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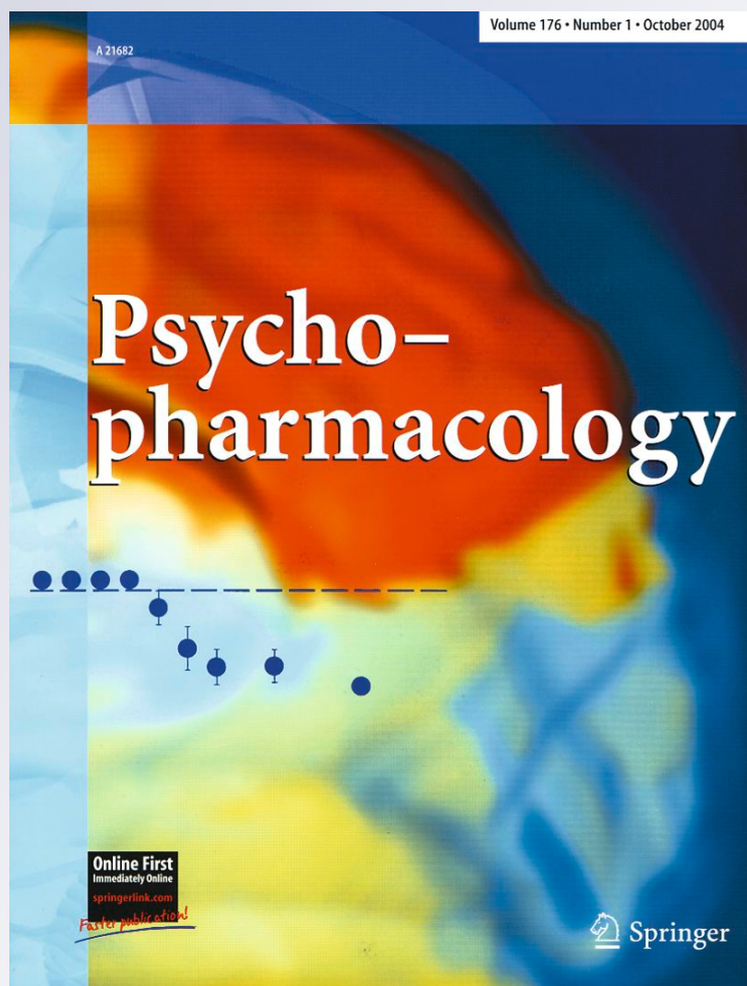
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Ketamine perturbs perception of the flow of time in healthy volunteers

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Abstract

Rationale Disturbances in the subjective experience of time have been observed both in schizophrenia and following acute administration of ketamine. However, effects of ketamine on more objective timing tasks have not yet been measured in humans, nor has it been established that timing effects are not merely secondary to a more general dysfunction in working memory (WM).

Objective and methods In a double-blind placebo-controlled crossover study, we characterised the effects of ketamine (100 ng/ml blood plasma level) on performance of perceptual timing and colour discrimination tasks, which were matched for WM and attentional demands. To test the ubiquity of ketamine's effects on timing, we also examined two distinct measures of temporal predictability.

Results Ketamine significantly distorted the subjective experience of time as measured by the Clinician-Administered Dissociative States Scales. Critically, ketamine also impaired accuracy on the perceptual timing task while having no effect on performance of the colour perception task. Although ketamine did not impair the ability to use prelearned temporal (or spatial) cues to predict target onset (or location), it did slow reaction times at long delays following non-informative neutral cues, suggesting an impaired ability to use the unidirectional flow of time itself to make temporal predictions.

Conclusions Ketamine induced selective impairments in timing, which could not be explained by more fundamental effects on the ability to hold information in WM. Rather our collected findings suggest that ketamine may disturb timing by selectively impairing the way in which information is temporally manipulated within WM.

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Keywords Ketamine · Timing · Duration · NMDA ·
Glutamate · Temporal expectation · Hazard function ·
Foreperiod · Temporal orienting · Attention

Introduction

The glutamatergic *N*-methyl-D-aspartate (NMDA) receptor antagonist ketamine is a useful pharmacological model of psychosis (Krystal et al. 1994; Corlett et al. 2007a). In healthy participants, it induces perceptual changes (Krystal et al. 1994; Duncan et al. 2001; Stone and Pilowsky 2006), thought disorder (Krystal et al. 2005; Malhotra et al. 1996; Duncan et al. 2001) and negative symptoms (Krystal et al. 2005; Duncan et al. 2001), as well as delusion-like thinking (Pomarol-Clotet et al. 2006). Moreover, ketamine induces selective impairments in cognitive function (Morgan and

Curran 2006) that are comparable to those observed in schizophrenia (Fletcher and Honey 2006). Notably, both ketamine (Honey et al. 2003, 2004) and schizophrenia (Perry et al. 2001; Kim et al. 2004) are associated with difficulties in manipulating, rather than maintaining, information in working memory (WM). While ketamine does not reproduce the signs and symptoms of established schizophrenia, it has been argued that the perceptual and cognitive changes it induces resemble those found during prodromal stages of the illness (Pomarol-Clotet et al. 2006; Corlett et al. 2006, 2007a, 2007b, 2011).

One feature of schizophrenia that has not been well-explored under ketamine is that of the experience of time. Numerous studies have demonstrated that patients with schizophrenia show reduced accuracy in estimating periods of time in the minutes-to-hours range (Rabin 1957; Johnson and Petzel 1971) and in timing of stimulus durations in the range of several hundreds of milliseconds to a few seconds (Lhamon and Goldstone 1956; Tysk 1983; Elvevåg et al. 2003; Davalos et al. 2003; Carroll et al. 2008, 2009). However, the effects of ketamine on timing are uncertain: in rats, ketamine had no direct effect on timing (Cheng et al. 2006) but instead ‘unlocked’ the clock-speeding effects of cocaine that had otherwise been masked by extensive training (Cheng et al. 2007a). Yet, subjective reports in healthy volunteers indicate that ketamine can distort the experience of time (Pomarol-Clotet et al. 2006; Stone and Pilowsky 2006), a phenomenon also observed in schizophrenia (Freedman 1974; Seeman 1976; Vogeley and Kupke 2007).

Although the effects of ketamine on timing have not yet been explored experimentally in humans, Rammsayer (2006) noted that memantine (a lower-affinity NMDA antagonist) impaired timing of auditory stimuli in the seconds (~1 s.) but not tens of milliseconds (~50 ms) range. Timing in the range of seconds, but not milliseconds, requires support from WM and sustained attention processes (Gibbon et al. 1984; Zakay and Block 1996): a seconds-range stimulus must be continually attended from stimulus onset to offset (as opposed to estimates of, e.g. pitch or colour, which can be estimated almost instantaneously), and the contents of WM must be continuously updated as stimulus presentation unfolds. Given the pattern of memantine's effects, Rammsayer (2006) concluded that its selective effects on seconds-range timing were secondary to its effects on WM. Given differences in their pharmacological specificity, we cannot presume that ketamine's effects would be similar to those of memantine nor, from a more cognitive point of view, that such effects necessarily reflect a more fundamental effect on WM. Timing of longer, seconds-range durations depends not only upon WM but also, of course, upon an index of elapsed time itself, for example accumulation of temporal pulses (Gibbon et al. 1984; Treisman 1963) or temporal integra-

tion of steadily climbing neuronal activity (Reutimann et al. 2004). Critically, psychophysical evidence suggests that this timing mechanism may be quite distinct for millisecond- and seconds-range timing (Gibbon et al. 1997; Buonomano et al. 2009). Thus, it is possible that a deficit in the seconds but not milliseconds, range reflects a specific temporal, rather than WM, deficit. We therefore sought to both establish whether ketamine, like memantine, is associated with a disturbance in an objective measure of time perception in the seconds time range and, crucially, to dissociate such an effect from concurrent mnemonic disturbance by directly comparing the effects of ketamine on performance of timing and colour discrimination tasks that were matched for WM demands (Coull et al. 2004, 2008). A ketamine-induced impairment of the timing, but not colour, task would strongly suggest a selective timing impairment over and above any effects on WM.

However, ‘timing’ is an imprecise term (Michon 1985; Pöppel 1997; Coull and Nobre 2008) that can refer equally well to how long a current event lasts (duration estimation) or when a future event is likely to occur (temporal prediction). Since ketamine has previously been shown to modulate neural indices of prediction error (e.g. Corlett et al. 2007a; 2011), we examine performance not only on time perception but also on measures of temporal prediction. Specifically, we tested the impact of ketamine on two distinct indices of temporal prediction derived from the temporal-orienting paradigm (Coull and Nobre 1998; Nobre 2001), a temporal analogue of the classic Posner covert orienting of spatial attention task (Posner et al. 1980). In the task, visual cues provide valid, invalid or neutral information concerning the likely delay before an imminent target is presented. The behavioural benefit of temporally informative cues is indexed by faster reaction times during valid compared to invalid trials (Coull and Nobre 1998; Griffin et al. 2001). Yet, even when the onset of the target cannot be predicted by temporal cues (i.e. in the neutral condition), the unidirectional flow of time itself, or ‘time's arrow’ (Eddington 1928), provides a degree of predictive power: since time flows inexorably forward an event that we expect to occur, but has not yet occurred, must do so at some time in the future. The ‘hazard function’ measures the increasing conditional probability (and, hence, increasing sense of temporal expectation) over time that an event will occur, given that it has not already occurred (Elithorn and Lawrence 1955; Luce 1986). Increasing temporal certainty of target presentation over time allows the response plan to be updated dynamically, as the trial unfolds, which ultimately translates into faster RTs at longer delays (Niemi and Näätänen 1981). The orienting paradigm can therefore index temporally predictive information that is conveyed either by valid, prelearned cues (temporal condition) or by the very passage of time itself (neutral condition). In order

to control for possible, though unlikely (Morgan and Curran 2006), effects of ketamine on non-specific attentional processes, we also examined effects of ketamine on the more classic, spatial, version of the task.

Methods

Participants

We collected data from 13 (6 male and 7 female; mean age 23.1 years) right-handed, healthy volunteers. Although three further volunteers had been recruited, their data could not be included in the final analysis due either to technical difficulties or subject dropout. Research was approved by Addenbrookes NHS Trust Research Ethics Committee. Participants spoke fluent English, had normal or corrected-to-normal vision with no colour blindness, and reported no history of clinical drug or alcohol abuse or of psychiatric illness.

Experimental design

Each participant was tested on two occasions, with a minimum of 1 week between sessions. A within-subject, double-blind crossover design was used such that participants received ketamine in one session and placebo in the other. Participants also completed two learning tasks during the testing session, and performed all tasks in the same order from one session to the next. Here, we report

data for the two timing tasks (data for other tasks will be reported elsewhere).

Ketamine infusion protocol

Racemic ketamine (2 mg/ml solution) or saline solution was administered intravenously by a computerised target-controlled infusion system, which calculated infusion rates necessary to achieve the ‘target’ blood concentration: 100 ng/ml plasma for 75 min immediately followed by a further 40 min at 200 ng/ml plasma. The timing tasks reported here were performed during the 100 ng/ml infusion period only. The infusion pump (Graseby 3500, Smiths Medical, Ashford, UK) was controlled by a personal laptop computer running Stanpump software, programmed to use a two-compartmental pharmacokinetic model for ketamine (Absalom et al. 2007) to calculate infusion rates. The order of infusion was counterbalanced across participants.

Cognitive tasks

Both tasks were performed at 100 ng/ml blood plasma level of ketamine. Twelve participants completed the discrimination tasks and 11 completed the orienting tasks.

Temporal and colour discrimination

Task The trial structure and stimuli used in the temporal and colour tasks were identical (Fig. 1), with the only

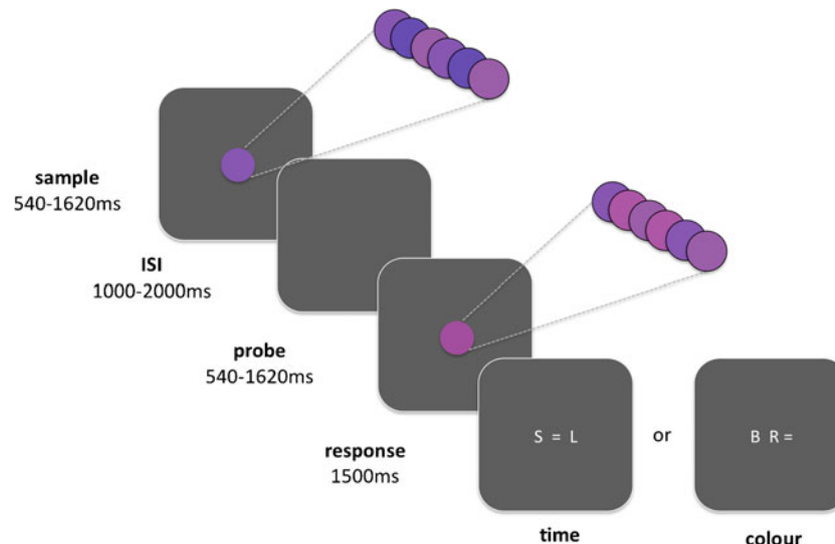


Fig. 1 Temporal and colour discrimination task. Participants estimated either the duration or colour of two consecutive stimuli. The first (*sample*) and second (*probe*) stimuli were presented for one of three durations (540, 1,080, 1,620 ms) and had an overall percept of one of three shades of purple (*maroon*, *violet* or *indigo*). Stimuli were not of a uniform colour but instead comprised rapidly flickering (90 ms)

presentations of three of five different shades of purple (see inset) to give the overall colour percept. Participants indicated whether the probe was shorter (*S*), longer (*L*) or the same (=) duration as the sample ('time' condition) or redder (*R*), bluer (*B*) or the same (=) shade of purple as the sample ('colour' condition) using a three-choice button press at the onset of the response screen

difference being whether participants were asked to make time or colour judgements. In the time condition, participants estimated whether the duration of the second (probe) stimulus was shorter (S), equal to (=), or longer (L) than the previous (sample) stimulus. In the colour condition, participants estimated whether the probe was redder (R), equal to (=), or bluer (B) than the sample. The response screen presented three alphanumeric characters, indicating each of the three possible responses for each condition. Their spatial position could vary on a trial-by-trial basis. Participants pressed one of the three corresponding response buttons (left, middle or right) to indicate their estimate of relative duration or colour. The response screen appeared for 1,500 ms, during which time the subject gave their response, and any responses slower than 1,500 ms were not recorded. Inter-trial intervals varied pseudo-randomly from 1 to 2 s. Participants performed 36 trials per condition, which comprised equal numbers of trials in which the probe was shorter than/longer than/equal to the sample in the time condition, or bluer than/redder than/equal to the sample in the colour task. The time and colour conditions were presented twice each in a blocked manner, with each block comprising 18 trials of either the time condition or the colour condition.

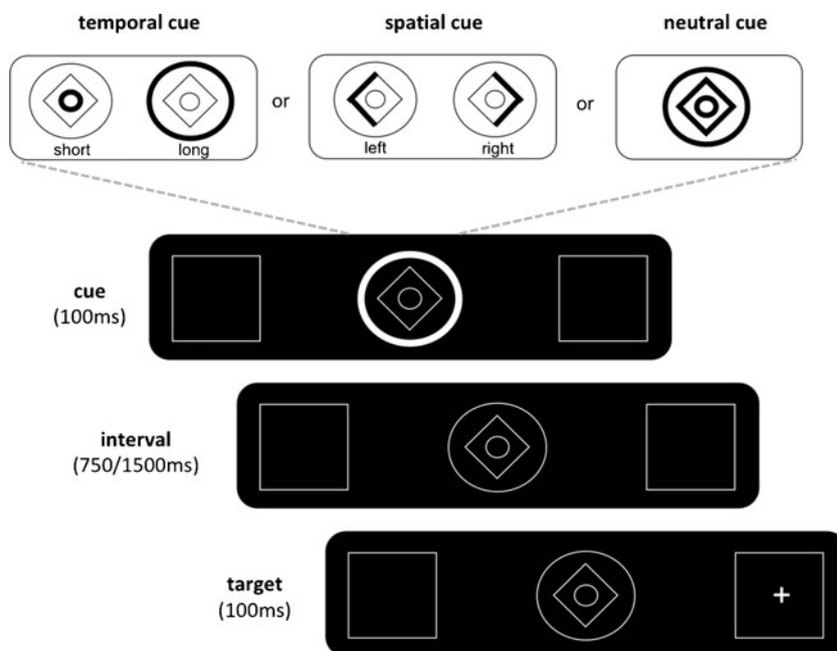
Stimuli The sample and probe stimuli were presented consecutively, separated by a pseudo-random inter-stimulus interval of 1–2 s. Each stimulus was presented for one of three durations (540, 1,080 or 1,620 ms), and had an overall percept of one of three shades of purple (maroon, violet or indigo) ranging from a reddish to a

bluish hue. Duration and colour stimulus attributes were counterbalanced such that any of the three stimulus durations could be paired with any of the three colours. The stimuli to be estimated were not a uniform colour for the entire duration of stimulus presentation. Rather, rapidly alternating presentations (90 ms) of five different shades of purple across the entire stimulus duration gave an overall percept of maroon, violet or indigo. During the colour task, the participant estimated the average shade of purple by amalgamating all shades presented during the flickering percept. This rather unusual colour task was designed to ensure that subjects maintained attention, and continually updated their colour estimate, throughout the entire stimulus presentation. This manipulation is crucial for equating sustained attention and working memory demands across the time and colour tasks since (1) time perception necessitates attention being sustained for the entire duration of the stimulus (~1 s.), whereas perception of a static colour would occur within the first hundred milliseconds or so, and (2) time perception requires stimulus onset to be held in working memory and then continually updated as time elapses, whereas perception of a static colour would require no such updating of working memory.

Temporal and spatial orienting of attention

Task The task manipulated participants' expectations of *where* or *when* target stimuli would appear within an experimental display. A central cue (inset Fig. 2) predicted either the location (spatial condition) or onset (temporal

Fig. 2 Temporal and spatial-orienting task. Participants detected a target appearing at one of two peripheral locations (*left/right*) after one of two cue-target intervals (*short/long*) as quickly as possible. In the spatial or temporal condition, the visual cue predicted where or when (respectively) the target was likely to appear, with 80% validity. In the neutral condition, the visual cue carried neither spatial nor temporal information



condition) of an imminent target or else it provided neither temporal nor spatial information (neutral condition). In the spatial and temporal conditions, cues could either correctly (valid trials) or incorrectly (invalid trials) predict where or when the target would appear, with an overall ratio of 80:20 valid/invalid trials. In the neutral conditions, the participants could predict neither where nor when the target would appear. The participants' task in all three conditions was simply to detect the target as rapidly as possible, while avoiding mistakes. Participants indicated covert detection of the target by pressing a response button with their right index finger. Inter-trial intervals varied between 600 and 1,000 ms. The computer recorded reaction times to target stimuli. Subjects were thoroughly familiarized with the cues prior to performing 120 experimental trials of each of the three conditions (360 in total). For the temporal and spatial conditions, this corresponded to 96 valid trials and 24 invalid trials.

Stimuli The basic visual display consisted of a central cueing stimulus (1° eccentricity) and two peripheral boxes (7° eccentricity), inside which the target ('x' or '+') appeared (Fig. 2). The central cue was a compound stimulus consisting of a diamond and two concentric circles. One part of the cue was briefly (100 ms) highlighted to inform the subject whether to attend to the location or time of onset of the target. During the spatial condition, the left (or right) side of the diamond brightened to inform the subject that the target was likely to appear in the left (or right) peripheral box. During the temporal condition, brightening of the inner or outer circle indicated that the target would appear after a short (750 ms) or long (1,500 ms) delay, respectively. During the neutral-cue condition, the entire cue brightened, providing no spatial or temporal information. Targets appeared for 100 ms in one of the two peripheral boxes and after one of two time intervals, according to the nature of the cue: in the spatial condition, targets appeared randomly at the short or long time delay, but their spatial location was determined by the nature of the preceding cue (left/right; valid/invalid); in the temporal condition, targets appeared randomly in the left or right box but their temporal onset was determined by the nature of the preceding cue (short/long; valid/invalid). Spatial locations and temporal onsets were counterbalanced across trials for all three experimental conditions.

Subjective rating scales

Subjective experience induced by the drug was recorded using a series of clinician-administered questionnaires: the Clinician-Administered Dissociative States Scales

(CADSS, Bremner et al. 1998), Brief Psychiatric Rating Scale, Startup and Startup and a Rating Scale for Psychiatric Symptoms. The time perception subscale of the CADSS (Krystal et al. 1994; 1998) indexed the subjective experience of time more specifically (see Appendix).

Data analysis

Temporal and colour discrimination

Mean accuracy (percentage of trials correct), omissions (percentage of missed responses) and reaction times (RTs) were analysed in $2 \times 2 \times 2$ repeated measures ANOVAs, with task (time/colour) and treatment (ketamine/placebo) as within-subjects factors, and treatment order (ketamine–placebo/placebo–ketamine) as a between-subjects factor. In addition, proportion of under or overestimates in the time condition were analysed in a $2 \times 3 \times 2 \times 2$ repeated measures ANOVA, with error type (under/overestimate), trial type (probe shorter/equal to/longer than sample) and treatment as within-subjects factors, and treatment order as a between-subjects factor.

Temporal and spatial orienting of attention

Mean RTs for temporal and spatial orienting conditions were analysed in $2 \times 2 \times 2 \times 2$ repeated measures ANOVAs with treatment, cue validity (valid/invalid) and either target onset (short/long) or target location (left/right) (for the temporal and spatial conditions, respectively) as within-subject factors and treatment order as a between-subject factor. Mean RTs for the neutral condition were analysed in a $2 \times 2 \times 2 \times 2$ repeated measures ANOVA with target onset, target location and treatment as within-subject factors and treatment order as a between-subject factor.

Subjective rating scales

Overall CADSS and CADSS time perception subscale scores were analysed in separate 2×2 repeated measures ANOVAs with treatment as a within-subject factor and treatment order as a between-subjects factor. Spearman's rank correlations further explored the relationship between these subjective CADSS scores and objective discrimination task performance. Specifically, the association between task performance and (1) global dissociative experience and (2) subjective experience of time was tested by correlating drug-induced change (ketamine–placebo) in mean timing or colour discrimination accuracy with drug-induced change in (1) overall CADSS and (2) CADSS time perception subscale scores, respectively.

Results

Temporal and colour discrimination

A significant main effect of task [$F(1, 10)=11.80, p<0.01$] revealed that the colour task was performed less accurately than the timing task. A significant main effect of treatment [$F(1, 10)=7.34, p<0.05$] was qualified by a significant interaction between treatment and task [$F(1, 10)=8.73, p<0.05$]. Post-hoc analysis with paired t tests indicated that ketamine, as compared to placebo, significantly reduced accuracy in the timing task ($t(11)=3.35, p<0.01$) but had no effect on performance of the colour task ($t(11)=-0.44, ns$) (Fig. 3a). Ketamine had no significant effect on reaction time or number of omissions. There were no significant main effects of, or interactions with, treatment order.

Given the selective effect of ketamine on timing, we explored these timing data further to examine whether the drug-induced impairments in accuracy were caused by a consistent bias to under or overestimate probe duration. First, each trial was classified according to whether the

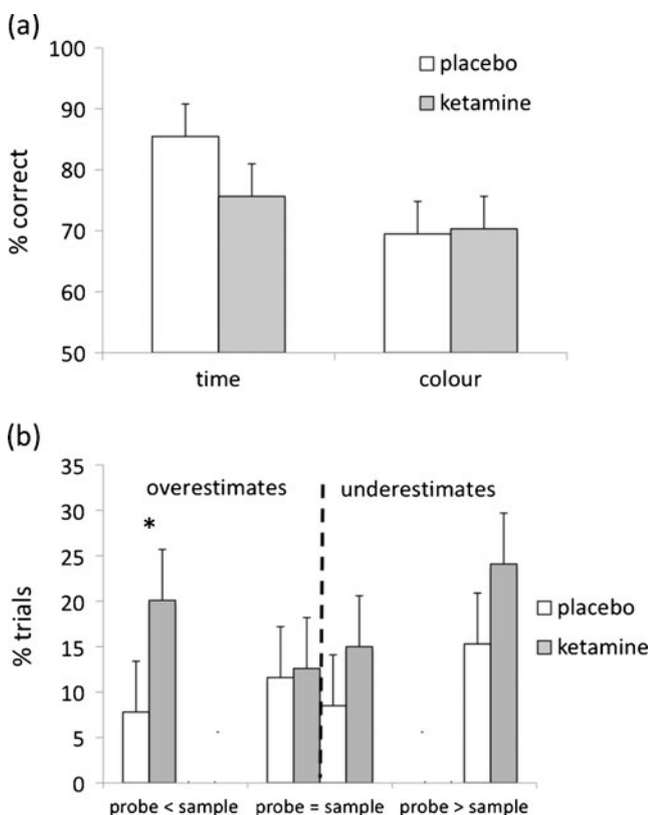


Fig. 3 Ketamine impairs perceptual timing. **a** Ketamine selectively impaired accuracy (% correct) of time discrimination, while having no effect on accuracy of colour discrimination. **b** During the temporal discrimination task, ketamine significantly increased the likelihood (% trials) that the probe stimulus would be overestimated when the probe was shorter than (<) the sample. Error bars represent the standard error of the mean

second stimulus (probe) was longer, shorter or of equal duration to the first stimulus (sample). Then, for each of these three trial types, we calculated the proportion of trials in which a subject under or overestimated the probe compared to the sample. If the probe and sample were of the same duration but the subject judged the probe to be shorter, then this was classed as an underestimate, whereas if the probe was judged to be longer, then this was classed as an overestimate. If the probe was longer than the sample but the participant judged it to be shorter or of equal length, then this was classed as an underestimate. Conversely, if the probe was shorter than the sample but the subject judged it to be longer or of equal length, then this was classed as an overestimate. A main effect of treatment showed that ketamine significantly increased the proportion of both over- and underestimates [$F(1, 10)=11.50, p<0.01$], but did not differentially induce one type of error more than the other (treatment \times error-type interaction [$F(1, 10)=0.09, ns$]). However, when data were broken down into trial type (probe shorter/equal to/longer than sample), a significant interaction between treatment, error type and trial type [$F(2, 20)=7.34, p<0.005$] indicated that ketamine, as compared to placebo, induced an increased incidence of overestimates, particularly when the probe was shorter than the sample ($t(11)=4.14, p<0.005$) (Fig. 3b). Although there was a corresponding increase in underestimates when the probe was longer than the sample (Fig. 3b), this trend merely approached statistical significance ($t(11)=1.84, p=0.09$).

Temporal and spatial orienting of attention

Any RTs faster than 100 ms were considered anticipatory and removed from the analysis. All remaining RTs were log transformed to normalize distributions (resulting RTs non-significantly different from a normal distribution as assessed by the Shapiro–Wilks test).

Temporal orienting of attention

A main effect of cue validity [$F(1, 9)=11.54, p<0.01$] indicated that targets appearing when expected were detected more quickly than targets appearing at an unexpected moment (Table 1). But, as predicted by the results of previous studies using the temporal orienting paradigm (e.g. Coull and Nobre 1998; Coull et al. 2000; Correa et al. 2006), this main effect was qualified by a significant validity \times onset interaction [$F(1, 9)=18.24, p<0.005$]. Specifically, the benefit of temporally valid cues was significant at the short ($t(10)=-4.71, p=0.001$) but not long ($t(10)=-0.48, ns$) interval (Table 1): when a target does not appear after the short interval as expected it must, by process of elimination, appear at the longer one. Such temporal predictability effectively removes the cost of

Table 1 Mean (\pm standard error) reaction times (RT) to detect targets in a cued simple RT task

	Temporal orienting				Spatial orienting			
	Short		Long		Left		Right	
	Valid	Invalid	Valid	Invalid	Valid	Invalid	Valid	Invalid
Placebo	310.06 (\pm 14.37)	341.84 (\pm 17.57)	290.88 (\pm 15.87)	295.50 (\pm 13.78)	311.31 (\pm 19.58)	339.03 (\pm 18.21)	308.30 (\pm 15.78)	346.53 (\pm 22.06)
Ketamine	316.68 (\pm 11.00)	346.75 (\pm 10.68)	301.53 (\pm 14.45)	301.82 (\pm 14.81)	321.00 (\pm 14.93)	361.01 (\pm 19.41)	314.79 (\pm 17.54)	354.18 (\pm 24.58)

In separate blocks, the cue predicted either the onset (temporal-orienting task) or location (spatial-orienting task) of the upcoming target. RTs are shown for targets appearing after either a short (750 ms) or long (1,500 ms) delay in the temporal-orienting task or on the left or right-hand side in the spatial-orienting task. The predictive information conveyed by the cue could be either valid (80% of trials) or invalid (20% of trials)

targets appearing unexpectedly late. Crucially however, ketamine did not interact with either cue validity [$F(1, 9)=0.25$, ns], temporal onset [$F(1, 9)=0.06$, ns], nor the interaction between the two [$F(1, 9)=0.05$, ns]. Nor did ketamine have any overall effect on performance (main effect of drug [$F(1, 9)=0.43$, ns]). Ketamine, therefore, had no effect on the use of temporal cues to predict stimulus onset.

Spatial orienting of attention

As predicted by prior studies (Posner et al. 1980), a main effect of cue validity ($F(1, 9)=29.24$, $p<0.0001$) confirmed that targets were detected more quickly when they appeared in expected versus unexpected spatial locations (Table 1). Cue validity did not interact with any other factor, including treatment [$F(1, 9)=0.05$, ns]. Ketamine, therefore, had no effect on the use of spatial cues to predict stimulus location.

Hazard function

As predicted by numerous previous studies (see Niemi and Näätänen 1981 for a review), a main effect of onset [$F(1, 9)=30.81$, $p<0.0001$] confirmed that RTs in the neutral cue condition were significantly faster at long versus short intervals. This speeding effect is due to the increasing conditional probability of target occurrence as a function of the passage of time, i.e. the hazard function, which allows for improved response preparation over time, and thus faster RTs. However, a significant treatment by onset interaction [$F(1, 9)=5.83$, $p<0.05$] revealed that, as compared to placebo, the relative speeding of RTs at the long interval was attenuated by administration of ketamine (Fig. 4). Therefore, although ketamine did not significantly slow performance in a global manner (main effect of drug [$F(1, 9)=1.71$, ns]), it selectively compromised the beneficial effects of the hazard function on performance.

Subjective rating scales

Consistent with numerous previous reports (e.g. Krystal et al. 1994, 1998; Fu et al. 2005) a main effect of treatment on the overall CADSS score [$F(1, 10)=10.95$, $p<0.01$] confirmed that ketamine induced a general dissociative state in healthy volunteers, though this effect tended [$F(1, 10)=4.61$, $p<0.06$] to be more pronounced in subjects receiving ketamine in the first session and placebo in the second (Table 2). More specifically, ketamine significantly [$F(1, 10)=7.22$, $p<0.05$] increased scores on the CADSS time perception subscale (see also Krystal et al. 1998), indicating a drug-induced distortion in the subjective experience of time. Again, this effect was greater

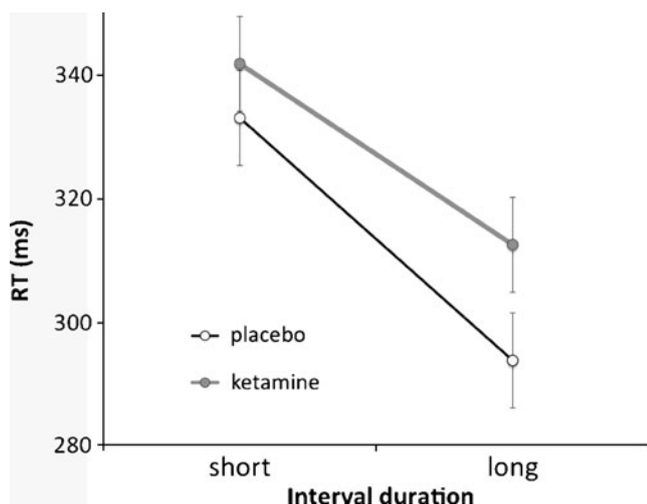


Fig. 4 Ketamine attenuates the beneficial effect of the hazard function. The neutral cue condition of the orienting task produces faster RTs for targets appearing after long rather than short intervals due to the “hazard function”. Ketamine slowed RTs selectively at the long, not short, intervals, thus significantly attenuating the benefit that is normally afforded by long delays. Error bars represent the standard error of the mean

[$F(1, 10)=5.09, p<0.05$] in subjects receiving ketamine in the first session and placebo in the second (Table 2).

We used Spearman's rank correlations to search for a link between these subjective measures of perceptual experience and more objective measures of visual perception. Neither the overall CADSS score, nor the time perception subscale, correlated with overall accuracy (% correct) of the time (Spearman's $\rho=-0.02$; Spearman's $\rho=0.29$, respectively, ns) or colour (Spearman's $\rho=-0.30$; Spearman's $\rho=-0.06$ respectively, ns) discrimination task. However, the overall time accuracy score incorporates errors of both over- and underestimates of probe duration. Prior reports of biases to either over- (Johnson and Petzel 1971; Tysk 1983) or under- (Rabin 1957; Elvevåg et al. 2003) estimation of duration in patients with schizophrenia prompted us to explore our data further for potential links between the subjective experience of time and the direction of objective temporal error (i.e. proportion of over- or underestimates). We

found a significant correlation between scores on the CADSS time perception subscale and the proportion of overestimates (Spearman's $\rho=-0.60, p<0.05$), though not underestimates (Spearman's $\rho=0.25, ns$) in the time discrimination task.

Discussion

Administration of an acute dose of ketamine (100 ng/ml blood plasma level) selectively impaired perception of temporal, but not colour, attributes of visual stimuli in healthy volunteers. As compared to placebo, ketamine reduced accuracy of perceptual timing while having no effect on accuracy of a colour perception task that was matched to the timing task for accessory cognitive processes, such as visual processing, sustained attention and working memory (WM). This is the first time that selective effects of ketamine on a perceptual timing task have been shown. Despite our relatively small sample size ($n=12$), the selectivity of the results, demonstrated by a significant task by drug interaction, is striking. Although the colour task was more difficult than the timing task, the lack of drug effect on the colour task is unlikely to merely reflect a floor effect since participants' performance in both sessions (~70% accuracy) was well above chance (33%), allowing sufficient margin for further deterioration in performance. Selective impairment on performance of the timing task was reflected in participants' subjective assessment of time. Ketamine significantly altered the inner experience of time and drug-induced changes in subjective ratings correlated with objective measures of duration overestimation in the perceptual timing task.

Further evidence of timing disturbances were noted during performance of a cued reaction time (RT) task with variable cue-target intervals. In this task, RTs are generally faster for targets appearing at long, rather than short, intervals due to the increasing conditional probability of target appearance over time (Niemi and Näätänen 1981). However, ketamine slowed RTs selectively at the long interval, thereby attenuating the increase

Table 2 Mean (\pm standard error) scores on the CADSS index of dissociative experience

	CADSS overall		CADSS time perception	
	K/P	P/K	K/P	P/K
Placebo	0.04 (\pm 0.04)	0.10 (\pm 0.06)	0	0.28 (\pm 0.18)
Ketamine	1.06 (\pm 0.39)	0.32 (\pm 0.13)	1.28 (\pm 0.40)	0.39 (\pm 0.18)

Higher scores represent a greater sense of dissociation and disturbed perception. Overall CADSS score represents the average score of all 27 questions, whereas the time perception subscale is calculated as the mean of the three questions relating specifically to the subjective experience of time. Scores are shown as a function of whether the participant received ketamine in the first session and placebo in the second (K/P) or vice versa (P/K)

in speed that is normally seen at long delays. By contrast, ketamine had no significant effect on RT when temporal pre-cues could be used to predict in advance when the target would appear. Similarly, ketamine had no effect on the ability to use spatial cues to predict target location, consistent with previous results in schizophrenic patients (Carter et al. 1992).

Are effects of ketamine on timing due to WM deficits?

Rammsayer (2006) concluded that effects of the NMDA receptor antagonist memantine on timing in the seconds (~1 s.) but not tens of milliseconds (~50 ms) range were due to a primary effect on the additional WM processes that are needed to process seconds-range durations. We aimed to disentangle drug effects on timing from effects on WM by examining the effects of ketamine on two tasks that were matched for WM demands, in terms of both manipulation and maintenance in WM. During the 500–1,500 ms stimulus presentation time in both tasks, subjects had to maintain attention and integrate (i.e. manipulate) either duration or colour information in WM: for the timing task, duration estimates had to be updated incrementally throughout stimulus presentation; for the colour task, colour estimates had to be averaged across stimulus presentation. Both tasks also required that a sample stimulus be stored (i.e. maintained) in WM for a delay of 2–4 s for comparison with a later probe, and for the resulting decision to be maintained in WM until appearance of the response screen. Since mnemonic (and attentional) load were thereby matched across tasks, selective effects of ketamine on timing, but not colour, discrimination, suggests that ketamine-induced impairments do not simply reflect a functional side effect of the drug on the WM processes that are necessary for processing seconds range rather than millisecond timing.

However, one difference between the timing and colour task was the specific way in which information was manipulated in WM. For the timing task, information was incrementally accumulated, whereas for the colour task, it was averaged. Accumulation implies a unidirectionality, a fundamental feature of the flow of time itself ('time's arrow', Eddington 1928). Averaging does not imply this unidirectionality. It is possible therefore, that ketamine influences timing behaviour by selectively impairing the ability to manipulate information in WM in a particular direction or order. A related possibility is that subjects were accumulating not time, but a numerical count of the flickering stimulus. However, this is unlikely given that (1) each flicker was presented for a duration of only 80 ms necessitating an abnormally fast rate of counting and (2) the vast majority of our trials contained at least one stimulus whose duration was below the critical 1,200 ms value at

which counting begins to become a useful strategy (Grondin et al. 1999).

Temporal expectations and the flow of time

As expected, both spatial (Posner et al. 1980) and temporal (Coull and Nobre 1998) cues speeded RTs when targets appeared at expected, rather than unexpected, locations or delays, respectively. These RT effects were not modulated by ketamine, in either the spatial or temporal domain. Although the glutamatergic system has previously been implicated in signaling mismatch between expected and actual outcome (also known as the prediction error) (Corlett et al. 2007a; 2011), we found no evidence in this study for a selective effect of ketamine on RTs to unexpected target locations or onsets. This null effect could represent a lack of effect of ketamine on the processing of unexpected events. Alternatively, it may represent the insensitivity of our behavioural measures: the behavioural consequences of prediction error are rather elusive, and it has previously proven easier to demonstrate the underlying neural responses to prediction error, and their modulation by ketamine, using, for example electrophysiological recordings (e.g. Schultz 1998; Umbricht et al. 2002) or fMRI (e.g. McClure et al. 2003; O'Doherty et al. 2003; Corlett et al. 2006). Future studies examining the effects of ketamine on neural responses to unexpected locations or onsets could help resolve this issue. A final possibility is that these null effects simply reflect a lack of power in the data. However, the predicted behavioural advantages of valid versus invalid cues were clearly present in both the temporal and spatial conditions, itself suggestive of sufficient power. Moreover, although ketamine had no effect on speed of responding in the temporal or spatial conditions, it selectively slowed RTs to long delay targets in the neutral condition. Together, these patterns of data suggest a selectivity of effect rather than insufficient power.

In the orienting task, temporal cues allow subjects to predict event onset, thus generating temporal expectations that help speed performance (Coull and Nobre 1998; Griffin et al. 2001). Yet, even when the onset of the imminent event cannot be predicted by arbitrary, pre-learned cues, the unidirectional flow of time itself (Eddington 1928) provides a degree of predictive power via the hazard function. In our study, the neutral cue condition of the orienting task, in which participants did not know in advance, whether the target would appear at the short or long delay, provides an index of the hazard function. We observed the expected speeding of RTs for targets appearing at long versus short delays following administration of placebo. Ketamine, however, significantly attenuated this increase in response speed (Fig. 4), slowing RTs selectively at the longer delay. Previous studies using a fixed foreperiod paradigm have also

reported ketamine-induced slowing selectively at long delays (Guillemain et al. 2001; Micallef et al. 2004). This was interpreted as reflecting an effect of ketamine on motor preparation processes due to particularly impaired temporal resolution at long delays. However, this is an unlikely explanation for our own results since ketamine did not slow RTs at long intervals when participants could use arbitrary temporal cues to predict in advance when the target would appear (i.e. the temporal cue condition of the orienting task), suggesting they were perfectly able to estimate and prepare for the long interval as long as it had been cued beforehand. Ketamine did not therefore affect the ability to use prelearned cues to predict in advance when the target would occur but, instead, interfered with the ability to use the very passage of time itself to predict target occurrence when no other sources of temporal information were available. Ketamine effects have previously been interpreted within a Bayesian framework (Corlett et al. 2009; 2011), in which information processing is contingent upon prior expectations. In Bayesian terms, our data show that ketamine does not disrupt the ability to make use of prior expectations to interpret sensory events, but does interfere with the ability to generate expectations online as a function of sensory evidence accumulated over time.

Relevance to schizophrenia

Temporal predictability in the temporal cue condition of the orienting task is driven by external, visual cues. Conversely, temporal predictability in the neutral cue condition, where visual cues are uninformative, depends upon an internal sense of the passage of time. After administration of ketamine, participants made good use of external visual cues to make temporal predictions, but were less able to use internal estimates of time to make predictions. This psychopharmacological dissociation is reminiscent of findings in patients with schizophrenia who rely more upon external sensory stimuli to interpret the consequences of their actions, rather than internal sensorimotor processes (Synofzik et al. 2010; Voss et al. 2010). Moreover, while the temporal cue condition requires straightforward maintenance of a fixed a priori temporal prediction in WM, the neutral cue condition requires updating (i.e. manipulation) of the prediction as a function of elapsing time. Notably, both ketamine and schizophrenia have previously been associated with deficits in manipulation, rather than maintenance in WM (Fletcher and Honey 2006). Our orienting data, taken together with the results on the temporal discrimination task discussed above, support the proposal that ketamine interferes with the way in which the unidirectional flow of time is integrated within WM. Intriguingly, disturbances in the temporal flow (past–present–future) of causal experience have previously been

suggested to underpin certain symptoms of schizophrenia (Fuchs 2007). The timing deficits observed in schizophrenia may therefore be more accurately characterised as impairments in temporal flow, such that the beginning, middle and end of the event to be timed are not necessarily processed (or retained in WM) in the correct order, thus compromising duration estimates. Further experiments are clearly needed to test this hypothesis more directly.

Patients with schizophrenia have previously been reported to both over- (Johnson and Petzel 1971; Tysk 1983) and underestimate (Rabin 1957; Elvevåg et al. 2003) stimulus duration. We therefore examined whether ketamine produced a consistent temporal error in one direction or the other by measuring the incidence of over- or underestimation of probe duration. Overall, ketamine induced both types of error, with no significant bias in one direction or the other (see also Cheng et al. 2006). However, a particularly high incidence of drug-induced overestimation was evident whenever the probe stimulus was shorter than the sample. This result is unlikely to reflect a ketamine-induced increase in speed of an internal clock (Gibbon et al. 1984), because any change in clock speed would apply equally to both sample and probe stimuli. Instead, ketamine appears to have either selectively elongated perception of the current (probe) event, or contracted the memorized representation of the sample. This issue could be addressed by future experiments in which the sample duration is encoded either before or after administration of ketamine (see Malapani et al. (2002) for an example of this approach in patients with Parkinson's disease).

Finally, patients with schizophrenia (Carroll et al. 2008) or individuals at high risk for schizophrenia (Penney et al. 2005) have previously been reported to show larger than normal differences in the relative accuracy of auditory versus visual timing, potentially due to a preferential impairment of timing in the auditory modality (Carroll et al. 2008). If ketamine can be considered a useful model of schizophrenia, then we may expect to replicate this 'modality effect' with ketamine and, furthermore, to predict that its effects on timing would be even more evident when tested in the auditory domain.

Possible neural substrates

We have shown that ketamine impairs accuracy on a temporal discrimination task that has previously been shown to engage Supplementary Motor Area, right prefrontal cortex and basal ganglia (Coull et al. 2004; 2008). Although ketamine had no effect on the ability to use visual cues to predict stimulus onset in the temporal orienting task, a process previously shown to engage left inferior parietal cortex and intraparietal sulcus (Coull and Nobre 1998; Coull et al. 2011), it did impair the ability to use the

flow of time to make temporal predictions (the hazard function), a process that has consistently been shown to depend upon right prefrontal cortex (Vallesi et al. 2007a, 2007b, 2009; Bueti et al. 2010). Taken together with data from the temporal discrimination task, we suggest that ketamine may impair timing by modulating right prefrontal cortex function. This hypothesis is supported by functional imaging data showing that ketamine modulates activity in prefrontal cortex, among other areas, during tasks of WM and executive function (Fu et al. 2005; Honey et al. 2008) or prediction error (Corlett et al. 2006).

These data might indicate therefore that timing depends upon glutamatergic function in prefrontal cortex. Alternatively, ketamine is known to increase prefrontal dopamine in rats (Bowers and Morton 1994; Verma and Moghaddam 1996), suggesting that its effect on timing may instead be a result of drug-induced modulation of prefrontal dopamine. Therefore, just as dysregulation of dopaminergic innervation of prefrontal cortex has been suggested as the common underlying mechanism for both schizophrenia and ketamine-induced impairments in WM (Fletcher and Honey 2006), it may also explain the timing deficits associated with these two states. Notably, the dopaminergic system has repeatedly been implicated in timing (Meck 1996), with converging evidence from the animal and clinical literatures (Coull et al. 2011). Yet dopaminergic manipulations provoke consistent changes in clock speed, with enhanced dopaminergic activity leading to overestimates of stimulus duration (Maricq and Church 1983; Meck 1983; Matell et al. 2004, 2006) and reduced activity inducing underestimates (Maricq and Church 1983; Meck 1983, 1986; MacDonald and Meck 2005) and results of the present study, as well as prior work in rats (Cheng et al. 2006, 2007a), suggest ketamine has no effect on clock speed. Instead, Cheng et al. (2007a, 2007b) suggest ketamine ‘unlocks’ the clock-speeding effects of dopaminergic agonists that are otherwise attenuated by extensive training. Specifically, although administration of dopaminergic agonists is known to speed the internal clock, extensive training on the task abolishes this effect (Cheng et al. 2007a, 2007b). The hypothesis is that as performance on the timing task becomes more and more automated through training, the dopaminergic system becomes less critical for timing, and instead, the glutamatergic system takes over to mediate a more habitual form of timing behaviour. If a glutamatergic antagonist, such as ketamine, is administered, then control reverts to the dopaminergic system, meaning dopaminergic agents will once more produce their clock-speeding effects. So why, in contrast to Cheng et al. (2006, 2007a), does ketamine modulate timing in our relatively naive participants? The answer may lie, once more, with the hazard function. Keeping track of the unidirectional flow of time in WM in order to time current

events, as well as to predict the onset of future events, is an overlearned, automated process that, as suggested above, appears to be subject to disruption by ketamine. Therefore, the results of this study appear not to reflect a dopamine-dependent clock-speed effect, but may instead indicate a glutamate-dependent modulation of the unidirectional nature of ‘time’s arrow’. Alternatively, in line with ketamine’s effects on the manipulation of information in WM (Honey et al. 2003, 2004), our results may reflect dopamine-dependent modulation of the way in which temporal information is integrated within WM. Future studies combining functional neuroimaging and psychopharmacological methodologies could ascertain more directly the role of prefrontal dopamine or glutamate in the effects of ketamine on timing or, more specifically, on the ability to integrate the unidirectional flow of time in WM.

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Appendix

The Clinician Administered Dissociative States Scales (CADSS, Bremner et al. 1998) comprises 19 self-report questions and 8 observer ratings, rated from 0 (not at all) to 4 (extremely), and incorporates five subscales (Krystal et al. 1994): body perception, environmental perception, feelings of unreality, memory impairment and time perception. The overall CADSS score (mean response to all questions) provides a global measure of dissociative experiences, while the time perception subscale (Krystal et al. 1994; 1998) indexes the subjective experience of time more specifically by averaging the three CADSS questions that relate directly to the perception of time, namely:

Q1: Do things seem to be moving in slow motion?

Q12: Does this experience seem to take much longer than you would have expected?

Q13: Do things seem to be happening very quickly, as if there is a lifetime in a moment?

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