Ketamine as a promising prototype for a new generation of rapid-acting antidepressants

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The discovery of ketamine’s rapid and robust antidepressant effects opened a window into a new generation of antidepressants. Multiple controlled trials and open-label studies have demonstrated these effects across a variety of patient populations known to often achieve little to no response from traditional antidepressants. Ketamine has been generally well tolerated across patient groups, with transient mild-to-moderate adverse effects during infusion. However, the optimal dosing and route of administration and the safety of chronic treatment are not fully known. This review summarizes the clinical effects of ketamine and its neurobiological underpinnings and mechanisms of action, which may provide insight into the neurobiology of depression, relevant biomarkers, and treatment targets. Moreover, we offer suggestions for future research that may continue to advance the field forward and ultimately improve the psychopharmacologic interventions available for those individuals struggling with depressive and trauma-related disorders.

Keywords: ketamine; depression; antidepressant; treatment mechanisms; neurobiology

Introduction

Depressive disorders are a leading cause of disability affecting millions worldwide, often causing chronic recurrent symptoms, increased morbidity, heightened risk of suicide, poor functional outcomes, and profound socioeconomic burden. However, the past five decades of research focusing on antidepressant development has unfortunately seen little success in creating fundamentally novel psychopharmacologic interventions. Over 20 antidepressant medications are currently available, all targeting the monoaminergic system. However, the efficacy of these medications is limited, with a substantial proportion of patients experiencing residual symptoms, persistent functional impairment, and failure to achieve sustained remission even when symptoms do improve. Furthermore, the full clinical benefit of these traditional antidepressants is only achieved following weeks to months of treatment. A large study by the National Institute of Mental Health found that, regardless of the primary or adjunctive antidepressant medication selected for treatment, less than one-third of depressed patients achieved remission within 12 weeks and 33% of patients did not achieve remission at all despite trials of four antidepressant medications over a 1-year period, leaving clinicians with few therapeutic options for treatment-refractory patients. Thus, there is a clear and urgent need for the development of novel, rapid-acting antidepressants with robust efficacy. Mounting evidence suggests that low doses of ketamine may possess both of these properties, acting rapidly and robustly to treat severely treatment-resistant depressed patients.

Ketamine, a glutamate N-methyl-D-aspartate receptor (NMDA-R) antagonist, is a U.S. Food and Drug Administration (FDA)-approved sedative medication originally developed for the induction and maintenance of anesthesia in adults. Ketamine...
has a short plasma half-life of 4 min and a 2.5-h plasma terminal half-life. It can be administered intravenously (most common), intramuscularly (93% bioavailability (BA)), intranasally (50% BA), intrarectally (25% BA), and orally (20% BA). In addition to its role as an anesthetic and as a pharmacological model of the core symptoms of schizophrenia, ketamine has received considerable attention in psychiatric research as a prototype for a new generation of antidepressants after the discovery of its profound and rapid effects on depressive symptoms. This review provides a discussion of clinical trials investigating ketamine’s rapid antidepressant effects, safety, and tolerability. In addition, we summarize some recent research examining the effect of ketamine on symptoms of posttraumatic stress disorder (PTSD) and neurocognitive functioning. We then briefly present mechanisms thought to underlie the rapid antidepressant effects of ketamine, including the neurobiology and potential clinical biomarkers of depression, neurocircuitry critical to affective regulation, and molecular pathways in synaptogenesis. We conclude by discussing the prospect for next-generation rapid-acting antidepressants and the clinical implications of these findings on ketamine.

Robust and rapid antidepressant effects

In the early 1990s, researchers conducting preclinical studies found that NMDA-R antagonists showed promising antidepressant qualities. Later in the 1990s, in a pilot study in patients with severely treatment-resistant depression, we discovered that a single subanesthetic dose of ketamine had striking, robust, rapid antidepressant effects within 4 h of intravenous administration. This finding has since been well replicated in multiple controlled trials, underscoring the rationale of targeting the glutamatergic system and the feasibility of developing truly novel rapid-acting antidepressant agents. These noticeable rapid antidepressant effects have also been reported in patient groups known to respond poorly to traditional antidepressant interventions, such as those with bipolar depression, anxious depression, and patients who are resistant to treatment with electroconvulsive therapy (ECT). In addition, ketamine showed efficacy in rapidly reducing suicidal ideation within hours of administration. The antidepressant effects of ketamine are evident within 2–4 h of treatment and are sustained for 3–7 days, with a response rate of approximately 25–85% within 24 h and 14–70% at 72 h postinfusion (see Tables 1 and 2 for selected controlled trials and open-label investigations). Early clinical trials primarily administered 0.5 mg/kg of ketamine infused intravenously over approximately 40 min, in a medication-free population (Fig. 1). More recently, several studies reported rapid antidepressant effects following various routes and regimens (Tables 1 and 2). Of note, repeated intravenous doses prolonged treatment response and intranasal administration exerted rapid antidepressant effects comparable to intravenous administration but with minimal psychotomimetic and dissociative effects.

To date, three meta-analyses have been published that consistently confirmed the efficacy of ketamine’s antidepressant effects and drew attention to the current limitations and areas for future investigations. Each study supports the association of ketamine with a superior clinical response and clinical remission relative to comparators (e.g., saline, midazolam) in both unipolar and bipolar depression, in those with and without treatment-resistant depression, and in both medicated and unmedicated study participants. These meta-analyses also highlight that additional research is needed to evaluate optimal dosing, route of administration, and treatment schedules; to further characterize the duration of effects and the long-term safety, tolerability, and efficacy of ketamine; and to explore the potential effects of other glutamatergic agents that may have fewer or less extreme side effects and lower addiction potential. Other limitations of ketamine trials are the short duration of trials, the short-term efficacy of a single infusion of ketamine, and the lack of complete blinding to treatment, because of the functional unblinding of treatment status by the acute adverse effects of ketamine. This latter limitation was partially addressed in a recent two-site controlled trial that demonstrated the rapid antidepressant effects of ketamine compared to midazolam (an anesthetic benzodiazepine) as an active comparator to optimize blinding to treatment status. In addition, the true biological effect of the drug is further supported by the relatively consistent time curve of response to ketamine across studies (i.e., improvement within 4 h, peak
### Table 1. Selected randomized controlled studies examining the rapid antidepressant effect of ketamine

<table>
<thead>
<tr>
<th>Study</th>
<th>Subjects</th>
<th>Treatment</th>
<th>ADE (4 h to 3 days)</th>
<th>ADE (7 days)</th>
<th>Response rate</th>
<th>Additional findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Abdallah et al.</td>
<td>N6-MDD,</td>
<td>Thiopental</td>
<td>NO</td>
<td>—</td>
<td>13% at 1–3 days</td>
<td>ADE in both groups after 6th ECT; ECT and/or thiopental appears to attenuate the rapid ADE of ketamine</td>
</tr>
<tr>
<td></td>
<td>N2-BD, MF</td>
<td>3.5 mg/kg + ECT 3×/week for 2 weeks</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Ketamine</td>
<td>NO</td>
<td>—</td>
<td>0% at 1–3 days</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.5 mg/kg + thiopental 3.5 mg/kg + ECT 3×/week for 2 weeks</td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>Berman et al.</td>
<td>N8-MDD,</td>
<td>Placebo</td>
<td>NO</td>
<td>—</td>
<td>12% at 3 days</td>
<td></td>
</tr>
<tr>
<td></td>
<td>N1-BD, MF</td>
<td></td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Ketamine</td>
<td>YES</td>
<td>—</td>
<td>50% at 3 days</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.5 mg/kg</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Diazgrandos et al.</td>
<td>N18-BD</td>
<td>Placebo</td>
<td>NO</td>
<td>NO</td>
<td>6% at 1 day; 71% at 1 day</td>
<td>Significant ADE at 40 min</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Ketamine</td>
<td>YES</td>
<td>NO</td>
<td>71% at 1 day</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.5 mg/kg</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ghasemi et al.</td>
<td>N9-MDD</td>
<td>Ketamine</td>
<td>YES</td>
<td>NO</td>
<td>42% at 1 day; 7% at 7 days</td>
<td>Response rates are following the last infusion</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.5 mg/kg</td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td></td>
<td></td>
<td>1×/2 days for 6 days</td>
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</tr>
<tr>
<td></td>
<td>N9-MDD</td>
<td>ECT 1×/2 days for 6 days</td>
<td>NO</td>
<td>YES</td>
<td>12% at 1 day; 25% at 7 days</td>
<td></td>
</tr>
<tr>
<td>Kudoh et al.</td>
<td>N35-MDD</td>
<td>Propofol</td>
<td>NO</td>
<td>—</td>
<td>—</td>
<td>Lower pain scores among ketamine group</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1.5 mg/kg</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>+ fentanyl 2 μg/kg</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Ketamine</td>
<td>YES</td>
<td>—</td>
<td>—</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>1.0 mg/kg + propofol 1.5 mg/kg + fentanyl 2 μg/kg</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lapidus et al.</td>
<td>N18-MDD</td>
<td>Placebo</td>
<td>NO</td>
<td>—</td>
<td>6% at 1 day</td>
<td>Significant improvement of anxiety symptoms @ 1 day; dose equivalent to 0.15–0.34 mg/kg intravenous ketamine</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Ketamine</td>
<td>YES</td>
<td>—</td>
<td>44% at 1 day</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>50 mg</td>
<td></td>
<td></td>
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</tr>
</tbody>
</table>

*Continued*
response around 24 h, and efficacy for about 1 week, as well as the maintenance of efficacy through repeated treatment (Tables 1 and 2)).

Posttraumatic stress disorder
Preliminary evidence supports the utility and safety of ketamine in treating PTSD symptoms, which are often highly comorbid and sometimes overlapping with depressive disorders. In an early case report, a young military veteran with highly treatment-resistant PTSD showed rapid improvement following a single infusion of a subanesthetic dose of ketamine treatment, with a 56% reduction in his PTSD symptoms, and this treatment response was maintained for 15 days. Ketamine was well tolerated and no adverse effects were reported.

Another case series reported a marked reduction in flashbacks in three women with PTSD treated with ifenprodil—an NMDA-R antagonist that selectively binds to the GluN2B subunit. The safety of subanesthetic doses of ketamine was reviewed in patients with PTSD or trauma history who were treated with ketamine for comorbid depression. Ketamine was well tolerated, with no evidence of worsening in PTSD symptoms.

More recently, Feder and colleagues provided promising pilot evidence supporting the utility and safety of ketamine in treating PTSD symptoms. This study randomized a cohort of PTSD patients to a single infusion of ketamine (0.5 mg/kg infused over 40 min) or midazolam in a double-blind crossover design. Compared to midazolam, ketamine showed a significant improvement in PTSD symptoms 24 h postinfusion. Similar to depression studies, dissociative symptoms occurred during infusion, peaked at 40 min, and were generally

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**Table 1. Continued**

<table>
<thead>
<tr>
<th>Study</th>
<th>Subjects</th>
<th>Treatment</th>
<th>ADE (4 h to 3 days)</th>
<th>ADE (7 days)</th>
<th>Response rate</th>
<th>Additional findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Murrough et al.</td>
<td>N47-MDD</td>
<td>Ketamine(^b) 0.5 mg/kg</td>
<td>YES</td>
<td>YES</td>
<td>64% at 1 day; 45% at 7 days</td>
<td>—</td>
</tr>
<tr>
<td></td>
<td>N25-MDD</td>
<td>Midazolam 0.045 mg/kg</td>
<td>YES</td>
<td>YES</td>
<td>28% at 1 day; 16% at 7 days</td>
<td>—</td>
</tr>
<tr>
<td>Valentine et al.</td>
<td>N10-MDD, MF</td>
<td>Placebo Ketamine(^b) 0.5 mg/kg</td>
<td>NO</td>
<td>NO</td>
<td>—</td>
<td>(^{1})H-MRS: Occipital amino acid levels did not correlate with ADE</td>
</tr>
<tr>
<td>Zarate et al.</td>
<td>N15-MDD, MF</td>
<td>Placebo Ketamine(^b) 0.5 mg/kg</td>
<td>NO</td>
<td>NO</td>
<td>0% at 1 day; 0% at 7 days</td>
<td>Depressed mood, guilt, work, interests, and psychic anxiety improved significantly among treatment group</td>
</tr>
<tr>
<td>Zarate et al.</td>
<td>N15-BD</td>
<td>Placebo Ketamine(^b) 0.5 mg/kg</td>
<td>NO</td>
<td>NO</td>
<td>64% at 40 min; 50% at 4 h; 43% at 1 day</td>
<td>Significant ADE at 40 min; decreased suicidal ideations</td>
</tr>
</tbody>
</table>

\(^a\)Intravenous ketamine bolus.  
\(^b\)Intravenous ketamine infusion over 40 min.  
\(^c\)Intranasal ketamine.  
—, not reported.

Abbreviations: ADE, antidepressant effect; N, number of subjects; MDD, major depressive disorder; BD, bipolar disorder; MF, medication free; \(^{1}\)H-MRS, proton magnetic resonance spectroscopy; ECT, electroconvulsive therapy.
### Table 2. Selected open-label studies examining the rapid antidepressant effects of ketamine

<table>
<thead>
<tr>
<th>Study</th>
<th>Subjects</th>
<th>Treatment</th>
<th>ADE (4 h to 3 days)</th>
<th>ADE (7 days)</th>
<th>Response rate</th>
<th>Additional findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Carlson et al.</td>
<td>N20-MDD</td>
<td>Ketamine&lt;sup&gt;a&lt;/sup&gt; 0.5 mg/kg</td>
<td>YES</td>
<td>—</td>
<td>30% at 4 h</td>
<td>Reduced activity in the right habenula</td>
</tr>
<tr>
<td>Chilukuri et al.</td>
<td>N9-MDD</td>
<td>Ketamine&lt;sup&gt;a&lt;/sup&gt; 0.5 mg/kg</td>
<td>YES</td>
<td>—</td>
<td>59% at 2 h</td>
<td>Improvement sustained at 3 days in all groups; two subjects attempted suicide—both were nonresponders to ketamine&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td>N9-MDD</td>
<td>Ketamine&lt;sup&gt;b&lt;/sup&gt; 0.5 mg/kg</td>
<td>YES</td>
<td>—</td>
<td>60% at 2 h</td>
<td></td>
</tr>
<tr>
<td></td>
<td>N9-MDD</td>
<td>Ketamine&lt;sup&gt;b&lt;/sup&gt; 0.25 mg/kg</td>
<td>YES</td>
<td>—</td>
<td>57% at 2 h</td>
<td></td>
</tr>
<tr>
<td>Diazgrandos, et al.</td>
<td>N33-MDD, MF</td>
<td>Ketamine&lt;sup&gt;a&lt;/sup&gt; 0.5 mg/kg</td>
<td>YES</td>
<td>—</td>
<td></td>
<td>Decreased suicidal ideations</td>
</tr>
<tr>
<td>Ibrahim et al.</td>
<td>N17-MDD-ER, MF</td>
<td>Ketamine&lt;sup&gt;a&lt;/sup&gt; 0.5 mg/kg</td>
<td>YES</td>
<td>—</td>
<td></td>
<td>Lower effect size with ECT-resistant (MDD-ER) group</td>
</tr>
<tr>
<td></td>
<td>N23-MDD, MF</td>
<td>Ketamine&lt;sup&gt;a&lt;/sup&gt; 0.5 mg/kg</td>
<td>YES</td>
<td>—</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ibrahim et al.</td>
<td>N42-MDD, MF</td>
<td>Ketamine&lt;sup&gt;a&lt;/sup&gt; 0.5 mg/kg</td>
<td>YES</td>
<td>YES</td>
<td>62% at 4–6 h</td>
<td>Riluzole following ketamine had no effects; ADE maintained at 28 days</td>
</tr>
<tr>
<td>Larkin and Beautrais</td>
<td>N14-MDD-SI</td>
<td>Ketamine&lt;sup&gt;c&lt;/sup&gt; 0.2 mg/kg</td>
<td>YES</td>
<td>YES</td>
<td>85% at 10 days</td>
<td>Decreased suicidal ideations</td>
</tr>
<tr>
<td>Machado et al.</td>
<td>N23-MDD, MF</td>
<td>Ketamine&lt;sup&gt;a&lt;/sup&gt; 0.5 mg/kg</td>
<td>YES</td>
<td>—</td>
<td>48% at 4 h</td>
<td>Serum BDNF did not correlate with ADE.</td>
</tr>
<tr>
<td>Mathew et al.</td>
<td>N26-MDD, MF</td>
<td>Ketamine&lt;sup&gt;a&lt;/sup&gt; 0.5 mg/kg</td>
<td>YES</td>
<td>YES</td>
<td>65% at 1 day; 54% at 3 days</td>
<td>Lamotrigine prior to ketamine and riluzole following ketamine had no effects</td>
</tr>
<tr>
<td>Murrough et al.</td>
<td>N24-MDD, MF</td>
<td>Ketamine&lt;sup&gt;a&lt;/sup&gt; 0.5 mg/kg</td>
<td>YES</td>
<td>—</td>
<td>71% at 24 h after last infusion</td>
<td>Significant ADE within 2 h; decreased suicidal ideations in ketamine responders as well as nonresponders; median time to relapse was 18 days</td>
</tr>
<tr>
<td>Okamoto et al.</td>
<td>N11-MDD</td>
<td>Ketamine&lt;sup&gt;c&lt;/sup&gt; 0.86 mg/kg + ECT 2×/w for 4w</td>
<td>—</td>
<td>YES</td>
<td></td>
<td>ADE in both groups; ketamine group had lower depression scores at 2nd and 4th ECT sessions and longer seizure at 1st and 6th ECT session.</td>
</tr>
<tr>
<td></td>
<td>N20-MDD</td>
<td>Propofol 0.94 mg/kg + ECT 2×/w for 4w</td>
<td>—</td>
<td>YES</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

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Continued
Table 2. Continued

<table>
<thead>
<tr>
<th>Study</th>
<th>Subjects</th>
<th>Treatment</th>
<th>ADE (4 h to 3 days)</th>
<th>ADE (7 days)</th>
<th>Response rate</th>
<th>Additional findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Phelps et al. 88</td>
<td>N26-MDD, MF</td>
<td>Ketamine&lt;sup&gt;a&lt;/sup&gt; 0.5 mg/kg</td>
<td>YES</td>
<td>—</td>
<td>43% at 4 h</td>
<td>Plasma ketamine/norketamine did not correlate with ADE</td>
</tr>
<tr>
<td>Rasmussen et al. 51</td>
<td>N10-MDD</td>
<td>Ketamine&lt;sup&gt;a&lt;/sup&gt; 0.5 mg/kg</td>
<td>YES</td>
<td>—</td>
<td>30% at 1 day; 80% at 4 days</td>
<td>Decreased suicidal ideations</td>
</tr>
<tr>
<td>Salvador et al. 68</td>
<td>N11-MDD, N11-HS, MF</td>
<td>Ketamine&lt;sup&gt;a&lt;/sup&gt; 0.5 mg/kg</td>
<td>YES</td>
<td>—</td>
<td>—</td>
<td>Repeated exposure to fearful faces: increased rACC activity in MDD but not HS; rACC activity correlated with ADE; rAG negatively correlated ADE</td>
</tr>
<tr>
<td>Salvador et al. 69</td>
<td>N15-MDD, MF</td>
<td>Ketamine&lt;sup&gt;a&lt;/sup&gt; 0.5 mg/kg</td>
<td>YES</td>
<td>—</td>
<td>—</td>
<td>N-back test (working memory): pgACC activity negatively correlated with ADE</td>
</tr>
<tr>
<td>Salvador et al. 89</td>
<td>N15-MDD, MF</td>
<td>Ketamine&lt;sup&gt;a&lt;/sup&gt; 0.5 mg/kg</td>
<td>YES</td>
<td>—</td>
<td>—</td>
<td>Frontal glx/glutamate negatively correlated with ADE, whereas glutamate positively correlated with anxiety improvement</td>
</tr>
</tbody>
</table>

<sup>a</sup>Intravenous ketamine infusion over 40 min.
<sup>b</sup>Intramuscular ketamine.
<sup>c</sup>Intravenous ketamine bolus.

—, not reported.

Abbreviations: ADE: antidepressant effect; N, number of subjects; MDD, major depressive disorder; MDD-SI, major depressive disorder with suicidal ideation; MDD-ER, ECT-resistant major depressive disorder; BD, bipolar disorder; MF, medication free; HS, healthy subjects; FH<sup>+</sup>/-, family history of alcohol abuse positive/negative; <sup>1</sup>H-MRS, proton magnetic resonance spectroscopy; MEG, magnetoencephalography; rACC, rostral anterior cingulate cortex; pgACC, pregenual anterior cingulate cortex; rAG, rostral amygdala; Glx, a <sup>1</sup>H-MRS measure reflecting glutamate and glutamine; ECT, electroconvulsive therapy.

Well tolerated, and subsided within 80 min following ketamine administration. 30 Also, analogous to ketamine studies in depressed and healthy subjects, physical adverse effects were transient. 30

Neurocognitive functioning
Cognitive deficits cardinal to depressive disorders contribute significantly to disability, risk for suicide, treatment noncompliance, and refractory treatment response. 31–37 As further described later, ketamine’s antagonism of the glutamatergic NMDA-R appears to be the first step in a cascade of events that converge to produce enhanced activity in excitatory networks and marked changes in synaptic plasticity and strength. 38–44 This enhanced synaptic plasticity occurs approximately 24 h after ketamine administration, the peak time of antidepressant response. Considering the critical role of synaptic plasticity in cognitive function, the question arises as to whether the demonstrated ketamine-induced synaptogenesis would translate into enhanced cognition 24 h posttreatment.

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Although ketamine studies have consistently demonstrated transient cognitive deficits during infusion, the hypothesized procognitive effects of ketamine 24 h posttreatment and the relationship between cognitive functions and treatment response have not been fully studied. In a controlled trial in which participants were randomized to ketamine or midazolam, there were no differential effects of treatment on cognitive performance and no correlation with antidepressant response. However, low baseline cognitive-processing speed uniquely predicted depression improvement at 24 h post-ketamine.45 Another recent study of ketamine’s effect on the neurocognitive performance of patients with bipolar depression revealed procognitive effects approximately 72 h postinfusion.46 These results appeared to be independent of depression severity or improvement, suggesting that ketamine may possess procognitive effects in addition to its influence on mood symptoms.46 These two investigations provide promising preliminary evidence. However, both studies have a relatively small sample size and did not include any long-term evaluation of cognitive changes. Future studies with larger sample sizes and longer-term follow-up are required to further characterize the cognitive effects of ketamine.

Safety and tolerability
Ketamine is generally well tolerated across both patient groups and healthy controls, with mild to moderate transient adverse side effects observed within the first 2 h of treatment. Ketamine administration produces transient perceptual disturbances, dissociation, dysphoria, euphoria, and/or anxiety during infusion. Physical adverse effects include dizziness, nausea, and mild increases in blood pressure and heart rate. There is also potential risk of ulcerative cystitis,47 although this is typically associated with chronic use at higher doses than what is generally administered in psychiatric clinical investigations. Given the short half-life of the drug, these adverse effects abate within a few minutes of stopping ketamine infusion and generally fully remit within 2 hours.

Open-label and placebo-controlled trials have supported the safety and tolerability of a single infusion of ketamine.21 However, the optimal dose, route of administration, and the safety of repeated or chronic treatment are still not fully known. This is particularly important given the addiction and excitotoxicity potential of ketamine. Case reports suggest that repeated ketamine infusions may safely and effectively extend the antidepressant effects of the drug for several months.48,49 Up to six infusions of repeated ketamine administered once, twice, or three times per week were found to be safe and efficacious in maintaining treatment response.22,50–53 Other studies reported safety and efficacy using a single administration of various routes and doses of ketamine, including 0.2 mg/kg intravenous bolus,54 50 mg intranasal,23 and 0.5 mg/kg or 0.25 mg/kg intramuscular injection.55

Mechanism of action
Convergent evidence implicates synaptic plasticity in the pathophysiology of depression.56,57 Sustained stress and depression precipitates neuronal atrophy and overall synaptic depression, particularly in the prefrontal cortex (PFC).58–60 These synaptic deficits are the result of reduced neurotrophic function (e.g., brain-derived neurotrophic factor (BDNF))61 and of the inhibition of the mammalian target of rapamycin (mTOR) pathway.62 Reduction of BDNF or inhibition of mTOR signaling leads to depressive-like behavior and blocks the effects of antidepressants in rodents.61,62 Conversely, increasing BDNF or enhancing mTOR signaling

![Figure 1](https://example.com/f1.png)

**Figure 1.** Ketamine administration and time course. (A) Comprehensive screening and evaluation; (B) ketamine 0.5 mg/kg infused over 40 min; (C) immediate antidepressant effect; and (D) rapid antidepressant effects sustained for up to 28 days after a single ketamine infusion.
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Figure 2. Mechanisms underlying the rapid antidepressant effects of ketamine. Changes in GluR1 appear to be region specific, with evidence for increased GluR1 in the medial prefrontal cortex but not the hippocampus. Abbreviations: ADE, antidepressant-like effects; NMDA-R, N-methyl-D-aspartate glutamate receptor; AMPA-R, \(\alpha\)-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid receptor; GluR1, R1 subunit of AMPA-R; eEF2k, eukaryotic elongation factor 2 kinase; BDNF, brain-derived neurotrophic factor; pmTOR, mammalian target of rapamycin; GSK3, glycogen synthase kinase 3. \(\oplus\), enhance/stimulate; \(\oslash\), inhibit/reduce; \(\Theta\), blockade.

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produces antidepressant-like effects.\(^{61,62}\) Therefore, it is proposed that activation of BDNF and mTOR signaling is a required step for efficacious antidepressant treatment. Ketamine is believed to exert its rapid antidepressant effect by increasing BDNF function and activating mTOR signaling.

High doses of ketamine inhibit overall NMDA-R activity, leading to sedation and to reduced glutamate signaling, as evident by reduced extracellular glutamate.\(^{39}\) In contrast, low doses of ketamine have a paradoxical effect on glutamatergic activity, leading to an increase of glutamate release.\(^{39,63}\) This paradoxical effect is hypothesized to be the consequence of a preferential blockade of the NMDA-R on a subpopulation of gamma-aminobutyric acid (GABA)ergic interneurons.\(^{64,65}\) Inhibition of these interneurons disinhibits prefrontal glutamatergic cells, leading to a glutamate surge, which appears to play a critical role in the antidepressant effects of ketamine.\(^{13}\)

Briefly, mounting evidence suggests that ketamine exerts its antidepressant effect through two pathways (Fig. 2).\(^{38,66}\) The “stop pathway” involves at-rest inhibition of the NMDA-R,\(^{66}\) presumably extrasynaptic NMDA-R, which is known to promote synaptic atrophy and neuronal death.\(^{57}\) At-rest inhibition of the NMDA-R culminates in increasing BDNF function by disinhibiting eEF2.\(^{66}\) The “go pathway” engages a cascade of events that includes (1) disinhibition of glutamatergic neurons, leading to a glutamate surge;\(^{39,63}\) (2) stimulation of synaptic glutamate receptors, particularly \(\alpha\)-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid (AMPA) receptors;\(^{38}\) and (3) activation of mTOR signaling and synaptic formation.\(^{38}\) The increase in BDNF function and mTOR signaling
converges to promote synaptogenesis and reversal of the synaptic deficits induced by prolonged stress and depression.\textsuperscript{13}

Clinical biomarkers

The rapid and robust antidepressant effects of ketamine offer a unique opportunity for clinical biomarker development to better understand the mechanism underlying rapid-acting antidepressants and to gain insight into the neurobiology of depression.\textsuperscript{13} To date, studies on biomarkers can largely be grouped into two categories: studies examining prefrontal excitability and synaptic strength or studies that investigated the role of neurotrophins, primarily BDNF.

An enhanced response to ketamine was associated with (1) pretreatment high medial PFC (mPFC) activity in response to fearful faces,\textsuperscript{68} (2) pretreatment low mPFC activity during a working memory task,\textsuperscript{69} (3) pretreatment low connectivity between the mPFC and amygdala,\textsuperscript{69} (4) posttreatment increases in synaptic strength,\textsuperscript{70,71} and (5) low pretreatment mPFC (glutamate + glutamine)/glutamate ratio,\textsuperscript{72} although not without inconsistencies.\textsuperscript{73}

In addition, although early studies failed to demonstrate a significant correlation between peripheral BDNF and response to ketamine,\textsuperscript{74,75} more recent studies confirmed a relationship between increased BDNF and enhanced treatment following ketamine administration.\textsuperscript{71,76} These findings were supported by a study showing a positive relationship between the functional variant of BDNF (Val66Met) and response to ketamine treatment.\textsuperscript{77}

Conclusions

The discovery of the rapid antidepressant effects of ketamine opened a window into a new generation of antidepressants and a better understanding of the biological underpinnings of depression. Ketamine studies showed us that rapid improvement in complex mood states such as depression, PTSD, and suicidality are possible with therapeutic interventions. It also highlighted the utility of targeting the glutamatergic system, a truly novel antidepressant mechanism. However, it is important to emphasize that ketamine remains primarily an investigational drug at this stage, with abuse liability and unknown safety and efficacy following chronic treatment.

Excitement in the field has led to concerted preclinical and clinical effort over the past decade by many research groups across the nation and throughout the world. Significant discoveries and major pathways have been implicated in the pathophysiology and treatment of depression. Novel NMDA-R modulators are being tested to mimic the rapid antidepressant effects of ketamine while minimizing adverse effects (for a review, see Ref. 78). However, several questions remain unanswered and require further investigation: (1) the optimal dose and preferable route of administration of ketamine; (2) the frequency and dose at which ketamine administration becomes harmful instead of beneficial; (3) the regimen to maintain treatment response; (4) the relationship between cognitive function and ketamine treatment; (5) the sustainability of antisuicidal properties and generalizability to the general population; and (6) the role of combined ketamine treatment and cognitive behavioral therapy.

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Easton Associates, Gilead Sciences, GlaxoSmithKline, Janssen Pharmaceuticals, Lundbeck Research USA, Medivation, Merz Pharmaceuticals, MK Medical Communications, Hoffmann–La Roche, SK Holdings, Sunovion Pharmaceuticals, Takeda Industries, and Teva Pharmaceutical Industries. He is on the Scientific Advisory Board for the following companies: Abbott Laboratories, Bristol-Myers Squibb, Eisai, Eli Lilly, Forest Laboratories, Lohoca Research Corporation, Mnemosyne Pharmaceuticals, Naurex, Pfizer Pharmaceuticals, and Shire Pharmaceuticals. He holds less than $150 in exercisable warrant options with Tetragenex Pharmaceuticals. He is also on the Board of Directors of the Coalition for Translational Research in Alcohol and Substance Use Disorders and was the principal investigator of a multicenter study in which Janssen Research Foundation provided drug and some support to the Department of Veterans Affairs. He is Editor of Biological Psychiatry. In addition, he has a patent on dopamine and noradrenergic reuptake inhibitors in the treatment of schizophrenia (patent number 5447948) and is a coinventor with Dr. Gerard Sanacora on a filed patent application by Yale University related to targeting the glutamatergic system for the treatment of neuropsychiatric disorders (PCTWO06108055A1). He has a patent pending on intranasal administration of ketamine to treat depression.

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