Noradrenergic neuromodulation of human attention for emotional and neutral stimuli

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Abstract

Introduction Norepinephrine (NE) has a regulatory role in human attention. Objective To examine its role in emotional modulation of attention, we used an attentional blink (AB) paradigm, in the context of psychopharmacological manipulation, where targets were either emotional or neutral items. Results and discussion We report behavioural evidence that β-adrenergic blockade with propranolol impairs attention independent of target valence. Furthermore, this effect is centrally mediated as administration of the peripheral β-adrenergic antagonist nadolol did not impair attention. By contrast, increasing NE tone, using the selective NE reuptake inhibitor reboxetine, improves detection of emotional stimuli. Conclusion In line with theoretical and animal models, these findings provide human behavioural evidence that the adrenergic system has a modulatory influence on selective attention that in some instances depends on item valence.

Keyword Norepinephrine · Attentional blink paradigm · Locus coeruleus

Introduction

Norepinephrine (NE) is a monoaminergic neurotransmitter released by ascending projections of the locus coeruleus (LC). NE plays a critical role in regulating cortical function and is implicated in the pathophysiology of major psychiatric disorders, such as depression, schizophrenia, and more recently attention deficit disorder (Beane and Marrocco 2004; Pliszka et al. 1996). While the role of NE system in arousal is well established (Jouvet 1969; Robinson and Berridge 1993), recent data from animal models suggest a role in facilitating processing of relevant, or salient, information (Berridge and Waterhouse 2003; Yu and Dayan 2005). Electrophysiological studies in monkeys indicate that there are two modes by which LC activity modulates attention (Aston-Jones and Cohen 2005). Phasic LC neuronal activation is evoked by salient or goal-relevant stimuli during vigilance tasks, a response positively associated with outcomes in task-related decisions and coupled with highly accurate responses (Aston-Jones et al. 1994). Phasic responses occur against the background of, and are in turn modulated by, tonic LC activity that correlates with general arousal levels (Aston-Jones et al. 1991, 1994).

We have recently shown (Strange and Dolan 2007) that the cortical circuitry engaged by salient stimuli is inhibited by pharmacological blockade of the noradrenergic system. Specifically, we demonstrated a role for β-adrenergic receptors in this process. This differs from previous psychopharmacological experiments addressing NE modulation of attention, which have typically examined the effects of α2-adrenergic receptor agonists such as clonidine (Coul et al. 2001). Subjects receiving clonidine, compared to placebo, are impaired at target discrimination (Clark et al. 1986) but also report subjective withdrawal and difficulties with concentration. Clonidine acts on presynaptic α2-adrenergic auto-receptors to inhibit NE release; thus, clonidine-induced impaired attention may reflect decreased arousal or, due to presynaptic inhibition of NE release, decreased stimulation of predominantly postsynaptic α1-
β-adrenergic receptors. Given that the $\beta_1$$\beta_2$-adrenergic antagonist propranolol is non-sedating (Harmer et al. 2001), its administration enables investigation of the adrenergic role in attention without sedation and localising specific attentional effects to a specific receptor subtype.

To investigate the noradrenergic modulation of human attention, we used a modification of the RSVP attentional blink (AB) paradigm in which subjects view a rapid sequence of visual stimuli (RSVP; Raymond et al. 1992) with a task requirement being to identify embedded targets. A classical finding using this type of manipulation is that identifying a first target stimulus (T1) causes transient impairment in detecting a subsequently presented second target (T2), an effect reduced with increasing temporal lag between the two targets. This paradigm allowed us to investigate how emotional stimuli modulate the allocation of limited attentional resources in time. Thus, the present study was designed to investigate the time dynamic of the interaction between attention and emotion.

Of particular interest to the study of a adrenergic role in attentional control is the recent finding that the AB can be modulated by the emotional significance of stimuli. The crucial observation in this study is that an arousing T2 stimulus is detected more often than a neutral T2 (Anderson and Phelps 2001), an effect abolished by amygdala damage (Anderson and Phelps 2001). These observations parallel those seen during studies of human episodic memory where memory is enhanced for emotional events, an effect not seen in patients with amygdala lesions (Cahill et al. 1994; Strange et al. 2003). Enhanced emotional memory effect are abolished by propranolol (Cahill et al. 1994; Strange et al. 2003), which leads to an hypothesis that the emotional modulation of the AB would also be abolished by β-adrenergic blockade. However, in light of our finding (Strange and Dolan 2007) that propranolol modulates the cortical circuitry involved in salience detection, the effects of propranolol on the AB might be more complex, i.e., there may be a generic effect of β-adrenergic blockade on T2 detection.

Thus, within the current RSVP task, each trial consisted of 15 words, 2 targets (bright green) and 13 distractors (black), presented for 130 ms each and immediately followed by the subsequent stimulus (Fig. 1). In half the trials, the T2 stimuli consisted of arousing words (e.g. rape and incest) and in the other half T2 were neutral words (e.g. pepper and omit). The neutral and emotional stimuli were matched for average word length, word frequency and interletter frequency. Order or presentation of emotional and neutral trials was random. The temporal lag between the first target (T1) and the second target (T2) was variable. At early lag (<500 ms, SOA), the detection of target T2 is more susceptible to the attentional blink compared with late lag (>500 ms, SOA). The subjects’ task was to monitor the RSVP stream and report both coloured green targets (T1 and T2) by typing them at the end of a stimulus sequence.

![Fig. 1 Schematic diagram of the AB task. Each trial consisted of 15 words [2 targets (bright green) and 13 distractors (black)] each presented for 130 ms (Experiment 1) or 110 ms (Experiment 2) and immediately followed by the subsequent stimulus. In half the trials, the T2 stimuli consisted of an arousing word (e.g. rape and incest) and in the other half T2 were neutral words (e.g. pepper and omit). The neutral and emotional stimuli were matched for average word length, word frequency and interletter frequency. Order or presentation of emotional and neutral trials was random. The temporal lag between the first target (T1) and the second target (T2) was variable. At early lag (<500 ms, SOA), the detection of target T2 is more susceptible to the attentional blink compared with late lag (>500 ms, SOA). The subjects’ task was to monitor the RSVP stream and report both coloured green targets (T1 and T2) by typing them at the end of a stimulus sequence.](image)

Materials and methods

Ninety-six healthy right-handed native English-speaking subjects took part in our studies. All volunteers gave informed consent, were free of neurological, psychiatric and physical illness and had not been on any medication for >3 months. The study was approved by the National Hospital for Neurology and Neurosurgery and the Institute of Neurology Joint Research Ethics Committee.
Experiment 1

Subjects In a double-blind experimental design, 36 subjects [18 male, mean age (SD)=24.2 (3.9); 18 female, mean age=23.1 (4.0)] received either a 40-mg oral dose of propranolol hydrochloride (a β1 and β2 adrenergic receptor antagonist) or a 100 mg oral dose of ascorbic acid (placebo pill). Drug allocation, in this and subsequent experiments, was balanced for gender. In view of the kinetics of propranolol’s peak plasma concentration (1–2 h), the attentional blink task commenced 90 min after drug administration. Blood pressure (BP) was measured immediately before drug administration (time 0 min) and the attention task (time +90 min).

Stimuli We used a modification of the RSVP paradigm (Anderson and Phelps 2001; Raymond et al. 1992) where each trial consisted of 15 words [2 targets (bright green) and 13 distractors (black)] each presented for 130 ms and immediately followed by the subsequent stimulus (Fig. 1). Six trial lags were introduced from lag 2 [1 distractor presented between the two targets (T1–T2) SOA=260 ms] to lag 7 [6 distractors presented between the two targets (T1–T2) SOA=910 ms]. In half the trials, the T2 stimuli consisted of an arousing word (e.g. rape and incest), and in the other half, T2 were neutral words (e.g. pepper and omit). The neutral and emotional stimuli were matched for average word length, word frequency and inter-letter frequency. The presentation order of emotional and neutral trials was random. The subjects’ task was to monitor the RSVP stream and report both coloured green targets (T1 and T2) by typing them at the end of a stimulus sequence (trials in which a T1 target was not reported correctly were discarded from the analysis of T2 effects).

Statistics Following the analysis of Anderson and Phelps (Anderson and Phelps 2001), data were segregated into early (lags 2–3, 260–390 ms) and late lag (lags 6–7, 780–910 ms). We performed a 2×2×2 drug (propranolol 40 mg, placebo) × valence of T2 stimulus (emotional, neutral) × lag (early vs late) analysis of variance (ANOVA). In this and all subsequent experiments, ANOVAs were followed by post-hoc two sample unequal variance t tests to test for simple effects.

Experiment 2

Subjects In a double-blind experimental design, 30 subjects [15 male, mean age (SD)=24.7 (2.8); 15 female, mean age=23.3 (3.1)] were allocated to one of three equal sized groups and received either a 40-mg oral dose of propranolol, a 4-mg oral dose of reboxetine methansulphonate (a selective norepinephrine reuptake inhibitor) or a 100-mg oral dose of ascorbic acid (placebo pill). In view of the kinetics of propranolol and reboxetine’s peak plasma concentration (1–2 and 1.5 h, respectively), the attentional blink task commenced 120 min after drug administration. One subject in the propranolol group was excluded from further analysis because behavioural performance was more than two standard deviations below the group average.

Stimuli The task used was identical to Experiment 1 except for two critical modifications; the time of the stimulus presentation was decreased from 130 ms in Experiment 1 to 110 ms, and targets were separated by six different time lags ranging from one to nine distractors presented between the two targets (lags 2–3–4–7–8–9) [e.g. lag 2 (T1–T2) SOA=220 ms; lag 9 (T1–T2) SOA=990 ms].

Statistics Data were collapsed into early (lags 2–3–4, 220–440 ms) and late lags (lags 7–8–9, 770–990 ms) and a drug (propranolol 40 mg, reboxetine, placebo) × T2 valence (emotional, neutral) × lag (early, late) 3×2×2 ANOVA performed. For each drug group, we also performed a 2×2 drug (drug, placebo) × T2 valence (emotional, neutral) ANOVA.

Experiment 3

Subjects In a double-blind experimental design, 30 subjects [15 male, mean age (SD)=25.2 (3.7); 15 female, mean age=25.5 (3.9)] were allocated to one of three equal size groups and received either a 20-mg oral dose of propranolol, a 40-mg oral dose of nadolol (a β1 and β2 adrenergic receptor antagonist that does not cross the BBB) or a 100 mg oral dose of ascorbic acid (placebo pill). In view of the kinetics of propranolol and nadolol peak plasma concentration (1–2 h), the attentional blink task commenced 120 min after drug administration. One subject in the propranolol group was excluded from further analysis because performance was more than two standard deviations below the group average (task outlier). Three subjects in the placebo group were excluded (two on the grounds of being task outliers and one on subsequent disclosure of not being a native English speaker).

Stimuli The task used was identical to Experiment 2 using the same equipment.

Statistics After exclusion criteria, the size of the placebo group was drastically reduced (seven subjects). Thus, to retain sufficient statistical power, we collapsed data from
the current placebo group with the placebo group from Experiment 2. Data were collapsed into early (lags 2–3–4, 220–440 ms) and late lags (lags 7–8–9, 770–990 ms) and separate analyses performed for each drug group (propanolol 20 mg, nadolol 40 mg) vs placebo, i.e. separate group (drug × placebo) × T2 valence (emotional, neutral) × lag (early, late) 2×2×2 ANOVAs.

Results

Experiment 1 This psychopharmacological experiment demonstrated that β-adrenergic blockade by propranolol significantly impaired detection of T2 targets independently of their emotional valence (i.e. emotional and neutral; see Fig. 2). In both placebo and drug groups, we observed enhanced reporting of emotional, relative to neutral, T2 stimuli and a proportional increase in detecting both types of T2 stimuli with increasing T1-T2 lag, in agreement with previous observations (Anderson and Phelps 2001). In the placebo group, the percentage of T2 correctly reported at early lag was for the neutral T2 (NT2) 86.99%±2.42 and for the emotional T2 (ET2) 93.93%±1.43. At late lag, these values increased to NT2=92.62%±1.28 and ET2=95.8%±1.13. Conversely, in the 40-mg propranolol group, the percentage of T2 correctly reported at early lag was NT2=80.69%±1.90 and ET2=91.26%±1.93; performance at late lag increased to NT2=88.18%±1.90 and ET2=93.84%±1.47.

A drug (propranolol, placebo) × T2 valence (neutral, emotional) × lag (early, late) 2×2×2 ANOVA yielded significant main effects of treatment \( [F(1,34)=4.98, P<0.05] \), T2-valence \( [F(1,34)=51.31, P<0.0001] \) and lag \( [F(1,34)=26.33, P<0.0001] \). In line with the observation of Anderson and Phelps that affective modulation of the attentional blink is most pronounced at early lags, we demonstrated a lag × T2-valence interaction significant at trend level \( [F(1,34)=3.72, P=0.062] \). Critically, there was no significant interaction between T2-valence and drug manipulation \( [F(1,34)=2.65, P=0.29] \) indicating that a propranolol effect was not influenced by T2 target valence (Fig. 2). The simple effect of group (placebo, propranolol) on T2 reporting was significant for neutral T2 \( [t(1,19)=2.04, P<0.05\text{ one-tailed}] \) but not for emotional T2 \( [t(1,19)=1.13, P=0.13] \).

Furthermore, no significant interaction was found between lags (early, late) and drug manipulation (propranolol, placebo) \( [F(1,34)=0.51, P=0.476] \) indicating that the effect of propranolol is present across all lags. Figure 2c shows
the percentage (%±SE) of correctly reported emotional and neutral T2 for early (260–390 ms, SOA) and late lags (780–910 ms, SOA). Importantly, performance on T1 target detection was not statistically different between placebo (96.27%) and propranolol (96.18%) groups \( [t(1,19) = 0.17, P > 0.05 \) one-tailed independent test].

As seen in Fig. 2, performance in both placebo and drug groups at late lags approached ceiling levels, which may have obscured differential effects at these later lags. Thus, in Experiment 2, we reduced the interstimulus interval, making T2 detection more difficult. Removing this ceiling effect enabled us to test the additional hypothesis that increasing NE levels, with the SNRI reboxetine, would enhance T2 detection.

Finally, the mean BP at the time of placebo/drug (0 min) administration for the placebo group was 90 mmHg and at the start of the experiment (+90 min) was 88.5 mmHg \( [t(1,17) = 0.95, P = 0.35 \) two-tailed paired t test]. For the propranol group at 0 min was 86.2 mmHg and showed a significant reduction at (+90 min) 81.9 mmHg \( [t(1,17) = 2.16, P < 0.05 \) two-tailed paired t test].

**Experiment 2** Three different drug groups (placebo, propranolol and reboxetine) performed an attentional blink task identical to Experiment 1 except for a reduction in stimulus presentation time from 130 to 110 ms and a subtle change in lag range. This experiment confirms the previous finding from Experiment 1 of a propranol-induced impaired detection of T2 targets, independently of whether these are emotional or neutral stimuli. It also demonstrates that reboxetine selectively boosts the detection of emotional T2 with no effect on the neutral T2 (Fig. 3). In the placebo group, the percentage of T2 correctly reported at early lag was NT2 = 64.02%±4.67 and ET2 = 74.26%±5.65, and at late lag NT2 = 77.87%±4.56 and ET2 = 84.5%±4.99. In the 40 mg propranolol group, the percentage of T2 correctly reported at early lag was NT2 = 46.1%±5.28 and ET2 = 61.6%±5.93 conversely at late lag NT2 = 62.68%±7.69 and ET2 = 70.48%±7.62. In the reboxetine group, the percentage of T2 correctly reported at early lag was NT2 = 62.8%±3.65 and ET2 = 82.64%±4.18 at late lag NT2 = 80.95%±2.05 and ET2 = 90.23%±1.92.

Figure 3c shows the percentage (%±SE) of correctly reported emotional (E) and neutral (N) T2 for early (220–440 ms, SOA) and late lags (770–990 ms, SOA). A drug (placebo, propranolol, reboxetine) × T2 valence (neutral, emotional) × lag (early, late) 3×2×2 ANOVA yielded a significant main effect of drug \( F(1,27) = 5.44, P < 0.01 \), T2-valence \( F(1,27) = 73.8, P < 0.0001 \) and lag \( F(1,27) = 93.54, P < 0.0001 \). The three-way interaction of drug × T2-valence × lag was also significant \( F(2,27) = 3.46, P < 0.05 \).

To investigate the effects of drug relative to placebo, we performed two further 2×2×2 ANOVAs. The effect of propranolol in a drug (placebo, propranolol) × T2-valence (neutral, emotional) × lag (early, late) 2×2×2 ANOVA yielded a significant main effect of drug manipulation \( F(1,18) = 4.58, P < 0.05 \), T2-valence \( F(1,18) = 24.87, P < 0.0001 \) and lag \( F(1,18) = 86.51, P < 0.0001 \). The interaction between T2-valence and drug manipulation was not significant \( F(1,18) = 0.62, P = 0.44 \). The simple effects of group (placebo, propranolol) on T2 detection was significant for neutral T2 \( [t(1,19) = 2.53, P < 0.01 \) and at trend level for emotional T2 \( [t(1,19) = 1.53, P = 0.07 \). These data replicate the results of Experiment 1, confirming that propranolol impairs T2 detection independently of T2 emotional valence. As in Experiment 1, the degree of affective modulation was most pronounced at early lags, evident in a significant interaction of lag × T2-valence \( F(1,18) = 6.59, P < 0.05 \). Again, no significant interaction was found between lags (early, late) and drug manipulation (propranolol, placebo) \( F(1,18) = 0.05, P = 0.862 \). Finally, in this more difficult version of our task, T1 detection was impaired in the propranolol group relative to placebo \( [t(1,19) = 1.84, P < 0.05 \) one-tailed independent-samples t test].

By contrast, reboxetine evoked a selective enhancement in emotional T2 detection. A drug (placebo, reboxetine) × T2 valence (neutral, emotional) × lag (early, late) 2×2×2 ANOVA yielded a significant main effect of T2 valence \( F(1,18) = 107.89, P < 0.0001 \) and lag \( F(1,18) = 54.39, P < 0.0001 \), but the main effect of reboxetine relative to placebo was not significant \( F(1,18) = 0.57, P = 0.46 \). However, the interaction between T2-valence and drug manipulation was significant \( F(1,18) = 10.61, P < 0.005 \). The simple effect of group (placebo, reboxetine) on T2 detection was not significant for neutral T2 \( [t(1,19) = 0.44, P = 0.66 \) and was at trend level for emotional T2 \( [t(1,19) = 1.54, P = 0.07 \). These data demonstrate that enhanced NE levels selectively improves detection of an emotional T2 stimulus with no effect on neutral T2. Again, detection of emotional relative to neutral T2 stimuli was greatest at early lags for both groups indexed by a significant lag × T2-valence \( F(1,18) = 22.8, P < 0.001 \) interaction. By contrast, a lag (early, late) × drug manipulation (reboxetine, placebo) interaction was not significant \( F(1,18) = 0.036, P = 0.852 \), indicating that the behavioural effect of reboxetine is present across all lags \( [t(1,17) = 0.95, P = 0.35 \) two-tailed paired t test].

The mean BP at the time of placebo/drug (0 min) administration for the placebo group was 90.9 mmHg and showed a significant reduction at (+90 min) 82.9 mmHg \( [t(1,19) = 2.79, P < 0.05 \) two-tailed paired t test]. For the reboxetine group at (0 min) was 91.3 mmHg and showed a non-significant reduction at (+90 min) 90.4 mmHg.
Finally, in the propranolol group, the mean BP at (0 min) was 85.1 and showed a significant reduction at (+90 min) 78.9 mmHg ($t_{(1,9)}=3.97$, $P<0.005$ two-tailed paired $t$ test).

Experiment 3 To further investigate the effects of propranolol, we conducted a third experiment in which three different drug groups (placebo, nadolol 40 mg and propranolol 20 mg) performed the AB task identical to Experiment 2. The first drug group received 40 mg of nadolol, a beta blocker with the same mechanism of action of propranolol (i.e. selective blockers of $\beta_1$ and $\beta_2$ adrenergic receptors) but that critically does not pass the BBB (Hardman et al. 2001). This manipulation controlled for the potential contribution of peripheral $\beta$-blockade to the attentional effects observed with propranolol. The second group received a dose of propranolol (20 mg) that was half that used in Experiments 1 and 2. This treatment was aimed at determining whether a lower dose of propranolol would result in an attentional effect that was specific for target valence. As shown in Fig. 4, correct T2 detection at early lag in the placebo group was NT2=59.19%±3.44 and ET2=75.32%±3.58, and at late lag NT2=76.45%±3.13 and ET2=86.68%±3.21. In the 20 mg propranolol group the percentage of T2 correctly reported at early lag was NT2=65.40%±6.40 and ET2=78.20%±6.31, whereas at late lag NT2=80.43%±5.02 and ET2=87.62%±4.27. In the 40 mg nadolol group, the percentage of T2 correctly reported at early lag was NT2=62.24%±4.76 and ET2=76.82%±5.57, and at late lag NT2=76.59%±4.08 and ET2=88.07%±3.69.

Thus, T2 detection did not differ between nadolol and placebo groups (Fig. 4). The drug (placebo, nadolol) × T2 arousal. Conversely, reboxetine significantly enhances detection of emotional T2 with no effect on neutral T2. D T2 target detection in placebo (solid lines), propranolol (small dashed lines) and reboxetine (large dashed lines) groups on early lag (220–440 ms, SOA), when the attentional demand is higher, compared with late lags (770–990 ms, SOA) for emotional (red lines) and neutral (blue lines) T2 stimuli.
valence (neutral, emotional) × lag (early, late) 2×2×2 ANOVA yielded a significant main effect of T2 valence \( F(1,25)=89.16, P<0.0001 \) and lag \( F(1,25)=12.45, P<0.005 \), but no significant main effect of drug relative to placebo \( F(1,25)=0.06, P=0.80 \) or drug × T2-valence interaction \( F(1,25)=1.48, P=0.23 \) (Fig. 4). These results confirm that the modulatory effects of propranolol observed in Experiments 1 and 2 are due to central, and not peripheral, effects.

Performance in the low dose propranolol group (20 mg) was equivalent to that of the placebo group. The drug (placebo, propranolol low dose 20 mg) × T2 valence (neutral, emotional) × lag (early, late) 2×2×2 ANOVA yielded a significant main effect of T2 valence \( F(1,24)=49.33, P<0.0001 \) and lag \( F(1,24)=68.99, P<0.0001 \). Although the low dose propranolol group (20 mg) performed the task slightly better than the placebo group, this difference was not statistically significant [main effect of drug relative to placebo \( F(1,24)=0.42, P=0.52 \)]. Critically, there was no significant interaction between drug and T2-arousal \( F(1,24)=1.08, P=0.31 \) (Fig. 4). Thus, only the higher propranolol dose (40 mg) impairs target detection, an effect that is independent of target arousal content. A lower propranolol dose does not impair detection of neutral or emotional T2 detection.

![Fig. 4 Behavioural results of Experimental 3. Subpanels a–c show the percentage of T2 stimuli that are correctly reported for each T1–T2 temporal lag respectively in the placebo, nadolol and propranolol low dose (20 mg) groups. In all groups, T2 detection improved with increasing temporal lags and is significantly enhanced for emotional (red squares) relative to neutral (blue diamonds) T2 detection. Both propranolol 20 mg (b) and nadolol 40 mg (c) do not show significant differences from the placebo group. d T2 target detection in placebo (solid lines), nadolol 40 mg (small dashed lines) and propranolol 20 mg (large dashed lines) groups on early lag (220–440 ms, SOA), when the attentional demand is higher, compared with late lags (770–990 ms, SOA) for emotional (red lines) and neutral (blue lines) T2 stimuli.

The mean BP at the time of placebo/drug (0 min) administration was for the placebo group was 90.5 mmHg and showed a non-significant reduction at (+90 min) 86.0 mmHg \( t(1,9)=1.8, P=0.102 \) two-tailed paired \( t \) test. For the nadolol group at (0 min) was 90 mmHg and showed a non-significant reduction at (+90 min) 88.6 mmHg \( t(1,9)=0.4, P=0.69 \) two-tailed paired \( t \) test. Finally, in the propranolol group, the mean BP at (0 min) was 92.3 and showed a significant reduction at (+90 min) 85.8 mmHg \( t(1,9)=2.79, P<0.05 \) two-tailed paired \( t \) test.

Discussion

The critical new observation we report is that \( \beta \)-adrenergic blockade by propranolol (40 mg) impairs detection of T2 stimuli regardless of the emotional arousal of these stimuli.
(Experiment 1). This is the first demonstration that a manipulation of the adrenergic system affects the AB. We note in this regard that a recent study found no effect of clonidine on neutral T2 detection (Nieuwenhuis et al. 2007). The robustness of our reported β-adrenergic effect is evident in a replication within Experiment 2. Furthermore, these data also replicate previous observations regarding emotional modulation of the attentional blink (Anderson and Phelps 2001). Specifically, placebo and drug groups, in Experiments 1 and 2, showed enhanced T2 detection with increasing T1–T2 lag for both emotional and neutral T2, while emotional T2 stimuli were more frequently detected than neutral T2, an effect more pronounced at early lags.

A recent comprehensive account of the role of NE in attention is derived from monkey electrophysiological experiments, which indicate two modes of LC activity (Aston-Jones and Cohen 2005). Phasic LC neuronal activation is evoked by salient or goal-relevant stimuli during vigilance tasks (Aston-Jones et al. 1994). Phasic responses are evoked against the background of, and in turn modulated by, tonic LC activity that correlates with general arousal levels (Aston-Jones et al. 1991; Aston-Jones et al. 1994). Whereas the LC phasic response itself is relatively brief in duration (typically of 50–100 ms), the ensuing neuromodulatory effects of NE on target cortical areas are known to be sustained relative to the LC phasic response.

One recent hypothesis regarding the neurobiological underpinnings of the attentional blink suggests that it is a product of the temporal dynamics of the LC–NE system (Nieuwenhuis et al. 2005). Although NE may potentiate processing in cortical areas, local NE release within LC is thought to be auto-inhibitory, due to effects of NE at presynaptic and dendritic α2 autoreceptors (Egan et al. 1983; Washburn and Moses 1989; Williams et al. 1985). This autoinhibition results in a refractory-like period after a LC phasic response, during which a subsequent LC phasic discharge is rarely observed (Aston-Jones et al. 1994; Usher et al. 1999). This refractoriness peaks at approximately 50–100 ms after the LC phasic response, typically 200–250 ms after the eliciting stimulus, and usually lasts 200 ms or until about 400–450 ms post-stimulus. The length of the refractory period coincides with the T1 and T2 temporal lag in which the blink for the second stimulus T2 is most marked (Nieuwenhuis et al. 2005). Thus, this formulation predicts an effect of NE modulation on both emotional and neutral T2 detection.

Our behavioural effects can be explained via an extension of the aforementioned AB–NE model (Nieuwenhuis et al. 2005) in which we also manipulated the salience level of the T2 target (arousing or neutral). Previous studies demonstrate that arousing stimuli presented in different modalities induce a robust phasic discharge of NE neurons in LC (Aston-Jones and Bloom 1981; Foote and Morrison 1987). Thus, the enhanced detection of emotional relative to neutral T2 stimuli, demonstrated by Anderson and Phelps (Anderson and Phelps 2001) and replicated here in Experiments 1 and 2, can be attributable to an increase in phasic NE release elicited by emotional, relative to neutral, stimuli. Propranolol has been demonstrated to reduce the phasic response elicited by novel stimuli in rats (Kitchigina et al. 1997). The effects we report after propranolol administration may consequently be explained in terms of a reduced impact of LC phasic response to targets. A critical observation in our data is that propranolol-treated subjects show impaired T2 detection, relative to placebo, for both neutral and emotional T2. Importantly, emotional T2 items, eliciting an increase in NE release, are still detected more frequently than neutral T2 even in the presence of propranolol. According to the model we propose, the behavioral effect in target detection is relative to the magnitude of NE release, assumed to be larger for an arousing T2. Note that our task required subjects to report both T1 and T2 stimuli correctly. The NE–AB hypothesis (Nieuwenhuis et al. 2005) states that initial T1 detection is mediated by phasic LC activity, with the subsequent LC refractory period leading to impaired T2 detection. If propranolol inhibits phasic LC activity, it follows that T1 detection should also be impaired by beta-blockade. In Experiment 2, in which an increase in task difficulty prevented T1 detection ceiling effects, we indeed showed a significant impairment for T1 detection for the propranolol compared with the placebo group. Previous evidence (Jolicoeur 1999) provides an interpretation that the increased difficulty in T1 detection under propranolol may also contribute to the increased AB magnitude that is independent of T2 emotionality, as demonstrated in this study.

One of the aims of Experiment 3 was to determine whether a lower dose (20 mg) of propranolol would yield arousal-specific effects on T2 detection. In other words, we tested whether less β-adrenergic blockade could be sufficient to abolish enhanced detection of emotional T2 targets, without affecting neutral T2 detection. We did not, however, observe any difference in T2 detection for either neutral or emotional targets between the propranolol 20 mg group and placebo. Experiment 3 also served to demonstrate that administration of nadolol, a peripherally acting β-adrenergic antagonist, has no effect on T2 detection, thus indicating that the attentional effects of propranolol are centrally mediated.

In addition to demonstrating attentional impairment evoked by blocking the β-adrenergic system, administration of a SNRI in Experiment 2 also enabled examination of the attentional effects of increased NE concentration at the
synaptic level. We provide evidence for reboxetine-evoked enhanced attention, which, in contradistinction to the impairment observed with propranolol, is dependent on the emotional salience of the target. That reboxetine failed to improve neutral T2 detection accords with previous data showing no effect of reboxetine on a neutral continuous performance attention task (Plewnia et al. 2006).

Reboxetine-induced enhanced emotional T2 detection can potentially be explained in terms of the NE–AB hypothesis (Nieuwenhuis et al. 2005). Previous studies demonstrate increased concentration of cortical NE, particularly in the frontal cortex, after acute reboxetine administration (Saccomi et al. 1999). By contrast, in the LC, NE reuptake inhibitors attenuate the firing activity of LC–NE neurons via α2-adrenergic autoreceptor overactivation (Beique et al. 2000; Kasamo et al. 1996; Lacroix et al. 1991). Thus, SNRIs mediate two opposite effects: at a cortical level, they potentiate attentional gain; but at the level of LC, SNRIs increase the refractory period via presynaptic α2 receptors. These two NE influences could potentially produce opponent effects on T2 detection involving facilitation and impairment in the AB task. In Experiment 2, we found that reboxetine selectively boosted detection of emotional T2 stimuli with no effect on neutral T2. These findings can be reconciled with the proposed model if, for neutral T2 detection, the facilitatory and impairing effects of reboxetine are in relative equilibrium, resulting in no net effect on neutral target detection. Conversely, the boost in NE elicited by the arousing T2 positively interacts with this state of equilibrium thereby significantly improving target detection for emotional items.

Enhanced processing of emotional stimuli might also, in theory, result in a lowered perceptual threshold for the arousing stimuli that arises epigenetically (Weinberger 1999). This framework predicts that propranolol would not have a selective effect on the perception of emotional stimuli, as the lower threshold for processing arousing stimuli is ‘hard-wired’ and therefore not dependent on phasic release of noradrenaline. This model could explain the findings from Experiments 1 and 2 showing that propranolol does not have a selective effect on emotional T2 processing.

Our results show a dissociation in the effects of propranolol on attention from effects previously reported on episodic memory (Strange et al. 2003). Whereas the effect of propranolol is selective for the emotional modulation of memory, we demonstrate for the first time in humans (Experiments 1 and 2) that engagement of β-adrenergic receptors in the central nervous system is essential for optimal target detection independent of stimulus arousal content. Moreover, we demonstrate that the increase in NE levels (Experiment 2) selectively improves detection of arousing stimuli. These attentional effects of pharmacological manipulation of the NE system in human subjects support and extend current theoretical models of the role of NE in attention.

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