**The influence of anxiety symptoms on clinical outcomes during baclofen treatment of alcohol use disorder: A systematic review and meta-analysis**

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**ABSTRACT** *(170 words)*

Given the high coexistence of anxiety symptoms in people with alcohol use disorder (AUD), we aimed to determine the influence of anxiety symptoms on outcomes in patients with AUD treated with GABAB receptor agonist baclofen. A meta-analysis of 13 comparisons (published 2010-2020) including baseline and outcome data on alcohol consumption and anxiety after 12 weeks was undertaken. There were significantly higher rates of abstinent days in patients treated with baclofen compared to placebo (p=0.004; high certainty evidence); specifically in those with higher baseline anxiety levels (p<0.00001; high certainty evidence) compared to those with lower baseline anxiety levels (p=0.20; moderate certainty evidence). The change in anxiety ratings over 12 weeks did not differ between those treated with baclofen or placebo (p=0.84; moderate certainty evidence). This may be due to different anxiety constructs being measured by scales not validated in this patient group, or that anxiety is not a biobehavioral mechanism by which baclofen may reduce alcohol drinking. Given the prevalence of anxiety symptoms in AUD all these factors warrant further research.

**Key words:** Alcohol, anxiety, baclofen, GABAB, clinical trials, meta-analysis, measurement, rating scales

1. **INTRODUCTION**

Baclofen (β-chlorophenyl-GABA) is a GABAB receptor full agonist, approved and long used in the treatment of muscle spasticity, which may also have analgesic properties independent of its myorelaxant effect [1,2]. The role of the GABAB receptor as a pharmacological target in the treatment of alcohol use disorder (AUD) has received increased attention in recent decades, with some clinical trials indicating that baclofen may reduce alcohol craving and drinking and promote abstinence [for review see 3]. However, human data are inconsistent [see meta-analysis studies: 4-7], which may be due, at least in part, to the heterogeneity of the enrolled samples, and differences in study designs and dosage regimes. Baclofen has been approved for AUD treatment in France, in patients who have not responded to other approved treatments [3].

Whilst the myorelaxant properties of baclofen are related to a ‘dampening’ of the spinal motor reflex [8], its potential mechanism of action in AUD is less clear. Central GABAB receptors are involved in the regulation of a large number of systems and functions, including several neurotransmitter systems (e.g. dopamine, serotonin, norepinephrine, glutamate), transduction pathways, memory, and other cognitive functions [8].

The optimal dosage of baclofen required to treat patients with AUD is also uncertain, with some patients being able to tolerate much higher doses compared to others [9]. Recent work suggests there may be a blunted sensitivity to baclofen in patients with AUD relative to controls, with no difference in pharmacokinetics suggesting a lower GABAB receptor sensitivity [10] and possibly a cross-tolerance between the effects of alcohol and baclofen on the GABAB receptor [11,12].

Anxiety disorders and AUD are among the more prevalent mental disorders worldwide [13]. Anxiety symptoms and disorders are frequent in patients with AUD [14], especially among women [15]. Social anxiety and post-traumatic stress disorder (PTSD) frequently pre-date AUD by many years and are an established risk factor in its aetiology [16-18]; panic attacks often occur in established AUD at high levels of daily drinking [19]; and anxiety symptoms are also common during alcohol withdrawal. Despite the high prevalence of co-occurrence, relatively little is known about the underlying neurobiological mechanisms involved, and even less about the effective management of this common co-morbid state [20].

Preclinical and clinical findings indicate a key role of the GABAB receptor in depression and anxiety disorders [21,22] and the presence of psychiatric comorbidity may influence the response to baclofen treatment in patients with AUD [3]. A recent narrative review of clinical trials of baclofen in patients with AUD and comorbid mental disorders was not able to draw any definitive conclusions [23].

We speculated that a meta-analytic approach might better elucidate the role of anxiety symptoms in determining outcomes to treatment with baclofen in patients with AUD; in terms of alcohol consumption, anxiety symptoms, and the potential interaction between them. The different scales used to measure anxiety also require further exploration, as each is based on a different construct and there is little in the literature validating their use in patients with AUD. Accordingly, the aims of this study were to review how anxiety symptoms were measured in trials of patients with AUD treated with baclofen; and, using a meta-analytic approach, to evaluate whether, compared to placebo, baclofen induced significant changes in anxiety symptoms and drinking outcomes over the course of treatment, and how these were related.

1. **METHODS**

**Search strategy**

We conducted a systematic review and meta-analysis in line with PRISMA guidelines, following an *a priori*–defined protocol, registered with PROSPERO [24]. We searched PsycINFO; Embase; MEDLINE; PubMed; Cochrane Library, with no date limitation, using the following search strategy: “alcohol related terms” AND “anxiety related terms” AND “baclofen related terms” (see 24 for full details). The last search was performed on February 26, 2019. References from included and excluded articles were reviewed to identify additional studies, and authors were contacted to identify studies in progress.

**Inclusion criteria**

*Types of studies*

Controlled clinical trials measuring the effects of baclofen on anxiety symptoms and alcohol consumption in patients with AUD. When a study reported more than one comparison, e.g. two different doses of baclofen and placebo, data for each comparison were included.

*Types of participants*

Men and women over 18 years with AUD, in whom levels of anxiety were also measured.

*Types of interventions*

Pharmacological treatment with baclofen alone or in combination with other medications or with psychosocial intervention.

*Comparator(s)/control*

Comparators initially included placebo, no intervention, other pharmacological interventions, and/or any psychological intervention.

* 1. **Selection of the outcomes to evaluate alcohol consumption**

There is a substantial range of outcome measures used in studies evaluating the efficacy of baclofen in the treatment of AUD [3]. Given the difficulty of comparing different outcome measures across studies, we selected a single measure (level of alcohol consumption) to evaluate the severity of alcohol use of participants at baseline.

Level of alcohol consumption at baseline (expressed in grams per week) was selected as a composite measure incorporating number of drinks per drinking day, and the number of drinking days per week. We used World Health Organization (WHO) criteria to classify the level of alcohol consumption into different categories of risk. WHO criteria consider ‘high risk’ as 60 -100g/day for men (40-60g/day for women), and ‘very high risk’ as greater than 100 g/day for men (60 g/day for women) [25]: therefore, a week with seven drinking days at these levels would constitute 420g to >700g alcohol week for men (280g->420g for women). Rate of abstinent days was used to evaluate the efficacy of baclofen at the end of the studies.

**Outcomes and their measures**

*Primary outcomes*

Severity of anxiety expressed as

Final score in continuous interviewer-rated and/or self-administered scales

Difference between baseline and final scores in continuous interviewer-rated and/or self-administered scales

* Alcohol consumption levels expressed as rate of abstinent days.

*Secondary outcomes*

* Use of other substances with abuse liability
* Craving as measured by validated scales, e.g. Visual Analog Scale (VAS)
* Severity of AUD as measured by validated scales, e.g. Clinical Global Impression scale (CGI)
* Severity of other psychiatric symptoms diagnosed using standard criteria, e.g. Diagnostic and Statistical Manual of Mental Disorders (DSM) or measured by validated scales, e.g. Positive and Negative Syndrome Scale (PANSS).

**Exclusion criteria**

Studies were excluded if: (a) they did not report information on anxiety symptoms; (b) patients were not affected by AUD; (c) patients were < 18-years-old; (d) they were published in papers in languages other than English, Italian, Portuguese, or Spanish.

**Data collection**

Screening and data extraction were independently conducted by two authors (H.A. and R.A.; J.S. and R.A.). Disagreements were resolved by discussions with a third author (D.S.B. for topics related to anxiety; L.L. for topics related to AUD). From each selected study or comparison (when there was more than one comparison per study), the following information was abstracted: (a) type of study (comparative vs placebo controlled trial), (b) number of participants assigned to baclofen and control group, (c) number of males and females, (d) mean age, (e) diagnostic criteria used for AUD and anxiety disorder, (f) duration of AUD and anxiety, (g) severity of AUD and anxiety at baseline, (h) dose of baclofen administered, (i) type of administration (fixed dose vs. titration), (j) scheduled duration of pharmacological treatment, (k) setting (inpatient or outpatient or a combination), (l) alcohol consumption at baseline (actively drinking or duration of abstinence), (m) presence of co-morbid mental or physical diseases (e.g. liver disease or mood disorders), (n) other pharmacological treatment offered, (o) other psychosocial treatment offered, (p) type of scale used to measure anxiety symptoms, and (q) outcome measures (see previous sections).

**Assessment of risk of bias**

Risk of bias of studies included in the meta-analysis was evaluated by two authors (J.S. and R.A.) and disagreements were discussed with a third author (D.S.B. or L.L.) according to the criteria indicated in Cochrane Reviews Handbook [26].

**Data analysis**

We analyzed continuous outcomes by calculating the mean difference (MD) with 95% confidence interval (CI). When different scales were used to measure the severity of anxiety symptoms, the standardized mean difference (SMD) was used. We considered that participants differed in their response to baclofen compared to placebo when CIs excluded 0 (p-values < 0.05) and there was a lack of difference when CIs included 0 (p-values ≥ 0.05). The presence of significant heterogeneity was defined as I² value > 50% or p-value for the chi-squared test ≤ 0.1 [26]. Random effects models were used for all the analyses. We assessed the overall quality of the evidence for the primary outcomes using the Grading of Recommendations Assessment, Development, and Evaluation (GRADE) system [27] reporting the main findings in the 'Summary of findings' tables (4a and 4b) [28]. Analyses were conducted using Review Manager 5.3.5.

1. **RESULTS**
	1. **Studies selected and comparisons extracted**

Two authors (RA, HA) independently screened the titles and abstracts of search hits to select studies of interest. The search identified 427 articles (4 from Cochrane reviews, 29 from Cochrane trials, 233 from Embase, 52 from Medline, 55 from PsycInfo and 54 from PubMed), of which 141 were duplicates. Six additional articles were identified through references (see Figure 1).

Screening the titles and abstracts of these records provided information that led to the exclusion of 208 articles (see References S1), while full text-review of the remaining 84 articles excluded a further 66 articles which did not meet all of the inclusion and none of the exclusion criteria. Nineteen studies were initially included in the qualitative synthesis, but one was excluded [29] as although stating it measured anxiety, it only reported data from a broader scale of psychological distress.

Eighteen studies [30-47] evaluated the effects of baclofen on anxiety symptoms and alcohol consumption in patients with AUD. As these studies varied in the duration of pharmacological treatment from less than 1 week to 12 months (see Table 1), studies with a duration of less than 12 weeks (a standard duration for many RCTs in AUD) were excluded from the meta-analysis to decrease the variability in response related to differences in duration [30,31,33,36,37,39,41,42].

Comparisons of baclofen with another drug (i.e. diazepam, naltrexone, chlordiazepoxide, lorazepam, and benfothiamine), rather than placebo were also excluded [31,35 (comparison 3, see below)-39]. Studies comparing baclofen to naltrexone [35] or benfothiamine [36] were excluded because there was no placebo comparison. Despite benfothiamine being defined by the authors as a ‘placebo’ [38] we included it in the comparator drug category, as benfothiamine (a nutritional supplement) has been found to modify anxiety symptoms in patients with AUD [48]. This is consistent with FDA guidelines for the development of medications for AUD, which highlight that, while an active comparator of proven efficacy can *also* be included, in general a placebo comparator *should* be used [49]. Studies comparing baclofen to diazepam [31,39b], chlordiazepoxide [36] or lorazepam [37] were excluded both due to lack of a placebo arm in addition to their short duration (see above) as a treatment for alcohol withdrawal, which is a distinct clinical treatment beyond the scope of this work.

Study characteristics are shown in Table 1. As three studies used two different doses of baclofen [32,43,44], two different comparisons were extracted from each. Patients assigned to placebo were divided into two groups, one for each comparison. One study compared baclofen treatment with naltrexone and their combination [35], enabling three potential comparisons [(1) baclofen (n=10) *vs* placebo (n=10); (2) the combination baclofen plus naltrexone (n = 10) *vs* the combination placebo plus naltrexone (n=10); (3) baclofen (n=10) *vs* naltrexone (n=10)].

* 1. **Scales used to evaluate anxiety symptoms**

Table 2 shows the main characteristics of the scales used to evaluate anxiety symptoms [50-55]. The majority of studies [30,32-35,39-41,43] used Spielberger's State-Trait Anxiety Inventory (STAI); three studies [40,45,38] used the Hamilton Anxiety Rating Scale (HAM-A or HARS); three studies [31,36,37],not ultimately included in the meta-analysis, only used the single anxiety question in the Clinical Institute Withdrawal Assessment for Alcohol, revised scale (CIWA-Ar). Two studies [46,47] used the Hospital Anxiety and Depression Scale (HADS-A), and one study [44] used the Depression, Anxiety and Stress Scale (DASS). One final study [29] stated that ‘symptoms of anxiety were measured by self-report using the Brief symptom Inventory (BSI)’: however, only the full BSI (rather than the anxiety sub-scale) findings were reported in the paper (see section below), and so after contacting the authors to confirm that it was not the anxiety measure that was presented, and they could not provide specific data for this, it was excluded from the analysis.

* + 1. *Spielberger State-Trait Anxiety Inventory (STAI) [55]*

There is limited evidence for the validity of the STAI as a measure of categorical diagnoses of anxiety in general population samples. Mean values of both State and Trait STAI range between 30-40 [56]. One recent study demonstrated that, among patients with epilepsy, a cut-off score ≥ 52 for both State and Trait STAI showed the best psychometric properties for the diagnoses of anxiety disorders [57]. As far as we are aware, no study has evaluated ‘cut-off’ values for the diagnoses of anxiety disorders among patients with AUD. Therefore, consistent with Wiglusz et al. [57], we used ≥ 52 as the cut-off value to categorize if samples in the present meta-analysis reached the threshold for an anxiety disorder.

* + 1. *Hamilton Anxiety Rating Scale (HAM-A or HARS) [54]*

The HAM-A is well validated as a symptom rating scale in patients with anxiety disorders. Scores of 14 and higher are often taken to designate a case [58]. Accordingly, we used the cut-off value ≥ 14 to categorize if samples compared in the present meta-analysis reached the threshold for an anxiety disorder.

* + 1. *Depression, Anxiety and Stress Scale (DASS) [52]*

The DASS comprises three separate scales, well validated in clinical samples, to evaluate depression (DASS-D), anxiety (DASS-A), and stress (DASS-S) [58]. The cut-off value suggestive for the presence of anxiety disorders is ≥ 8 for the anxiety scale.

* + 1. *Hospital Anxiety and Depression Scale (HADS-A) [53]*

The HADS-A is validated for screening for anxiety and depression in medically unwell patients [58]. Scores of >8 higher on the anxiety sub-scale are often taken to designate a case including in one report of patients with rheumatological conditions [59]. To our knowledge, no study has evaluated cut-off values for probable anxiety disorder among patients with AUD; therefore we used the standard cut-off value to categorize if samples compared in the present meta-analysis reached the threshold for an anxiety disorder

* + 1. *Brief Symptom Inventory (BSI) [50]*

The BSI [50] is a shortened, 53-item version of the Symptom Checklist-90. Participants have to rate from 0 (not at all) to 4 (extremely) the items corresponding to different subscales investigating anxiety, phobic anxiety, somatization, obsessive-compulsive, sensitivity, depression, hostility, paranoid ideation, and psychoticism. The cut-off for clinical ‘caseness’ for psychological distress is ≥ 63.

* + 1. *Clinical Institute Withdrawal Assessment for alcohol, revised scale (CIWA-Ar) [51]*

The CIWA-Ar is a 10-item scale used clinically to measure the severity of alcohol withdrawal and response to medication. There is a single question relating to anxiety (scored 0-7), which is observer-rated. We excluded all studies in which only the CIWA single item question was used as the measure of anxiety symptoms.

* 1. **Scales and outcomes used to evaluate alcohol consumption**

The measures used to evaluate AUD severity varied widely including the Alcohol Dependence Scale (ADS), Alcohol Use Disorder Identification Test (AUDIT), and CAGE screening questions. The outcomes selected by the different studies to evaluate alcohol consumption or abstinence from alcohol comprised: the number or rate of Cumulative Abstinent Days (CAD), the number of Drinks per Drinking Day (DDD), the number or rate of Heavy Drinking Days (HDD), the Total Alcohol Consumption (TAC) expressed in grams of alcohol per day or grams per week, the number or rate of abstinent participants, the rate of heavy drinkers, the time to first relapse, the time to first lapse, and time to first Heavy Drinking Day.

* 1. **Comparisons included in the meta-analysis**

We included 13 comparisons in the meta-analysis (32HD, 32LD, 34, 35(1), 35(2), 40, 43HD, 43LD, 44HD, 44LD, 45, 46, 47; 1143 participants) between baclofen (610 participants) and placebo (533 participants) extracted from studies lasting 12 weeks or longer (see Table 3).

* + 1. *Risk of bias in included comparisons (see Figures 2 and 3)*

Random sequence generation: eleven comparisons were considered to have low risk of bias. No information about methods of sequence random generation was available in two comparisons, so were judged as at unclear risk of bias.

Allocation concealment: eight comparisons were considered at low risk of bias, three comparisons did not report methods of allocation concealment, so were judged as at unclear risk of bias. Two other comparisons, during the trial, the inclusion of patients in a scheduled group was interrupted, and it is unclear that it may have constituted a risk of bias [32HD,32LD].

Blinding of participants and personnel: seven comparisons were judged at low risk of bias; three comparisons did not report details about blinding so were judged at unclear risk of bias; two comparisons, during the study, the inclusion of patients in a scheduled group was interrupted and it is unclear that it may have constituted a risk of bias [32HD,32LD]; in another it was unclear if the high dose of baclofen may constitute a risk of bias [47].

Blinding of outcome assessment: only one comparison was judged at low risk of bias [40]; for the other comparisons information about blinding was not available so were judged as at unclear risk of bias. Incomplete outcome data: five comparisons were considered low risk of bias; five comparisons did not report numbers and reasons of dropouts or missing data, so were judged as at unclear risk of bias. Three comparisons were judged to be at high risk of bias: one study as patients who relapsed were excluded [40], one reported the primary outcome as “an estimated percentage of abstinent patients during 20 consecutive weeks” without providing an explanation of how these figures were reached (e.g. how the authors deal with drop out [46]), and one as a high number of patients discontinued before the end of the study [47].

Selective reporting: nine comparisons were considered at low risk of bias; three comparisons did not report enough information, so were judged as at unclear risk of bias. In one comparison a high number of patients discontinued before the end of the study and were moved to an open-label baclofen study, and so considered as at high risk of bias [47].

Other biases: eight comparisons were considered as at low risk of bias; four comparisons did not report enough information, so were judged as at unclear risk of other bias. In one comparison not all patients had an AUD, and so was considered as at high risk of bias [47].

* + 1. *Severity of anxiety at baseline*

Only three comparisons [44HD,44LD,47] met the criteria at baseline (based on mean scores on anxiety scales) for the sample to be considered within the clinically anxious range. One comparison [32LD] reported mean baseline anxiety scores that spanned the threshold for the presence of anxiety disorders based on the defined cut-off levels (see Table 3), and so was not included in the analyses divided according to the presence or not of anxiety disorders at baseline. In all other comparisons, mean baseline scores were lower than the cut-off values suggestive of clinically significant pathology of anxiety.

* + 1. *Alcohol consumption at baseline*

As study samples varied considerably in terms of proportions of men and women included, it was not possible to differentiate between high and very high risk levels of alcohol consumption at baseline within samples (see above). However, using WHO drinking risk level criteria, seven comparisons [32HD, 32 LD, 34, 43HD, 43LD, 44HD, 44LD] were in participants whose drinking level at baseline were within or above the high risk range, regardless of gender (see Table 3). No data were available to calculate the drinking risk level for three comparisons [35(1), 35(2), 47], and one study [40] recorded a very low drinking risk level at baseline.

1. **Meta-analysis**
	1. **Abstinent days (%, studies = 12 weeks)**

The rate of abstinent days at 12 weeks was available for 8 comparisons (34, 35(1), 35(2), 40, 43HD, 43LD, 44HD, 44LD), involving 298 patients. The analysis found a significantly higher rate of abstinent days among patients treated with baclofen (p=0.03; moderate certainty evidence) (see Tables 3 and 4a).

One comparison had a high risk of bias (40): the analysis found a higher rate of abstinent days among patients treated with baclofen after exclusion of this study (p=0.02), without evidence of heterogeneity [7 comparisons (34, 35(1), 35(2), 43HD, 43LD, 44HD, 44LD) , 266 patients, MD: 12.96, 95% CI 1.78 to 24.13; Tau² = 97.61; Chi² = 11.00, df = 6 (p=0.09); I² = 45%].

* + 1. Studies divided into subgroups according to anxiety

Two comparisons (44HD,44LD), involving 104 patients reported mean baseline anxiety scores suggestive of reaching the threshold for an anxiety disorders based on the cut-off values of the scale used to evaluate anxiety symptoms. The analysis found a significantly higher rate of abstinent days among patients with levels of anxiety reaching the threshold for an anxiety disorder treated with baclofen rather than placebo (p=0.0010; moderate certainty evidence): none of them had a high risk of bias (see Table 3 and 4a).

Six comparisons (34, 35(1), 35(2), 40, 43HD, 43LD) involving 194 patients reported mean baseline anxiety scores **not** reaching the threshold for an anxiety disorders according to the cut-off values of the scales used to evaluate anxiety symptoms. The analysis found no difference in the rate of abstinent days between baclofen and placebo in the low anxiety group (p=0.28; moderate certainty evidence) (see Table 3 and 4a). One comparison had a high risk of bias (40). The analysis found no difference after its exclusion (p=0.24) without evidence of heterogeneity [5 comparisons (34, 35(1), 35(2), 43HD, 43LD), 162 patients, MD: 7.56, 95% CI -5.18 to 20.31; Tau² = 72.75; Chi² = 6.12, df = 4 (p=0.19); I² = 35%].

* 1. **Abstinent days (%, studies ≥ 12 weeks)**

The rate of abstinent days at 12 weeks or longer was available for 12 comparisons (32HD, 32LD, 34, 35(1), 35(2), 40, 43HD, 43LD, 44HD, 44LD, 45, 47), involving 825 patients. The analysis found high certainty evidence (see Table 4b) for a significantly higher rate of abstinent days among patients treated with baclofen (p=0.004). Two comparisons had a high risk of bias (40, 47). The analysis found a higher rate of abstinent days among patients treated with baclofen after their exclusion (p=0.03), without evidence of heterogeneity [10 comparisons (32HD, 32LD, 34, 35(1), 35(2), 43HD, 43LD, 44HD, 44LD, 45), 473 patients, MD: 8.50, 95% CI 0.99 to 16.01; Tau² = 47.87; Chi² = 13.73, df = 9 (p=0.13); I² =34%].

* + 1. Studies divided into subgroups according to anxiety

Three comparisons in studies with outcome data at 12 weeks or longer (44HD, 44LD, 47) involving 424 patients reported mean baseline anxiety scores suggestive of anxiety disorders based on the cut-off values of the scale used to evaluate anxiety symptoms. The analysis found high certainty evidence (see Table 4b and Figure 4) for a significantly higher rate of abstinent days among patients treated with baclofen (p<0.00001). One comparison had high risk of bias (47). After its exclusion, the analysis still found a significantly high rate of abstinent days among patients treated with baclofen (p=0.0010), without evidence of heterogeneity [2 comparisons (44HD,44LD), 104 patients, MD: 23.47, 95% CI 9.53 to 37.41; Tau² = 0.00; Chi² = 0.08, df = 1 (p=0.78); I² = 0%).

Eight comparisons (32HD, 34, 35(1), 35(2), 40, 43HD, 43LD, 45), involving 339 patients, reported mean baseline anxiety scores that **did not** reach the defined threshold for an anxiety disorder (see Table 3 and Figure 4). There was no difference in the rate of abstinent days between baclofen and placebo (p=0.20; moderate certainty evidence). One comparison had high risk of bias (40). After its exclusion, the analysis still did not find any difference in the rate of abstinent days among patients (not reaching the defined threshold for an anxiety disorder) treated with baclofen and placebo (P=0.18), without evidence of heterogeneity [7 comparisons (32HD, 34, 35(1), 35(2), 43HD, 43LD, 45), 307 patients, MD: 5.19, 95% CI -2.33 to 12.70; Tau² = 10.64; Chi² = 6.67, df = 6 (p=0.35); I² = 10%].

* 1. **Anxiety: Final score (studies = 12 weeks)**

The outcome rating on an anxiety scale at 12 weeks was available (or we calculated it) for 7 comparisons (34, 40, 43HD, 43LD, 44HD, 44LD, 47), involving 477 patients. The analysis found no difference between groups treated with baclofen or placebo for anxiety outcome rating (p=0.48, without heterogeneity, high certainty evidence, see Tables 3 and 4a). Two comparisons had high risk of bias (40,47). After their exclusion, the analysis still did not find any difference between anxiety outcome ratings (p=0.23), without evidence of heterogeneity, [5 comparisons (34, 43HD, 43LD, 44HD, 44LD), 226 patients, SMD: -0.17, 95% CI -0.44 to 0.10; Tau² = 0.00; Chi² = 1.61, df = 4 (p=0.81); I² = 0%].

* + 1. Studies divided into subgroups according to anxiety

Three comparisons (44HD, 44LD, 47), involving 323 patients, with mean baseline anxiety scores that reached the defined threshold for an anxiety disorder (see Table 3) reported an outcome anxiety rating at 12 weeks. The analysis of these comparisons found no difference between baclofen and placebo on the outcome anxiety rating (p=0.63, without evidence of heterogeneity, moderate certainty evidence; see Table 4a). One of them had high risk of bias (47). After its exclusion, the analysis still found no difference in outcome anxiety rating (p=0.46) [2 comparisons (44HD, 44LD), 104 patients, MD: -2.17, 95% CI -7.91 to 3.57; Tau² = 0.00; Chi² = 0.00, df = 1 (p=0.98); I² = 0%).

Four comparisons (34, 40, 43HD, 43LD), involving 154 patients, with mean baseline anxiety scores that **did not** reach the defined threshold for an anxiety disorder (see Table 3) reported an outcome anxiety rating at 12 weeks. The analysis found no difference in outcome anxiety rating between baclofen and placebo (p=0.58, without evidence of heterogeneity, moderate certainty evidence, see Table 4a). One comparison had a high risk of bias (40). After its exclusion, the analysis still found no difference in outcome anxiety rating (p=0.54, without evidence of heterogeneity) [3 comparisons (34, 43HD, 43LD), 122 patients, MD: -1.63, 95% CI -6.83 to 3.57; Tau² = 0.72; Chi² = 2.06, df = 2 (p=0.36); I² = 3%].

* 1. **Anxiety: Final score (studies ≥ 12 weeks)**

The outcome score on an anxiety rating at 12 weeks or longer was available (or could be calculated) for 10 comparisons (32HD, 32LD, 34, 40, 43HD, 43LD, 44HD, 44LD, 45, 47), involving 684 patients. The analysis found no difference between groups treated with baclofen or placebo for anxiety outcome rating (p=0.79, without heterogeneity, high certainty evidence, see Tables 3and 4b). Two studies had high risk of bias (40,47). After their exclusion, the analysis still did not find any difference between anxiety outcome ratings (P=0.80, without evidence of heterogeneity), [8 comparisons (32HD, 32LD, 34, 43HD, 43LD, 44HD, 44LD, 45), 433 patients, SMD: 0.03, 95% CI -0.19 to 0.25; Tau² = 0.02; Chi² = 8.51, df = 7 (p=0.29); I² = 18%].

* + 1. Studies divided into sub-groups according to anxiety

Three comparisons (44HD, 44LD, 47), involving 323 patients, with mean baseline anxiety scores that reached the defined threshold for an anxiety disorder (see Table 3) reported an outcome anxiety rating at 12 weeks. The analysis of these three comparisons found no difference between baclofen and placebo on the outcome anxiety rating (p=0.63, without evidence of heterogeneity, moderate certainty evidence, see Table 4b). One of them had a high risk of bias (47). After its exclusion, the analysis still found no difference in outcome anxiety rating (p=0.46) after its exclusion [2 comparisons (44HD,44LD), 104 patients, MD: -2.17, 95% CI -7.91 to 3.57; Tau² = 0.00; Chi² = 0.00, df = 1 (p=0.98); I² = 0%).

Six comparisons (32HD, 34, 40, 43HD, 43LD, 45), involving 299 patients, with mean baseline anxiety scores that **did not** reach the defined threshold for an anxiety disorder (see Table 3) reported an outcome anxiety rating at 12 weeks. The analysis found no difference in outcome anxiety rating between baclofen and placebo (p=0.70, without evidence of heterogeneity, moderate certainty evidence, see Table 4b). One comparison had high risk of bias (40). After its exclusion, the analysis still found no difference in outcome anxiety rating (p=0.75) [5 comparisons (32HD, 34, 43HD, 43LD, 45), 267 patients, SMD: 0.05, 95% CI -0.27 to 0.37; Tau² = 0.05; Chi² = 6.10, df = 4 (p=0.19); I² = 34%].

* 1. **Anxiety: Delta score (studies = 12 weeks)**

The difference between baseline and final score at 12 weeks was available (or we calculated it) for 7 comparisons (34, 40, 43HD, 43LD, 44HD, 44LD, 47), involving 578 patients. The analysis found no difference between baclofen and placebo for the change in anxiety level over 12 weeks (p=0.82, without heterogeneity, high certainty evidence, see Tables 3and 4a). Two comparisons had high risk of bias (40,47), but after their exclusion, the analysis still found no difference in change of anxiety score (p=0.45, without evidence of heterogeneity) [5 studies (34, 43HD, 43LD, 44HD, 44LD), 226 patients, SMD: 0.10, 95% CI -0.17 to 0.38; Tau² = 0.00; Chi² = 1.00, df = 4 (p=0.91); I² = 0%].

* + 1. Studies divided into sub-groups according to anxiety

Three comparisons (44HD, 44LD, 47), involving 424 patients, with mean baseline anxiety scores that reached the defined threshold for an anxiety disorder (see Table 3) reported a change in anxiety score at 12 weeks. The analysis of these comparisons found no difference between baclofen and placebo for the change in anxiety score over 12 weeks (p=0.99, without evidence of heterogeneity, high certainty evidence, see Table 4a). One study had high risk of bias (47). After its exclusion, the analysis still found no difference in change of anxiety score (p=0.31) without evidence of heterogeneity [2 comparisons (44HD,44LD), 104 participants, MD: 4.87, 95% CI -4.45 to 14.18; Tau² = 0.00; Chi² = 0.11, df = 1 (p=0.75); I² = 0%).

Four comparisons (34, 40, 43HD, 43LD), involving 154 patients, with mean baseline anxiety scores that **did not** reach the defined threshold for an anxiety disorder (see Table 3) reported a change in anxiety score at 12 weeks. The analysis found no difference between baclofen and placebo for the change in anxiety score over 12 weeks (P=0.66, without evidence of heterogeneity, moderate certainty evidence, see Table 4a). One comparison had high risk of bias (40). After its exclusion, the analysis still found no difference in change of anxiety score (p=0.97) [3 comparisons (34, 43HD, 43LD), 122 patients, MD: 0.11, 95% CI -9.97 to 10.20; Tau² = 0.00; Chi² = 0.24, df = 2 (p=0.88); I² = 0%].

* 1. **Anxiety: Delta score (studies ≥ 12 weeks)**

The difference between baseline and final score at 12 weeks or longer was available (or we calculated it) for 10 comparisons (32HD, 32LD, 34, 40, 43HD, 43LD, 44HD, 44LD, 45, 47), involving 785 patients. The analysis found no difference between baclofen and placebo for the change in anxiety score over 12 or more weeks (p=0.84, without heterogeneity, moderate certainty evidence, see Table 3 and 4b). Two comparisons had high risk of bias (40,47), but after their exclusion, the analysis still found no difference in change of anxiety score (p=0.99) without evidence of heterogeneity [8 comparisons (32HD, 32LD, 34, 43HD, 43LD, 44HD, 44LD, 45), 433 participants, SMD: -0.00, 95% CI -0.20 to 0.19; Tau² = 0.00; Chi² = 4.85, df = 7 (p=0.68); I² = 0%].

* + 1. Studies divided into sub-groups according to anxiety

Three comparisons (44HD, 44LD, 47), involving 424 patients, with mean baseline anxiety scores that reached the defined threshold for an anxiety disorder (see Table 3) reported a change in anxiety score at 12 weeks or longer*. The analysis found no difference between baclofen and placebo for the change in anxiety score over 12 or more weeks* (p=0.99, without evidence of heterogeneity, high certainty evidence, see Table 4b). One study had high risk of bias (47). After its exclusion, the analysis still found no difference in change of anxiety score (p=0.31) without evidence of heterogeneity [2 comparisons (44HD,44LD), 104 participants, MD: 4.87, 95% CI -4.45 to 14.18; Tau² = 0.00; Chi² = 0.11, df = 1 (p=0.75); I² = 0%) without evidence of heterogeneity.

Six comparisons (32HD, 34, 40, 43HD, 43LD, 45), involving 299 patients, with mean baseline anxiety scores that **did not** reach the defined threshold for an anxiety disorder (see Table 3) reported a change in anxiety score at 12 weeks or longer. The analysis found no difference between baclofen and placebo for the change in anxiety score over ≥12 weeks (p=0.78, without evidence of heterogeneity, moderate certainty evidence, see Table 4b). One study had a high risk of bias (40). After its exclusion, the analysis still found no difference in change of anxiety score [5 comparisons (32HD, 34, 43HD, 43LD, 45), 267 patients, SMD: -0.08, 95% CI -0.32 to 0.17; Tau² = 0.00; Chi² = 3.12, df = 4 (p=0.54); I² = 0%].

1. **DISCUSSION**

This study is the first to analyze systematically the impact of anxiety symptoms on the outcome of treatment with baclofen in patients with AUD, as well as explore the range and validity of the tools used to assess anxiety. We focused the analysis on those studies which had outcome data at 12 or more weeks, to enhance the stability of the results found, and limit the impact of acute alcohol withdrawal on measuring anxiety symptoms. There were limited differences between baclofen and placebo in the change on outcome measure at 12 weeks or more than 12 weeks suggesting that baclofen-placebo differences in anxiety symptoms at this point are relatively stable, as found in other treatment studies in patients with generalized anxiety disorder, social anxiety disorder and panic disorder [60].

There were significantly higher rates of abstinent days among patients treated with baclofen compared to placebo after 12 or more weeks (with moderate and high certainty of evidence, respectively). This remained so in the small number of comparisons where patients had high levels of anxiety at baseline (moderate and high certainty of evidence, respectively); however, the analysis found moderate certainty of evidence that patients treated with baclofen did not differ in the rate of abstinent days, after 12 or more weeks treatment, from those treated with placebo among those with low anxiety levels. We also found that, regardless of levels of anxiety at baseline, patients treated with baclofen did not differ from those treated with placebo in final anxiety rating expressed as final score and difference between basal and final score after 12 or more weeks (with moderate to high certainty of evidence).

These findings suggest that baclofen *may* be more effective in patients with higher baseline anxiety levels compared to those with lower baseline anxiety levels. There has been much speculation that a better response to the anti-alcohol effects of baclofen may be related to its anxiolytic actions [3,23,61-79]. However, our results demonstrate that this apparently better response among patients with higher baseline anxiety levels is not related to an anxiolytic effect induced by baclofen, as we found no differences in the final anxiety levels between patients treated with baclofen and placebo. As such, it is conceivable to speculate that reduction in anxiety is not a biobehavioral mechanism by which baclofen may reduce alcohol drinking. It is plausible that higher levels of anxiety represent an indirect proxy of a specific endophenotype of patients with AUD who may better respond to baclofen in terms of alcohol-related outcomes (e.g. patients with higher severity of AUD).

We also note the wide confidence intervals around the baseline anxiety levels which suggests significant inter-personal variation within the samples we analysed. We also observed that the measurement of anxiety symptoms was highly variable. Of those included in the meta-analysis, four studies used the Spielberger State anxiety subscale, two the Spielberger Trait anxiety subscale, two used the DASS, and one each the HAM-A and HADS-A. Of these the DASS, HAM-A and HADS-A have some evidence of sensitivity to change. Although the most widely used, the Spielberger Trait scale scores were reported as either State or Trait in different studies. Trait anxiety measures although constructed as a more stable phenomenon showed reductions over time in the samples in which it was used, and so its usefulness as a construct in clinical populations with AUD is hard to assess. Overall the STAI was not constructed for use in clinical samples or treatment studies and the only validation study against a diagnostic measure of anxiety was undertaken in patients with epilepsy [57], and so this reference point was used as the cut off to define clinically significant anxiety in this analysis.

Levels of anxiety at baseline were also highly variable, and using the cut off points for potential ‘caseness’ from the literature only two studies included patients who (at a group level) scored sufficiently highly to meet the criteria for an anxiety disorder [44,47]. Despite this variety, all but one study [43] showed a reduction in anxiety symptoms from baseline over time in both cases and controls, suggesting that the reduction of anxiety symptoms in early abstinence is a consistent phenomenon in need of further research.

The study is limited by the fact that anxiety levels could only be measured at the sample mean level and not individual level, making it difficult to draw firm conclusions about the impact of baclofen treatment in individual patients based on their level of anxiety.

Anxiety symptoms are frequently seen as a factor across many mental disorders (including depression, PTSD and OCD, autistic spectrum disorder) [80-83]. Their presence is frequently associated with higher suicide risk, longer duration of illness, and greater likelihood of treatment nonresponse [84]. Given anxiety symptoms are an important predictor of outcome across a range of disorders validated measures of severity and change are needed. Specifically, in AUD, the substantial coexistence of anxiety symptoms, and the potential impact of this on outcomes, requires further research to establish the optimal method for measuring anxiety constructs in this patient group. The STAI has been the most frequently used measure of anxiety symptoms in patients with AUD; however, given that both state and trait measures reduced over the 12 weeks of the study,  its utility is questionable and future studies are needed to investigate the value of using the STAI to assess anxiety levels in patients with AUD. Of note the General Anxiety Disorder -7 (GAD-7) questionnaire [85] has not been used in any of the studies in this analysis. It is a specific measure of anxiety symptom severity and has been shown to be sensitive to change in clinical populations [86,87], and consisting of only 7 questions (unlike 40 for the STAI) it would be less burdensome to use in clinical practice and research trials than the STAI, but needs validation in this population.

Given the putative mechanism of action of baclofen and the findings of this meta-analysis which suggest that levels of anxiety may be a predictor of outcome, there is a real need to reach consensus on how best to measure anxiety levels in patients with AUD. Improving the consistency of measures would enhance the quality of gathered evidence, and extend understanding of the potential role of anxiety in patients treated with baclofen, and help to clarify the role of anxiety on treatment outcomes in AUD.

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**FIGURES AND TABLES WITH LEGENDS**

Figure 1. Flow diagram of study selection

No legend

Figure 2. Risk of bias

No legend

Figure 3. Risk of bias summary

No legend

Figure 4. Forest plot of the comparison “baclofen *vs* placebo” in the rate of abstinent days

Legend

CI: confidence interval

HD: high dose

IV: inverse variance

ID: low dose

SD: standard deviation

Table 1. Main characteristics of the studies in which baclofen was administered to patients with AUD and anxiety symptoms were evaluated

Legend

ADS: Alcohol Dependence Scale

CAD: Cumulative Abstinent Days

CIWA-Ar: Clinical institute withdrawal assessment for alcohol scale – Revised

DASS: Depression, Anxiety and Stress Scale

DDD: Drinks per Drinking Day

HADS-A: Hospital Anxiety and Depression Scale

HAM-A or HARS: Hamilton Anxiety Rating Scale

HD: High Dose Baclofen

HDD: Heavy Drinking Day

LD: Low Dose Baclofen

NR: Not Reported

TAC: Total Alcohol Consumption

Table 2. Scales used to evaluate anxiety in studies of Baclofen in AUD

Legend

BDI: Beck Depression Inventory

BSI: Brief Symptom Inventory

SCL-90-R: Revised 90 item Symptom checklist from

DASS: Depression, Anxiety and Stress Scale

GHQ: General Health Questionnaire

HADS-A: Hospital Anxiety and Depression Scale

HAM- A: Hamilton Rating Scale for Anxiety

HARS: Hamilton Anxiety Rating Scale

STAI: State-Trait Anxiety Inventory

Table 3. Comparisons included in the meta-analysis in which baclofen was compared to placebo for at least 12 weeks

Legend Mean (SD)

ADS: Alcohol Dependence Scale

B: Baclofen

B+N: Baclofen + Naltrexone

BSI: Brief Symptoms Inventory

CAD: Cumulative Abstinent Days

DASS: Depression, Anxiety and Stress Scale

HADS-A: Hospital Anxiety and Depression Scale

HAM-A: Hamilton Anxiety

HARS: Hamilton Anxiety Rating Scale

HD: High Dose Baclofen

HDD: Heavy Drinking Day

LD: Low Dose Baclofen

N: Naltrexone

N.A.: Not Available

P: Placebo

SD: Standard Deviation

SE: Standard Error

T: Total

TAC: Total Alcohol Consumption

Table 4a: Summary of findings: Baclofen compared to placebo for patients with AUD in studies in which anxiety symptoms were evaluated (studies of 12 weeks)

Legend

a. Downgraded one level for imprecision: fewer than 400 participants included in the analysis

CI: confidence interval

MD: mean difference

SMD: standardized mean difference

Table 4b: Summary of findings: Baclofen compared to placebo for patients with AUD in studies in which anxiety symptoms were evaluated (studies of 12 weeks or longer)

Legend

a. Downgraded one level for imprecision: fewer than 400 participants included in the analysis

CI: confidence interval

MD: mean difference

SMD: standardized mean difference

**Supporting information**

**References S1**

References of excluded articles

Figure 1. Flow diagram of study selection

* 4 studies excluded because baclofen was compared only to another drug (no placebo arm)
* 5 studies excluded because the duration was < 12 weeks

Records excluded
(n = 208)

Studies included in qualitative synthesis
(n = 18)

Records after duplicates removed
(n = 286)

## Included

## Eligibility

Records screened
(n = 292)

Records identified through database searching
(n = 427)

## Identification

Additional records identified through reference searches

(n = 6)

## Screening

Full-text articles excluded, with reasons (e.g review article, observational study)
(n = 65)

Full-text articles assessed for eligibility
(n = 84)

At data extraction study excluded as anxiety not measured using anxiety scale

(n = 1)

Studies included in quantitative synthesis (meta-analysis)
(n = 9)









**Table 1. Main characteristics of the studies in which baclofen was administered to patients with AUD and anxiety symptoms were evaluated**

|  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- |
| **#** | **Study** | **Baclofen group** | **Control group** | **Baclofen** | **Treatment** | **Data available during and/or at the end of the study** |
| **Type of administration** | **Daily dose** | **Duration** | **Setting** | **Anxiety Symptoms** | **Alcohol**  |
| 1 | Addolorato 2002 [30]1 Drink = 12 g | N = 20 | **Placebo**N =19 | Fixed | 30 mg | 4 weeks | Outpatient | State STAI (baseline, 1,2,3,4 weeks) | CAD, DDD ((baseline, 1,2,3,4 weeks)Abstinent participants (4 weeks)  |
| 2 | Addolorato 2006 [31] | N = 18 | **Diazepam**N = 19 | Fixed | 30 mg | 10 days | Outpatient | CIWA-Ar (anxiety item: day 1, 2, 3, 4, 5 and 10 days) | NR |
| 3 | Beraha 2016 [32HD] | N = 58 | **Placebo** N = 31 | Titration | Up to 150 mg | 16 weeks | Inpatient and outpatient | Trait STAI (baseline, 4 weeks, and 16 weeks) | TAC, CAD, Abstinent participants, Heavy drinkers (16 weeks) |
| Beraha 2016 [32LD] 1 Drink =?  | N = 31 | **Placebo** N = 31 | Fixed | 30 mg | 16 weeks | Inpatient and outpatient | Trait STAI (baseline, 4 weeks, and 16 weeks) | TAC, CAD, Abstinent participants, Heavy drinkers (16 weeks) |
| 4 | Farokhnia 2017[33]1 Drink = 14 g | N = 18 | **Placebo** N = 16 | Fixed | 30 mg | 8 days | Outpatient laboratory | State and trait STAI (baseline and 8 days) | Alcohol self-administered Mini drinks self-administered  |
| 5 | Garbutt 2010a[34]1 Drink =14 g  | N = 40 | **Placebo** N = 40 | Fixed | 30 mg | 12 weeks | Outpatient | State STAI (baseline, 1, 2, 3, 4, 6, 8, 10 and 12 weeks) Trait STAI (baseline and 12 weeks)  | Abstinent days, HDD (baseline, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11 and 12 weeks) |
| 6 | Garbutt 2010b[35(1)]1 Drink =14 g  | N =10 | **Placebo** N = 10 | Fixed | 30 mg | 12 weeks | Outpatient | State STAI (baseline, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11 and 12 weeks; graph) | HDD, Abstinent days (baseline and at 12 weeks)Time to first HDD (baseline, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11 and 12 weeks; graph) |
| Garbutt 2010b[35(2)]1 Drink =14 g | Baclofen +NaltrexoneN = 10 | Naltrexone **+ placebo** N = 10 | Fixed | 30 mg | 12 weeks | Outpatient | State STAI (baseline, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11 and 12 weeks; graph) | HDD, Abstinent days (baseline and at 12 weeks)Time to first HDD (baseline, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11 and 12 weeks; graph) |
| Garbutt 2010b[35(3)]1 Drink =14 g | N =10 | **Naltrexone** N = 10 | Fixed | 30 mg | 12 weeks | Outpatient | State STAI (baseline, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11 and 12 weeks; graph) | HDD, Abstinent days (baseline and at 12 weeks)Time to first HDD (baseline, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11 and 12 weeks; graph) |
| 7 | Girish 2016 [36] | N = 30 | **Chlordiazepoxide**N = 30 | Fixed | 30 mg | 9 days | Inpatient | CIWA-Ar (anxiety item: baseline, 1, 2, 3, 4, 5, 6, 7, 8, and 9 days) Symptom free days | NR |
| 8 | Gulati 2019 [37] | N = 34 | **Lorazepam** N = 32 | Fixed | 30 mg | 8 days | Inpatient | CIWA-Ar (baseline, 1, 2, 3, 4, 5, 6, 7, and 8 days; data on anxiety subscale NR) | NR |
| 9 | Gupta 2016 [38] | N = 72 | **Benfothiamine**N = 50 | Fixed | 30 mg | 12 weeks | Outpatient | Decrease in HARS (12 weeks) | HDD, Time to relapse, CAD, Abstinent participants (12 weeks) |
| 10 | Hauser 2017[29]1 Drink = 14 g | N = 88 | **Placebo**N = 92 | Fixed | 30 mg | 12 weeks | Outpatient | BSI and PLC (baseline, 6, and 12 weeks)BSI and PLC (diff between baseline and 12 weeks) | D and HDD per 2 weeks (graph) |
| 11 | Krupitsky 1993[39a] | N = 29 | **Placebo**N = 23 | Fixed | 37.5 mg | 3 weeks | Inpatient | State, Trait STAI and MMPI –Taylor at 3 weeks | NR |
| Krupitsky 1993 [39b] | N = 29 | **Diazepam**N = 23 | Fixed | 37.5 mg | 3 weeks | Inpatient | State, Trait STAI and MMPI –Taylor at 3 weeks | NR |
| 12 | Krupitsky 2017[40] | N = 16 | **Placebo**N = 16 | Fixed | 50 mg | 12 weeks | Outpatient | State, Trait STAI and HARS at 12 weeks | Time to relapse at 12 weeks |
| 13 | Leggio 2013 [41]1 Drink =? | N = 7 | **Placebo**N = 7 | Fixed | 30 mg | 7 days | Outpatient laboratory | Trait and State STAI at 7 days | NR (lab) |
| 14 | Lyon 2011 [42]1 Drink =?  | N = 25 | **Placebo**N = 19 | Fixed | 30 mg | 3 days | Inpatient | NR | NR |
| 15 | Morley 2014 [43HD]1 Drink =10 g | N = 14 | **Placebo**N = 7 | Fixed | 60 mg | 12 weeks | Outpatient | State STAI at 12 weeks | DDD, HDD, Time to first lapse, Time to first relapse at 12 weeks |
| Morley 2014 [43LD]1 Drink =10 g | N = 14 | **Placebo**N = 7 | Fixed | 30 mg | 12 weeks  | Outpatient | State STAI at 12 weeks | DDD, HDD, Time to first lapse, Time to first relapse at 12 weeks |
| 16 | Morley 2018 [44HD]1 Drink =10 g | N = 35 | **Placebo**N = 16 | Fixed | 75 mg | 12 weeks | Outpatient | DASS (final score and diff betw baseline and final score) at 12 weeks | DDD, HDD, Time to first lapse, Time to first relapse, Abstinent days, Abstinent participants at 12 weeks |
| Morley 2018 [44LD]1 Drink =10 g | N = 36 | **Placebo**N = 17 | Fixed | 30 mg | 12 weeks | Outpatient | DASS (final score and diff betw baseline and final score) at 12 weeks | DDD, HDD, Time to first lapse, Time to first relapse, Abstinent days, Abstinent participants at 12 weeks |
| 17 | Muller 2015 [45] | N = 28 | **Placebo**N = 28 | Titration | Up to 270 mg | 24 weeks  | Outpatient | HAM-A (baseline, and 24 weeks) | CAD, Abstinent participants (baseline and 24 weeks) |
| 18 | Reynaud 2017 [46]  | N = 158 | **Placebo**N = 162 | Titration | Up to 180 mg | 26 weeks | Outpatient | NR | Abstinent participants for 20 consecutive weeks from day 28TAC & HDD (diff betw baseline and 6 months) |
| 19 | Rigal et al., in press [47]1 Drink =10 g | N = 162 | **Placebo**N = 158 | Titration | Up to 300 mg | 12 months | Outpatients | HADS-A (baseline, 3, 6, and 12 months)  | TAC, HDD, Abstinent days, Null or Low-risk alcohol drinkers, Heavy drinkers (baseline and 12 months) |

**Table 2. Scales used to evaluate anxiety in studies of baclofen in AUD**

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
| **Scale** | **Construction** | **Form** | **Rating** | **Primary samples** | **Use in Baclofen studies** |
| BSI [50] | Adapted from SCL-90-RGood psychometric properties for whole scaleSensitive to change | Nine subscales, including one on ‘anxiety’6 items on anxiety sub-scale (Q1,12,19,38,45,49)Timeframe: last 7/7 | 0 (not at all) - 4 (extremely)Max score on anxiety subscale = 24.***Or 63+ cut off for case in whole BSI*** | Designed to reflect the psychological distress and symptom patterns of psychiatric and medical patients, as well as community samples | Hauser 2017 [29] (Used full BSI, not anxiety subscale) |
| CIWA-Ar [51]  | Clinical rating scale for the monitoring of the severity of alcohol withdrawal | Ten questionsSingle question on anxiety 'Do you feel nervous?'Time frame: currently | 0 (no anxiety – at ease)4 (Moderately anxious, or guarded, so anxiety is inferred)7 (Equivalent to acute panic states as seen in severe delirium) | Patients in alcohol withdrawal | Addolorato 2002 [30]Girish 2016 [36]Gulati 2019 [37] |
| DASS [52] | Three scales to measure depression, anxiety and stress Sensitive to change | Each scale contains 14 items, including Anxiety scaleBurden: <2 minutesTimeframe: ‘over the past week’ | 0 (not at all) – 3 (very much)Range 0-420-7 normal;8-9 mild;10-14 moderate15-19 severe20+ extremely severe***Or 8+ cut off for case*** | Range of populations, developed in non-clinical setting | Morley 2018 [44] |
| HADS-A [53] | Not confounded by physical illnessSome sensitivity to changeGood correlation with BDI and GHQ | Self-assessment/ or interviewerBurden: <5 mins7 items (anxiety)Time frame: currently | 0 (not present) – 3 (severely)Range 0-210-7 normal;8-10 mild;11-14 moderate15-21 severe***Or 8+ cut off for case*** | Screening for significant anxiety in medically unwell patients. | Reynaud 2017 [46]Rigal et al., in press [47] |
| HAM-A or HARS [54] | More accurate than self-reportGood inter-rater reliabilityReasonable sensitivity to change | Clinician ratedBurden: 10-15 mins14 itemsTime frame: in the last week | 0 (not present)- 4 (severe)Range 0-560-5: no anxiety6-14: minor anxiety15-56: major anxiety***Or 14+ cut off for case*** | Based on clinical anxious samplesSeverity rating not diagnosis/ screeningIncludes depressive symptoms | Gupta 2016 [38]Krupitsky 2017 [40]Muller 2015 [45] |
| STAI Y [55]  | S (State) has lower reliability than T (Trait)T not sensitive to changeLimited discrimination from depressive states | Self-reportBurden: 20 minsS-20 items State – ‘now’T- 20 items Trait – ‘in general’ | 0 (almost never) – 4 (almost always)Range 0-80***Or 52+ cut off for case in Epilepy sample*** | Used extensively in patients with chronic medical conditionsMost commonly used, but developed for non-clinical conditions | Addolorato 2002 [30]Beraha 2016 [32]Faroknia 2017 [33]Garbutt 2010 [34]Garbutt 2010b [35]Krupitsky 1993 [39]Krupitsky 2017 [40]Leggio 2013 [41]Morley 2014 [43] |

**Table 3. Comparisons included in the meta-analysis in which baclofen was compared to placebo for at least 12 weeks**

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
|  | **Study** | N patients | **Males and Females** | **Baseline values** | **Data available at 12 weeks or later** |
| **Anxiety** | **Alcohol** | **Anxiety** | **Alcohol** |
| **Scale used and mean baseline score** | **Cut-off values for cases** | **Anxiety disorders** | **Mean****consumption in g/week** | **Scale used and** **mean final score** | **Scale used and mean difference between basal and final scores** | **Reduction in anxiety symptoms (% of basal score)** | **Abstinent days (%)** |
| 1 | Beraha 2016 [32HD] | B: 58P: 31 | M: 62 (69.66%)F: 27 (30.34%) | Trait STAIB: 49.7 (12.4)P: 48.6 (10.6) | 52 | NO | B: 896.7P: 864.4 | Trait STAI at 16 weeksB: 37.3 (12.1)P: 38.0 (11.1) | Trait STAI at 16 weeksB: 12.40 (24.50)P: 10.60 (21.70) | At 16 weeksB: 24.9%P: 21.8% | At 16 weeksB: 71.0 (34.94)P: 69.8 (28.98) |
| 2 | Beraha 2016 [32LD] | B: 31P: 31 | M: 42 (67.74%)F: 20 (32.26%) | Trait STAIB: 52.1 (9.5)P: 48.6 (10.6) | 52 | Unclear | B: 808.2P: 864.4 | Trait STAI at 16 weeksB: 42.2 (14.1)P: 38.0 (11.1) | Trait STAI at 16 weeksB: 9.90 (23.60)P: 10.60 (21.70) | At 16 weeksB: 19.0%P: 21.8% | At 16 weeksB: 67.7 (34.94)P: 69.8 (28.98) |
| 3 | Garbutt 2010a [34] | B: 40P: 40 | M: 44 (55.00%)F: 36 (45.00%) | State STAIB: 34.1 (10.8)P: 38.5 (12.7)Trait STAIB: 38.3 (8.7)P: 42.1 (11.4) | 52 | NO | B: 623.4P: 598.9 | State STAI at 12 weeksB: 27.5 (13.5)P: 32.2 (16.7) | State STAI at 12 weeksB: 6.6 (24.30)P: 6.3 (29.40) | At 12 weeksB: 19.3%P: 16.4% | At 12 weeksB: 49.9 (27.9)P: 50.6 (25.9) |
| 4 | Garbutt 2010b [35(1)] | B: 10P: 10 | M: 12 (60.00%)F: 8 (40.00%) | State STAIin a graph | 52 | NO | N.A. | N.A. | N.A. | N.A. | At 12 weeksB: 69.8 (34.94)P: 33.4 (28.98) |
| 5 | Garbutt 2010b [35(2)] | B + N: 10P + N: 10 | M: 9 (45.00%)F: 11 (55.00%) | State STAIin a graph | 52 | NO | N.A. | N.A. | N.A. | N.A. | At 12 weeksB: 55.4 (34.94)P: 58.9 (28.98) |
| 6 | Krupitsky 2017 [40] | B: 16P: 16 | M: 26 (81.25%)F: 6 (18.75%) | State STAIB: 43.2 (SE: 1.8)P: 37.2 (SE: 2.3)Trait STAIB: 45.5 (1.7)P: 41.7 (1.9) | 52 | NO | B: 8.5 (SE: 1.3)P: 9.7 (SE: 1.4) | State STAI at 12 weeksB: 34.8 (SE: 2.4)P: 33.8 (SE: 1.0) | State STAI at 12 weeksB: 8.4 (16.80) P: 3.4 (13.20) | At 12 weeksB: 19.4%P: 9.1% | At 12 weeksB: 79.1 (34.94)P: 78.7 (28.98) |
| 7 | Morley 2014 [43HD] | B: 14P: 7 | M: 7 (33.33%)F: 14 (66.67%) | State STAIB: 35.93 (30.33-41.53)P: 36.62 (30.68-42.55)Trait STAIB: 39.50 (33.79-45.21)P: 43.08 (35.71-50.44) | 52 | NO | B: 594.7 (475.7)P: 765.1 (655.8) | State STAI at 12 weeksB: 36.61 (28.24-44.98)P: 32.44 (22.59-42.29) | State STAI at 12 weeksB: -0.68 (26.67) P: 4.18 (30.13)  | At 12 weeksB: -1.9%P: 11.4% | At 12 weeksB: 33.2 (31.2)P: 22.6 (29.1) |
| 8 | Morley 2014 [43LD] | B: 14P: 7 | M: 12 (57.14%)F: 9 (42.86%) | State STAIB: 42.23 (33.39-51.07)P: 36.62 (30.68-42.55)Trait STAIB: 46.92 (40.20-53.65)P: 43.08 (35.71-50.44) | 52 | NO | B: 617.0 (466.4)P: 765.1 (655.8) | State STAI at 12 weeksB: 33.18 (24.13-42.22)P: 32.44 (22.59-42.29) | State STAI at 12 weeksB: 9.05 (33.95) P: 4.18 (30.13)  | At 12 weeksB: 21.4%P: 11.4% | At 12 weeksB: 30.9 (34.7)P: 22.6 (29.1) |
| 9 | Morley 2018 [44HD] | B: 35P: 16 | M: 36 (70.59%)F: 15 (29.41%) | DASSB: 11.82 (8.92)P: 10.56 (8.58) | 8 | YES | B: 720.3 (581.3)P: 708.2 (509.5) | DASS at 12 weeksB: 7.83 (7.55)P: 9.91 (15.74) | DASS at 12 weeksB: 3.99 (16.47) P: 0.65 (24.32) | At 12 weeksB: 33.8%P: 6.2% | At 12 weeksB: 64.56 (SE: 7.69)P: 43.35 (SE: 7.60) |
| 10 | Morley 2018 [44 LD] | B: 36P: 17 | M: 38 (71.70%)F: 15 (28.30%) | DASSB: 14.72 (10.00)P: 10.56 (8.58) | 8 | YES | B: 934.4 (764.7)P: 708.2 (509.5) | DASS at 12 weeksB: 7.65 (9.72)P: 9.91 (15.74) | DASS at 12 weeksB: 7.07 (19.72)P: 0.65 (24.32) | At 12 weeksB: 48.0%P: 6.2% | At 12 weeksB: 68.54 (SE: 5.90)P: 43.35 (SE: 7.60) |
| 11 | Muller 2015 [45] | B: 28P: 28 | M: 39 (69.64%)F: 17 (30.36%) | HAM-AB: 2.1 (2.3)P: 3 (3.5) | 14 | NO | N.A. | HAM-A at 24 weeksB: 1.9 (5)P: 0 (0) | HAM-A at 24 weeksB: 0.2 (7.30) P: 3.00 (3.50)  | At 24 weeksB: 9.5%P: 100% | At 24 weeksB: 49.3 (34.94)P: 39.8 (28.98) |
| 12 | Reynaud2017 [46] | B: 158P: 162 | M: 225 (72.58%)F: 85 (27.42%) | HADS-AB: 5.8 (1.5)P: 5.7 (1.6) | 8 | NO | N.A. | N.A. | N.A. | N.A. | N.A. |
| 13 | Rigal in press [47] | B: 162P: 158 | M: 224 (70.00%)F: 96 (30.00%) | HADS-AB: 10.4 (4.4)P: 10.6 (4.6) | 8 | YES | N.A. | HADS-A at 12 weeksB: 9 (4)P: 9 (5) | HADS-A at 12 weeksB: 1.4 (8.4) P: 1.6 (10.0)  | At 12 weeksB: 13.5%P: 15.1% | At 48 weeksB: 53.6P: 39.3 |

**Table 4a**

| **Summary of findings:**  |  |
| --- | --- |
| **Baclofen compared to placebo for patients with AUD in studies in which anxiety symptoms were evaluated (studies of 12 weeks)** |  |
| **Patient or population**: Patients with AUD (studies of 12 weeks) **Setting**: Outpatient **Intervention**: baclofen **Comparison**: placebo |  |
| **Outcomes** | **Type of patients** | **N° of participants****(comparisons)** | **Certainty of****the evidence****(GRADE)** | **Anticipated absolute effects\* (95% CI)** | **Heterogeneity** |
| **Risk with placebo** | **Risk difference** **with baclofen** |  |
| **Rate of abstinent days** at the end of treatment | All patients | 298(8 comparisons) | ⨁⨁⨁◯MODERATE a | The mean rate of abstinent days ranged from 22.6 to 78.7 | MD **11.24 higher**(1.29 higher to 21.19 higher)  | Tau² = 78.44; Chi² = 11.69, df = 7 (p=0.11); I² = 40% |
| Anxious patients | 104(2 comparisons) | ⨁⨁⨁◯MODERATE a | The mean rate of abstinent days ranged from 43.35 to 43.35 | MD **23.47 higher**(9.53 higher to 37.41 higher)  | Tau² = 0.00; Chi² = 0.08, df = 1 (p=0.78); I² = 0% |
| Not anxious patients | 194(6 comparisons) | ⨁⨁⨁◯MODERATE a | The mean rate of abstinent days ranged from 22.6 to 78.7 | MD 5.54 higher(4.56 lower to 15.64 higher)  | Tau² = 32.63; Chi² = 6.25, df = 5 (p=0.28); I² = 20% |
| **Anxiety symptoms** **expressed as final score** at the end of treatment | All patients | 477(7 comparisons) | ⨁⨁⨁⨁HIGH | The mean final score of anxiety symptoms ranged from 9 to 33.8 | SMD 0.07 SD lower(0.25 lower to 0.12 higher)  | Tau² = 0.00; Chi² = 2.71, df = 6 (p=0.84); I² = 0% |
| Anxious patients | 323(3 comparisons) | ⨁⨁⨁◯MODERATE a | The mean final score of anxiety symptoms ranged from 9 to 9.91 | SMD 0.05 SD lower (0.28 lower to 0.17 higher)  | Tau² = 0.00; Chi² = 0.56, df = 2 (p=0.75); I² = 0% |
| Not anxious patients | 154(4 comparisons)  | ⨁⨁⨁◯MODERATE a | The mean final score of anxiety symptoms ranged from 32.2 to 33.8 | SMD 0.09 SD lower(0.41 lower to 0.23 higher)  | Tau² = 0.00; Chi² = 2.11, df = 3 (p=0.55); I² = 0% |
| **Anxiety symptoms** **expressed as difference between basal and final score** at the end of treatment | All patients | 578(7 comparisons)  | ⨁⨁⨁⨁HIGH | The mean difference between basal and final score of anxiety symptoms ranged from 0.65 to 6.3 | SMD 0.02 SD higher(0.15 lower to 0.19 higher)  | Tau² = 0.00; Chi² = 2.66, df = 6 (p=0.85); I² = 0% |
| Anxious patients | 424(3 comparisons) | ⨁⨁⨁⨁HIGH | The mean difference between basal and final score of anxiety symptoms ranged from 0.65 to 2 | SMD 0.00 SD (0.19 lower to 0.19 higher)  | Tau² = 0.00; Chi² = 1.67, df = 2 (p=0.43); I² = 0% |
| Not anxious patients | 154(4 comparisons)  | ⨁⨁⨁◯MODERATE a | The mean difference between basal and final score of anxiety symptoms ranged from 3.4 to 6.3 | SMD 0.07 SD higher(0.25 lower to 0.39 higher) | Tau² = 0.00; Chi² = 0.86, df = 3 (p=0.84); I² = 0% |
|  | \***The risk in the intervention group** (and its 95% confidence interval) is based on the assumed risk in the comparison group and the **relative effect** of the intervention (and its 95% CI). **CI:** Confidence interval; **MD:** Mean difference; **SMD:** Standardised mean difference  |  |
|  | **GRADE Working Group grades of evidence****High certainty:** We are very confident that the true effect lies close to that of the estimate of the effect**Moderate certainty:** We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different**Low certainty:** Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect**Very low certainty:** We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect  |  |

#### Explanations

a. Downgraded one level for imprecision: fewer than 400 participants included in the analysis

**Table 4b**

| **Summary of findings:**  |  |
| --- | --- |
| **Baclofen compared to placebo for patients with AUD in studies in which anxiety symptoms were evaluated (studies of 12 weeks or longer)** |  |
| **Patient or population**: Patients with AUD (studies of 12 weeks or longer) **Setting**: Outpatient **Intervention**: baclofen **Comparison**: placebo |  |
| **Outcomes** | **Type of patients** | **N° of participants****(comparisons)** | **Certainty of****the evidence****(GRADE)** | **Anticipated absolute effects\* (95% CI)** | **Heterogeneity** |
| **Risk with placebo** | **Risk difference** **with baclofen** |
| **Rate of abstinent days** at the end of treatment | All patients | 825(12 comparisons) | ⨁⨁⨁⨁HIGH | mean rate of abstinent days ranged from 22.6 to 78.7 | MD **9.01 higher**(2.85 higher to 15.16 higher) | Tau² = 36.19; Chi² = 16.78, df = 11 (p=0.11); I² = 34% |
| Anxious patients | 424(3 comparisons) | ⨁⨁⨁⨁HIGH | mean rate of abstinent days ranged from 39.3 to 43.35 | MD **16.5 higher**(9.67 higher to 23.32 higher)  | Tau² = 0.00; Chi² = 1.34, df = 2 (p=0.51); I² = 0% |
| Not anxious patients | 339(8 comparisons) | ⨁⨁⨁◯MODERATE a | mean rate of abstinent days ranged from 22.6 to 78.7 | MD 4.35 higher(2.25 lower to 10.95 higher)  | Tau² = 0.00; Chi² = 6.80, df = 7 (p=0.45); I² = 0% |
| **Anxiety symptoms** **expressed as final score** at the end of treatment | All patients | 684(10 comparisons) | ⨁⨁⨁⨁HIGH | mean final score of anxiety symptoms ranged from 0 to 38 | SMD 0.02 SD higher(0.13 lower to 0.17 higher)  | Tau² = 0.00; Chi² = 8.64, df = 9 (p=0.47); I² = 0% |
| Anxious patients | 323(3 comparisons) | ⨁⨁⨁◯MODERATE a | mean final score of anxiety symptoms ranged from 9 to 9.91 | SMD 0.05 SD lower(0.28 lower to 0.17 higher)  | Tau² = 0.00; Chi² = 0.56, df = 2 (p=0.75); I² = 0% |
| Not anxious patients | 299(5 comparisons)  | ⨁⨁⨁◯MODERATE a | mean final score of anxiety symptoms ranged from 0 to 38 | SMD 0.05 SD higher(0.21 lower to 0.32 higher)  | Tau² = 0.02 Chi² = 6.18, df = 5 (p=0.29); I² = 19% |
| **Anxiety symptoms** **expressed as difference between basal and final score** at the end of treatment | All patients | 785(10 comparisons)  | ⨁⨁⨁◯MODERATE a | mean difference between basal and final score of anxiety symptoms ranged from 0.65 to 10.6 | SMD 0.01 SD lower(0.16 lower to 0.13 higher)  | Tau² = 0.00; Chi² = 5.96, df = 9 (p=0.74); I² = 0% |
| Anxious patients | 424(3 comparisons) | ⨁⨁⨁⨁HIGH | mean difference between basal and final score of anxiety symptoms ranged from 0.65 to 2 | SMD 0 SD (0.19 lower to 0.19 higher)  | Tau² = 0.00; Chi² = 1.67, df = 2 (p=0.43); I² = 0% |
| Not anxious patients | 299(6 comparisons)  | ⨁⨁⨁◯MODERATE a | mean difference between basal and final score of anxiety symptoms ranged from 3 to 10.6 | SMD 0.03 SD lower(0.27 lower to 0.2 higher) | Tau² = 0.00; Chi² = 4.24, df = 5 (p=0.52); I² = 0% |
|  | \***The risk in the intervention group** (and its 95% confidence interval) is based on the assumed risk in the comparison group and the **relative effect** of the intervention (and its 95% CI). **CI:** Confidence interval; **MD:** Mean difference; **SMD:** Standardised mean difference  |  |
|  | **GRADE Working Group grades of evidence****High certainty:** We are very confident that the true effect lies close to that of the estimate of the effect**Moderate certainty:** We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different**Low certainty:** Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect**Very low certainty:** We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect  |  |

#### Explanations

a. Downgraded one level for imprecision: fewer than 400 participants included in the analysis