Ketamine for the acute treatment of severe suicidal ideation: double blind, randomised placebo controlled trial

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ABSTRACT

OBJECTIVE

To confirm the rapid onset anti-suicidal benefits of ketamine in the short term and at six weeks, overall and according to diagnostic group.

DESIGN

Prospective, double blind, superiority, randomised placebo controlled trial.

SETTING

Seven French teaching hospitals between 13 April 2015 and 12 March 2019.

ELIGIBILITY CRITERIA FOR PARTICIPANTS

Aged 18 or older with current suicidal ideation, admitted to hospital voluntarily. Exclusion criteria included a history of schizophrenia or other psychotic disorders, substance dependence, and contraindications for ketamine.

PARTICIPANTS

156 participants were recruited and randomised to placebo (n=83) or ketamine (n=73), stratified by centre and diagnosis: bipolar, depressive, or other disorders.

INTERVENTION

Two 40 minute intravenous infusions of ketamine (0.5 mg/kg) or placebo (saline) were administered at baseline and 24 hours, in addition to usual treatment.

MAIN OUTCOME MEASURES

The primary outcome was the rate of patients in full suicidal remission at day 3, according to the scale for suicidal ideation total score ≤3. Analyses were conducted on an intention-to-treat basis.

RESULTS

More participants receiving ketamine reached full remission of suicidal ideas at day 3 than those receiving placebo: 46 (63.0%) of 83 participants in the ketamine arm and 25 (31.6%) of 73 in the placebo arm (odds ratio 3.7 (95% confidence interval 1.9 to 7.3), P<0.001). This effect differed according to the diagnosis (treatment: P<0.001; interaction: P=0.02): bipolar (odds ratio 14.1 (95% confidence interval 3.0 to 92.2), P<0.001), depressive (1.3 (0.3 to 5.2), P=0.6), or other disorders (3.7 (0.9 to 17.3, P=0.07)). Side effects were limited. No manic or psychotic symptom was seen. Moreover, a mediating effect of mental pain was found. At week 6, remission in the ketamine arm remained high, although non-significantly versus placebo (69.5% v 56.3%; odds ratio 0.8 (95% confidence interval 0.3 to 2.5), P=0.7).

CONCLUSIONS

The findings indicate that ketamine is rapid, safe in the short term, and has persistent benefits for acute care in suicidal patients. Comorbid mental disorders appear to be important moderators. An analgesic effect on mental pain might explain the anti-suicidal effects of ketamine.

TRIAL REGISTRATION

ClinicalTrials.gov NCT02299440.

Introduction

Around 700 000 people worldwide die from suicide annually, and 10 to 20 times this number attempt suicide.1 Suicide is the second most important cause of death in adolescents and young adults.2 The 12 month prevalence of suicidal ideas is 2% in the adult population globally, including 0.5% with a suicidal plan.3 Although most suicidal ideas will not lead to a suicidal act, all suicidal acts are preceded by suicidal ideas. Thus rapid resolution of a suicidal crisis before it is acted on might prevent many deaths. Moreover, reducing the intensity of the suicidal pain could facilitate psychosocial intervention.

Only limited evidence based options are available to treat suicidal crises. Antidepressants might reduce the risk of suicide, particularly in individuals aged over 25, but onset of beneficial effects is delayed by several weeks and a trial of several drugs is often necessary.4 Moreover, antidepressants are not recommended for people with bipolar disorders. Similarly, clozapine and lithium might be effective anti-suicidal drugs in schizophrenia and mood disorders, respectively, but not in the short term.5 Psychotherapy takes several sessions to be efficient,6 and the evidence for electroconvulsive therapy remains weak.7 Admission to hospital, anxiolytics, and hypnotics
are commonly used despite limited scientific evidence. Moreover, suicide occurs at high rates in psychiatric units during admission and after discharge, questioning the efficacy of this procedure.

Recently, ketamine has been shown to have a rapid effect on depressive symptoms and suicidal ideation after a single dose.\textsuperscript{5-11} A meta-analysis\textsuperscript{12} showed a beneficial effect on suicidal ideation scores within 4 hours after infusion, lasting for at least the first 72 hours. Of note, a recent review of the literature suggested that intravenous ketamine has a better effect than intranasal esketamine.\textsuperscript{13} Previous studies, however, were subject to several methodological limitations. Firstly, the suicidal risk was often poorly measured (eg, with one single item of a depression scale). Secondly, response (usually defined as a 50% score reduction on a scale) was the most common outcome rather than remission (that is, complete absence of suicidal ideas). Thirdly, samples were usually small. Fourthly, most studies were conducted in unipolar disorder with limited knowledge about the effect in bipolar disorder, despite the high suicide risk,\textsuperscript{14} and in non-mood disorders. Finally, the psychophysiological mechanisms of action remain poorly understood. Notably, it is now established that mental pain contributes to an increased risk of suicidal ideas and acts,\textsuperscript{15} suggesting that suicidal acts aim to put an end to unbearable mental pain. Whether the antalgic effects of ketamine contribute to its anti-suicidal effects remains to be tested.

We aimed in this study to examine full remission of suicidal ideas 72 hours after two infusions of ketamine versus placebo in a large sample. Three groups of patients were a priori selected: those with a bipolar disorder, or a depressive disorder, or another main diagnosis. Furthermore, we tested whether ketamine acted on suicidal ideas through an analgesic effect on mental pain. Finally, we examined the persistence of the effect of ketamine at six weeks. We hypothesised that (a) ketamine will be better than placebo for inducing full suicidal remission at day 3; (b) this effect will vary according to the diagnostic group; (c) this effect will be mediated through alleviating mental pain; (d) this effect will persist over six weeks.

Methods

Study design

This six week, double blind, randomised placebo controlled study (named KETIS) was conducted in seven academic hospitals in metropolitan France. Ethical approval was obtained on 18 July 2014 from the research ethics board “Comité de Protection des Personnes (CPP) Sud Méditerranée III” (ref: 2014.06.03 bis). This study was prospectively registered on 20 November 2014 on https://www.clinicaltrials.gov/ (NCT02299440), and is listed on EudraCT: 2014-001324-30. All patients gave their informed, written, and signed consent before inclusion.

Patients

Participants were recruited during admission to hospital in psychiatry for suicidal ideation. Inclusion criteria were patients aged 18 or older, with a clinician rated scale for suicidal ideation (SSI\textsuperscript{16}) total score >3; voluntarily admitted to hospital; French speaking; able to provide informed consent; insured or beneficiary of a health insurance plan. Exclusion criteria were a history of schizophrenia or other psychotic disorders based on Diagnostic and Statistical Manual of Mental Disorders, fourth edition (DSM-IV) criteria; schizoid or schizotypic personality disorders; presence of psychotic symptoms at initial interview; substance dependence during the preceding month (except nicotine or caffeine); positive urine screening for illicit substances (except cannabis); pregnancy (known or positive at baseline urine test) or breastfeeding; unstable somatic condition; known or suspected contraindication for ketamine, including hypersensitivity to ketamine, hypertension, class IV cardiac insufficiency, history of stroke, hepatic or cutaneous porphyria, history of intracranial hypertension; clinically important anomalies found during clinical examination, biological tests or electrocardiogram; non-stabilised hypertension or hypertension >180/100; concomitant electroconvulsive therapy; current participation or participation within the past three months in another interventional study; patients under judicial protection or guardianship.

Randomisation and masking

Patients were randomised 1:1 to placebo or ketamine after inclusion, before perfusion preparation. Patients were further randomised by blocks of random size.
Stratified by centre and by diagnostic category based on the Mini-International Neuropsychiatric Interview (MINI) 5.0,7 using a programme developed specifically for the study (SAS; Cary, NC). Participants were assigned a unique identification code. The three diagnostic categories were bipolar disorder; major depressive disorder; and any psychiatric disorder with no mention of a bipolar disorder, major depressive disorder, or any exclusion diagnosis (notably psychotic disorders and substance dependence; see exclusion criteria in “Patients” section).

Investigators and patients were blinded to study arm. The perfusion product was prepared within each participating department by a designated nurse, who was the only person with knowledge of randomisation results. This nurse did not participate in other aspects of the study and kept the arm assignments secret. Both ketamine and placebo perfusions were transparent and visually similar.

Procedures

At baseline, a sociodemographic and clinical assessment was conducted comprising the MINI 5.07 for psychiatric diagnoses according to DSM-IV criteria; the SSI (both clinician rated and self-rated versions18); the clinician rated Columbia suicide severity rating scale (CSSRS)15; the self-rated physical and psychological pain-visual analogue scale (PPPVAS)16; the self-rated Beck hopelessness scale (BHS)17; the 30 item inventory of depressive symptomatology (IDS-C30), clinician rated version22; and the clinical global impression scale (CGI) to assess the global impression according to the clinician.

Then, patients received a first 40 minute intravenous infusion of ketamine (0.5 mg/kg) or placebo 0.9% (saline solution) in addition to their current treatment.

This procedure has been used in most previous studies.12 A second administration was performed 24 hours later (including during weekends or bank holidays). The choice of two infusions was based on available data at the time of protocol writing in 2013/2014.8 23 Moreover, at that time, most protocols used a placebo, whereas later studies used midazolam.12 Usual care for these patients included a combination of admission to hospital, regular meetings with the healthcare staff (physicians, nurses), medication, individual and group psychotherapy, and family meetings.

Clinical evaluations were conducted at 40 minutes, 2 hours, 4 hours, day 1 (before the second infusion), day 2, and day 3. They comprised the SSIs, CSSRS, PPP-VAS, BHS, IDS-C30, CGI, and an assessment of safety and side effects with the Young mania rating scale (YMRS),24 patient rated inventory of side effects (PRISE),25 and brief psychiatric rating scale (BPRS).26 Patients were then followed up until the end of week 6, with assessments at day 4, week 2, week 4, and week 6.

Outcomes

The primary outcome was the rate of patients with a clinician rated SSI total score ≤3 (that is, in current full suicidal remission) at day 3 for each treatment arm and in each diagnostic group. SSI is a scale assessing suicidal ideation based on 19 items scored 0 to 2 (maximum score 38).18 Here, we used the level of suicidal ideation on the day of assessment. Secondary outcomes at day 3 were, for each arm and in each diagnostic group: rates of remission at intermediate time points; changes in SSI, BHS, PPP-VAS, IDS-C30, and CGI mean scores between baseline and day 3; rates of suicide attempts (CSSRS) during the three day period; and treatment side effects in each arm (YMRS, PRISE, and BPRS). At week 6, secondary outcomes were the rates of full suicidal remission in both arms.

Protocol amendments

Details of changes to the original protocol are presented in table S1. Briefly, changes were minor, including changes in investigators, biobank procedures (for future studies), extension of the inclusion period to reach the targeted sample size, and stopping the use of the self-rated quick 16 item Inventory of Depressive Symptomatology to save time, in addition to several questionnaires at 40 minutes, 2 hours, and 4 hours to reduce the burden of assessment.

Statistical analysis

This was a superiority trial. Calculation of sample size was done with nQuery software version 8.7.2.0 (table S2). Based on available literature at the time of the study conception, the hypothesis was an absolute difference of 45% resolution of suicidal ideation at day 3 between the two arms in each of the diagnostic groups (60% in the ketamine arm vs 15% in the placebo arm). To test this difference with a power of 85%, a two sided risk of 5% was taken into account the risk of α inflation associated with multiple comparisons (n=3), 52 patients were required for each diagnostic category (26 per arm), for a total of 156. A data monitoring committee oversaw the study.

Quantitative data were expressed as mean and standard deviation or median and interquartile range, according to their distribution. Qualitative data were expressed as absolute number and frequency (%). Comparison between groups used, as appropriate, Student’s t, Wilcoxon’s, X2, or Fisher’s tests. The rates of patients with SSI ≤3 at day 3 were compared between the two arms by a logistic regression model to take into account an arm by diagnostic group interaction to test heterogeneity. If the interaction term was significant, the between arm comparison was performed within each of the diagnostic categories. Holm’s corrections were performed to adjust for multiple comparisons. The associated odds ratios were estimated with 95% confidence interval with the profile likelihood method.

Linear mixed models were used to examine the effect of ketamine compared with placebo over time, on the different scores used as secondary endpoints, with patient considered as random effect. Time, drug, and time by drug interaction were tested. Analyses were performed within each of the diagnostic groups
when interaction with arm was significant. The associations between treatment, suicidal ideas, and psychological pain were explored with a mediation model. All included patients were analysed according to the intention-to-treat principle. A P value ≤0.05 was considered as statistically significant. Statistical analysis was performed with R 3.5.1 software (R Development Core Team, 2018). R Foundation for Statistical Computing, Vienna, Austria).

Patient and public involvement
Patients and the public were not involved in the design or conduct of this study, because their involvement in the design of scientific studies is recent in France and was not customary when the study was started in 2013. As the benefits of public involvement are obvious, this approach will be prioritised in our future studies. Moreover, patients will be involved in the discussion and dissemination of the findings of this study.

Results
Patient characteristics
Between 13 April 2015 and 12 March 2019, 156 patients were recruited and randomised to ketamine (n=73) or placebo (n=83), stratified into three groups: bipolar disorder (n=26/26, respectively); depressive disorder (26/30); and other diagnoses (21/27; fig 1). One centre (Centre Hospitalier Universitaire, Nîmes) included 75.0% of all patients.

Details of the participants’ characteristics are shown in table 1 (and tables S3-S5 for characteristics of each diagnostic group). Of note, patients had a past history of a suicidal act in 67 (93.1%) of 72 participants (data for one patient missing) in the ketamine arm, and in 70 (85.4%) of 82 participants (data for one patient missing) in the placebo arm. At inclusion, patients were severely suicidal in 71 (97.3%) of 73 participants in the ketamine arm and in 71 (86.6%) of 82 participants (data for one patient missing) in the placebo arm, based on the suicidality section of MINI 5.0 (10 or more points on a 32 point scale). All patients remained in hospital at day 3. At weeks 2, 4, and 6, rates of admission to hospital in the placebo and ketamine arms were, respectively 53.4%/41.8%, 34.4%/31.7%, and 22.7%/11.7%.

Main outcomes at day 3
By day 3 (primary endpoint), two patients withdrew consent and one was lost to follow-up (after the second infusion) in the placebo arm. On day 3, 46 (63.0%) of 73 patients in the ketamine arm and 25 (31.6%) of 79 patients (data for one patient missing) in the placebo arm had reached full remission of suicidal ideas (odds ratio 3.7 (95% confidence interval 1.9 to 7.3), P<0.001). These results were unchanged after adjustment for sex and presence of severe suicidal ideas (odds ratio 3.9 (2.0 to 7.9), P<0.001) or antalgic use (3.7 (1.9 to 7.6), P<0.001). Change over time in rates of suicidal
remission occurred by 60 minutes after infusion and persisted over the three day period (fig 2).

This effect differed according to the diagnostic group, with a significant interaction between arm and group, after adjustment for sex and presence of severe suicidal ideas (treatment arm: t=−4.7; df=1, P<0.001; diagnostic group: t=−3.3; df=2; P=0.2; interaction: t=−7.1; df=2; P=0.03); 84.6% (n=22) vs 28.0% (n=7) in the bipolar disorder group (odds ratio 14.1 (3.0 to 92.2), P<0.001), 42.3% (n=11) vs 35.7% (n=10) in the depressive disorder group (1.3 (0.3 to 5.2), P=0.6), and 61.9% (n=13) vs 30.8% (n=8) in the other diagnoses group (3.7 (0.9 to 17.3), P=0.07). Adding the main effect of recruitment centre into the analyses did not modify the outcomes.

Secondary outcomes at day 3
Table S6 in the supplemental materials shows information on the detailed outcomes. We found significant differences (after adjustment for centre

and diagnostic group) between the ketamine and placebo arms at day 3 in SSI median scores—both for the clinician rated (1.0 (interquartile range 0.8-0.0) vs 8.0 (2.0-15.5), respectively; unstandardised regression coefficient β=−5.0 (95% confidence interval −7.7 to −2.3), P<0.001) and patient rated versions (median score 7.0 (4.0-12.0) vs 11.5 (7.0-16.2); β=−2.5 (−4.5 to −0.4), P=0.02)—and in scores of depression (17.4 (standard deviation 12.1) vs 24.2 (12.7); β=−6.5 (−10.5 to −2.4), P=0.002), psychological pain (3.7 (interquartile range 0.3-6.3) vs 5.0 (2.0-8.0); β=−1.2 (−2.2 to −0.1), P<0.001), hopelessness (9.0 (4.0-15.0) vs 13.0 (8.0-17.0); β=−2.5 (−4.4 to −0.6), P=0.01), and global clinical improvement, but not physical pain (0.1 (0.0-3.0) vs 0.5 (0.0-3.5); β=−0.1 (−1.0 to 0.8), P=0.8). During the first three days, one suicide attempt occurred in the ketamine arm and none in the placebo arm.

No increase in YMRS and BPRS scores was seen in any patient from any arm (table S6). Table 2 reports the main side effects during the first three days. All side effects were rated as minor, and all symptoms reported in table 2 reduced significantly between the first assessment and day 4 (all P<0.05). Seventeen patients (23.3%) experienced at least one side effect in the ketamine group versus seven (8.4%) in the placebo group. The most common side effects in the ketamine group were sedation (11.0%), depersonalisation (9.6%), and nausea (6.8%).

Finally, we tested the hypothesis that the improvement of suicidal ideation was mediated by its effect on psychological pain, as assessed by the patient rated version of the SSI. The treatment was associated with a significant reduction in SSI scores after 72 hours (t=−1.9, P=0.05, accounting for centre), but this association was weaker (t=−1.8, P=0.07) after accounting for psychological pain, while psychological pain remained highly associated with SSI scores (t=7.2; P<0.001), suggesting a mediation effect.

Outcomes at week 6
Between day 4 and week 6 (study end), seven patients withdrew consent and seven patients were lost to follow-up in the placebo arm. In the ketamine arm, one patient died from suicide (determined by the oversight committee to be unrelated to the intervention), four withdrew consent, and eight were lost to follow-up. Therefore, 60 (82.2%) patients of the 73 in the ketamine group and 66 (79.5%) patients of 83 in the placebo group completed the study.

Over the study, eight patients (9.8%) in the placebo arm (data for one patient missing) and six patients (8.2%) in the ketamine arm attempted suicide (two v zero, respectively, in the bipolar disorder group; one v five in the depressive disorder group; and five v one in the other diagnosis group). Between day 4 and week 6, the ketamine arm continued to have better full suicidal remission than the placebo arm (69.5 v 56.3% at week 6), although this was not significant at week 6 owing to reduced suicidality in the placebo group over time (odds ratio 0.8 (95% confidence interval 0.3 to 2.5).
Results in individuals not fulfilling the criteria for a full suicidal remission (as indicated by depression and/or suicidal ideation scores; and (c) some of them finally reached remission of suicidal ideas.30 Therefore, this group might be particularly heterogeneous, with both more patients sensitive to a placebo effect and more patients requiring repeated ketamine infusions.

The persistence of the ketamine effect at six weeks of intake is not in line with previous studies.8 10 11 31 and the related meta-analysis,12 but those three studies together analysed only 63 patients. More long term studies are needed.

Over the six week period, 8.2% of patients in the ketamine arm and 9.8% in the placebo arm attempted suicide, including one fatal act. Detailed examination of these events in the intervention group showed that (a) none of them had exacerbated depression or suicidal ideation scores after infusion, suggesting that ketamine had no direct negative effect; (b) all these patients were poor responders to ketamine during the first three days as indicated by depression and suicidal ideation scores; and (c) some of them finally reached remission of suicidal ideas after several days, which might have led to decreased vigilance. It must be remembered that the resolution of a suicidal crisis necessitates more than medication alone. Psychological, social, and family care and support should always be combined with pharmacotherapy. Finally, this study was not designed to assess the benefits of ketamine for prevention of a suicidal act, and larger studies and meta-analyses will be necessary.

Discussion
Principal findings
This study confirmed in a large randomised controlled trial that ketamine is a fast acting, efficient treatment of suicidal ideation. In this population at very high suicidal risk, 63.0% reached full remission at three days after two infusions in the ketamine group in comparison with 31.6% in the placebo group. This effect was rapid, with 43.8% remission only two hours after the first infusion versus 7.3% in the placebo group. Ketamine was well tolerated without severe side effects. Main side effects, including sedation, depersonalisation/derealisation, nausea, and dizziness, were of short duration and occurred in around 10% or fewer participants. In addition, the effect persisted at six weeks in 69.5% of individuals treated with ketamine (versus 56.3% in those receiving placebo).

This study highlights a major moderating effect on primary mental disorders. A strong effect of ketamine versus placebo was found in the group with bipolar disorder, whereas the effect was moderate and did not quite reach significance in the group with “other psychiatric disorders,” and was non-significant in major depressive disorders. Results in bipolar disorder—a disorder associated with a high suicidal risk and limited options to treat depression—are highly encouraging and support two previous small studies.8 28 Notably, no mood switch was seen in the 26 patients with bipolar disorder treated with ketamine. Results in individuals not fulfilling the criteria for a full major depressive episode (whether bipolar or major depressive disorders) also support the observation that ketamine is efficient independently from depressive episodes, as suggested by previous authors.29 This group comprised a substantial number of individuals with post-traumatic stress disorder, dysthymia, and anxiety disorders (panic disorder, agoraphobia, generalised anxiety disorder). These patients would probably also have had personality disorders, although this was not formerly measured.

The non-significant outcomes in depressive disorders are more challenging to interpret. This group showed the highest placebo effect (36% vs 28% in bipolar disorder and 31% in other diagnoses). This placebo effect is also higher than that found in a meta-analysis of 10 trials analysing 157 suicidal patients with overall remission rates (clinician rated measures) of around 30% in the control group.27 Moreover, the effect of ketamine (42%) was lower than in the two other groups in our study (84% and 62%) and in the meta-analysis (around 55%).29 Our study, therefore, might have lacked power to detect an effect in this particular group with depressive disorders. Additionally, one study of treatment resistant depression suggests that repeated doses of ketamine might be necessary for some patients to achieve remission of severe suicidal ideas.30 Therefore, this group might be particularly heterogeneous, with both more patients sensitive to a placebo effect and more patients requiring repeated ketamine infusions.

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Table 2 | Reports of side effects in each treatment arm within the 72 hour period

| Side effect                      | Ketamine (n=73) N (%) | Placebo (n=83) N (%) |
|----------------------------------|-----------------------|----------------------|
| Sedation                         | 8 (11.0)              | 2 (2.4)              |
| Depersonalisation/derealisation   | 7 (9.6)               | 2 (2.4)              |
| Nausea                           | 5 (6.8)               | 2 (2.4)              |
| Dizziness                        | 3 (4.1)               | 1 (1.2)              |
| Agitation                        | 2 (2.7)               | 1 (1.2)              |
| Tremor                           | 2 (2.7)               | 1 (1.2)              |
| Blurred vision                   | 2 (2.7)               | 1 (1.2)              |
| Anger                            | 1 (1.4)               | 1 (1.2)              |
| Hallucination                    | 1 (1.4)               | 1 (1.2)              |
| Sweating                         | 1 (1.4)               | 1 (1.2)              |
| Hypotension                      | 1 (1.4)               | 1 (1.2)              |
| Tachycardia                      | 1 (1.4)               | 1 (1.2)              |
| Vomiting                         | 1 (1.4)               | 1 (1.2)              |
| Dry mouth                        | 1 (1.4)               | 1 (1.2)              |
| Diarrhoea                        | 1 (1.4)               | 1 (1.2)              |
| Vagal syncope                    | 1 (1.4)               | 1 (1.2)              |

P=0.7; fig 3). Results for each intermediate endpoint are reported in table S6.

Fig 2 | Change in rates of suicidal remission over time within 72 hours of the first injection. Here, suicidal remission corresponds to a score ≤ 3 on the Beck scale for suicidal ideation, clinician rated version.
Of note, a recent review of 15 studies suggested that in the short term no more suicidal acts occurred in the ketamine group than in the placebo group.

Overall, the tolerance of ketamine was good, as three quarters of patients had no side effects, and side effects were largely minor and of short duration. This result is in line with a recent review of literature emphasising that tolerance of ketamine is good.32

Additional findings

Our study suggests that the beneficial effect of ketamine on suicidal ideation could be mediated by an effect on psychological pain. Although mental pain does not necessarily lead to suicidal ideas, recent studies suggest that individuals with severe suicidal ideas (notably those with a plan) also have high levels of mental pain.33 Ketamine might therefore exert its effects through analgesic mechanisms that reduce mental pain. Indirect support for this suggestion is the observation that the effects of ketamine on depression might involve the opioid system34 (although this is controversial35), that buprenorphine—a μ opioid partial agonist—is also effective on suicidal ideas,36 and that mental pain has been associated with the nociceptin system.33

Limitations

Results should be interpreted in light of several limitations. Firstly, although this is a large study and sufficiently powered, analyses within diagnostic groups were on smaller samples, which might explain both the large effect size of ketamine in bipolar disorder, and the lack of significant differences in the depressive disorder group. Secondly, as ketamine can induce recognisable effects (depersonalisation, dizziness), masking might have been compromised for both the patients and the investigators, but this was not formally measured. It should, however, be noted that only 9.6% of patients in the ketamine group experienced depersonalisation and 4.1% dizziness (ν 2.4% in the placebo group) while other side effects were unspecific and found in the placebo group. Midazolam has been used instead of placebo in a few studies and should be considered as a suitable control in the future. Thirdly, the rapid resolution of suicidal ideas after receiving ketamine does not equate to a reduced risk of suicidal acts, notably after hospital discharge. Indeed, the rates of suicide attempts during follow-up were similar between the groups. Moreover, ketamine is a drug with a potential for abuse. Longer follow-up of larger samples will be necessary to examine benefits on suicidal behaviours and long term risks.

Conclusion

This large trial confirms that ketamine rapidly induces remission of severe suicidal ideation in adults, an effect persisting over six weeks in two thirds of patients. This effect seems to be dependent upon comorbid mental disorders. The tolerance was good. Long term benefits and safety of ketamine must be examined and drugs with different mechanisms of action will have to be investigated for non-responders.

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Ethical approval: Ethical approval was obtained on 18 July 2014 from the research ethics board “Comité de Protection des Personnes (CPP) Sud Méditerranée III” (ref. 2014.06.03 bs). This study was prospectively registered on the 20 November 2014 on https://www.clinicaltrials.gov/ (NCT02299440) and is listed on EudraCT. 2014-001234-30. All patients gave their informed, written and signed consent before inclusion.

Data sharing: Individuals’ participant data that underline the results reported in this article (after de-identification) will be available. The study protocol, statistical analysis plan, and informed consent form will also be provided on request. Data will be immediately available after publication, with no end date. Data will be provided to investigators whose proposed use of the data has been approved by an independent review committee identified for this purpose for individual participant data meta-analysis or review. Proposals should be directed to pascale.FABBRO@chu-nimes.fr

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Web appendix: Supplementary materials