Current and emerging medications for borderline personality disorder: is pharmacotherapy alone enough?

Highlights

- Many psychotropic agents have been tested for the treatment of BPD symptom domains.

- In most cases, the studies are open-label trials or affected by severe limitations. There is little evidence of efficacy.

- Evidence supports the use of mood stabilizers, second-generation antipsychotics, and omega-3 fatty acids to treat BPD.

- Combining drugs with psychotherapy demonstrates better results than single therapies on several core symptoms of BPD.

- Data on the tolerability of psychotropic agents in BPD are scarce. Specific examinations of each drug’s adverse effects are needed.
Abstract

Introduction
The treatment of borderline personality disorder (BPD) remains an open question for clinicians. There is scarce evidence available. The guidelines' conclusions diverge. Together with these factors, the complexity of BPD generates uncertainty in day-to-day practice. This narrative review aims to provide an overview of advances in BPD treatment and posit a critical opinion based on clinical evidence and practice.

Areas covered
We reviewed clinical trials of the efficacy of the main class of drugs in BPD: antidepressants, mood stabilizers, first-, second-, and third-generation antipsychotics, and other agents (opiate antagonists, clonidine, oxytocin, omega-3 fatty acids). We also included in this review studies on combinations of drugs and psychotherapies.

Expert Opinion
An individualized, tailored pharmacotherapy for BPD that targets the prominent symptom clusters can improve relevant aspects of the clinical picture. However, no medication is indicated to treat the global psychopathology of BPD. Polypharmacy should be avoided or strictly limited. To date, pharmacotherapy alone does not suffice to manage the complexity BPD. Combining medication with psychotherapy may improve specific BPD symptom dimensions. In particular, it may help those aspects that respond slowly or not at all to monotherapy.

Keywords
Borderline personality disorder; pharmacotherapy; psychotherapy; combined therapy; antidepressants, mood stabilizers; antipsychotics
1. Borderline Personality Disorder (BPD): common recommendations for the complex clinical entity

Borderline personality disorder (BPD) is a severe and heterogeneous mental disorder. It is characterized by a pattern of identity diffusion, impulsive behaviors, and chronic instability in affectivity and relationships [1]. Self-injuries and suicidal conduct may occur during critical phases of the disorder. Patients with BPD exhibit impairment in a wide range of functions [2], to a greater extent than patients with other psychiatric disorders [3]. They have long-term difficulties over their lifespans [4,5]. BPD does not suddenly emerge in adulthood. In fact, early precursors, prodromal signs, and processes that confer vulnerability to later personality pathology appear in youth [6,7,8]. A full diagnosis of BPD is as valid in adolescence as it is in adulthood. Borderline personality disorder is the most common personality disorder in clinical populations [9,10]. The lifetime prevalence of BPD has been estimated at around 1.4% in the community [11] and 25% in psychiatric inpatient settings [12]. Two studies based on data from the National Epidemiologic Survey on Alcohol and Related Conditions found higher rates, at 2.7% [13] and 5.9%, respectively [14]. The high prevalence significantly impacts mental health services [15,16]. The complexity of borderline personality disorder requires particular care in diagnostic processes and early therapeutic interventions.

Although several sets of guidelines for the treatment of BPD have been published in the last eighteen years, no medication has obtained an official indication for this mental disorder.

The American Psychiatric Association (APA) [17,18] guidelines supported the role of pharmacotherapy that is oriented at specific symptom domains to treat stated symptoms during periods of acute decompensation and trait vulnerabilities. It identified three symptom dimensions as the targets of drugs: affective dysregulation (including depressed mood, anxiety, anger, and mood lability), impulsive–behavioral dyscontrol, and cognitive-perceptual symptoms. Antidepressants and mood stabilizers were recommended as first- and second-line interventions for affective dysregulation and impulsive behavioral dyscontrol. Antipsychotics (first- and second-generation
antipsychotics) represented the first-line of treatment for cognitive-perceptual symptoms and the third choice for impulsive–behavioral dyscontrol symptoms. Similarly, in the practical and biological guidelines published by the World Federation of Societies of Biological Psychiatry (2007) [19], the main symptomatic targets of pharmacotherapy with different evidence levels were affective dysregulation, cognitive-perceptual symptoms, impulsivity, and anger. Moreover, the authors stated that the effects of pharmacotherapy in these patients may have increased responses in combination with psychosocial interventions. The guidelines from the National Institute for Health and Clinical Excellence (NICE) [20,21] reached rather different conclusions from the preceding guidelines. They stated that drug therapy was not recommended other than to treat mental disorders in comorbidity or to control specific acute symptoms during a crisis and with a short-term prescription [22]. NICE supported the central role of psychotherapy in treating BPD. The Australian National Health and Medical Research Council guidelines (2012) [23] took an intermediate approach. It recognized the role of medications as second-line or adjunctive treatment and confirmed psychotherapy as the first-line treatment. In substantial accordance with the treatment guidelines, two Cochrane systematic reviews [24,25] underlined the importance of combining psychotherapy and medications. Among pharmacotherapy, some evidence focused on second-generation antipsychotics, mood stabilizers, and omega-3 fatty acids. The data did not support the role of antidepressants.

This review aims to provide an overview of the advances in pharmacotherapy and combined therapy for BPD and express a critical opinion based on clinical evidence and practice.

1.1 Antidepressants

In recent years, antidepressants have been widely prescribed in patients affected by BPD, despite a lack of high-quality evidence of their efficacy in the scientific literature [26]. Tricyclic antidepressants (TCAs) and monoamine oxidase inhibitors (MAOIs) have been prescribed progressively less for all psychiatric indications. They have significant side effects, and there is a
high risk of a potentially fatal overdose. These medications have been replaced by selective serotonin reuptake inhibitors (SSRIs) and serotonin and noradrenaline reuptake inhibitors (SNRIs). The overall usage of antidepressants in BPD has decreased recently, and now the prescription of these agents is recommended only for comorbid affective disorder [25].

To our knowledge, no placebo-controlled trials on the efficacy of antidepressants in BPD have been performed since 2010. In general, we observed a decrease in the number of investigations on medications in patients with a diagnosis of BPD in the last nine years. The existing studies often showed significant methodological limitations [27].

Among SSRIs, most RCTs investigated the efficacy of fluoxetine (5 trials) [28-32]. Three studies compared fluoxetine with placebo [28-30]. One study compared fluoxetine with dialectical behavioral therapy (DBT) [31], and one trial compared fluoxetine with the antipsychotic olanzapine and a combination of fluoxetine and olanzapine [32]. A more recent study evaluated the efficacy of fluoxetine in monotherapy in comparison with a combined therapy of fluoxetine and interpersonal psychotherapy adapted to BPD (IPT-BPD) [33]. The results of this study will be discussed in the paragraph on combining drugs and psychotherapy. In the initial trials, fluoxetine was found efficacious in decreasing anger, irritability, and impulsive aggression and in improving global functioning. Unexpectedly, the effect on depression was controversial and questionable. Two studies did not find any effect on depression [28,31], while one study [29] stated a beneficial effect of fluoxetine on depression and anxiety. This discrepancy may be due to a difference in the dose of fluoxetine used in the three samples. In the two studies published by Salzmann et al. and Simpson et al., the maximum dose was 60 mg/day, while in Markovitz’s study, the maximum dose was 80 mg/day. The only RCT on fluvoxamine showed a significant effect on affective instability with a decrease in rapid mood shifts but not on aggressiveness and impulsivity [34]. More recently, sertraline was compared with olanzapine in one RCT [35]. The authors observed that sertraline decreased depression, obsessive symptoms, and hypersensitivity in interpersonal relationships,
while olanzapine was more effective on aggression, paranoid ideation, and self-mutilating behaviors.

There have been only two open trials of SNRIs in the treatment of BPD. The first evaluated the efficacy of venlafaxine [36], and the second investigated the effects of duloxetine [37]. Both drugs showed a reduction of somatic symptoms in BPD [36,37], and duloxetine improved depressive symptoms, impulsivity, outbursts of anger, and affective instability [37].

In summary, the findings of controlled studies of the efficacy of antidepressants in BPD are outdated as they have not been replicated after 2010. The main results were that SSRIs positively impacted affective symptoms. In particular, they helped a depressed mood, anxiety, anger, and impulsive behaviors in BPD patients. Nevertheless, the available evidence is weak on whether to support or exclude the use of SSRIs. Controlled trials of SNRIs are lacking, and there have been no new studies of new antidepressants, like vortioxetine.

The results of RCTs are displayed in Table 1.

1.2 Mood stabilizers

Clinicians have considered mood stabilizers an efficient therapeutic option in treating BPD patients since the 1990s. This class of medications has been found particularly useful in the short-term control of anger and impulsivity in BPD [38]. Although a Cochrane review (2010) [24] found some evidence of the efficacy of mood stabilizers (valproate and lamotrigine) in BPD, since 2010, few trials have been conducted to consolidate the data and reinforce their use in clinical practice [27,22].

Lithium was studied in only one RCT [39] that compared it with desipramine and placebo in BPD patients receiving concomitant psychotherapy. Although lithium showed a major efficacy in improving anger, irritability, and self-injuries, the risk of toxicity and the need for regular blood level controls represented significant limitations to its prescription in BPD. Nowadays, no more evidence is available about lithium carbonate, and there has been no study of lithium sulfate in BPD.
Carbamazepine was tested in one RCT [40] and found not to be significantly superior to placebo on measures of affective symptoms, behavioral dyscontrol, and global assessment. Due to the limited evidence of therapeutic benefits and the risk of adverse effects, carbamazepine is a questionable treatment option for this disorder.

Over the years, the most widely studied mood stabilizer in BPD patients has been valproate. Four double-blind placebo-controlled trials and one unblinded controlled study have been published. Two studies [41,42] reported the efficacy of valproate in reducing impulsive and aggressive behaviors of BPD patients. One study [43] showed the superiority of valproate on interpersonal sensitivity, anger, hostility, and aggressiveness in a sample of BPD patients with comorbid bipolar disorder. A more recent study [44] evaluated the effects of extended-release divalproex in BPD patients treated with four weeks of dialectical behavioral therapy (DBT). The results of this trial are less encouraging because no significant differences between the two groups were observed in the global assessment. The unblinded controlled trial [45] compared valproate with the association of valproate and omega-3 fatty acids (docosahexaenoic acid and eicosapentaenoic acid). The combined therapy was found superior to the single therapy for impulsivity, self-harm, and outbursts of anger.

Another mood stabilizer that was tested in BPD patients was topiramate. Five RCTs were performed on this medication [46-51]. Two short-term [46,47] and two follow-up [48-50] studies showed that topiramate significantly reduced irritability and anger. One study [51] reported the improvement of global functioning, hostility, interpersonal sensitivity, and somatization symptoms in patients affected by BPD and comorbid mood disorders.

The efficacy of lamotrigine was evaluated in four RCTs [52-55]. The findings revealed significant effects on the outcomes of anger, impulsivity, affective instability, and interpersonal dysfunctions [52-54]. However, one study performed by Crawford and colleagues [55] showed that adding lamotrigine to usual care did not result in symptomatic relief and cost-effectiveness balance.
To our knowledge, oxcarbazepine has been tested in only one BPD open-label study [56]. This study reported improvements in anxiety, interpersonal relationships, impulsivity, affective fluctuations, and anger.

In our opinion, more studies on this class of drugs are needed to draw more reliable conclusions. Large-scale double-blind placebo trials should be conducted. Drug-drug comparison trials between different mood stabilizers would clarify their effects on symptom domains and establish whether there are significant differences in efficacy between the drugs. Nonetheless, the available evidence suggests that valproate, topiramate, and lamotrigine are a therapeutic option in treating impulsivity, anger, and affective instability in BPD patients.

The results of RCTs are displayed in Table 2.

1.3 Antipsychotics

Cognitive-perceptual symptoms represent a psychopathological dimension of BPD. In particular, stress-related transient psychotic-like symptoms (paranoid ideation and dissociative symptoms) can frequently occur over the course of the disorder.

Individuals with BPD have increased plasma and cerebrospinal fluid levels of dopamine metabolites, which could explain the transient psychotic and dissociative symptoms.

Before 1990, several studies on the use of first-generation antipsychotics in BPD resulted in controversial findings. These studies concerned flupentixol decanoate [57], loxapine versus chlorpromazine [58], thiothixene [59], and trifluoperazine [60]. After 1990, two RCTs were performed to evaluate the efficacy of haloperidol versus placebo [61] and phenelzine [62]. The results showed that haloperidol promoted the reduction of hostility, irritability, and impulsive-aggressive behaviors in BPD, but it failed to improve global severity and psychoticism. Moreover, haloperidol seemed to be responsible for the onset of depressive symptoms during prolonged treatment.
More recently, more investigations have been published on second (clozapine, risperidone, olanzapine, paliperidone, and ziprasidone) and third generation (aripiprazole) antipsychotics in the treatment of BPD. The use of newer antipsychotics has been preferred in clinical practice because of these medications are better tolerated than first-generation drugs. In fact, new antipsychotics are associated with fewer and milder extrapyramidal adverse effects, a lower risk of tardive dyskinesia, and a considerable improvement in cognitive functions [63]. This class of drugs showed a moderate effect on negative affectivity, disinhibition, and interpersonal dysfunction by improving dorsolateral and orbital frontal cortex activation [64,65].

Among new antipsychotics, clozapine was tested in four open-label studies of patients with BPD [66-69]. All the studies concluded that this medication was useful in reducing impulsivity, affective instability, self-mutilating behaviors, and cognitive-perceptual symptoms. However, these trials had a diagnostic bias due to the inclusion of patients with psychotic symptoms or disorders, which significantly affected the results. In a 2-year observational study that was recently published [70], clozapine was associated with a significant decrease in psychiatric admissions of patients with BPD without cases of serious adverse effects. However, generally, the risk of adverse effects, such as agranulocytosis, is considerable with clozapine. It is particularly dangerous in BPD patients because of their low adherence to treatment prescriptions and blood chemistry tests.

Four open-label studies [71-74] and one double-blind controlled trial [75] evaluated the efficacy of risperidone. The findings from the uncontrolled trials suggested an effect of this drug on impulsivity, aggressiveness, affective instability, and anxiety. The results of the only RCT highlighted the efficacy on psychoticism, paranoid ideation, phobic anxiety, and interpersonal sensitivity [22,75]. Two open-label studies were performed on the active metabolite of risperidone, paliperidone: One concerned the extended-release oral formulation [76], and one was on the long-acting intramuscular formulation [77]. Although their data need to be replicated, they showed that paliperidone improved global and specific BPD symptoms, such as impulsive behavioral dyscontrol and cognitive-perceptual disturbance.
Olanzapine was studied in nine RCTs and one open-label trial. In fact, it is the most thoroughly studied second-generation antipsychotic in BPD. Among RCTs, four trials evaluated the efficacy of this drug versus placebo [78-81]. Two studies compared olanzapine plus DBT psychotherapy versus placebo plus DBT [82,83]. Three drug-to-drug studies investigated the efficacy of olanzapine to sertraline [84], haloperidol [85], and asenapine, respectively [86]. The studies generally agreed that olanzapine improves global functioning, affective instability, anxiety and anger, interpersonal sensitivity, and impulsive-aggressive behaviors in BPD patients. The comparison of olanzapine with asenapine showed olanzapine had a superior effect on paranoid ideation [86]. The comparison with haloperidol did not find a difference between the two drugs on generic behavioral symptoms [85]. However, Shafti and Shahveisi administered olanzapine at a low mean dose (7 mg/day) and haloperidol at a relatively high mean dose (7mg/day). In the open-label randomized trial [87], olanzapine was compared with aripiprazole. Both medications were found effective in reducing global symptoms. Olanzapine did not show any superior effect on symptoms of hostility. It must be noted that a Cochrane systematic review (2010) [24] identified an increase in self-harming behavior when using olanzapine in BPD [24]. In contrast, BPD patients were found less likely to engage in self-mutilating acts when treated with aripiprazole [88-90].

Aripiprazole was found useful in the management of affective dysregulation, depression, impulse control, and paranoid symptoms in three RCTs of BPD patients [87-89] and one open-label trial [91], which tested the efficacy of the drug in augmentation of sertraline in a group of non-responders. The recent protocol for the “Verbal Experiences Response in Borderline Personality Disorder to Aripiprazole Trial Medication – Verbatim” randomized controlled trial [92] aims to inform BPD treatment decisions when the patients present with auditory verbal hallucinations.

Several studies on quetiapine mostly agreed and shared promising findings. Unfortunately, most of these studies were open-label [93-99], and only two RCTs were published more recently [100,101]. Quetiapine showed effects on a wide range of BPD symptoms, including impulsive-aggressive behaviors, anger, and affective instability, and it had positive effects on anxious-depressive
symptoms and general psychopathology [22,25]. The recently approved antipsychotic asenapine was studied in one open-label study [102] and one controlled trial, which compared asenapine with olanzapine [86]. Asenapine was found effective in improving affective instability, impulsivity, and cognitive symptoms, and it was superior to olanzapine on affective instability [86]. Ziprasidone was tested in one open-label and one controlled study [103,104]. Neither investigation found any evidence of the efficacy of this antipsychotic on symptoms of BPD. Concerning inhaled loxapine, there were only case reports [105,106] and one retrospective analysis [107] available. The results suggested that inhaled loxapine may be useful for the emergency treatment of agitated patients with BPD and other personality disorders [108]. To date, we have no data on the newest antipsychotics, lurasidone, brexpiprazole, and cariprazine, in BPD sample.

In conclusion, first-generation antipsychotics can be administered to BPD patients during acute states with impulsive-aggressive behaviors and psychotic-like symptoms. However, there is little evidence of the efficacy. The treatment should be prescribed in low doses and for short periods because of its remarkable side effects and elevated dropout rate [109,22]. Among second-generation antipsychotics, there was evidence to support the therapeutic effect of olanzapine on cognitive symptoms, aggressiveness, outbursts of anger, and anxiety in BPD patients. Further controlled investigations are needed to confirm the preliminary, promising data on aripiprazole and quetiapine. Data on the newest medications have not been collected yet.

The results of RCTs are displayed in Table 3.

**1.4 Other psychotropic agents**

In recent years, opiate antagonists, oxytocin, clonidine, and omega-3 fatty acids have garnered growing attention as treatments for BPD symptoms.

Among opiate antagonists, naloxone and naltrexone were not found to be superior to placebo in treating acute dissociative states in patients with BPD in two small placebo-controlled studies [110,111]. An open-label trial reported the efficacy of nalmefene in reducing BPD global
symptomatology, self-injurious behavior, binge eating, and alcohol consumption in a group of BPD patients with alcohol use disorder [112].

The effects of intranasal oxytocin on social cognition in BPD were observed in one open-label study [113] and five randomized placebo-controlled trials [114-118]. The findings showed an improvement in interpersonal hypersensitivity, social behavior, and dysphoric emotional response to stress, with better regulation of abnormal behavioral and neural patterns in response to social context. The studies of oxytocin mainly examined laboratory outcomes related to prosocial behaviors instead of providing measures of clinical outcomes.

The α2 adrenergic agonist clonidine, usually administered to reduce blood pressure, was studied in two RCTs on its potential effect on BPD symptoms. The first study [119] compared two dosages of clonidine (75 or 150 mcg/day) and found that both resulted in a significant decrease in aversive inner tension, dissociative symptoms, the urge to commit self-injurious behaviors, and suicidal ideation. In the second placebo-controlled trial among BPD patients with or without post-traumatic stress disorder (PTSD) [120], BPD-related psychopathology (self-destruction and self-perception) improved selectively in the PTSD-positive subgroup.

In recent decades, omega-3 fatty acids have received increasing interest for their effects on psychiatric disorders, particularly depression. In BPD, three placebo-controlled studies compared the efficacy of omega-3 fatty acids with placebo [121-123], and one RCT compared the association of omega-3 fatty acids and valproate with single valproate [45]. The association of eicosapentaenoic acid and docosahexaenoic acid was found efficacious in treating depressive symptoms, aggressive behaviors, impulsivity, anger, and self-injury. The only study that evaluated the long-term efficacy of these agents in BPD (six months after discontinuation of omega-3 fatty acids with ongoing valproate) [124] suggested a long-lasting effect in terms of anger control.

Tonic glutamate overactivity due to chronic stress was found in subjects affected by BPD [125]. Therefore, recent investigations have focused on the role of N-methyl-D-aspartate (NMDA) receptor activity in sustaining BPD symptoms. Excessive stimulation of glutamate receptors
(NMDA) causes excitotoxicity. Memantine is an NMDA antagonist and a potential neuroprotective agent. The only RCT available [126] compared the effect of memantine with usual treatment in a group of BPD patients. Memantine was found useful in improving delusions, aggression, disinhibition, irritability, and depressed mood. Furthermore, memantine presented minimal abuse potential and was demonstrated to inhibit the abuse of addictive drugs [127].

The results of RCTs are displayed in Table 4.

1.5 Side effects
The real effects of psychotropic agents in the treatment of BPD symptoms are strongly affected by several issues concerning patient adherence to the prescription of medications and their sensitivity to the risk of side effects. Inadequate compliance with pharmacotherapy, sudden spontaneous changes of drugs and doses, weak and unstable therapeutic alliance, phenomena of pharmacophobia, or exaggerated intolerance to even mild adverse effects are all relevant problems that clinicians must deal with when treating BPD.

Concerning the tolerability of medications, adverse events have not been assessed in most studies of BPD patients. Only about 30% of the studies included in this review reported data on adverse effects. The adverse effects of antidepressants, mild sedation, akathisia, headache, nausea, and insomnia were found with fluoxetine [32,33]. Nausea was observed with fluvoxamine [34]. Headache and nausea were found with duloxetine [37].

Among mood stabilizers, the most common adverse effects were: tremors, diarrhea, nausea, dyspepsia, and transient hepatic transaminases increase with valproic acid [43,45]. Weight loss, fatigue, dizziness, paresthesia, and subjective cognitive impairment were found with topiramate [47,49,51]. Rash, pruritus, irritability, confusion, headache, and dizziness were found with lamotrigine [53-55]. Sedation, dizziness, headache, and nausea were found with oxcarbazepine [56]. Concerning first-generation antipsychotics, studies reported the incidence of extrapyramidal effects, in particular, with haloperidol [61,62,85]. Investigations that assessed the side effects of
second-generation antipsychotics revealed that clozapine may induce granulocytopenia and agranulocytosis (a concentration of granulocytes below 100 cells/mm$^3$ of blood) in about 1% of treated patients. Other common adverse effects are excessive salivation and weight gain [66,67]. Sleepiness, fatigue, hyperprolactinemia, and significant weight gain were observed with both risperidone [71-73] and olanzapine [32,81,82,85,86]. One Cochrane systematic review highlighted an increased risk of suicidal ideation and self-mutilating behaviors with olanzapine. The main adverse effects of aripiprazole were headache, nausea, insomnia, anxiety, and inner restlessness [87-89,91]. Quetiapine was responsible for sedation, dry mouth, and dizziness [96-100]. Patients treated with asenapine reported akathisia, restlessness, anxiety, and irritability [86,102]. Paliperidone was associated with insomnia [76], while ziprasidone was linked to gastrointestinal adverse effects [104].

The only data available among other therapeutic agents stated that omega-3 fatty acids were well-tolerated in BPD patients. Nalmefene was associated with dizziness, nausea, low appetite, and unsteady movement [112]. Oxytocin may cause headache, whereas clonidine induces xerostomia and orthostatic reactions (Ziegenhorn et al., 2009).

### 1.6 Combination of medications and psychotherapy

The conclusions of recent systematic reviews [128,25] and guidelines [20,129] underlined the role of psychotherapy as a first-line treatment of patients with BPD and confirmed the importance of combining psychotherapy and medications.

Several psychotherapeutic approaches were developed or adapted specifically for BPD patients. Most investigated models of psychotherapy were dialectical behavior therapy (DBT) [130] among cognitive-behavioral therapies and mentalization-based therapy [131] among psychodynamic treatments. An increasing amount of evidence has been obtained about the adaptation of interpersonal psychotherapy to BPD (IPT-BPD) [132-137].
In this review, we included only the trials that compared the combination of psychotherapy and medication with single psychotherapy or single medication to answer to our initial question: “is pharmacotherapy alone enough?”

Three RCTs evaluated the advantages of combining DBT and pharmacotherapy in comparison with DBT and placebo [31,82,83]. The association of fluoxetine with DBT did not provide additional benefits over single DBT [31]. The combined therapy of olanzapine and DBT showed a significant advantage over DBT plus placebo in treating depression, anxiety, and impulsive-aggressive behaviors [82]. In addition, olanzapine with DBT promoted a faster decrease in irritability and aggression than did DBT plus placebo [83].

Three other controlled trials investigated whether a combined treatment of IPT adapted to BPD (IPT-BPD) and fluoxetine was superior to single pharmacotherapy with fluoxetine [33,136,137]. The combination of pharmacotherapy and psychotherapy was found more efficacious than pharmacotherapy alone to treat three BPD core symptoms: disturbed interpersonal relationships, affective instability, and impulsivity. It was also more effective in reducing anxiety and improving the subjective perception of the quality of life [33]. The addition of IPT-BPD to the antidepressant showed advantages on impulsivity, interpersonal dysfunctions, and quality of life obtained that were maintained during two years of follow-up after the termination of psychotherapy [137]. One study of predictive factors of response to the addition of psychotherapy indicated that patients who presented more severe BPD psychopathology and a higher degree of fear of abandonment, affective instability, and identity disturbance had a higher chance of responding to combined therapy [136]. These symptoms, particularly identity disturbance, should be considered with particular care, as they are often refractory to usual BPD treatment.

In summary, there is insufficient data on the combination of pharmaco- and psychotherapy to draw any conclusions. However, most of the studies suggested that a combined approach may be useful
for improving specific BPD symptom dimensions, in particular those that respond slowly or not at all to monotherapy.

The results of RCTs are displayed in Table 5.

2. Meta-analytic studies

Several meta-analyses on BPD treatment have been published in recent years. A meta-analytic study performed by Nosè et al. [138] suggested that antidepressants and mood stabilizers are useful on affective instability and anger, whereas antipsychotics show significant effects on impulsive–aggressive behaviors. In a meta-analysis of anger and depression in BPD, Mercer et al. [38] found that mood stabilizers are more beneficial than SSRIs in treating affective symptoms (both depressed mood and anger), while antipsychotics show a moderate effect on anger, but not depression. Ingenhoven and collaborators published two meta-analyses [139,140] with a symptom dimensions approach. In the first study [139], the authors concluded that impulsive–behavioral dyscontrol, anger, and anxiety show more improvement with mood stabilizers, and cognitive-perceptual symptoms and anger show a moderate response with antipsychotics. None of the dimensions or mood symptoms improve with antidepressants. In the second meta-analysis [140], the authors focused on the efficacy of antipsychotic drugs (including both classic neuroleptics and atypical antipsychotics). The antipsychotics showed weak effectiveness on cognitive-perceptual symptoms and mood lability (with a more pronounced effect on anger), and they were entirely ineffective on impulsive behavioral dyscontrol [22,140]. These findings contrast with another meta-analysis that included RCTs and open trials [141]. In the latter study, mood stabilizers and antipsychotics are found to be useful for treating affective dysregulation and impulsive behavioral dyscontrol. These two classes of drugs and antidepressants prove effective for treating affective dysregulation, while only antipsychotics improve cognitive-perceptual symptoms. Saunders and Silk [142] analyzed data from placebo-controlled trials on the efficacy of antipsychotics on the following symptom dimensions: affective instability, anxiety, inhibition,
cognitive-perceptual disturbance, and impulsivity aggression. All these dimensions benefit from treatment with antipsychotics.

Cochrane Systematic Reviews [24,25] of pharmacological interventions for BPD found evidence that supported the efficacy of mood stabilizers, second-generation antipsychotics, and omega-3 fatty acids on BPD symptoms. Nevertheless, the authors emphasized the need for higher quality evidence as most of the current recommendations are based on single study effect estimates [22].

3. Conclusions
We can conclude that a careful examination of trials published in recent years provides only limited evidence of the efficacy of some mood stabilizers (valproic acid), new generation antipsychotic (olanzapine and, to lesser degree, aripiprazole), and omega-3 fatty acids.

The small number of controlled studies, the heterogeneity of the results, and the methodological shortcomings in methods impede reaching any reliable conclusion on the utility of these drugs in clinical practice. Even less is known about specific adverse effects in BPD. Therefore, we suggest using these psychotropic agents only when a comorbid condition is present, or an acute crisis has to be addressed.

4. Expert opinion
Drugs are routinely prescribed in clinical practice for the treatment of BPD, although no psychotropic medication has been licensed for this disorder. Why are medications so frequently used, often in polypharmacy, for this clinical condition? Data from the literature shows that the percentage of BPD patients who have been treated with medications ranges between 90 and 99% [143], and multiple drugs were used in 67-84% of those cases [144]. Low-quality evidence, divergent guidelines recommendations, and differences between experimental settings and real-world clinical practice contributed to confusion and contrasting findings. At the moment, there is a generally low level of evidence to support the efficacy of psychotropic agents in BPD patients.
There are four evidence levels (from strongest to weakest). Level A signals that at least three RCTs are available. Level B means that at least two RCTs or one RCT and ≥1 prospective, large, open-label study have been published. Level C connotes that one RCT and one prospective, open-label study/case series or at least two prospective, open-label study/case series have been found. Level D means that the efficacy of the medication is only supported by expert opinions. In the guidelines of the WFSBP International Task Force [19], the evidence on patients with BPD was systematically estimated. Antidepressants and mood stabilizers showed Level B evidence of efficacy (fair), while second-generation antipsychotics had Level C evidence (minimal).

Although more controlled studies on these medications in BPD have become available in recent years, there is general agreement among the reviews in this field that the evidence is weak because of the severe methodological limitations of most studies. Small sample sizes, different gender proportions, heterogeneous selection criteria and assessment instruments, and high rates of dropouts were the most relevant limitations. In addition, the mean duration of the trials is too short (32 days- to 24 weeks) and insufficient to draw conclusions about the long-term efficacy and tolerability of medications in a long-lasting disorder such as BPD. For many drugs, most studies were open-label trials, which had limited clinical meaning, because there is significant evidence of spontaneous changes in BPD symptoms over time [145,146].

Current treatment guidelines diverge on the role of medications in BPD, which produces uncertainty among clinicians. APA guidelines [17,18] and WFSBP guidelines [19] support the use of drugs with psychotherapy, when possible, to treat symptoms of BPD. In fact, the APA concluded that BPD is a complex condition that can be effectively treated with a targeted pharmacological approach and suggested that single classes of drugs may affect specific symptom domains. Taking a rather similar approach, WFSBP guidelines concluded that the off-label use of psychotropic agents may help individuals with BPD to improve the cluster of affective symptoms, including depressed mood, anxiety, mood swings, and impulsivity. Despite considering approximately the same evidence as the previous guidelines, the NICE and ANHMRC [20,23] expressed substantially
divergent opinions and were strongly in favor of psychotherapy with much more limited use of medications. In fact, neither source recommended drug therapy other than to treat mental disorders in comorbidity (depression) or control specific acute symptoms during a crisis and with a short-term administration. One reason for this discordance is that the English and Australian expert groups considered more parameters to draw clinical recommendations, such as evidence of cost-effectiveness, not only for medications but also for clinical management and psychological therapies. In addition, these guidelines were developed in light of the peculiar care pathway for personality disorders in Britain and Australia.

However, a series of clinical factors must be considered to decide the most appropriate treatment approach for these patients. BPD is a challenging and often treatment-resistant clinical condition [144] with a high risk of suicidal behaviors. Patients frequently present an acute exacerbation of symptoms with impulsive and self-harming conduct, agitation, and depressive or transient psychotic symptoms. Adequate psychotherapy for BPD is not always available, and patients may refuse or discontinue it. These problematic situations lead clinicians to prescribe medications. There is a high risk of administering too many drugs for too long to manage different domains of impairing and refractory symptoms. Clinicians should choose the fewest possible medications based on the predominant psychopathological factors and design pharmacotherapy in terms of individualized, tailored intervention rather than applying unspecific polypharmacy.

Among pharmacotherapies from the last decade, we observed a shift from a high predominance of prescribing antidepressants to the present-day preference for mood stabilizers and second-generation antipsychotics. Since 2010 there have been no trials among BPD patients to assess the efficacy and tolerability of antidepressants (except for one follow-up study). Recent systematic reviews stated that there is insufficient evidence to prescribe these drugs to BPD patients, except in the case of comorbidity with depression or anxiety disorders. In contrast, more recent controlled trials indicated that mood stabilizers (valproate, topiramate, and lamotrigine) and second- and third-generation antipsychotics (olanzapine and quetiapine, and aripiprazole, respectively) can
significantly affect core symptoms of the disorder, such as impulsive-behavioral dyscontrol, affective instability, and cognitive-perceptual symptoms. Nevertheless, the only antipsychotic for which there is sizable positive evidence is olanzapine. Classical neuroleptics have failed to demonstrate effects on BPD symptoms, except for reducing impulsive-aggressive behaviors during acute states, and adverse effects strongly limit their use. Dietary omega-3 supplementation and oxytocin have garnered growing interest among researchers due to promising findings in terms of beneficial effects, particularly on the symptom domain of anger. They may represent two interesting therapeutic options. Omega-3 fatty acids have a high level of tolerability with very mild or absent side effects. Oxytocin seems to address the domain of social cognition, which tends to respond poorly to other pharmacological treatments.

In our opinion, an individualized, tailored pharmacotherapy for BPD, with the fewest medications needed to target the prominent symptom clusters could improve relevant aspects of the clinical picture. However, no medication is actually indicated to treat the global psychopathology of this personality disorder. Polypharmacy is still common in clinical practice. It is not supported by data and should be avoided or strongly limited. We conclude that at the moment, chemical pharmacotherapy alone cannot manage a complex, severe, impairing condition such as BPD. Several core symptom domains, such as fears of abandonment, chronic feelings of emptiness, and identity disturbances, are refractory to pharmacotherapy. They could be the focus of specific psychotherapies or combined therapies. As we have reported in this review, the initial data appear in favor of the efficacy of these interventions on resistant BPD symptoms. The number of controlled, well-designed trials to test medications in BPD has generally decreased over the last few years probably because preceding investigations had fewer encouraging results than expected. Based on available findings, psychotherapy appropriately adapted for severe personality disorders or associated with select mood stabilizers or new antipsychotics appears to be the most promising therapeutic tool for treating BPD patients.
Further issues about pharmacotherapy have resulted from new diagnostic categories proposed by ICD 11 [147]. In particular, there has been a debate over whether the new ICD-11 diagnostic category of complex PTSD (CPTSD) can be interpreted as substantially equivalent to PTSD comorbid with Borderline Personality Disorder (BPD). It is likely that a group of patients receiving a diagnosis of BPD with DSM-5 criteria develop their symptoms on a post-traumatic basis and present both symptoms of PTSD and symptoms of disturbances in self-organization, which overlap with several core symptoms of BPD [148]. The ICD-11 diagnostic criteria for CPTSD reflect a structure model consisting of two higher-order factors. The first factor includes typical symptoms of PTSD, such as prolonged feelings of terror, marked preoccupation with a relationship with a perpetrator, isolation, and disruption in intimate relationships, and search for a rescuer. The second factor includes disturbances in self-organization, such as problems in affect regulation, feelings of shame, guilt, or failure, and difficulties in sustaining relationships. Of course, these two factors’ structure is not recognizable in the majority of BPD patients. Moreover, a history of trauma is not necessary for a diagnosis of BPD, while it is an essential requirement for the diagnosis of CPTSD [149]. Nevertheless, there is abundant evidence of the trauma of early life for BPD patients, and a subgroup of patients may have been misdiagnosed as BPD. They might be better classified in the new category of CPTSD. The crossover between CPTSD and BPD opens up the field’s consideration of trauma therapies and medications. At the moment, no controlled trials of pharmacotherapy are available in samples of CPTSD patients. A recent systematic review and meta-analysis of available trials in this clinical population considered psychosocial interventions, such as cognitive-behavioral therapy, exposure alone, and eye movement desensitization and reprocessing. It found initial evidence of partial efficacy [150]. Determining whether to use medication as a single or combined therapy is interesting, and the topic needs to be explored in its entirety.
References


24
90. Bellino S, Bozzatello P, Brignolo E, Bogetto F. Antipsychotics in the Treatment of Impulsivity in Personality Disorders and Impulse Control Disorders. Current Psychopharmacology, 2013, 2, 5-17. 2211-5579/13 $58.00+0.00


140. Ingenhoven TJ, Duivenvoorden HJ. Differential effectiveness of antipsychotics in borderline personality disorder: meta-analyses of placebo-controlled, randomized clinical trials on


** These studies provide important data in prominent fields of research: an accurate review and a meta-analysis of available evidences of efficacy of pharmacotherapy and the first findings about clinical predictors of response and long-term efficacy of combined therapy in BPD.

* These studies are also of particular interest for the topic of this review: the first is not so recent, but is the only available set of treatment guidelines for BPD provided by the Federation of Societies of Biological Psychiatry with a special focus on the use of drugs; the second is a narrative review of literature on BPD pharmacotherapy with an attempt to address some core issues and open questions.
Table 1: Randomized controlled trials of antidepressants for borderline personality disorder.

<table>
<thead>
<tr>
<th>Study (year) [Ref.]</th>
<th>Interventional arm(s)</th>
<th>Comparison arm(s)</th>
<th>Sample</th>
<th>Treatment duration</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Salzamann et al. (1995) [28]</td>
<td>Fluoxetine (20–60 mg/day)</td>
<td>Placebo</td>
<td>27 BPD patients</td>
<td>12 weeks</td>
<td>Decrease in anger, improvement in global functioning. No effect on depression</td>
</tr>
<tr>
<td>Markovitz (1995) [29]</td>
<td>Fluoxetine (20–80 mg/day)</td>
<td>Placebo</td>
<td>17 BPD patients with concomitant affective and anxiety disorders</td>
<td>14 weeks</td>
<td>Decrease of all symptoms</td>
</tr>
<tr>
<td>Coccaro and Kavoussi (1997) [30]</td>
<td>Fluoxetine (20–60 mg/day)</td>
<td>Placebo</td>
<td>40 patients with PDs (33% BPD) and comorbidity with dysthymia, anxiety disorders or substance abuse</td>
<td>12 weeks</td>
<td>Improving of irritability, impulsive-aggression, and global functioning</td>
</tr>
<tr>
<td>Simpson et al. (2004) [31]</td>
<td>Fluoxetine (40 mg/day) + DBT</td>
<td>Placebo + DBT</td>
<td>25 female BPD patients</td>
<td>12 weeks</td>
<td>No effects</td>
</tr>
<tr>
<td>Zanarini et al. (2004) [32]</td>
<td>Fluoxetine (15 mg/day)</td>
<td>Fluoxetine (15 mg/day) + olanzapine (2.5 mg/day) and olanzapine (2.5 mg/day)</td>
<td>45 female BPD patients</td>
<td>8 weeks</td>
<td>Both the comparison arms resulted to be superior to fluoxetine monotherapy to decrease all symptoms</td>
</tr>
<tr>
<td>Rinne et al. (2002) [34]</td>
<td>Fluvoxamine (150–250 mg/day)</td>
<td>Placebo</td>
<td>38 female BPD patients, comorbidity with mood and anxiety disorders</td>
<td>6 weeks</td>
<td>Decrease of rapid mood shifts. No effects on aggression and impulsivity</td>
</tr>
<tr>
<td>Jariani et al. (2010) [35]</td>
<td>Sertraline (50–100 mg/day)</td>
<td>Olanzapine (5–10 mg/day)</td>
<td>120 BPD patients on methadone therapy</td>
<td>12 weeks</td>
<td>Sertraline &gt; olanzapine in decreasing depression, hypersensitivity in interpersonal relationship, and obsessions. Olanzapine &gt; sertraline in decreasing anxiety, aggression, paranoia, and self-harming. Sertraline = olanzapine in decreasing Somatization symptoms</td>
</tr>
<tr>
<td>Markovitz et al. (1995) [36]</td>
<td>Venlafaxine</td>
<td></td>
<td></td>
<td></td>
<td>Effective in reducing somatic symptoms</td>
</tr>
<tr>
<td>Bellino et al. (2010) [37]</td>
<td>Duloxetine (60 mg/day)</td>
<td></td>
<td>18 BPD patients</td>
<td>12 weeks</td>
<td>Improvement of depressive symptoms, impulsivity, outbursts of anger, and affective instability</td>
</tr>
</tbody>
</table>

Ref.: reference; BPD: borderline personality disorder; DBT: dialectical behavioral therapy
Table 2: Randomized controlled trials of mood stabilizers for borderline personality disorder.

<table>
<thead>
<tr>
<th>Study (year) [Ref.]</th>
<th>Interventional arm(s)</th>
<th>Comparison arm(s)</th>
<th>Sample</th>
<th>Treatment duration</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Links et al. (1990) [39]</td>
<td>Lithium (986 mg/day) and Desipramine placebo</td>
<td>17 BPD patients receiving concomitant psychotherapy</td>
<td>6 weeks</td>
<td>Lithium was superior to increase anger, irritability and self-mutilation. No effects on mood symptoms</td>
<td></td>
</tr>
<tr>
<td>De la Fuente and Lotstra (1994) [40]</td>
<td>Carbamazepine (plasma level)</td>
<td>Placebo</td>
<td>20 BPD outpatients</td>
<td>4.5 weeks</td>
<td>No effects</td>
</tr>
<tr>
<td>Hollander et al. (2001) [41]</td>
<td>Valproate sodium (plasma level)</td>
<td>Placebo</td>
<td>16 BPD patients</td>
<td>10 weeks</td>
<td>No effects (probably because of high drop-out rate)</td>
</tr>
<tr>
<td>Hollander et al. (2005) [42]</td>
<td>Valproate sodium (plasma level)</td>
<td>Placebo</td>
<td>96 cluster B PDs (52 BPD) with impulsive-aggression</td>
<td>12 weeks</td>
<td>Improvement of global symptomatology, impulsive-aggression, and irritability</td>
</tr>
<tr>
<td>Frankenburg and Zanarini (2002) [43]</td>
<td>Valproate sodium (plasma level)</td>
<td>Placebo</td>
<td>30 female BPD patients with comorbid bipolar II disorder</td>
<td>6 months</td>
<td>Improvement of interpersonal sensitivity, anger, hostility, and aggressiveness</td>
</tr>
<tr>
<td>Moen et al. (2012) [44]</td>
<td>Divalproex ER (plasma level)</td>
<td>Placebo</td>
<td>15 BPD patients after 4 weeks of “condensed” DBT</td>
<td>4 weeks</td>
<td>No differences between the two arms</td>
</tr>
<tr>
<td>Bellino et al. (2014) [45]</td>
<td>EPA and DHA with valproic acid (plasma level)</td>
<td>Placebo</td>
<td>43 BPD outpatients</td>
<td>12 weeks</td>
<td>Combined therapy &gt; single therapy for impulsivity, self-harm, and outbursts of anger</td>
</tr>
<tr>
<td>Nickel et al. (2004) [46]</td>
<td>Topiramate (50–250 mg/day)</td>
<td>Placebo</td>
<td>31 female BPD patients</td>
<td>8 weeks</td>
<td>Improvement of irritability and anger</td>
</tr>
<tr>
<td>Nickel et al. (2005) [47]</td>
<td>Topiramate (50–250 mg/day)</td>
<td>Placebo</td>
<td>42 male BPD patients</td>
<td>8 weeks</td>
<td>Improvement of irritability and anger</td>
</tr>
<tr>
<td>Loev et al. (2008) [51]</td>
<td>Topiramate (25–200 mg/day)</td>
<td>Placebo</td>
<td>56 female BPD patients with concomitant mood disorders</td>
<td>10 weeks</td>
<td>Improvement of somatization, interpersonal sensitivity, hostility and global functioning</td>
</tr>
<tr>
<td>Tritt et al. (2005) [52]</td>
<td>Lamotrigine (50–200 mg/day)</td>
<td>Placebo</td>
<td>24 female BPD patients</td>
<td>8 weeks</td>
<td>Improvement of anger</td>
</tr>
<tr>
<td>Reich et al. (2009) [53]</td>
<td>Lamotrigine (25–275 mg/day)</td>
<td>Placebo</td>
<td>28 BPD patients with major depression and anxiety disorders</td>
<td>12 weeks</td>
<td>Improvement of impulsivity and affective instability</td>
</tr>
<tr>
<td>Leiberich et al., (2008) [54]</td>
<td>Lamotrigine (50–200 mg/day)</td>
<td>Placebo</td>
<td>27 female BPD patients</td>
<td>18 months follow-up</td>
<td>Improvement of aggression in the long-term treatment</td>
</tr>
<tr>
<td>Crawford et al., (2015) [55]</td>
<td>Lamotrigine (25–400 mg/die)</td>
<td>Placebo</td>
<td>28 BPD patients</td>
<td>52 weeks</td>
<td>No effect in terms of symptomatic reliefs and cost-effectiveness balance</td>
</tr>
</tbody>
</table>
Ref.: reference; BPD: borderline personality disorder; DBT: dialectical behavioral therapy

Table 3: Randomized controlled trials and open trials of antipsychotics for borderline personality disorder.

<table>
<thead>
<tr>
<th>Study (year) [Ref.]</th>
<th>Interventional arm(s)</th>
<th>Comparison arm(s)</th>
<th>Sample</th>
<th>Treatment Duration</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cornelius et al. (1993) [61]</td>
<td>Haloperidol (&lt;6 mg/day)</td>
<td>Placebo</td>
<td>54 BPD outpatients from Soloff et al. (1993) study</td>
<td>16 weeks</td>
<td>Improvement of hostility and impulsive-aggression</td>
</tr>
<tr>
<td>Soloff et al. (1993) [62]</td>
<td>Haloperidol (4 mg/day)</td>
<td>Phenelzine (60 mg/day) and placebo</td>
<td>108 BPD inpatients with concomitant major depression</td>
<td>5 weeks</td>
<td>Improvement of irritability</td>
</tr>
<tr>
<td>Szigety &amp; Schultz (1998) [75]</td>
<td>Risperidone (2.5 mg/day)</td>
<td>Placebo</td>
<td>27 BPD patients</td>
<td>8 weeks</td>
<td>No difference with regard to global functioning. Risperidone &gt; placebo in reducing psychoticism, paranoid ideas, phobic anxiety and interpersonal sensitivity</td>
</tr>
<tr>
<td>Zanarini and Frankenburg (2001) [78]</td>
<td>Olanzapine (5.33 mg/day)</td>
<td>Placebo</td>
<td>28 female BPD patients</td>
<td>6 months</td>
<td>Improvement of anger, interpersonal sensitiv, anxiety, paranoid ideation and global functioning</td>
</tr>
<tr>
<td>Bogenschutz and Nurnberg (2004) [79]</td>
<td>Olanzapine (5–10 mg/day)</td>
<td>Placebo</td>
<td>40 BPD outpatients</td>
<td>12 weeks</td>
<td>Improvement of anger and global symptoms</td>
</tr>
<tr>
<td>Schulz et al. (1998) [80]</td>
<td>Olanzapine (2.5–20 mg/day)</td>
<td>Placebo</td>
<td>314 BPD patients</td>
<td>12 weeks</td>
<td>No differences between two groups. Faster amelioration in olanzapine group.</td>
</tr>
<tr>
<td>Zanarini et al. (2011) [81]</td>
<td>Olanzapine low doses (2.5 mg/day) and moderate doses (5–10 mg/day)</td>
<td>Placebo</td>
<td>451 BPD outpatients</td>
<td>12 weeks</td>
<td>Olanzapine moderate doses &gt; placebo in global symptoms</td>
</tr>
<tr>
<td>Soler et al. (2005) [82]</td>
<td>Olanzapine (5–20 mg/day) + DBT</td>
<td>Placebo + DBT</td>
<td>60 BPD patients</td>
<td>12 weeks</td>
<td>Improvement of anxiety, depression and impulsive-agression</td>
</tr>
<tr>
<td>Linehan et al. (2008) [83]</td>
<td>Olanzapine (5 mg/day) + DBT</td>
<td>Placebo + DBT</td>
<td>24 female BPD patients</td>
<td>21 weeks</td>
<td>No differences of general symptoms. Faster decrease of irritability and aggression in olanzapine group</td>
</tr>
<tr>
<td>Jariani et al. (2010) [84]</td>
<td>Olanzapine (5-10 mg/day)</td>
<td>Sertraline (50-100 mg/day)</td>
<td>120 BPD patients on methadone maintenance therapy</td>
<td>12 weeks</td>
<td>Both drugs improving depression, anxiety, aggression, sensitivity in interpersonal relationships, obsessive symptoms, pessimistic behaviors and somatization disorders</td>
</tr>
<tr>
<td>Shafti and Shahveisi (2010) [85]</td>
<td>Olanzapine (7 mg/day)</td>
<td>Haloperidol (7 mg/day)</td>
<td>28 female BPD inpatients</td>
<td>8 weeks</td>
<td>No differences</td>
</tr>
<tr>
<td>Bozzatello et al. (2017) [86]</td>
<td>Olanzapine (5-10 mg/day)</td>
<td>Asenapine (5-10 mg/day)</td>
<td>51 BPD patients between 18 and 50 years</td>
<td>12 weeks</td>
<td>Asenapine &gt; Olanzapine in treating affective instability Olanzapine &gt; Asenapine in treating paranoid ideation and dissociation</td>
</tr>
<tr>
<td>Study</td>
<td>Treatment 1</td>
<td>Treatment 2</td>
<td>Participants</td>
<td>Duration</td>
<td>Outcome</td>
</tr>
<tr>
<td>------------------------------</td>
<td>----------------------</td>
<td>----------------------</td>
<td>--------------</td>
<td>----------</td>
<td>--------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Shafti and Kaviani (2015) [87]</td>
<td>Olanzapine (6.4 mg/day)</td>
<td>Aripiprazole (7 mg/day)</td>
<td>24 female BPD inpatients</td>
<td>8 weeks</td>
<td>Olanzapine &gt; Aripiprazole on uncooperativeness and excitement</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Aripiprazole &gt; Olanzapine on suspiciousness and unusual thought content</td>
</tr>
<tr>
<td>Nickel et al. (2006) [88]</td>
<td>Aripiprazole (15 mg/day)</td>
<td>Placebo</td>
<td>57 BPD patients</td>
<td>8 weeks</td>
<td>Improvement of depression, anxiety, anger, aggressiveness, paranoia and global functioning</td>
</tr>
<tr>
<td>Nickel et al. (2007) [89]</td>
<td>Aripiprazole (15 mg/day)</td>
<td>Placebo</td>
<td>52 BPD patients</td>
<td>18 months</td>
<td>Effective and relatively safe agent in the long-term treatment</td>
</tr>
<tr>
<td>Chanen et al. (2018) [92]</td>
<td>Aripiprazole (2-30 mg/day)</td>
<td>Placebo</td>
<td>154 BPD patients</td>
<td>12 weeks + 27-week follow-up</td>
<td>Improvement of general psychopathology, borderline personality pathology, social and occupational functioning</td>
</tr>
<tr>
<td>Black et al. (2014) [100]</td>
<td>Quetiapine low (150 mg/day) and moderate doses (300 mg/day)</td>
<td>Placebo</td>
<td>95 BPD patients (70.5% female)</td>
<td>8 weeks</td>
<td>Quetiapine low doses &gt; placebo in all the outcomes measures (in particular affective instability, cognitive-perceptual symptoms and aggressiveness). Faster response in both the treatment groups</td>
</tr>
<tr>
<td>Lee et al. (2016) [101]</td>
<td>Quetiapine low (150 mg/day) and moderate doses (300 mg/day)</td>
<td>Placebo</td>
<td>95 BPD patients (70.5% female)</td>
<td>8 weeks</td>
<td>Improvement of all the outcomes measures with both quetiapine doses</td>
</tr>
<tr>
<td>Pascual et al. (2008) [104]</td>
<td>Ziprasidone (84 mg/day)</td>
<td>Placebo</td>
<td>60 BPD patients</td>
<td>12 weeks</td>
<td>No effects</td>
</tr>
<tr>
<td>Bellino et al. (2009) [91]</td>
<td>Sertraline (50-100 mg/day) + Aripiprazole (10-15 mg/day)</td>
<td>Open label trial</td>
<td>21 BPD patients between 18 and 50 years</td>
<td>12 weeks</td>
<td>Improvement of impulsivity and dissociation/paranoid ideation</td>
</tr>
<tr>
<td>Perrella et al. (2007) [97]</td>
<td>Quetiapine (400-800 mg/day)</td>
<td>Open label trial</td>
<td>29 BPD outpatients</td>
<td>12 weeks</td>
<td>Improvement of mood symptoms and aggression</td>
</tr>
<tr>
<td>Bellino et al. (2011) [76]</td>
<td>Paliperidone ER (3-6 mg/day)</td>
<td>Open label trial</td>
<td>18 BPD outpatients</td>
<td>12 weeks</td>
<td>Improvement of impulsive dyscontrol, anger, and cognitive-perceptual disturbances</td>
</tr>
<tr>
<td>Rocca et al. (2002) [71]</td>
<td>Risperidone (3.27 mg/day)</td>
<td>Open label trial</td>
<td>15 BPD outpatients</td>
<td>8 weeks</td>
<td>Improvement of depressive symptoms, energy and global functioning</td>
</tr>
</tbody>
</table>

BPD: borderline personality disorder; DBT: dialectical behavioral therapy; >: superior of.
Table 4: Randomized controlled trials of other psychotropic agents and for borderline personality disorder.

<table>
<thead>
<tr>
<th>Study (year) [Ref.]</th>
<th>Interventional arm(s)</th>
<th>Comparison arm(s)</th>
<th>Sample</th>
<th>Treatment Duration</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Philipsen et al. (2004) [110]</td>
<td>Naloxone (0.4 mg/day)</td>
<td>Placebo</td>
<td>9 BPD patients</td>
<td>15 min</td>
<td>No differences in dissociative symptoms</td>
</tr>
<tr>
<td>Schmahl et al. (2012) [111]</td>
<td>Naltrexone (50 or 200 mg/day)</td>
<td>Placebo</td>
<td>29 BPD patients</td>
<td>3 weeks</td>
<td>No differences in dissociative symptoms</td>
</tr>
<tr>
<td>Bertisch et al. (2013) [115]</td>
<td>Oxytocin (26 IU)</td>
<td>Placebo</td>
<td>40 female nonmedicated BPD patients</td>
<td>45 minutes</td>
<td>Improvement of social hypersensitivity, anger and aggressive behavior</td>
</tr>
<tr>
<td>Philipsen et al. (2004) [119]</td>
<td>Clonidine (75 µg/day)</td>
<td>Clonidine (150 µg/day)</td>
<td>14 female BPD patients</td>
<td>30,60 and 120 min</td>
<td>Decrease (maximum after 30-60 min) of all the symptoms with both doses of clonidine</td>
</tr>
<tr>
<td>Ziegenhorn et al. (2009) [120]</td>
<td>Clonidine (up to 0.3 mg/day)</td>
<td>Placebo</td>
<td>18 BPD patients, some of them with concomitant PTSD</td>
<td>6 weeks</td>
<td>Decrease of hyperarousal symptoms independently of PTSD comorbidity. Decrease of BPD specific and general symptoms mainly in the PTSD-positive subgroup</td>
</tr>
<tr>
<td>Zanarini and Frankenburg, (2003) [121]</td>
<td>EPA (1 g/day)</td>
<td>Placebo</td>
<td>30 female BPD patients</td>
<td>8 weeks</td>
<td>Improvement of Depressive symptoms and aggressive behaviors</td>
</tr>
<tr>
<td>Hallahan et al. (2007) [122]</td>
<td>EPA (1.2 g/day)+DHA (0.9 g/day)</td>
<td>Placebo</td>
<td>49 patients with self-harm behaviors (35 BPD)</td>
<td>12 weeks</td>
<td>Improvement of depression, suicidality and reaction to daily stress</td>
</tr>
<tr>
<td>Amminger et al. (2013) [123]</td>
<td>PUFAs (1.2 g/day)</td>
<td>Placebo</td>
<td>15 adolescent BPD patients with high risk of psychosis</td>
<td>12 weeks</td>
<td>Improvement of global functioning and schizophrenia-like symptoms</td>
</tr>
<tr>
<td>Bellino et al. (2014) [45]</td>
<td>EPA (1.2 g/day) + DHA (0.8 g/day) + valproate (800-1300 mg/day)</td>
<td>Valproate (800-1300 mg/day)</td>
<td>43 BPD outpatients</td>
<td>12 weeks</td>
<td>No differences with regard to global symptoms. Improvement of impulsivity, anger and self-mutilating conducts in omega-3 group</td>
</tr>
<tr>
<td>Bozzatello et al. (2018) [124]</td>
<td>EPA (1.2 g/day) + DHA (0.8 g/day) + valproate (800-1300 mg/day)</td>
<td>Valproate (800-1300 mg/day)</td>
<td>43 BPD outpatients</td>
<td>24 weeks follow-up</td>
<td>Combined therapy with omega-3 fatty acids showed long-lasting effects after discontinuation in terms of anger control.</td>
</tr>
<tr>
<td>Kulkarni et al. (2018) [126]</td>
<td>Memantine (20 mg/die)</td>
<td>Placebo</td>
<td>33 BPD outpatients</td>
<td>8 weeks</td>
<td>Memantine showed &gt; improvement on aggression, disinhibition, irritability and depressed mood</td>
</tr>
</tbody>
</table>

BPD: borderline personality disorder; PTSD: Posttraumatic Stress Disorder; EPA: Eicosapentaenoic acid; PUFAs: Polyunsaturated fatty acid; >: superior of.
Table 5: Randomized controlled trials of combined therapy (pharmacotherapy + psychotherapy) for borderline personality disorder.

<table>
<thead>
<tr>
<th>Study (year)</th>
<th>Interventional arm(s)</th>
<th>Comparison arm(s)</th>
<th>Sample</th>
<th>Treatment duration</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Simpson et al. (2004) [31]</td>
<td>Fluoxetine (40 mg/day) + DBT</td>
<td>Placebo + DBT</td>
<td>25 female BPD patients</td>
<td>12 weeks</td>
<td>No effects on depression, anxiety, anger, dissociation, and global functioning</td>
</tr>
<tr>
<td>Soler et al. (2005) [82]</td>
<td>Olanzapine (5–20 mg/day) + DBT</td>
<td>Placebo + DBT</td>
<td>60 BPD patients</td>
<td>12 weeks</td>
<td>Improvement of anxiety, depression, and impulsive-aggression</td>
</tr>
<tr>
<td>Linehan et al. (2008) [83]</td>
<td>Olanzapine (5 mg/day) + DBT</td>
<td>Placebo + DBT</td>
<td>24 female BPD patients</td>
<td>21 weeks</td>
<td>No differences in treatment of general symptoms. Faster decrease of irritability and aggression in the olanzapine group</td>
</tr>
<tr>
<td>Bellino et al. (2010) [33]</td>
<td>Fluoxetine (20–40 mg/day) + IPT-BPD</td>
<td>Fluoxetine (20–40 mg/day)</td>
<td>44 BPD patients</td>
<td>32 weeks</td>
<td>Combined therapy &gt; fluoxetine in improving BPD core symptoms (interpersonal relationships, affective instability, and impulsivity), anxiety and QoL</td>
</tr>
<tr>
<td>Bellino et al. (2015) [136]</td>
<td>Fluoxetine (20–40 mg/day) + IPT-BPD</td>
<td>Fluoxetine (20–40 mg/day)</td>
<td>44 BPD patients</td>
<td>32 weeks</td>
<td>More severe BPD psychopathology and higher degree of fear of abandonment, affective instability, and identity disturbance predicted response to combined therapy</td>
</tr>
<tr>
<td>Bozzatello and Bellino (2016) [137]</td>
<td>Fluoxetine (20–40 mg/day) + IPT-BPD</td>
<td>Fluoxetine (20–40 mg/day)</td>
<td>44 BPD patients</td>
<td>24 months</td>
<td>Persistence of advantages of combined therapy on impulsivity, interpersonal dysfunctions, and QoL</td>
</tr>
</tbody>
</table>

Ref.: reference; BPD: borderline personality disorder; DBT: dialectical behavioral therapy; IPT-BPD: interpersonal psychotherapy adapted to BPD; QoL: quality of life; > : superior to