (adjunctive rTMS and Withania somnifera), and 1 evaluated problem-solving (adjunctive rTMS). Among the 10 adjunctive trials, the number of patients, mean age, and percentage of male patients in each study ranged between 13 and 175, 28 and 49 years, and 32.4% and 60%, respectively. The timing of cognitive assessment varied between 2 and 24 weeks after baseline assessment. The widely used criteria defining euthymic BD among the studies were Young Maria Rating Score (YMRS) ≤ 14 or Montgomery Aberg Depression Rating Scale (MADRS) ≤ 18.

Figure 2 shows that positive outcomes were observed for working memory, verbal learning, and attention. The forest plots present the SMD for each study compared with TAU. Out of the interventions used in the 6 studies (209 patients) that evaluated working memory, Withania somnifera was the only one to demonstrate significant benefits in this domain relative to TAU (SMD = 0.58, 95% CI = 0.03–
1.13; Figure 2A). Out of the interventions used in the 9 studies (443 patients) that evaluated verbal learning, adjunctive galantamine was the only one to demonstrate benefits in this domain relative to TAU (SMD = 1.22, 95% CI = 0.10–2.35; Figure 2B). Out of the interventions used in the 5 studies (314 patients) that evaluated attention, only adjunctive EPO suggested improvements (SMD = 0.61, 95% CI = 0.00–1.23; Figure 2C) in this domain relative to TAU.

According to the SUCRA results for the remaining domains (processing speed, visual learning, problem-solving, social cognition, and executive function), we revealed the potential benefit of Withania somnifera and EPO in processing speed and executive function, respectively. In addition, rTMS also ranked highly in visual learning domain. However, for these five cognitive domains, none exhibited significant effects compared with TAU (Supplementary Figure S1 and Figure S2).

**Figure 2** Network analysis of adjunctive interventions for specific cognitive domains in euthymic bipolar disorder. (A) Outcome of working memory. (B) Outcome of verbal learning. (C) Outcome of attention. The size (area) of the nodes is proportional to the number of patients in each intervention. Line widths are proportional to the number of patients in trials providing direct comparisons between the nodes. The right parts of the forest plots indicate the standardized mean differences for each study in direct comparisons with treatment as usual.

Abbreviations: TAU, treatment as usual; rTMS, repetitive transcranial magnetic stimulation.
Through the systematic review, majority of other candidate pharmacological compounds cannot include in quantitative analysis revealed the disappointing results. For example, add-on melatonin did not seem to affect cognition in patients with severe mental illness including BD.\textsuperscript{22} One study suggested the beneficial effect of methylene blue in residual mood symptoms in BD, but the effects on cognitive symptoms were not significant.\textsuperscript{23} The trail for pramipexole in BD revealed negative findings in primary cognitive outcomes, and only suggested the potential benefit in subgroup of strictly defined euthymic subjects.\textsuperscript{24} The trials for atorvastatin and DHA for cognitive deficits in BD also revealed the negative results.\textsuperscript{26,27} The only promising findings came from the most recently published pilot study for rTMS targeting cognitive function in BD, which indicating a significant group by time interaction in verbal learning domain.\textsuperscript{25}

**Cognitive Outcomes for Network Development in Patients with BD Experiencing Depressive or Manic Episodes**

Adjuvant creatine monohydrate, high- and low-frequency rTMS, infliximab, tianeptine and mifepristone were the interventions used (compared with TAU) in the 6 studies including patients with BD experiencing depressive episodes.\textsuperscript{43–48} Across these studies, attention and social cognition were not measured, and only one evaluated problem-solving. Across the 5 domains included in the NMA (Supplementary Figure S3), none of the adjunctive therapies revealed significant benefits compared with TAU. A pilot study assessing effect of adjunctive tDCS on cognition in depressed BD patients revealed no changes in cognitive scores.\textsuperscript{28} Olanzapine augmentation therapy in a trial among major depressive disorder and BD patients under depression did not improve their cognitive tasks neither.\textsuperscript{29}

Trails of olanzapine and risperidone were the two medications that included patients with BD undergoing acute manic episodes with evaluated cognitive functions.\textsuperscript{49,50} Perlis et al reported no significant differences in Cognitive Test of Delirium between olanzapine and risperidone in patients with BD undergoing acute manic or mixed episodes.\textsuperscript{49} Shi et al noted improvements in the cognitive component of the Positive and Negative Syndrome Scale with olanzapine treatment compared with placebo,\textsuperscript{50} while this improvement was associated with improvement in acute mania state. Because the cognitive measurements were non-specific, the data were not pooled for analysis.

**Partially Ordered Set with Hasse Diagram**

Because our NMA contained multiple cognitive outcomes, we used the Hasse diagram for partially ordered set to resolve conflicting treatment rankings between different outcomes. Structured Hasse diagrams were generated only for euthymic BD with combined outcomes across certain domains (Figure 3). Compared with TAU, adjunctive intranasal insulin, NAC, and EPO ranked highly for combined processing speed, verbal learning, and executive function (Figure 3A). Similar rankings were noted when outcomes across 2 domains (among processing speed, verbal learning, and executive function) were considered (Figure 3B–D). Adjunctive galantamine had the lowest ranking for combined outcomes of processing speed and executive function (Figure 3C). Otherwise, adjunctive quetiapine had the lowest ranking compared with TAU, which suggests that add-on second-generation antipsychotics exacerbate cognitive deficits in BD under TAU.

**Risk of Bias Assessment**

The scores for bias risk for the 21 included RCTs are presented in Supplementary Table S2 and summarized in Figure 4. In general, the studies were of satisfactory quality, with an average of >4 of the 7 assessed categories having low risk of bias. Selective reporting and allocation concealment were the categories with the best and worst scores, respectively. However, in 3 studies\textsuperscript{35,45,46} the risk of bias was unclear in most categories. As none of our NMA contains treatment loop, there is no need for assessing consistency in the evidence.

**Discussion**

To the best of our knowledge, this NMA is the first to compare different interventions for cognitive function in BD and to find a common matrix demonstrating the rankings of interventions for various cognitive outcomes. The topic of cognitive deficits in BD is important, but current manuscript revealed from relatively small and heterogeneous dataset. We applied reasonable methodology, but ultimately conclusions still cannot be drawn. According to our findings, we suggested the adjunctive EPO, *Withania somnifera*, and galantamine benefited attention, working memory, and verbal learning compared with TAU, respectively. Besides, the SUCRA results indicated that rTMS ranked highly for visual learning compared with TAU, but its benefits were not significant. The Hasse diagrams
Figure 3 Hasse diagrams of adjunctive interventions for cognitive outcomes in euthymic bipolar disorder. (A) Outcomes of processing speed, verbal learning, and executive function. (B) Outcomes of processing speed and verbal learning. (C) Outcomes of processing speed and executive function. (D) Outcomes of verbal learning and executive function.

Abbreviations: TAU, treatment as usual; rTMS, repetitive transcranial magnetic stimulation; tDCS, transcranial direct current stimulation.

Figure 4 Summary of the risk of bias assessment.
showed that for combined outcomes in processing speed, verbal learning, and executive function, intranasal insulin, NAC, and EPO were ranked highly and adjunctive quetiapine was ranked less compared with TAU. Future homogeneous trials still need to build up the evidence.

Galantamine, which is approved for the treatment of Alzheimer disease, inhibits cholinesterase and potentiates nicotinic neurotransmission. Early case reports, when multiple domains were approached, which maintains the multidimensional characteristics of the underlying decision problem and can reveal the uncertainties, also presents indisputable evidence. Finding a treatment that is superior across all domains may be impossible. Instead, interventions can be tailored to specific domains of impairment.

As shown in Figure 3, when multiple domains were considered, intranasal insulin, NAC, and EPO had higher rankings, whereas quetiapine had a low ranking. Converging lines of evidence indicate that insulin has activity in the brain. Reduced cerebrospinal fluid insulin levels have been reported in patients with Alzheimer disease, and trials for cognitive impaired patients have reported that intranasal insulin demonstrated improvements on memory.

Evidence is emerging for the effectiveness of NAC in treating various psychiatric disorders. Its effectiveness may be due to its glutamatergic modulation and antioxidative activities. A study noted improvements in working memory in patients with psychosis (including schizophrenia and BD) under NAC treatment; however, the present analysis did not exert significant cognitive benefits of NAC among BD patients. Whether the benefits of NAC are limited to psychotic BD or whether larger sample sizes are necessary to reveal significant effects warrants further investigation.

Second-generation antipsychotic treatments for cognitive dysfunction in BD, which mainly work to reduce mood symptoms, may also have direct adverse cognitive effects mediated by their anticholinergic, extrapyramidal, or sedative effects. In the present study, quetiapine had a low ranking in the Hasse diagram, which was consistent with the exacerbated cognition in trial conducted by Rakofsky et al. Because of conflicting results on their efficacies between outcomes, the Hasse diagram rankings for adjunctive Withania somnifera, galantamine, memantine or neurostimulation indicated sparse representation.

This study has several limitations. First, all the included studies were compared with TAU, which consisted of various psychotropic medications; the uneven effects of TAU medications could not be assessed. Second, the small number of studies and small sample sizes prevented us from exploring potentially important factors, such as subtypes or cognitive stages of BD. Therefore, we could not determine the robustness of our findings across different categories of BD. Third, because all the studies came from
a single trial of the intervention, we adopted the fixed effects model. Consequently, the uncertainty of the effect sizes, such as the confidence interval, may be underestimated, and the generalizability of our results needs further validation. In addition, these effect sizes are based on single, generally small studies, and are thus subject to effect size inflation that can occur with smaller studies. Furthermore, because of the limited number of common measurements across studies, the outcome selection strategy was arbitrary. As various cognitive tests were used as endpoints, collapsing them into domain-specific groups introduced some inevitable validity bias. Future establishment of consensus for cognitive measurements in patients with BD is warranted. Finally, the reviewed studies included in our analysis had different lengths of follow-up, and mostly investigated immediate or short-term intervention effects. Those differences amongst studies could indicate as a source of heterogeneity. Future studies with long-term follow-up are necessary to determine whether such effects are sustained over time. As cognitive outcomes in BD is an emerging important issue, we believe that newly published trials will keep presenting new findings in the coming years. The ongoing trials may impact on the current findings of the NMA, especially the ranking of choices in each domain. Therefore, we suggest that updating search and re-analyzing new data should be done regularly in the future to renew the knowledge in this field.

Conclusion
No single intervention demonstrated improvements on cognitive function across all domains; therefore, interventions tailored to each patient’s specific cognitive deficits is preferable. As current review with the relatively small and heterogeneous dataset, future trials with consensus to explore strategies to alleviate cognitive impairment in BD should be given clinical priority.

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The Authors declare that there is no conflict of interest.

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