Efficacy and Safety of Esmethadone (REL-1017) in Patients With Major Depressive Disorder and Inadequate Response to Standard Antidepressants: A Phase 3 Randomized Controlled Trial

Maurizio Fava, MD; Stephen M. Stahl, MD, PhD; Luca Pani, MD; Sara De Martin, PhD; Andrew J. Cutler, MD; Vladimir Maletic, MD, MS; Charles W. Gorodetzky, MD, PhD; Frank J. Voci, PhD; Frank L. Sapienza, MS; Thomas R. Kosten, MD; Cornelia Kröger, PhD; Paggard Champas, PhD; Cedric O’Gorman, MD; Clotilde Guidetti, MD; Andrea Alimonti, MD; Stefano Comai, PhD; Andrea Mattarei, PhD; Franco Folli, MD; David Bushnell, MS; Sergio Traversa, PharmD; Charles E. Inturrisi, PhD; Paolo L. Manfredi, MD; and Marco Pappagallo, MD

Abstract

Objective: To test esmethadone (REL-1017) as adjunctive treatment in patients with major depressive disorder (MDD) and inadequate response to standard antidepressants.

Methods: In this phase 3, double-blind, placebo-controlled trial, outpatients with MDD (DSM-5) were randomized to daily oral esmethadone (75 mg on day 1, followed by 25 mg daily on days 2 through 28) or placebo between December 2020 and December 2022. The primary efficacy measure was change from baseline (CFB) to day 28 in the Montgomery-Asberg Depression Rating Scale (MADRS) score. The intent-to-treat (ITT) population included all randomized participants. The per-protocol (PP) population included completers without major protocol deviations impacting assessment. Post hoc analyses included participants with severe depression (baseline MADRS score ≥35).

Results: For the ITT analysis (n = 227), mean CFB was 15.1 (SD 11.3) for esmethadone (n = 113) and 12.9 (SD 10.4) for placebo (n = 114), with a mean difference (MD) of 2.3, which was not statistically significant (P = .154; Cohen effect size [ES] = 0.21). Remission rates were 22.1% and 13.2% (P = .076), and response rates were 39.8% and 27.2% (P = .044) with esmethadone and placebo, respectively. For the PP analysis (n = 198), mean CFB was 15.6 (SD 11.2) for esmethadone (n = 101) and 12.5 (SD 9.9) for placebo (n = 97), with an MD of 3.1 (P = .051; ES = 0.29). In post hoc analyses of patients with baseline MADRS ≥35 in the ITT population (n = 112), MD was 6.9; P = .0059; ES = 0.57, and for the PP population (n = 98), MD was 7.9; P = .0015; ES = 0.69. Adverse events (AEs) were predominantly mild or moderate and transient, with no significant differences between groups.

Conclusions: The primary end point was not met. Esmethadone showed stronger efficacy in PP than in ITT analyses, with the discrepancy not attributable to AEs impacting treatment adherence. Significant efficacy occurred in post hoc analyses of patients with severe depression. Esmethadone was well tolerated, consistent with prior studies.

Trial Registration: ClinicalTrials.gov identifier: NCT04688164

J Clin Psychiatry 2024;85(3):24m15265

Author affiliations are listed at the end of this article.
Clinical Points

- The majority of patients fail to achieve remission from first-line antidepressants, and new treatment options are needed. No N-methyl-D-aspartate receptor antagonists are US Food and Drug Administration approved as adjunctive treatment for major depressive disorder.
- This trial of esmethadone did not meet its primary outcome; however, some of the secondary and post hoc outcomes were promising. Esmethadone was overall safe and well tolerated.

The neurobiology of MDD is progressively disengaging from the classic serotonergic hypothesis, and the risk-benefit ratio of available antidepressants, which mostly target monoaminergic neurotransmissions, has been questioned. Alternative hypotheses for the neurobiology of MDD implicate impairment of neural plasticity. The pivotal role of N-methyl-D-aspartate receptors (NMDARs) in neural plasticity is well established and dysregulation of glutamatergic signaling via NMDARs is increasingly recognized as a potential pathological mechanism for neuropsychiatric disorders and a target for novel antidepressants.

Uncompetitive NMDAR antagonists have been approved by the Food and Drug Administration (FDA) for treating MDD. Intranasal esketamine has been FDA-approved for treatment-resistant depression (TRD) and for MDD with suicidal ideation; the oral twice daily dextromethorphan-bupropion combination has been approved for MDD. Recently, in silico and in vitro studies have advanced our understanding of the interactions of uncompetitive NMDAR antagonists and NMDARs and the comparative pharmacological affinity and activity of different NMDAR uncompetitive antagonists. Experimental models of depressive-like behavior suggest that NMDAR uncompetitive antagonists may improve depressive-like behavior via brain-derived neurotrophic factor–dependent restoration of neural plasticity. Recent reviews indicate that NMDAR uncompetitive antagonists, including esmethadone (REL-1017), are at the forefront among novel antidepressant candidates.

Esmethadone is a novel NMDAR uncompetitive antagonist antidepressant candidate with promising safety, tolerability, and efficacy results from phase 1 and phase 2 trials. In phase 1 studies, oral esmethadone was found to be safe and well tolerated at doses up to 150 mg, nausea and vomiting limited the use of higher doses, and oral esmethadone exhibits linear pharmacokinetics with dose proportionality. A phase 2 study with adjunctive oral once daily esmethadone confirmed the safety and tolerability seen in phase 1 and showed rapid and robust efficacy.

Chiral configuration is known to impart opioid activity to racemic opioid molecules; as a rule, after chiral separation, only one of the two chiral opioid enantiomers retains meaningful opioid agonist activity. Esmethadone, the dextro-isomer of racemic methadone, is a low affinity, low-potency NMDAR uncompetitive antagonist that binds to the phenylcyclohexyl site of the NMDAR at low-micromolar half-maximal inhibitory concentrations (IC$_{50}$). Esmethadone has 20- to 40-fold lower affinity for mu opioid receptors compared with levomethadone and does not contribute in a meaningful way to the opioid effects of racemic methadone, which are a result of its enantiomer, levomethadone. Esmethadone may even act as an opioid antagonist within the racemic mixture, attenuating the opioid agonist effects of levomethadone. Preclinical studies showed lack of self-administration in animal models predictive of abuse potential. Clinical studies in recreational substance users showed that tested doses of esmethadone up to 150 mg were statistically equivalent to placebo. These recent state-of-the-art studies to define abuse potential confirm prior literature indicating that esmethadone has no meaningful opioid agonist activity and no meaningful abuse potential.

In addition to opioid affinity and NMDAR affinity, esmethadone inhibits serotonin and norepinephrine transporters, with affinities in the micromolar range that are approximately 500-fold (serotonin transporters) and 100-fold (norepinephrine transporters) lower than those seen for duloxetine. These 100-plus fold differences in IC$_{50}$ compared with duloxetine suggest that a primary monoaminergic antidepressant mechanism of action for esmethadone is unlikely. Preclinical studies and the ongoing clinical use of NMDAR antagonists are advancing our understanding of the neurobiology of MDD. Esmethadone appears to have preferential activity on GluN2D subtypes, an NMDAR subtype implicated in MDD, triggered by chronic excitotoxicity at resting membrane potential.

Leaving aside the mechanism of action, oral once daily esmethadone showed efficacy in a phase 2 trial and confirmed a favorable tolerability and safety profile across multiple clinical studies without any signal for the metabolic, cardiovascular, and neurological side effects seen with atypical antipsychotic drugs currently approved as adjunctive treatment of MDD. Esmethadone does not cause Olney lesions in rats, a potential indicator of its safety compared to other uncompetitive NMDAR antagonists. Long-term human exposure to esmethadone in millions of patients treated for over half a century with racemic S,R-methadone for opioid use disorder and pain, generally at doses higher than the doses proposed for MDD, has been safe overall. We therefore evaluated the efficacy, safety, and
tolerability of esmethadone in a phase 3 study of patients with MDD unresponsive to monoaminergic antidepressants.

METHODS

This multicenter trial was conducted in accordance with the International Council on Harmonization guidelines for Good Clinical Practice, the principles of the Declaration of Helsinki, and all regulatory requirements. The study protocol was reviewed and approved by an institutional review board, and written informed consent was obtained from all participants after receiving a complete description of the study and prior to any study procedure.

Study Design

This was a 28-day double-blind, placebo-controlled, randomized phase 3 trial conducted in 43 centers in the United States from December 2020 to December 2022. The overall duration of the trial, including the screening period, was approximately 58 days (Supplementary Figure 1). During the screening period, clinicians from the Massachusetts General Hospital Clinical Trials Network and Institute (MGH-CTNI) independently assessed prior antidepressant treatment response and history using the MGH Antidepressant Treatment Response Questionnaire (ATRQ). Raters from the sites were required to obtain certification and training prior to rating study participants. The screening and rater review process are presented in Supplementary Appendix 1.

Participants

Adult patients ages 18–65 years were eligible if they met criteria for MDD defined by Diagnostic and Statistical Manual of Mental Disorders, Fifth Edition (DSM-5 criteria), if they had a 17-item Hamilton Depression Rating Scale (HAM-D) score ≥19 and did not show an increase in absolute value of >40% or a decrease >20% on the HAM-D score between screening and baseline, and if they had a body mass index (BMI) between 18 and 30 kg/m². Patients also had a current major depressive episode lasting 8 weeks to 36 months and had an inadequate response from 1–3 courses of antidepressant treatment during the same episode, as determined by a clinician from the MGH-CTNI who independently assessed each patient using the MGH-ATRQ. Patients were required to have a Montgomery-Asberg Depression Rating Scale (MADRS) total score at baseline ≥24 points. Patients were taking the same SSRI, serotonin-norepinephrine reuptake inhibitor, or bupropion for at least 8 weeks prior to screening and maintained the same adequate dose for the last 4 weeks. Exclusion criteria included use of opioids, anxiolytics, antipsychotics, anticonvulsants, mood stabilizers, stimulants, NMDAR antagonists, electroconvulsive therapy, vagus nerve stimulation, or repetitive transcranial magnetic stimulation. Any medication taken consistently by the patient for 30 days prior to screening that was not a prohibited medication was continued during the trial (see Supplementary Table 1 for list of prohibited medications). Initiation of medications during the trial was not allowed. Patients at risk for suicide and patients with history of bipolar disorder, psychosis or mania, substance use disorder or heavy alcohol use, and patients with positive results on urine test for alcohol or illicit drugs also were excluded. For a full list of inclusion/exclusion criteria, please see Supplementary Appendix 2.

Procedures

Patients were randomized by an unblinded pharmacist through an interactive web response system in a 1:1 ratio to esmethadone or placebo. The randomization code used in the interactive web response system was prepared by a statistician who was not involved in the study. On day 1, patients received a loading dose of esmethadone 75 mg or placebo. The loading dose of esmethadone was computed based on pharmacokinetic data from a phase 1 study to achieve steady-state concentrations by day 1. Oral 75 mg esmethadone is well tolerated without evidence of subjective effects in healthy volunteers, in patients with MDD, or in recreational substance users, making unblinding unlikely. On days 2–28, patients received esmethadone 25 mg or placebo. The first 200 patients underwent a 2-week safety-withdrawal assessment after discontinuation of study medication. Safety assessments included evaluation of frequency and severity of adverse events (AEs), and changes in clinical laboratory tests (chemistry, hematology, and urinalysis), electrocardiogram, physical examination, vital signs (blood pressure, heart rate, and respiratory rate), weight, and body temperature. Other safety assessments included suicidal ideation and behavior with the Columbia-Suicide Severity Rating Scale (C-SSRS; a higher score indicates a higher intensity of suicidal ideation); present state dissociative symptoms with the Clinician-Administered Dissociative States Scale (CADSS; a higher score indicates a higher likelihood of the presence of a dissociative state); and psychotic symptoms with the 4-item Positive Symptom Rating Scale. “Drug liking,” “drug high,” and “desire to take the drug again” were assessed with a 100-point visual analog scale (VAS). The Misuse, Abuse, and Diversion Drug Event Reporting System (MADDERS) was used to assess potentially abuse-related events. Potential withdrawal was assessed for 14 days after abrupt treatment discontinuation (days 28–42) using the Physician Withdrawal Checklist (PWC), Clinical Opiate Withdrawal Scale (COWS),
and Subjective Opiate Withdrawal Scale (SOWS). Abuse-related AEs were defined as AEs of special interest.

### Outcomes
The primary efficacy end point was the mean change from baseline (CFB) to day 28 for the MADRS total score. Remission (MADRS total score ≤10) and response (≥50% MADRS improvement from baseline) were key secondary end points.

### Statistical Analysis
The study was designed to achieve 90% power, with an overall 2-tailed alpha level of 0.05. The sample size was estimated based on a Cohen effect size (ES) assumption of 0.45 in the main estimand of the primary efficacy end point, defined as the absolute CFB to day 28 in the MADRS total score. Using the treatment policy for intercurrent events, a net sample size of 210 complete patients was estimated, which corresponded to approximately 220 randomized participants considering an early termination rate of 5%. The sample size of 210 total completers was computed using a 2-sided t-test assuming equal variances. The assumed ES was determined by taking a conservative approach from analysis of a phase 2 study and considering studies of adjunctive treatment of MDD with brexipiprazole and esketamine. Sample size calculations were performed with the software package nQuery 8, version 8.5.2.0.

The intent-to-treat (ITT) population included all randomized patients. The prespecified supportive analysis was the per-protocol (PP) population, which...
Table 1.
Baseline Characteristics—Safety Population

<table>
<thead>
<tr>
<th></th>
<th>All Patients (N = 227)</th>
<th>Esmethadone (N = 113)</th>
<th>Placebo (N = 114)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mean</td>
<td>SD</td>
<td>Mean</td>
</tr>
<tr>
<td>Age, y</td>
<td>42.5</td>
<td>14.6</td>
<td>43.3</td>
</tr>
<tr>
<td>Body mass index, kg/m²</td>
<td>26.0</td>
<td>3.0</td>
<td>25.8</td>
</tr>
<tr>
<td>N</td>
<td>%</td>
<td></td>
<td>%</td>
</tr>
<tr>
<td>Age ≥50 years</td>
<td>97</td>
<td>42.7</td>
<td>50</td>
</tr>
<tr>
<td>Female</td>
<td>169</td>
<td>74.4</td>
<td>82</td>
</tr>
<tr>
<td>Race</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Asian</td>
<td>13</td>
<td>5.7</td>
<td>6</td>
</tr>
<tr>
<td>Black/African American</td>
<td>30</td>
<td>13.2</td>
<td>16</td>
</tr>
<tr>
<td>White</td>
<td>175</td>
<td>77.1</td>
<td>85</td>
</tr>
<tr>
<td>Multiracial</td>
<td>6</td>
<td>2.6</td>
<td>4</td>
</tr>
<tr>
<td>Other</td>
<td>3</td>
<td>1.3</td>
<td>2</td>
</tr>
<tr>
<td>Hispanic or Latino</td>
<td>52</td>
<td>22.9</td>
<td>29</td>
</tr>
<tr>
<td></td>
<td>Mean</td>
<td>SD</td>
<td>Mean</td>
</tr>
<tr>
<td>MADRS total score</td>
<td>35</td>
<td>4.8</td>
<td>34.7</td>
</tr>
<tr>
<td>Time since first diagnosis, y</td>
<td>15.4</td>
<td>10.7</td>
<td>15.1</td>
</tr>
<tr>
<td>Lifetime depressive episodes</td>
<td>6.9</td>
<td>8.4</td>
<td>7.7</td>
</tr>
<tr>
<td>Depression episodes in past 5 years</td>
<td>2.2</td>
<td>1.2</td>
<td>2.3</td>
</tr>
</tbody>
</table>

Abbreviation: MADRS = Montgomery-Asberg Depression Rating Scale.

included all patients completing the 28-day treatment period without major protocol deviations impacting efficacy assessments. A post hoc analysis was performed in patients with severe depression, defined as baseline MADRS ≥35. The safety population comprised all randomized patients who received any dose of study drug and coincided with the ITT. Data for the primary end point were analyzed using mean difference (MD) in MADRS total score and using a mixed-effect model with repeated measures (MMRM), with consideration of repeated assessments of the MADRS 10 total score and with the independent variables of treatment, visit, the interaction of treatment and visit, and baseline MADRS total score. Comparisons of response rate and remission rate were analyzed using 95% CI for MD (Wilson confidence limits), χ² test (2-sided with α = 0.05), and odds ratio (OR) with 95% CI.

RESULTS

The ITT population comprised 227 randomized patients. The PP population comprised 198 patients completing treatment without major protocol deviations affecting efficacy assessments (Figure 1). Among the 29 patients included in the ITT population who were excluded pre-database lock from the PP analysis (17 placebo and 12 esmethadone), 19 patients (13 placebo and 6 esmethadone) did not complete treatment, and 11 patients (5 placebo and 6 esmethadone) had major protocol deviations affecting treatment outcome (1 patient did not complete treatment and had a major protocol deviation).

Patients in the esmethadone and placebo groups were generally comparable for baseline demographic and clinical characteristics (Table 1). Mean (SD) age was 43.5 (14.6) years, 74% were female, and 77% were white. Mean baseline MADRS score was 35.0 (4.8), and approximately 50% of patients (112/227) has a baseline MADRS score of ≥35, indicating severe depression. Mean (SD) duration of the current major depressive episode was 1.2 (2.2) years. Patients had an average of 6.9 (8.4) lifetime major depressive episodes and an average of 15.4 (10.7) years since the first MDD diagnosis, indicating that most patients had a long history of depression.

Efficacy

For the primary end point in the ITT population, mean (SD) CFB to day 28 for the MADRS total score was 15.1 (11.3) for esmethadone (n = 113) and 12.9 (10.4) for placebo (n = 114) (MD: 2.3 (10.9); P = .154; ES = 0.21 (Table 2, Figure 2A). In the PP population, mean (SD) CFB for MADRS was 15.6 (11.2) for esmethadone (n = 101) and 12.5 (9.9) for placebo (n = 97) (MD: 3.1 (10.6); P = .051; ES = 0.29 (Figure 2B).

Remission rate at day 28 was 22.1% with esmethadone and 13.2% with placebo (MD: 9.0%, 95% CI, −0.9 to 18.8; P = .076; OR: 1.88, 95% CI, 0.88 to 4.08). Response rate was 39.8% with esmethadone and 27.2% with placebo (MD: 12.6%, 95% CI, 0.5 to 24.8; P = .044; OR: 1.77, 95% CI, 0.98 to 3.23) (Figure 2C). In the PP population, remission rates at day 28 were 23.8% and 13.4%
Mean Change From Baseline to Day 28 for MADRS Total Score in the ITT Population, in the PP Population Prespecified Supportive Analysis and in Post Hoc Disease-Severity Population Enriched Analyses of Patients With Severe Depression (MADRS ≥35)

<table>
<thead>
<tr>
<th></th>
<th>Esmethadone</th>
<th>Placebo</th>
<th>Difference drug minus placebo</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>N</td>
<td>Mean</td>
<td>SD</td>
</tr>
<tr>
<td><strong>Primary end point</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ITT (n = 227)</td>
<td>113</td>
<td>-15.1</td>
<td>11.3</td>
</tr>
<tr>
<td>PP (n = 190)</td>
<td>100</td>
<td>-15.6</td>
<td>11.2</td>
</tr>
<tr>
<td><strong>Post hoc analyses</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ITT MADRS ≥35 (n = 112)</td>
<td>51</td>
<td>-18.5</td>
<td>13.3</td>
</tr>
<tr>
<td>PP MADRS ≥35 (n = 90)</td>
<td>45</td>
<td>-19.2</td>
<td>13.0</td>
</tr>
</tbody>
</table>

Bold values are significant.

Abbreviations: ITT = intent-to-treat; MADRS = Montgomery-Asberg Depression Rating Scale; PP = per-protocol.

(P = .062), and response rates were 42.6% and 29.9% (P = .064) for esmethadone and placebo, respectively.

In post hoc analyses of patients with severe depression (MADRS score ≥35 at baseline), significant improvement occurred with esmethadone vs. placebo in both the ITT and PP populations (MD CFB 6.9 and 7.9; P = .0059 and P = .0015; ES = 0.57 and 0.68, respectively) (Table 2 and Figure 3). In this post hoc disease-severity population analysis, remission rates at day 28 were 27.5% and 11.5% (MD: 16.0%, 95% CI, 1.3 to 30.6; P = .031), and response rates at day 28 were 43.1% and 21.3% (MD: 21.8%, 95% CI, 4.8 to 38.9; P = .013) for esmethadone and placebo, respectively, in the ITT population. Remission rates were 28.9% and 11.3% (MD: 17.6%, 95% CI, 1.8 to 33.3; P = .028), and response rates were 46.7% and 22.6% (MD: 24.0%, 95% CI, 5.6 to 42.4; P = .012) for esmethadone and placebo, respectively, in the PP population.

The MMRM analysis for the ITT, PP, and post hoc severe depression populations showed significant differences between esmethadone and placebo only for the severe population with ES of 0.51 and 0.64 for the ITT and PP populations (Supplementary Table 2).

Safety and Tolerability

The incidence of treatment-emergent adverse events (TEAEs) was comparable between esmethadone and placebo (Table 3). No serious AE or deaths related to study treatment were reported. Seven patients discontinued study treatment due to AE (5 placebo [anxiety, depression, panic attack, pregnancy, throat irritation, pruritus, urticaria] and 2 esmethadone [suicidal ideation, nausea, dizziness]). AEs were predominantly mild or moderate and transient. The most common TEAEs were headache, COVID-19, dizziness, and gastrointestinal complaints. There were 3 unrelated or unlikely related serious AEs in the esmethadone group.

No clinically significant findings were observed with esmethadone for vital signs, body weight/BMI, or clinical laboratory testing. For the esmethadone group, mean (SD) weight and BMI were 72.5 (11.0) kg and 25.8 (7.8) kg/m² at baseline and 73.0 (10.8) kg and 26.0 (2.7) kg/m² at end of treatment; for the placebo group, mean (SD) weight and BMI were 73.8 (13.2) kg and 26.3 (3.2) kg/m² at baseline and 75.4 (14.5) kg and 26.8 (3.4) kg/m² at end of treatment. One patient with esmethadone had an increase in alanine aminotransferase levels >3 times upper limit of normal, and 2 patients with esmethadone had an increase in aspartate aminotransferase levels >3 times upper limit of normal. These events were attributed to viral illness and use of acetaminophen in 1 patient, to HIV medications in 1 patient, and to excessive physical exercise in 1 patient. These 3 AEs were considered unlikely related to esmethadone. No abnormal liver function tests occurred with placebo.

Mean (SD) CFB to day 28 for the QT interval with Fridericia correction (QTcF) interval was 0.24 (13.5) ms for esmethadone and −3.1 (11.9) ms for placebo. Analysis of worst CFB at any time point for the QTcF interval showed no QTcF increase ≥60 ms and no QTcF >480 ms with either esmethadone or placebo. No differences between esmethadone and placebo were observed for shifts from baseline to worst value on the C-SSRS. No signal of abuse potential was observed from CADSS and VAS, and no signal for withdrawal was seen on PWC, COWS, or SOWS. No cases of withdrawal, misuse, abuse, or diversion were recorded in MADERS. In this study, mean esmethadone end-of-dose concentration at steady state day 7 was 192 ng/mL.

DISCUSSION

Approximately 50%–60% of patients with MDD fail to achieve an adequate response following their first antidepressant treatment.7 These patients are left with few satisfactory pharmacological options. Several atypical antipsychotics are FDA-approved for the treatment of depressed patients with inadequate response to first-line antidepressant therapy. However, atypical antipsychotics carry significant neurological,
Figure 2.
Mean Change From Baseline to Day 28 for Montgomery-Asberg Depression Rating Scale (MADRS) Total Score for the Primary Efficacy Analysis in the Intent-to-Treat (A) and Per-Protocol (B) Populations and Response and Remission Rates (C)*

A  Intent-to-Treat Population

- Esmethadone (n=113)
- Placebo (n=114)

Mean difference: –2.3
Effect size = 0.21
P = 0.154

B  Per-Protocol Population

- Esmethadone (n=101)
- Placebo (n=97)

Mean difference: –3.1
Effect size = 0.29
P = 0.051

C  Response and Remission Rates

- Esmethadone
- Placebo

P = .044
39.8
27.2
P = .076
22.1
13.2

*Response (P = .044) and remission (P = .076). Response was at least 50% reduction in MADRS from baseline; remission was MADRS score of 10 or less.
cardiovascular, and metabolic side effects. No NMDAR antagonists are FDA-approved as adjunctive treatment for MDD. Intranasal esketamine, approved for MDD with acute suicidal ideation and TRD, has limitations due to dissociative and psychotomimetic effects requiring clinical supervision. The oral twice daily dextromethorphan-bupropion combination has been approved for MDD.

In this multicenter trial, esmethadone did not meet the primary efficacy end point in the ITT analysis (MD = 2.3; P = .154; ES = 0.21). However, esmethadone showed a statistically significant improvement in response rate compared to placebo (P = .044) and an encouraging nonsignificant trend for improvement in remission rate (P = .076). In the PP prespecified supportive analysis, results trended toward a more favorable outcome (MD = 3.1;
P = .051; ES = 0.29). In this trial, the difference in efficacy between ITT and PP analyses is attributed to the exclusion of patients with protocol noncompliance due to reasons other than esmethadone-related AEs. The 29 patients who were not compliant with the protocol and were not included in the PP analysis are unlikely to inform on treatment efficacy, suggesting that in this study, the PP analysis of compliant patients may have provided an enhanced measure of efficacy. Arguably, if the sample size calculation had been based on more widely accepted assumed ES for standard antidepressants, ie, ES around 0.3, the trial may have potentially met its primary end point. Furthermore, in the analyzed populations (ITT, PP, severe depression population), the esmethadone-treated groups had a slope that was still declining at the conclusion of the 4-week treatment period, potentially suggesting that a longer study may have led to enhanced separation from placebo.

No biomarker consistently improves the accuracy of MDD diagnosis, and according to DSM-5 criteria, the assessment of severity is critical for diagnosis. A MADRS score of 35 or more indicates severe depression and may enhance diagnostic accuracy. We hypothesize that in this study, the subgroup of patients with severe depression may have included a lower number of patients with transient reactive depression and a lower number of “professional patients,” thus explaining the favorable results seen in the severity-enriched post hoc analysis.

Adverse events were mild or moderate and transient and were comparable in the two groups. No treatment-related serious AEs were observed. No withdrawal effects and no signals of potential abuse were observed on a broad battery of specialized measurements, confirming the lack of meaningful abuse potential seen in ad hoc studies. The effects on QT prolongation in these patients with MDD and concomitant antidepressants were mild and consistent with prior results and with results seen in drug-free healthy volunteers.

This study may have been underpowered relative to other antidepressant trials, and the 4-week treatment course may not have captured the full therapeutic effect of esmethadone. The safety/tolerability profile was consistent with previous phase 1 and phase 2 studies.

**CONCLUSION**

While neither the primary efficacy end point nor the secondary end point of remission was achieved, the statistically significant difference in the key secondary end point of response rate and results in the prespecified supportive PP analysis suggest meaningful antidepressant effects of esmethadone. Post hoc analyses of patients with severe MDD showed statistically significant efficacy with robust ES, consistent with results seen in the phase 2 trial. The overall results from this study strengthen prior evidence for the efficacy, safety, and tolerability of esmethadone as a promising antidepressant for the adjunctive treatment of MDD. The side effect profile of esmethadone compares favorably with the side effects of the currently FDA-approved adjunctive treatments for MDD. Further phase 3 studies will better characterize the potential efficacy of esmethadone as adjunctive treatment in patients with persistent MDD despite ongoing treatment with adequate doses of standard antidepressants.

**Article Information**

Published Online: June 17, 2024. https://doi.org/10.4088/JCP.24m15265
© 2024 Physicians Postgraduate Press, Inc.
Submitted: January 16, 2024; accepted March 25, 2024.
References


Esmethadone in Patients With MDD and Inadequate Antidepressant Response


Efficacy and Safety of Esmethadone (REL-1017) in Patients with Major Depressive Disorder and Inadequate Response to Standard Antidepressants: A Phase 3 Randomized Controlled Trial

Maurizio Fava, MD; Stephen M. Stahl, MD, PhD; Luca Pani, MD; Sara De Martin, PhD; Andrew J. Cutler, MD; Vladimir Maletic, MD, MS; Charles W. Gorodetzky, MD, PhD; Frank J. Vocci, PhD; Frank L. Sapienza, MS; Thomas R. Kosten, MD; Cornelia Kröger, PhD; Paggard Champasa, PhD; Cedric O’Gorman, MD; Clotilde Guidetti, MD; Andrea Alimonti, MD; Stefano Comai, PhD; Andrea Mattarei, PhD; Franco Folli, MD; David Bushnell, MS; Sergio Traversa, PharmD; Charles E. Inturrisi, PhD; Paolo L. Manfredi, MD; and Marco Pappagallo, MD

10.4088/JCP.24m15265

LIST OF SUPPLEMENTARY MATERIAL FOR THE ARTICLE

1. Appendix 1  Screening and Rater Review Process
2. Appendix 2  Inclusion and Exclusion Criteria
3. Figure 1    Study Design
4. Table 1     Time from Discontinuation of Prohibited Medications, Supplements, and Other Substances or Therapies
5. Table 2     MMRM Analysis for Intent-To-Treat, Per Protocol, and Post Hoc Severe Depression Population for Mean Change from Baseline to Day 28

DISCLAIMER

This Supplementary Material has been provided by the author(s) as an enhancement to the published article. It has been approved by peer review; however, it has undergone neither editing nor formatting by in-house editorial staff. The material is presented in the manner supplied by the author.
Supplementary Material

Appendix 1
Screening and Rater Review Process
A SAFER interview will be conducted off-site during the screening period and eligibility will be assessed based on the inclusion/exclusion criteria by the investigator and verified by an MGH-CTNI-certified clinician who will interview the participants using the HAMD17, SAFER/ATRQ, and eligibility criteria per the study protocol.
The SAFER interview assesses depression in a real-world setting and confirms that the participant’s illness is a specific state and excludes participants with any symptoms that are nonspecific or not readily assessable.

The ATRQ is administered at screening by a certified rater, the MGH-CTNI-certified clinician as part of the SAFER Interview. The ATRQ examines the efficacy and adequacy of any antidepressant treatment in a step-by-step procedure. This widely accepted questionnaire evaluates improvement (0% to 100%) and adequacy (adequate duration and dose) (Chandler 2010).

Appendix 2
Inclusion Criteria
To enroll in the clinical study, participants must meet the following inclusion criteria:

1. Must be able to read, speak, and understand English or Spanish and must provide written informed consent prior to the initiation of any protocol-specific procedures.
2. Male or female participant, aged 18 to 65 years, inclusive.
3. Body mass index (BMI) between 18.0 and 30.0 kg/m², at screening.
4. Participant is willing and able to commit to meet all study requirements, adhere to both approved ADT and study drug regimen, and complete all assessments and all scheduled visits, per investigator judgment.
5. Women of childbearing potential (WOCBP) and men whose sexual partners are WOCBP must use at least 1 highly effective method of contraception from screening and for at least 2 months after the last study drug administration. For men with female sexual partners of childbearing potential, examples of medically acceptable forms of contraception include vasectomy or male condom for participants, plus an additional method of contraception for their female partners. Highly effective methods of contraception are those which have a failure rate of <1% (when implemented consistently and correctly) and include:
   - Intratuterine device (IUD)
   - Bilateral tubal ligation, bilateral salpingectomy, or bilateral tubal occlusive procedure
   - Hormonal contraceptives (eg, oral, patch, or injectable)
   - A double-barrier protection method (eg, condom, sponge, or vaginal diaphragm with spermicide cream, foam, or gel)
   - Abstinence from heterosexual intercourse is accepted if this is the participant’s usual lifestyle and must be continued until at least 2 months after the last dose of
study drug.
Women who are not of childbearing potential must be congenitally or surgically sterile (hysterectomy and/or bilateral oophorectomy/salpingo-oophorectomy, as determined by the participant’s medical history) or must be post-menopausal. Post-menopausal is defined as being amenorrheic for at least 1 year without another cause and a follicle-stimulating hormone (FSH) level ≥40 mIU/mL as confirmation.

6. Diagnosed with MDD as defined by the Diagnostic and Statistical Manual, Fifth Edition (DSM-5), and confirmed by the SCID-5 MDD.

7. Hamilton Depression Rating Scale-17 (HAMD17) score ≥19 at screening and independently confirmed by SAFER assessment.

8. At baseline, before definitive admission and randomization of the participant, the MADRS10 scale will be administered and the participant must show a MADRS10 score of ≥24.

9. Diagnosed with a current MDE lasting from 8 weeks to 36 months as defined by the DSM-5 and confirmed by the SCID-5 MDD, as well as independent confirmation of HAMD17 score, SAFER/ATRQ, and contextual appropriateness to be a participant in this study, after evaluation by an MGH-CTNI clinician.

10. Treated for at least 6 weeks prior to screening and stabilized for at least 6 weeks prior to baseline on an approved dosing regimen of ADT (eg, SSRI, SNRI, or bupropion (a NDRI and nicotinic receptor antagonist) during the current MDE, and committed to remaining on the same stable dosing regimen for the screening period and for the entire study, at or above the minimally adequate dose in the ATRQ. Maximal doses and recommended doses for each ADT are at the discretion of the investigator and medical monitor, except for citalopram and escitalopram.

    Note: Discontinuation of any of the listed ADT must occur at least 6 weeks prior to baseline.

    Note: Participants taking trazodone and/or bupropion as secondary ADT are permitted.

    Note: A dosing eDiary will be used beginning at screening to document the stability of background antidepressant(s); only participants reporting a minimum of 80% adherence during screening will be randomized.

11. An appropriate and valid participant in the study, after independent MGH-CTNI SAFER/ATRQ assessment of the participant’s MDD condition to confirm the diagnosis of MDD, as well as the inadequate response to 1 to 3 valid courses of treatment with an antidepressant medication in the current MDE, defined as <50% improvement with an antidepressant medication at doses listed on the SAFER and ATRQ Interview Forms (Criteria: State versus trait; Assessability; Face validity; Ecological validity; and Rule of three Ps [pervasive, persistent, and pathological]).

Exclusion Criteria
Individuals meeting any of the following criteria at screening or baseline are ineligible to participate in this study:
1. History or presence of clinically significant abnormality as assessed by physical examination, medical history, 12-lead ECG, vital signs, or laboratory values, which in the opinion of the investigator would jeopardize the safety of the participant or the validity of the study results, including established QT prolongation, long QT syndrome, torsades de pointes, bradyarrhythmia, ventricular tachycardia, uncompensated heart failure (greater than NYHA Class 1 CHF), uncontrolled hypokalemia, or uncontrolled hypomagnesemia.

2. More than class 2 angina pectoris or a myocardial infarction (MI) or acute coronary syndrome within the past 3 months.

3. Any medical, psychiatric condition, or social context that, in the opinion of the investigator, is likely to unfavorably alter the risk-benefit of subject participation, to interfere with protocol compliance, or to confound safety or efficacy assessments.

4. Have any significant illness, of any nature, including possible SARS-COV-2 related fever and symptoms, requiring hospitalization, emergency treatment, or isolation (quarantine) within 4 weeks prior to screening or during the screening period, and as determined by the investigator.

5. History or first degree relative with history of unexplained sudden death or long QT syndrome.

6. Triplicate 12-lead ECG with average QTcF ≥450 msec, and/or a QRS interval ≥120 msec at screening.

7. Current or recent uncontrolled orthostasis or orthostatic hypotension necessitating treatment.

8. Poorly controlled diabetes as defined by a glycosylated hemoglobin (HbA1c) >7.5%, despite standard care.

9. Any use of long-term prescribed opioids (ie, >120 days in a 6-month period) within 6 months prior to screening or any recreational use of opioids.

10. More than 3 doses of opioids within 30 days prior to baseline.

11. Any use of benzodiazepines within 30 days prior to baseline and/or more than 3 doses of antipsychotics, when used for non-psychiatric indications, within 30 days prior to baseline.

12. Use of any anxiolytic, antipsychotic, anticonvulsant/antiepileptic, mood stabilizer, or stimulant medication(s) within 30 days prior to baseline. Note: Participant should be medically stable, the medication was appropriately tapered and participant has no withdrawal symptoms.

13. Use of St. John’s Wort, (Hypericum Perforatum) within 30 days prior to baseline.

14. Participated in a ketamine, esketamine, dextromethorphan or any other NMDAR-antagonist study, or who received esketamine at any time.

15. Received ketamine, memantine, and/or dextromethorphan treatment within 30 days prior to baseline.

16. History of allergy or hypersensitivity to methadone or related drugs.

17. Receiving new-onset psychotherapy (individual, group, marriage, or family therapy) within 2 months prior to screening, or planning to start psychotherapy at any time during participation in the study.

18. Any lifetime experience of electroconvulsive therapy (ECT) and/or vagus nerve stimulation (VNS) or any other type of physical brain stimulation.

19. Received repetitive transcranial magnetic stimulation (rTMS) less than 6 months prior to the screening visit.
20. Any current and primary psychiatric disorder (ie, a condition that is the primary focus of
distress and/or treatment other than MDD), as defined by the DSM-5 and confirmed by
psychiatric history and/or examination by the investigator. These disorders include, but
are not limited to, any psychotic disorder, post-traumatic stress disorder, borderline
personality disorder, antisocial personality disorder, obsessive-compulsive disorder,
intellectual disability, or pervasive developmental disorder.
21. Participants who, in the investigator’s judgment, are at significant risk for suicide. A
participant with a C-SSRS ideation score of 4 or 5 within the last 6 months or any suicide
attempt within the past year of either screening or baseline must be excluded.
22. Any lifetime history of bipolar I or II disorder, psychosis and/or mania as defined by the
DSM-5 and confirmed by psychiatric history and/or examination by the investigator.
23. Comorbid moderate to heavy alcohol or substance use disorder, as defined by DSM-5, at
screening or within the 12 months prior to screening. Heavy drinking is defined as an
average of 3 or more drinks per day, in the last month.
24. A positive result on the urine drug/alcohol screen within 30 days prior to baseline (Day
1). At investigator discretion, a retest is permitted.
25. HAMD17 score <19 at Baseline or an increase in absolute value of >40% or a decrease in
absolute value of >20% on the HAMD17 score between screening and baseline as
conducted by the certified site rater.
26. Evidence of clinically significant hepatic or renal impairment, including an estimated
glomerular filtration rate (eGFR) <60 mL/min/1.73 m² (CKD-EPI 2009 calculation),
alanine aminotransferase (ALT) or aspartate aminotransferase (AST) >2.0 × upper limit
of normal (ULN), bilirubin >1.5× ULN (participants with history of Gilbert’s syndrome
diagnosis may be included if approved by medical monitor), or clinically significant
abnormal endocrine laboratory values (including clinically significant abnormal thyroid
parameters, ie, thyroid stimulating hormone [TSH] < 0.9 x LLN or > 1.25 x ULN.
27. Diagnosed with and/or treated for any type of cancer (excluding basal cell carcinoma and
in situ melanoma) within 4 years prior to screening.
28. Any planned elective surgery requiring general anesthesia.
29. Participant has had gastric bypass surgery or has had any procedures or disorders that
interfere with gastrointestinal transit or absorption.
30. Participated in a clinical study with an investigational medication in the past 6 months, or
participated in more than 2 clinical studies with investigational medications in the past 2
years.
31. Females who are currently lactating.
Supplementary Figure 1. Study Design

28 DAYS OF DOSING (QD)

Screening confirms MDE with inadequate response to 1-3 ADT

1:1 RANDOMIZATION

ADT + 25 mg REL-1017 (Day 1 loading dose 75 mg)

ADT + matching placebo

Follow-up assessments & Reliance OLS

D 4  D 7  D 14  D 21  D28 End of treatment
**Supplementary Table 1. Time from Discontinuation of Prohibited Medications, Supplements, and Other Substances or Therapies**

<table>
<thead>
<tr>
<th>Prohibited Medications, Supplements, and Other Substances or Therapies</th>
<th>Minimum Time from Discontinuation to Screening&lt;sup&gt;a&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ketamine, esketamine, dextromethorphan, or any other NMDAR-antagonist administered as part of a clinical study</td>
<td>Lifetime</td>
</tr>
<tr>
<td>Esketamine</td>
<td>Lifetime</td>
</tr>
<tr>
<td>Electroconvulsive therapy (ECT) and/or vagus nerve stimulation (VNS) or any other type of physical brain stimulation</td>
<td>Lifetime</td>
</tr>
<tr>
<td>Repetitive transcranial magnetic stimulation (rTMS)</td>
<td>180 days</td>
</tr>
<tr>
<td>Long-term opiate use (i.e. &gt;120 days)</td>
<td>180 days</td>
</tr>
<tr>
<td>New-onset psychotherapy</td>
<td>60 days</td>
</tr>
<tr>
<td>Ketamine, memantine and/or dextromethorphan</td>
<td>30 days</td>
</tr>
</tbody>
</table>

<sup>a</sup> The medical monitor should be contacted for any questions regarding the potential for pharmacological interactions with concomitant medications used by participants during the study. These include off-label use of medications for depression.

<table>
<thead>
<tr>
<th>Prohibited Medications, Supplements, and Other Substances or Therapies</th>
<th>Minimum Time from Discontinuation to Baseline&lt;sup&gt;a&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anxiolytic drugs</td>
<td>30 days</td>
</tr>
<tr>
<td>Antipsychotic drugs</td>
<td>30 days</td>
</tr>
<tr>
<td>Anticonvulsants/Antiepileptic drugs</td>
<td>30 days</td>
</tr>
<tr>
<td>Mood stabilizers (including lithium and valproic acid)</td>
<td>30 days</td>
</tr>
<tr>
<td>Stimulants (including amphetamines)</td>
<td>30 days</td>
</tr>
<tr>
<td>More than 3 doses of opioids</td>
<td>30 days</td>
</tr>
<tr>
<td>Any doses of benzodiazepines</td>
<td>30 days</td>
</tr>
<tr>
<td>St. John’s Wort</td>
<td>30 days</td>
</tr>
</tbody>
</table>

<sup>a</sup> The medical monitor should be contacted for any questions regarding the potential for pharmacological interactions with concomitant medications used by participants during the study. These include off-label use of medications for depression.
Supplementary Table 2. MMRM Analysis for Intent-To-Treat, Per Protocol, and Post-Hoc Severe Depression Population for Mean Change from Baseline to Day 28.

<table>
<thead>
<tr>
<th></th>
<th>Placebo</th>
<th>Esmethadone</th>
<th>LS Mean Difference (esmethadone – placebo)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Intent to Treat</strong></td>
<td>N=114</td>
<td>N=113</td>
<td></td>
</tr>
<tr>
<td>Baseline, mean (SD)</td>
<td>35.3 (4.3)</td>
<td>34.7 (5.2)</td>
<td></td>
</tr>
<tr>
<td>LS Mean (SE)</td>
<td>-13.37 (1.09)</td>
<td>-15.10 (1.05)</td>
<td>-1.74 (1.52)</td>
</tr>
<tr>
<td>95% CI</td>
<td>-15.52, -11.22</td>
<td>-17.18, -13.02</td>
<td>-4.74, 1.26</td>
</tr>
<tr>
<td>p-value</td>
<td></td>
<td></td>
<td>0.255</td>
</tr>
<tr>
<td>Effect size</td>
<td></td>
<td></td>
<td>-0.16</td>
</tr>
<tr>
<td><strong>Per Protocol</strong></td>
<td>N=97</td>
<td>N=103</td>
<td></td>
</tr>
<tr>
<td>Baseline, mean (SD)</td>
<td>35.1 (4.4)</td>
<td>34.6 (5.3)</td>
<td></td>
</tr>
<tr>
<td>LS Mean (SE)</td>
<td>-12.69 (1.10)</td>
<td>-15.63(1.06)</td>
<td>-2.94 (1.53)</td>
</tr>
<tr>
<td>95% CI</td>
<td>-14.87, -10.51</td>
<td>-17.73, -13.54</td>
<td>-5.96, 0.08</td>
</tr>
<tr>
<td>p-value</td>
<td></td>
<td></td>
<td>0.057</td>
</tr>
<tr>
<td>Effect size</td>
<td></td>
<td></td>
<td>-0.28</td>
</tr>
<tr>
<td><strong>Severe Depression (MADRS 10 ≥35)</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Intent to Treat</strong></td>
<td>N=61</td>
<td>N=51</td>
<td></td>
</tr>
<tr>
<td>Baseline, mean (SD)</td>
<td>38.3 (2.9)</td>
<td>39.4 (3.3)</td>
<td></td>
</tr>
<tr>
<td>LS Mean (SE)</td>
<td>-11.83 (1.58)</td>
<td>-17.87 (1.70)</td>
<td>-6.04 (2.33)</td>
</tr>
<tr>
<td>95% CI</td>
<td>-14.97, 8.70</td>
<td>-21.24, 14.50</td>
<td>-10.65, -1.42</td>
</tr>
<tr>
<td>p-value</td>
<td></td>
<td></td>
<td>0.011</td>
</tr>
<tr>
<td>Effect size</td>
<td></td>
<td></td>
<td>-0.51</td>
</tr>
<tr>
<td><strong>Per Protocol</strong></td>
<td>N=53</td>
<td>N=45</td>
<td></td>
</tr>
<tr>
<td>Baseline, mean (SD)</td>
<td>38.2 (3.0)</td>
<td>39.4 (3.4)</td>
<td></td>
</tr>
<tr>
<td>LS Mean (SE)</td>
<td>-11.56 (1.58)</td>
<td>-18.81 (1.69)</td>
<td>-7.25 (2.32)</td>
</tr>
<tr>
<td>95% CI</td>
<td>-14.69, -8.43</td>
<td>-22.17, -15.45</td>
<td>-11.87, -2.64</td>
</tr>
<tr>
<td>p-value</td>
<td></td>
<td></td>
<td>0.002</td>
</tr>
<tr>
<td>Effect size</td>
<td></td>
<td></td>
<td>-0.64</td>
</tr>
</tbody>
</table>