Baseline reward processing and ventrostriatal dopamine function are associated with pramipexole response in depression

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The efficacy of dopamine agonists in treating major depressive disorder has been hypothesized to stem from effects on ventrostriatal dopamine and reward function. However, an important question is whether dopamine agonists are most beneficial for patients with reward-based deficits. This study evaluated whether measures of reward processing and ventrostriatal dopamine function predicted response to the dopamine agonist, pramipexole (ClinicalTrials.gov Identifier: NCT02033369). Individuals with major depressive disorder (n = 26) and healthy controls (n = 26) (mean ± SD age = 26.5 ± 5.9; 50% female) first underwent assessments of reward learning behaviour and ventrostriatal prediction error signalling (measured using functional MRI). 11C- (+)-PHNO PET before and after oral amphetamine was used to assess ventrostriatal dopamine release. The depressed group then received open-label pramipexole treatment for 6 weeks (0.5 mg/day titrated to a maximum daily dose of 2.5 mg). Symptoms were assessed weekly, and reward learning was reassessed post-treatment. At baseline, relative to controls, the depressed group showed lower reward learning (P = 0.02), a trend towards blunted reward-related prediction error signals (P = 0.07), and a trend towards increased amphetamine-induced dopamine release (P = 0.07). Despite symptom improvements following pramipexole (Cohen’s d ranging from 0.51 to 2.16 across symptom subscales), reward learning did not change after treatment. At a group level, baseline reward learning (P = 0.001) and prediction error signalling (P = 0.004) were both associated with symptom improvement, albeit in a direction opposite to initial predictions: patients with stronger pretreatment reward learning and reward-related prediction error signalling improved most. Baseline D2/3 receptor availability (P = 0.02) and dopamine release (P = 0.05) also predicted improvements in clinical functioning, with lower D2/3 receptor availability and lower dopamine release predicting greater improvements. Although these findings await replication, they suggest that measures of reward-related mesolimbic dopamine function may hold promise for identifying depressed individuals likely to respond favourably to dopaminergic pharmacotherapy.
Introduction

Although several treatments are available for major depressive disorder, response rates are modest and highly varied. Half of patients fail to respond to first-line antidepressants (Levkovitz et al., 2011), and there are no consistently replicated, clinically meaningful predictors of response to specific classes of antidepressant medications. Finding ways to tailor treatment to a given individual is therefore an important step towards reducing the global burden of depression.

One approach is to subtype patients based on symptoms associated with specific underlying neurobiological features, to which personalized treatments can be directed. A promising target for subtyping depression is motivational disturbance, particularly anhedonia, which has been linked to poorer response to selective serotonin reuptake inhibitor treatment (McMakin et al., 2012), psychotherapy (McMakin et al., 2012), and transcranial magnetic stimulation (Downar et al., 2014), suggesting that anhedonic individuals may require alternative treatment approaches. Translational research has linked the reward and motivation-related deficits that characterize anhedonia to mesolimbic dopamine system dysfunction (Berridge and Kringelbach, 2015). For example, manipulating phasic dopamine neuron firing in the ventral tegmental area, which projects to the ventral striatum, alters anhedonic behaviour in rodents (Chaudhury et al., 2013). In psychiatrically healthy humans, PET imaging has shown that blunted ventrostriatal dopamine release is associated with decreased motivation to work for rewards (Treadway et al., 2012). Furthermore, ventrostriatal deep brain stimulation has been found to reduce anhedonia severity (Bewernick et al., 2010). Collectively, these findings suggest that for a subset of depressed individuals with prominent anhedonia, a treatment that specifically targets ventrostriatal dopamine may be warranted. However, to achieve this level of treatment precision, valid indicators of anhedonia-related ventrostriatal dopamine dysfunction are required.

Reward learning is a measure that correlates with mesolimbic dopamine function (Steinberg et al., 2013) and may be useful for identifying individuals likely to benefit from dopaminergic pharmacotherapy. It is the process by which behaviour is updated based on prior reinforcement, and is guided by phasic dopamine neuron firing that encodes differences between anticipated and actual rewards, known as reward prediction errors (Glimcher, 2011). Reward learning is impaired in major depressive disorder, particularly among anhedonic individuals (Pizzagalli et al., 2008b; Fletcher et al., 2015). Similarly, individuals with depression display blunted prediction error signals to reward in the ventral striatum (Kumar et al., 2008, 2018; but see Rutledge et al., 2017) and the extent of this blunting correlates with anhedonia (Greenberg et al., 2015). Further support for the importance of phasic dopamine firing in reward learning comes from studies showing that pharmacological challenges assumed to reduce phasic dopamine signalling disrupt reward learning (Pizzagalli et al., 2008a), whereas administering drugs that enhance striatal dopamine signalling improves reward learning (Der-Avakian et al., 2013; Pergadia et al., 2014). Together, these findings suggest that reward learning and prediction error signalling are both closely linked to ventrostriatal dopamine function, and may be useful for identifying depressed individuals who would benefit from a dopamine-targeting medication.

Pramipexole is a high-affinity D2 receptor agonist that may be suitable for treatment of anhedonia, as several randomized controlled trials have found it to be efficacious in treating major depressive disorder (Goldberg et al., 2004; Fawcett et al., 2016) as well as motivational symptoms in Parkinson’s disease (Drijgers et al., 2012). Building on our prior report focusing on cross-sectional abnormalities in ventrostriatal dopamine function in medication-naive individuals with major depressive disorder (Schneier et al., 2018), we tested whether reward learning and ventrostriatal prediction error signalling prospectively predicted response to pramipexole. To directly assess the relationship between ventrostriatal dopamine function and response to pramipexole, we also examined whether baseline ventrostriatal dopamine release, measured using 11C-(+)-PHNO 11C-(+)-propyl-hexa-hydro-naphtho-oxazin, a D2/3 agonist PET imaging in conjunction with oral amphetamine, predicted response to pramipexole. Given pramipexole’s known effects on striatal dopamine (Mierau and Schingnitz, 1992), we hypothesized that individuals showing impaired reward learning and blunted ventrostriatal prediction errors to reward would disproportionally benefit from pramipexole treatment (i.e. show greater depressive and anhedonic symptom improvement). Consistent with links between reward learning, ventrostriatal prediction error signalling and ventrostriatal dopamine function, we also expected that lower...
ventrostriatal dopamine release would predict greater response to pramipexole.

Materials and methods

Participants

Individuals with major depressive disorder (n = 26) and healthy controls (n = 26) were recruited from clinics at the New York State Psychiatric Institute and Icahn School of Medicine at Mount Sinai. Inclusion and exclusion criteria are outlined in the Supplementary material. Procedures were approved by both institutional review boards, and participants provided written informed consent prior to participating, in accordance with the Declaration of Helsinki. The Clinical trials registration can be found at https://clinicaltrials.gov/ct2/show/NCT02033369.

Clinical measures

Three outcome measures assessing depressive symptoms, anhedonia and clinical global improvement were administered weekly across 6 weeks of treatment: the 17-item Hamilton Depression Rating Scale (HDRS) (Hamilton, 1960), the Snaith-Hamilton Pleasure Scale (SHAPS) (Snaith et al., 1995; Ameli et al., 2014), and the Clinical Global Impression-Change Scale (CGI) (Guy, 1976). Additional assessments are described in the Supplementary material.

Behavioural probabilistic reward task

Reward learning was assessed pre- and post-treatment using the Probabilistic Reward Task (PRT), which has been described in detail (Pizzagalli et al., 2008b). This task uses a differential reinforcement schedule to induce a response bias towards a more frequently rewarded (‘rich’) stimulus (see Supplementary material). Each trial began with a fixation cross (500 ms), followed by a schematic face without a mouth (500 ms). Next, a short (10 mm) or a long (11 mm) mouth was displayed (100 ms). Participants indicated whether the short or long mouth was presented. There were three blocks of 100 trials, and 40 correct trials in each block were followed by monetary reward (‘Correct! You won 20 cents’). Long and short mouths were presented with equal frequency; however, one of the lengths (the ‘rich stimulus’) was rewarded three times more frequently than the other (the ‘lean stimulus’). Participants were not informed of this contingency. Two versions were administered in a counterbalanced order from pre- to post-treatment: one where the length varied and another where the nose length varied.

After quality control, signal detection analysis (Macmillan and Creelman, 1991) was used to calculate response bias (the tendency to bias responding to the rich stimulus). Reward learning (defined as block 3 – block 1 response bias) was evaluated as a predictor of treatment response.

Computational model

To unravel the mechanisms driving any observed association between reward learning and treatment response, we used a reinforcement learning model to compute two parameters for each individual: reward sensitivity and learning rate (see Supplementary material) (Huys et al., 2013). Higher reward sensitivity indicates greater subjective value of a reward, whereas greater learning rate indicates greater weight of immediate prior rewards on future decisions.

Imaging acquisition and analysis

Functional MRI reinforcement learning paradigm

Full details of the functional MRI acquisition, learning paradigm and analysis can be found elsewhere (Schneier et al., 2018) and in the Supplementary material. Scanning was conducted on a GE SIGNA 3T scanner (GE Healthcare) with a 32-channel head coil. T1-weighted structural images (1 mm isotropic, 200 slices, field of view = 256 mm) and functional echo-planar images (repetition time = 2000 ms, echo time = 28 ms, flip angle = 77°, field of view = 19.2, 3 mm isotropic voxels, 40 slices) were acquired in six runs of 20 trials.

During functional MRI, participants performed a separate two-phase reinforcement learning task (Reinen et al., 2014) consisting of counterbalanced gain (winning money) and loss conditions (avoiding losing money from an endowment). On each trial, participants had to choose one of two shapes. After making a choice they received anticipatory feedback (‘correct’ or ‘incorrect’; 70/30 probability based on choice), followed by a monetary outcome. The trial staging allowed us to model prediction errors separately for anticipatory feedback and monetary outcomes. In the gain condition, ‘correct’ feedback triggered a $1 or $0.50 monetary gain (50/50 probability), whereas ‘incorrect’ feedback triggered a $0.50 or $0 monetary gain (50/50 probability). In the loss condition, correct feedback triggered a loss of $0 or $0.50 (50/50 probability), whereas incorrect feedback triggered a loss of $0.50 or $1 (50/50 probability). This design was used to equate the magnitude of both gain and loss prediction errors, while allowing for differences in motivational context.

Functional MRI analysis

A Q-learning model generated trial-by-trial prediction error values that were used as regressors for functional MRI analyses. Prediction error beta values generated from the general linear model were extracted from regions of interest in the left and right ventral striatum, defined by automated meta-analysis (neurosynth.org). A higher value for the gain prediction error beta indicates increased ventrostriatal activation for unexpected receipt of reward or better-than-expected feedback, and decreased activation for unexpected omission of reward or worse-than-expected feedback, in the gain condition. Conversely, a higher value for the loss prediction error beta indicates increased ventrostriatal activation for unexpected omission of loss or better-than-expected feedback, and decreased activation for unexpected receipt of loss or worse-than-expected feedback, in the loss condition. Eight prediction error variables were extracted: gain and loss prediction errors, under feedback and outcome conditions, in left and right ventral striatum. The four gain and four loss prediction errors were averaged to create a gain and a loss prediction error that were evaluated as predictors of treatment response.
PET imaging

The PET imaging methods are described in our prior report (Schneier et al., 2018). Subjects completed two 120-min \(^{11}C\)-(+)-PHNO PET scans (5-h apart), before and after 0.5 mg/kg of oral amphetamine. In contrast to functional MRI, which measures task-evoked changes in blood oxygen level-dependent activation, PET imaging calculates regional dopamine release as the difference in binding potential between two scans. Therefore, we chose an anatomical (rather than a functional) ventral striatum region of interest for PET analyses, which was drawn on each individual’s T1 image using criteria for ventral striatum boundary definitions defined in prior PET studies (Mawlawi et al., 2001; Martinez et al., 2003). Time-activity curves were calculated as the mean activity within the region of interest in each time frame. Reference tissue-based kinetic modelling yielded binding potential relative to non-displaceable compartment (BPND) (Innis et al., 2007). Percentage change from baseline BPND following amphetamine (BPND) was used as the measure of dopamine release (Martinez et al., 2003).

Pramipexole treatment

One day after behavioural testing and imaging, participants began 6 weeks of open-label pramipexole monotherapy. Doses (ranging from 0.5 to 2.5 mg/day) were adjusted weekly based on clinical response, and participant’s symptoms were assessed at each weekly visit via clinical interview.

Statistical analysis

Baseline group differences were assessed using the following: response bias: Group (control, depressed) × Block (1, 2, 3) ANOVA; functional MRI analyses: separate Group × Hemisphere (left, right) × Condition (feedback, outcome) ANOVAs for gain and loss prediction errors; PET analyses: paired samples t-tests for dopamine D2/3 receptor availability (BPND) and dopamine release (ΔBPND).

Predictors were then assessed for their ability to predict end-point symptom severity as well as rate of change in symptom improvement across the 6 weeks of treatment. This approach allowed us to examine potential biomarkers of overall versus rapid antidepressant effects (Supplementary material). First, multiple regression assessed whether measures of reward processing (reward learning, prediction error signals) and dopamine function (BPND and ΔBPND) predicted post-treatment symptom severity on the HDRS, SHAPS and CGI, controlling for baseline scores. Next, we used linear mixed effects models (implemented in STATA 13.1) to evaluate whether these measures of reward processing and dopamine function predicted the slope of symptom improvement across 6 weeks of treatment. Models included random intercepts and slopes. The predictors in the model were Baseline symptom scores, Predictor, Week, and a Predictor × Week interaction term. A significant Predictor × Week interaction indicated that the variable predicted the slope of symptom improvement across treatment.

Data availability

The data that support the findings of this study are available on request from the corresponding author.

Results

Sample characteristics

Twenty-four controls and 25 patients were considered because they had either valid behavioural, functional MRI or PET data (see CONSORT diagram, Supplementary Fig. 1). Sample characteristics are summarized in Table 1.

Baseline group differences in reward learning, prediction errors and ventrostriatal dopamine function

Reward learning on the Probabilistic Reward Task

A main effect of Block emerged \([F(2,80) = 5.62, P = 0.005, \eta_p^2 = 0.12]\), due to overall higher response bias in block 3 than in block 1 \((P = 0.03)\), indicating that the task effectively induced a response bias. Furthermore, a main effect of Group emerged \([F(1,40) = 5.65, P = 0.02, \eta_p^2 = 0.12]\) due to overall lower response bias in the depressed [mean ± standard deviation (SD) = 0.11 ± 0.15] than control \((0.20 ± 0.09)\) group (Cohen’s \(d = 0.73\); Fig. 1A). The main effect was not qualified by a Block × Group interaction \((P = 0.92)\). Groups did not differ in computationally-defined reward sensitivity \([t(40) = 0.40, P = 0.69]\) or learning rate \([t(40) = 0.50, P = 0.62]\) parameters.

Ventröstriatal prediction error signals

A trend-level main effect of Group emerged for the gain prediction error signal \([F(1,45) = 3.59, P = 0.07, \eta_p^2 = 0.07, d = 0.54]\). Averaged across conditions and hemispheres, the depressed group had blunted ventrostriatal prediction error responses when learning to gain rewards compared to controls (Fig. 1B). No group effects emerged for the loss prediction error signal (all \(P’s > 0.10)\).

Dopamine function

As previously reported (Schneier et al., 2018), there were no group differences in ventrostriatal dopamine D2/3 receptor availability (BPND) \([t(38) = -0.11, P = 0.92, d = 0.03]\) (Fig. 1C). In contrast, there was a trend for greater ventrostriatal dopamine release (ΔBPND) in the depressed relative to the control group \([t(38) = 1.85, P = 0.07, d = 0.58]\) (Fig. 1D).

Associations between reward learning, prediction error signals, ventrostriatal dopamine function, and symptom severity are reported in the Supplementary material.

Changes in reward learning and symptoms following pramipexole

Among 22 depressed patients who started pramipexole, 21 completed 6 weeks of treatment. The average maximum dose of pramipexole was 1.6 ± 0.7 mg/day. There were significant improvements across all measures from pre- to post-
treatment (Supplementary Table 1; see Supplementary Table 2 for treatment-emergent adverse events). Of those who completed treatment, 17 had valid PRT data at baseline and post-treatment. Despite significant improvements in symptoms, there were no changes in response bias, reward sensitivity or learning rate from pre- to post-treatment (all P’s > 0.10). However, after controlling for baseline HDRS or CGI scores, after controlling for baseline HDRS or CGI scores, respectively (all P’s > 0.10). However, after controlling for baseline SHAPS scores, baseline reward learning predicted post-treatment SHAPS scores (β = –0.77, P = 0.001); unexpectedly, better—rather than worse—baseline reward learning predicted lower post-treatment anhedonia (Fig. 2A). When the same analysis was run for the reward sensitivity and learning rate parameters (using a separate regression model for each parameter), only reward sensitivity emerged as a significant predictor of post-treatment SHAPS scores, β = –0.70, P = 0.004 (Fig. 2B). The reward learning and reward sensitivity predictors survived correction for multiple comparisons [corrected alpha = 0.05/(three PRT indices × three outcome measures) = 0.0056].

Greater baseline reward learning and reward sensitivity predict lower post-treatment anhedonia

Baseline PRT performance did not predict post-treatment HDRS or CGI scores, after controlling for baseline HDRS or CGI scores, respectively (all P’s > 0.10). However, after controlling for baseline SHAPS scores, baseline reward learning predicted post-treatment SHAPS scores (β = –0.77, P = 0.001); unexpectedly, better—rather than worse—baseline reward learning predicted lower post-treatment anhedonia (Fig. 2A). When the same analysis was run for the reward sensitivity and learning rate parameters (using a separate regression model for each parameter), only reward sensitivity emerged as a significant predictor of post-treatment SHAPS scores, β = –0.70, P = 0.004 (Fig. 2B). The reward learning and reward sensitivity predictors survived correction for multiple comparisons [corrected alpha = 0.05/(three PRT indices × three outcome measures) = 0.0056].

Linear mixed effects models examining predictors of change in HDRS, CGI and SHAPS scores across the 6 weeks of treatment failed to show a Reward learning × Week interaction or a Reward sensitivity × Week interaction (all P’s > 0.10).

Stronger ventrostriatal gain and weaker ventrostriatal loss prediction errors predict symptom improvement

Ventrostriatal gain prediction error signals
Gain prediction error signals did not predict post-treatment symptom scores. However, there was a significant Gain prediction error × Week interaction [B = −0.08, 95% confidence interval (CI) = −0.13 to −0.02, P = 0.004] for the model predicting CGI scores. Specifically, stronger gain prediction error signals predicted greater improvements in global illness severity across the 6 weeks of treatment (Fig. 3A). In addition, a trend-level Gain prediction error × Week interaction emerged for the model predicting HDRS scores [B = −0.32, 95% CI = −0.68 to 0.04, P = 0.08] where again, stronger gain prediction error signals predicted greater reductions in depressive symptoms across treatment. Contrary to initial predictions, gain prediction error did not predict improvement in SHAPS scores (P > 0.10).

Ventrostriatal loss prediction error signals
Loss prediction error signals did not predict post-treatment symptom scores on any outcome measure. However, significant Loss prediction error × Week interactions emerged for the model predicting SHAPS scores [B = 0.61, 95% CI = 0.20 to 1.02, P = 0.004] and for the model predicting HDRS scores [B = 0.39, 95% CI = 0.04 to 0.73, P = 0.03]. For both models, more blunted loss prediction errors (i.e. a reduced ventrostriatal response to monetary loss) predicted greater symptom improvement across treatment.

Table 1 Demographic and clinical characteristics of sample

<table>
<thead>
<tr>
<th></th>
<th>HC baseline (n = 24)</th>
<th>MDD baseline (n = 25)</th>
<th>P-value (HC versus MDD baseline)</th>
<th>MDD Week 6 (n = 21)</th>
<th>P-value (MDD baseline versus Week 6)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age, mean (SD)</td>
<td>26.9 (5.5)</td>
<td>26.5 (6.3)</td>
<td>0.84</td>
<td>26.2 (5.6)</td>
<td>0.47</td>
</tr>
<tr>
<td>Female, n (%)</td>
<td>12 (50)</td>
<td>13 (52)</td>
<td>0.89</td>
<td>11 (52)</td>
<td>0.64</td>
</tr>
<tr>
<td>Caucasian, n (%)</td>
<td>10 (42)</td>
<td>10 (40)</td>
<td>0.93</td>
<td>9 (43)</td>
<td>0.87</td>
</tr>
<tr>
<td>Years of education, mean (SD)</td>
<td>15.4 (1.5)</td>
<td>14.6 (1.4)</td>
<td>0.06</td>
<td>14.8 (1.6)</td>
<td>0.02</td>
</tr>
<tr>
<td>Income below $60 000 p.a., n (%)</td>
<td>14 (74)*</td>
<td>18 (86)*</td>
<td>0.34</td>
<td>19 (90)</td>
<td>0.40</td>
</tr>
<tr>
<td>HDRS age at onset, mean (SD)</td>
<td>17.4 (6.4)</td>
<td>2.4 (3.9)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>HDRS 17-item total, mean (SD)</td>
<td>0.2 (0.4)</td>
<td>20.2 (2.7)</td>
<td>&lt; 0.001</td>
<td>8.1 (5.4)</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>SHAPS, mean (SD)</td>
<td>18.8 (4.9)</td>
<td>32.0 (6.7)</td>
<td>&lt; 0.001</td>
<td>25.3 (6.9)</td>
<td>0.003</td>
</tr>
<tr>
<td>CGI, mean (SD)</td>
<td>3.0 (0.9)</td>
<td>1.8 (0.8)</td>
<td>&lt; 0.001</td>
<td></td>
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</tr>
<tr>
<td>MASQ Anhedonic Depression subscale, mean (SD)</td>
<td>37.7 (9.7)</td>
<td>82.6 (10.5)</td>
<td>&lt; 0.001</td>
<td>59.6 (18.5)</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>Apathy Evaluation Scale, mean (SD)</td>
<td>23.8 (5.0)</td>
<td>40.9 (8.8)</td>
<td>&lt; 0.001</td>
<td>31.7 (9.5)</td>
<td>0.001</td>
</tr>
<tr>
<td>TEPS subscale</td>
<td></td>
<td></td>
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<tr>
<td>Anticipatory, mean (SD)</td>
<td>49.0 (5.3)</td>
<td>36.2 (8.1)</td>
<td>&lt; 0.001</td>
<td>43.2 (7.9)</td>
<td>0.02</td>
</tr>
<tr>
<td>Consummatory, mean (SD)</td>
<td>38.4 (7.3)</td>
<td>30.2 (7.7)</td>
<td>&lt; 0.001</td>
<td>35.6 (6.5)</td>
<td>0.007</td>
</tr>
</tbody>
</table>

*Some participants chose not to report their income, therefore income totals are out of 19 healthy controls and 21 patients.

As the CGI change scale captures change in clinical impairment from one time point to the next, the ‘baseline’ mean and SD for this measure reflects ratings given at Week 1 (which capture changes in clinical impairment from baseline to Week 1).

HC = healthy control; MASQ = Mood and Anxiety Symptom Questionnaire; MDD = major depressive disorder; MDE = major depressive episode; TEPS = Temporal Experience of Pleasure Scale (greater scores on TEPS indicate less anhedonia).
The Gain prediction error × Week interaction for the model predicting CGI scores and the Loss prediction error × Week interaction for the model predicting SHAPS scores (Table 2) survived correction for multiple comparisons [corrected alpha = 0.05/(two prediction errors × three outcomes) = 0.0083].

Lower D2/3 receptor availability and dopamine release predict greater improvement in global illness severity

Dopamine D2/3 receptor availability (BPND)
Dopamine D2/3 receptor availability did not predict post-treatment symptom scores or slope of symptom improvement on the SHAPS or HDRS (P’s > 0.05). However, it did predict the slope of global illness severity improvement on the CGI. Specifically, a BPND × Week interaction emerged (B = 0.20, 95% CI = 0.04 to 0.37, P = 0.02) where lower dopamine D2/3 receptor availability predicted greater improvements in global illness severity across treatment (Fig. 3B).

Dopamine release (∆BPND)
Dopamine release did not predict post-treatment symptom scores or slope of symptom improvement on the SHAPS or HDRS (all P’s > 0.10). However, the ∆BPND × Week interaction for the model predicting CGI scores was marginally significant (B = –1.04, 95% CI = –2.08 to 0.01, P = 0.05) indicating that lower ventrostriatal dopamine
Figure 2 Baseline reward learning and reward sensitivity predict post-treatment anhedonia. Partial regression plots showing that (A) better baseline reward learning and (B) greater baseline reward sensitivity (as assessed using computational modelling) on the Probabilistic Reward Task (PRT) predicted lower post-treatment anhedonia (as assessed by the SHAPS) after controlling for baseline SHAPS scores. For visualization purposes, the grey dashed line shows the healthy control group mean and indicates that patients with scores equal to or greater than the control group mean (i.e. those with relatively more normative scores) showed the lowest post-treatment anhedonia.

Figure 3 Predictors of change in global illness severity across the 6 weeks of treatment. Figures show the moderating effect of baseline ventral striatal gain prediction error (A), ventral striatal dopamine D_{2/3} receptor availability (B) and the trend-level moderating effect of ventral striatal dopamine release (C) on the rate of global clinical improvement on the CGI across the 6 weeks of treatment. For visualization purposes, scores for values at the mean, 1 SD above the mean (`High`), and 1 SD below the mean (`Low`) are plotted. Scores for values equal to the healthy control group mean are also shown. Higher baseline gain prediction error signals, lower dopamine D_{2/3} receptor availability and lower dopamine release, predicted greater global clinical improvement. For the models involving the gain prediction error signal (A) and dopamine release (C) as predictors, patients with scores more similar to the healthy control group mean (i.e. those with relatively more normative scores), were those showing the greatest clinical improvement over the course of treatment. For the model involving ventral striatal dopamine D_{2/3} receptor availability as the predictor (B), the MDD group mean was equal to and overlapped with the healthy control group mean. DA = dopamine; HC = healthy control; PE = prediction error; VS = ventral striatal.
release predicted greater improvements in global illness severity across treatment (Fig. 3C). Neither of the PET predictors survived correction for multiple comparisons [alpha = 0.05 / (two PET indices × three outcomes) = 0.0083].

**Discussion**

Building on recent PET analyses on this sample (Schneier et al., 2018), which found no differences between controls and individuals with major depressive disorder on measures of striatal dopamine receptor availability or release, this study examined whether measures of reward-related mesolimbic dopamine system function (reward learning, functional MRI-based ventrostriatal prediction error signalling, PET-based ventrostriatal dopamine release) predicted clinical response to dopamine agonist treatment in major depressive disorder. Replicating prior findings (Kumar et al., 2008, 2018; Pizzagalli et al., 2008b), depression was characterized by significantly reduced reward learning and blunted ventrostriatal gain prediction error signals (trend) at baseline. Following pramipexole treatment, the depressed group showed significant reductions in depression, anhedonia and global illness severity. As hypothesized, baseline reward learning and ventrostriatal prediction error signalling were associated with post-treatment anhedonia severity and change in global illness severity, respectively, following 6 weeks of treatment with pramipexole. However, counter to the direction of predictions, individuals with better reward learning, greater reward sensitivity, and stronger ventrostriatal prediction error signalling to gains showed the greatest improvements in anhedonia (reward learning and sensitivity) or global illness severity (ventrostriatal prediction error signals). Although these findings await replication in a larger placebo-controlled study, the results suggest that depressed individuals with more normative reward learning and striatal prediction error signalling may respond favourably to a dopamine agonist. While unexpected, our results are consistent with an earlier literature suggesting that individuals with atypical depression (a subtype characterized by preserved reward sensitivity) may preferentially improve with dopaminergic pharmacotherapy (Stewart and Thase, 2007).

Direct measures of dopamine function also predicted clinical response to pramipexole. We hypothesized that individuals with more pronounced dopamine deficits (i.e. those with reduced ventrostriatal dopamine release), would show the greatest response to pramipexole. Results fit these predictions, where lower baseline ventrostriatal dopamine release predicted greater improvement in global illness severity. However, contrary to predictions, the depressed group did not show blunted ventrostriatal dopamine release relative to controls, but rather, showed a trend for increased ventrostriatal dopamine release. Accordingly, and consistent with the direction of effects observed for reward learning and ventrostriatal prediction error signalling, depressed patients with ventrostriatal dopamine release more similar to controls were those who responded more favourably to pramipexole.

Lower ventrostriatal dopamine D2/3 receptor availability also predicted greater improvement in global illness severity across treatment. Studies examining striatal dopamine receptor availability in depression have produced mixed findings, with nine PET studies reporting no difference, four reporting increases and one reporting decreases in receptor availability (for a review see Schneier et al., 2018). One explanation for higher receptor availability in depression is that depression-related dopamine deficits may cause a compensatory up-regulation of D2/3 receptors (Dunlop and Nemeroff, 2007). Accordingly, depressed individuals with lower ventrostriatal BPND (who showed the greatest global clinical improvement following pramipexole) might be those with more normative D2/3 receptor availability. However, PHNO binding is sensitive to competition with endogenous dopamine, with 42% of ventrostriatal BPND variance estimated to be attributable to endogenous dopamine (Caravaggio et al., 2016). Thus, lower ventrostriatal BPND at baseline could alternatively represent higher levels of endogenous dopamine.

Taken together, these findings suggest that measures of reward processing and striatal dopamine function are associated with lower post-treatment anhedonia severity and

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### Table 2 Linear mixed effect models showing the moderating effects of striatal prediction error signals on symptom improvement across the 6 weeks of treatment

<table>
<thead>
<tr>
<th>Model term</th>
<th>Coefficient</th>
<th>SE</th>
<th>Z</th>
<th>P</th>
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<tbody>
<tr>
<td><strong>Model 1: Ventral striatal gain prediction error predicts change in global illness severity</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Week</td>
<td>-0.26</td>
<td>0.04</td>
<td>-6.59</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Ventral striatal gain prediction error</td>
<td>0.34</td>
<td>0.12</td>
<td>2.96</td>
<td>0.003</td>
</tr>
<tr>
<td>Week × Ventral striatal gain prediction error</td>
<td>-0.08</td>
<td>0.03</td>
<td>-2.87</td>
<td>0.004</td>
</tr>
<tr>
<td><strong>Model 2: Ventral striatal loss prediction error predicts change in anhedonia</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Baseline SHAPS</td>
<td>0.82</td>
<td>0.17</td>
<td>4.79</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Week</td>
<td>-0.77</td>
<td>0.31</td>
<td>-2.50</td>
<td>0.012</td>
</tr>
<tr>
<td>Ventral striatal loss prediction error</td>
<td>-1.52</td>
<td>0.91</td>
<td>-1.67</td>
<td>0.095</td>
</tr>
<tr>
<td>Week × Ventral striatal loss prediction error</td>
<td>0.61</td>
<td>0.21</td>
<td>2.91</td>
<td>0.004</td>
</tr>
</tbody>
</table>

*The CGI-Change Scale measures changes in global illness severity and was therefore first administered after 1 week of treatment. Accordingly, models do not include a baseline CGI score term.*

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greater improvements in global illness severity, respectively, following treatment with a dopamine agonist in individuals with major depressive disorder. However, contrary to conventional assumptions, individuals with more normative rather than more disrupted reward and dopamine function, responded most favourably. These findings are consistent with a recent study showing that greater baseline ventrostriatal prediction error signalling predicted greater reductions in anhedonia in a naturalistic longitudinal study (Eckstrand et al., 2019). Furthermore, they align with studies showing links between better baseline reward processing and superior response to Behavioral Activation Therapy (Carl et al., 2016; Walsh et al., 2017), a therapy thought to specifically target anhedonia (Hopko et al., 2003). A critical next step is to determine whether baseline reward processing predicts superior response to treatments specifically targeting reward processing, or whether it predicts greater treatment responsiveness more generally.

Some limitations must be considered when interpreting the current findings (these are discussed further in the Supplementary material). First, a placebo group could not be included given the costs of intensive multimodal neuroimaging in this study (Schneier et al., 2018). Hence, the findings only point to a relationship between baseline measures of ventrostriatal reward function and pramipexole response at the group, rather individual patient, level. Future studies should test the specificity of our findings using placebo and/or a non-dopaminergic antidepressant control. Second, larger sample sizes are needed to test whether the differential predictive effects observed for reward learning (on anhedonia) and striatal prediction error signalling and ventrostriatal dopamine function (on global illness severity) are robust. Finally, functional MRI and PET imaging were only performed at baseline; therefore, we could not evaluate whether pramipexole altered ventrostriatal prediction error signalling or dopamine function. This is an important area for future research, as it remains unclear whether longer-term treatment with pramipexole may alter brain reward function via allostatic processes (Supplementary material).

Identifying ways to improve treatment precision for individuals with depression represents a major challenge. Using a multimodal approach, our findings suggest that measures of reward processing and ventrostriatal dopamine function may identify individuals with depression likely to respond favourably to a dopaminergic antidepressant. These findings pave the way for larger studies focused on improved antidepressant treatment precision, which is a critical step towards reducing the global burden of depression.

Acknowledgements
We thank Roberto Valdovinos and Danielle Moskow for assistance with data collection, and Page van Meter for assistance with data management.

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Competing interests
F.R.S. has received research support from Forest Laboratories/Allergan and Feelmore Labs. M.S. has received research support from Forest Laboratories, Pierre-Fabre, CHDI, and Otsuka; and has provided consultation for Amsen. D.V.I. has received consulting fees from Alkermes, Axsome, Centers of Psychiatric Excellence, Jazz, Lundbeck, MyndAnalytics (CNS Response), Otsuka, Precision Neuroscience, and Sundovion; and has received research support (through his academic institutions) from Alkermes, Astra Zeneca, Brainsway, LiteCure, Neosync, Roche, and Shire. A.A-D. has received research support from Takeda and Forest Pharmaceuticals and has served on advisory boards for Roche, Forum, and Otsuka. D.A.P. has received consulting fees from BlackThorn Therapeutics, Boehringer Ingelheim, Compass, Takeda and an honorarium from Alkermes for activities unrelated to the current research. D.A.P. has a financial interest in BlackThorn Therapeutics, which has licensed the copyright to the Probabilistic Reward Task through Harvard University. D.A.P. interests were reviewed and are managed by McLean Hospital and Partners HealthCare in accordance with their conflict of interest policies. All other authors report no biomedical financial interests or potential conflicts of interest.

Supplementary material
Supplementary material is available at Brain online.

References


Dunlop BW, Nemeroff CB. The role of dopamine in the pathophysiology of depression. Arch Gen Psychiatry 2007; 64: 327–37.


Supplementary Information

Supplementary Methods

Inclusion/exclusion criteria

Inclusion criteria for the depressed group were a current diagnosis of major depressive disorder, confirmed using the Structured Clinical Interview for DSM-IV (First et al., 2002); a score of 17 to 28 on the 17-item Hamilton Depression Rating Scale (HDRS) (Hamilton, 1960); and <2 weeks of lifetime psychiatric medication (and no medication in the past 3 months). Exclusion criteria for the depressed group were lifetime psychotic, bipolar, attention deficit, or substance use disorders (including nicotine). Inclusion criteria for the age-, gender- and race/ethnicity-matched healthy control group were an absence of lifetime psychiatric disorders or major medical illnesses. Exclusion criteria were tobacco or illicit substance use in the past 3 months; a family history of schizophrenia; pregnancy; breastfeeding; use of hormonal contraceptives.

Additional measures of hedonic function

The primary measure of anhedonia was the Snaith-Hamilton Pleasure Scale (SHAPS) (Snaith et al., 1995), which is one of the most widely used assessments of anhedonia in depressed samples. This scale consists of 14 items that inquire about the level of pleasure experienced in response to pleasant stimuli and situations. It is scored on a scale from 1 (Definitely Agree) to 4 (Definitely Disagree) with scores ranging from 14 to 56. Higher scores are indicative of more severe anhedonia. This scoring method preserves the continuous structure of the data and is modified from the original scoring using by Snaith and colleagues (Snaith et al., 1995), which recoded the four response categories into dichotomous categories (0=Agree, 1=Disagree) for the purposes of categorizing individuals as anhedonic or non-anhedonic. The modified scoring method, which has been used across numerous studies, has
been found to produce good to excellent internal consistency across both clinical and non-clinical populations (Franken et al., 2007). Additional self-report measures of hedonic function were administered at baseline and post-treatment: the Temporal Experience of Pleasure Scale (TEPS) (Gard et al., 2006), the anhedonic depression subscale of the Mood and Anxiety Symptom Questionnaire MASQ) (Watson et al., 1995), and the Apathy Evaluation Scale (AES) (Marin et al., 1991).

**PRT quality control criteria**

In order for response bias to be accurately interpreted, quality control criteria were applied to confirm that participants performed at a level that was above chance, and therefore high enough for them to be exposed to the asymmetrical reinforcement ratio. First, trials where the reaction time (RT) was < 150ms or > 2500ms were excluded, as were remaining trials with RT failing ±3SD from the mean. Cases where there were less than 80 valid trials per block, greater than 10 outlier trials per block, less 55% accuracy per block, or where the rich to lean reward ratio was lower than 2.5:1, were excluded from analyses. Next, response bias and discriminability scores were computed using these formulae:

**Response bias:**

$$ \log b = \frac{1}{2} \log \left( \frac{\text{Rich}_{\text{correct}} \cdot \text{Lean}_{\text{incorrect}}}{\text{Rich}_{\text{incorrect}} \cdot \text{Lean}_{\text{correct}}} \right) $$

**Discriminability:**

$$ \log d = \frac{1}{2} \log \left( \frac{\text{Rich}_{\text{correct}} \cdot \text{Lean}_{\text{correct}}}{\text{Rich}_{\text{incorrect}} \cdot \text{Lean}_{\text{incorrect}}} \right) $$

To compute response bias and discriminability for cases that had a zero in the formula, 0.5 was added to every cell in the formula matrix (Hautus, 1995). For regression analyses examining PRT performance as predictors of symptom improvement, the primary variable of interest was reward learning (block 3 response bias – block 1 response bias),
which was chosen in light of prior studies that have linked this variable to individual differences in striatal dopamine clearance (Kaiser et al., 2018).

**Computational modelling of the Probabilistic Reward Task (PRT)**

To separate the influence of reward sensitivity (which operationalizes reduction in consummatory pleasure) and learning rate (which operationalizes participants’ ability to learn from reward feedback) on PRT performance, we fitted a series of reinforcement learning models to the PRT choice data. In a prior, independent study, worse anhedonia has been linked to blunted reward sensitivity (Huys et al., 2013). Learning rate was also of interest given that this separate study showed that a single low dose of pramipexole altered learning rate in healthy individuals (Huys et al., 2013) (however, no study to date has examined the influence of pramipexole on these parameters in individuals with depression). These models tested whether participants associated rewards with stimulus-action pairs (‘Stimulus-Action’ model), with actions (‘Action’ model), or with a mixture of the two stimulus-action associations weighted by an uncertainty factor (‘Belief’ model). They also tested whether subjects treated zero outcomes as losses (‘Punishment’ model). The models were fitted using an empirical Bayesian random-effects approach and were compared using integrated group-level Bayesian Information Criterion factors following previously established procedures (Huys et al., 2013). Individual subject parameter inference was constrained by an empirical prior distribution and no further assumptions were made. We found that the ‘Belief’ model gave the most parsimonious account of the data (group-level log Bayes factor compared to the second-most parsimonious model = 23, which is >20 and represents very strong evidence in favor of the better fitting model). This model assumes uncertainty within subjects about the presented stimulus. As such, they might assign rewards to both stimuli with only a certain preference for the actual presented stimulus. Five parameters were derived: (i) reward sensitivity assessed the immediate behavioral impact of rewards; (ii) learning rate
represented subjects’ ability to accumulate rewards over time and learn from the rewards; (iii) belief indicated subjects’ uncertainty about which stimulus was actually presented; (iv) instruction sensitivity measured subjects’ ability to follow the instructions; (v) initial bias indicated subjects’ initial bias towards one response or the other. This study focused on the reward sensitivity and learning rate parameters, which were analyzed in the transformed space in order to prevent issues associated with non-Gaussianity. In the current study, reward sensitivity and learning rate parameters were negatively correlated at trend level in both the healthy control \((r= -0.41, p=0.07)\) and depressed groups \((r= -0.40, p=0.06)\).

**fMRI preprocessing and analysis**

Functional images were preprocessed with SPM8 and analyzed with NeuroElf software (http://neuroelf.net/). Images were slice-time corrected and realigned to the first image in each run, warped to the Montreal Neurological Institute template, and smoothed (6mm Gaussian kernel). Next, first-level analyses were conducted using a general linear model (GLM) that included six stick function regressors convolved with a hemodynamic response for choice, feedback, and outcome, each with trial-specific parametric regressors (choice value, feedback prediction error, and monetary outcome prediction error). A high-pass temporal filter (Fourier transform, 200sec) and motion parameters were included in the model as nuisance regressors. A second-level model was developed that included regressors for choice, feedback prediction error, and outcome prediction error, each separated by a jittered interval. Prediction error regressors for use with the fMRI GLM were generated using a Q-learning model as has been established for this task in prior work (for details, see Reinen et al., 2014). Gain and loss condition learning signals were analyzed using separate regressors in the same model.
Analyses focused on activation in a ventral striatal region of interest (See Supplementary Fig. 2). This region was defined using the automated meta-analysis Neurosynth. We used "ventral striatum" as the Neurosynth search term, identified the peak coordinates, and extracted activation for each participant in voxels within a 6mm sphere (radius) surrounding the region and its bilateral counterpart. There are currently 14,371 studies in the Neurosynth database and the term "ventral striatum" yielded 415 studies and 12,989 activations.

**PET imaging**

First, a 7sec computed tomography (CT) scan was conducted, followed by a 120-minute baseline scan. Next, 0.5 mg/kg of amphetamine was administered orally. Three hours later, another CT scan was conducted, followed by the second 120-minute scan. PET scanning was performed on a Biograph multispectral PET-CT (Siemens Healthineers, Knoxville, TN).

**Aspects of trial design**

Participants were recruited from the New York State Psychiatric Institute Division of Translational Imaging (healthy controls) and the New York State Psychiatric Institute Division of Clinical Therapeutics Anxiety Disorders Clinic and Depression Evaluation Service, and the Depression and Anxiety Center of Mount Sinai Icahn School of Medicine (MDD participants). Participants were initially screened via telephone and those who appeared eligible were invited to take part in a psychiatric and medical history evaluation (by a MD or PhD/PsyD), as well as a full SCID interview to confirm diagnosis of MDD. For those deemed eligible following the SCID interview, a physical examination was performed by a physician. This included a blood test for hematology, liver, thyroid and kidney function assessments, as well as a pregnancy test in women of child-bearing potential. Participant’s
height and weight were obtained, an EKG was performed and participants were screened for metal and other MR contraindications. Ineligible subjects with MDD or other disorders were referred for clinical treatment.

Participants who were deemed eligible based on the initial clinical and medical screen were scheduled to return to complete self-report measures of anhedonia, mood and anxiety symptoms, the PRT, MRI and PET scans. These procedures were scheduled for 2 separate days less than 1 week apart. Following these baseline assessments the MDD participants received 6 weeks of open-label treatment with pramipexole, during which time they returned for weekly visits with a psychiatrist to monitor treatment. Weekly independent evaluator assessments of symptom severity and side effects were also obtained. Following the six weeks of treatment, a second counterbalanced version of the PRT was administered behaviorally. Full details of the trial protocol can be found at https://clinicaltrials.gov/ct2/show/NCT02033369.

**Supplementary Results**

*Associations among PRT performance, fMRI and PET variables and baseline symptom severity*

There were no correlations among the PRT variables (i.e., reward learning, reward sensitivity, learning rate), the fMRI-based ventral striatal prediction errors, and the PET-based BP_{ND} or ΔBP_{ND} across the entire sample. When examining the groups separately, among controls there were trend-level correlations between ΔBP_{ND} and gain feedback prediction error in both the left ($r=-0.40$, $p=0.08$) and right ($r=-0.44$, $p=0.053$) ventral striatum, such that greater ventrostriatal dopamine release was associated with a stronger ventrostriatal response to unexpected correct feedback (note that negative values of ΔBP_{ND} indicate greater ventrostriatal dopamine release). There was also a trend-level correlation
between lower $\Delta BP_{ND}$ and greater gain outcome prediction error in the left ventral striatum ($r=0.40$, $p=0.09$). No significant associations emerged within the depressed group (all $ps>0.05$).

In terms of correlations between measures of reward processing, ventrostriatal dopamine function and symptoms in the depressed group, lower reward sensitivity (derived from the computational model) was associated with worse anhedonia on the SHAPS ($r=-0.46$, $p=0.03$), replicating prior work (Huys et al., 2013). In contrast, neither ventrostriatal prediction errors, $BP_{ND}$ nor $\Delta BP_{ND}$ correlated with baseline symptom severity (all $ps>0.05$).

**Baseline PRT reward learning and reward sensitivity as predictors of post-treatment anhedonia, controlling for changes in non-anhedonic symptoms and depressive episodes**

Highlighting specificity, both PRT reward learning and the reward sensitivity parameter remained significant predictors at an uncorrected threshold ($p<0.02$ for both predictors) of post-treatment SHAPS scores when controlling for changes in non-anhedonic depressive symptoms (MASQ GDD subscale scores) and anxiety (MASQ AA subscale scores). Furthermore, we confirmed that neither baseline reward learning nor baseline reward sensitivity predicted post-treatment MASQ GDD scores or MASQ AA scores, after controlling for baseline MASQ GDD and MASQ AA scores, respectively (all $ps>0.30$). In addition, baseline reward learning and baseline reward sensitivity remained significant predictors of post-treatment SHAPS scores when controlling for the number of lifetime major depressive episodes ($p<0.01$ for both predictors), indicating that the results were not driven by individual differences in baseline depressive illness severity.

Given that baseline reward learning and reward sensitivity predicted post-treatment SHAPS scores, we also examined whether changes in PRT performance from pre- to post-treatment predicted change in SHAPS scores. To do this we ran stepwise multiple regression
analyses that included baseline SHAPS scores and baseline PRT performance in the first step, post-treatment PRT performance in the second step, and post-treatment SHAPS scores as the dependent variable. Results showed that neither post-treatment reward learning ($\beta=0.17$, $p=0.37$) nor post-treatment reward sensitivity ($\beta=0.10$, $p=0.63$) emerged as significant predictors of post-treatment SHAPS scores when entered into the second step of the model. Furthermore, in both models, baseline reward learning ($\beta=-0.71$, $p=0.002$) and baseline reward sensitivity ($\beta=-0.61$, $p=0.008$) remained significant at the second step, confirming that baseline reward learning and sensitivity, rather than change in reward learning and sensitivity, predicted post-treatment anhedonia severity.

**Supplementary Discussion**

**Additional limitations and future directions**

*Lack of association between reward learning and measures of ventrostriatal dopamine function*

Given that reward learning has been hypothesized to be driven by phasic firing of dopamine in the striatum, we had expected that fMRI-based measures of ventrostriatal prediction error signaling and PET-based measures of ventrostriatal dopamine release and receptor availability would correlate with individual differences in behavioral reward learning. However, neither ventrostriatal prediction error signaling nor measures of dopamine release or receptor availability correlated with behavioral performance on the PRT (i.e., as indexed by response bias or the two computational parameters) (see Supplementary Results). There are several possible reasons for this. First, the sample size was relatively small, which may have reduced our power to observe significant associations between measures of ventrostriatal dopamine function and behavior. Second, although the PRT has been widely used to study reward learning and its relationship to frontostriatal function (e.g., Vrieze et al.,
2013; Kaiser et al., 2018), it is a relatively simple task that only involves learning actions in response to a single stimulus dimension that is associated with gains. Accordingly, the relationship between ventrostriatal dopamine release and ventrostriatal prediction error signaling with other aspects of learning (e.g., learning stimulus-outcome associations using more complex, multidimensional stimuli) was not examined. Future studies may benefit from using additional measures of reward learning in order to clarify the lack of association observed in the current study.

Possible reasons for the lack of changes in reward learning from pre- to post-treatment

Given evidence that a single low dose of pramipexole has been previously found to reduce reward learning in healthy humans (Pizzagalli et al., 2008) and rats (Der-Avakian et al., 2013; Lamontagne et al., 2018) as well as reduce phasic dopamine firing in rodents (Tokunaga et al., 2012), the lack of effects of pramipexole on reward learning was surprising. One possibility is that pramipexole improves symptoms via alterations in other facets of reinforcement learning. For example, Argyelan and colleagues (Argyelan et al., 2018) found that in Parkinson’s Disease, dopamine agonists (levodopa or pramipexole) modulated punishment learning and attenuated striatal responses to punishment, leaving reward learning and reward-related striatal activation unaltered. Although this resulted in a higher ratio of striatal activation to reward versus punishment, this effect was driven by dopaminergic attenuation of punishment-related neural activation. Future studies assessing changes in both punishment and reward learning pre- and post-pramipexole could provide further insights into pramipexole’s effects on other aspects of learning.
Potential for negative effects of long-term pramipexole treatment on reward function

One important consideration that warrants further investigation is whether treatment with pramipexole may lead to a worsening of depression and anhedonia when used over the long term, in a similar manner to that observed in individuals who abuse recreational substances that have dopaminergic effects, such as cocaine. Specifically, although these recreational drugs have pro-hedonic effects over the short term, with continuous long-term exposure their use leads to compensatory changes in the reward system via the process of allostasis, which can lead to the emergence of depression (Koob & Le Moal, 2001). Although we only examined the effects of a relatively short course of pramipexole treatment (i.e., 6 weeks) some studies suggest that longer-term treatment (e.g., 16 weeks) may be relatively safe and effective in the management of treatment-resistant depression. For example, Lattanzi et al. (2002) reported a 68% response rate to pramipexole augmentation in a sample of individuals with treatment-resistant depression who were treated with pramipexole for up to 16 weeks. In a follow-up to this study that tracked rates of sustained remission in patients who received treatment with pramipexole for up to 1 year, Cassano and colleagues (2004) found that 60.9% of patients experienced sustained remission of their major depressive episode during the follow-up period. Similar findings have been observed in individuals with bipolar depression, where pramipexole given as an adjunct to a mood stabilizer for an average of 6.7 ± 9.0 months was found to improve depressive symptoms significantly within four weeks and this improvement was maintained for over 9 months (El-Mallakh et al., 2010). This suggests that unlike drugs of abuse, pramipexole does not appear to cause a worsening of depressive or anhedonic symptoms over the long term. However, future studies are needed to fully understand the potential adverse effects of long-term treatment with pramipexole, particularly since long-term treatment with the drug or other dopamine agonists...
has been associated with the emergence of impulse control disorders in subsets of individuals with Parkinson’s disease undergoing dopamine replacement therapy (Weintraub et al., 2010; Garcia-Ruiz et al., 2014).
Supplementary References


### Supplementary Table 1. Changes in reward learning and symptoms from pre- to post-treatment in the depressed (MDD) group

<table>
<thead>
<tr>
<th>Measure, M (SD)</th>
<th>MDD baseline</th>
<th>MDD week 6</th>
<th>P Value</th>
<th>Cohen’s d</th>
</tr>
</thead>
<tbody>
<tr>
<td>HDRS 17-item total</td>
<td>20.0 (2.6)</td>
<td>8.1 (5.4)</td>
<td>&lt;0.001</td>
<td>2.16</td>
</tr>
<tr>
<td>SHAPS</td>
<td>31.6 (6.6)</td>
<td>25.3 (6.9)</td>
<td>0.003</td>
<td>0.75</td>
</tr>
<tr>
<td>CGI</td>
<td>3.0 (0.9)¹</td>
<td>1.8 (0.8)</td>
<td>&lt;0.001</td>
<td>1.00</td>
</tr>
<tr>
<td>MASQ Anhedonic Depression subscale</td>
<td>82.9 (10.7)</td>
<td>59.6 (18.5)</td>
<td>&lt;0.001</td>
<td>1.14</td>
</tr>
<tr>
<td>MASQ General Distress Depression subscale</td>
<td>40.2 (11.8)</td>
<td>23.9 (12.9)</td>
<td>&lt;0.001</td>
<td>1.38</td>
</tr>
<tr>
<td>MASQ General Distress Anxiety subscale</td>
<td>23.6 (7.1)</td>
<td>18.0 (5.4)</td>
<td>0.001</td>
<td>0.84</td>
</tr>
<tr>
<td>MASQ Anxious Arousal subscale</td>
<td>25.3 (7.5)</td>
<td>21.7 (6.0)</td>
<td>0.03</td>
<td>0.51</td>
</tr>
<tr>
<td>Apathy Evaluation Scale</td>
<td>41.0 (9.2)</td>
<td>31.7 (9.5)</td>
<td>0.001</td>
<td>0.85</td>
</tr>
<tr>
<td>TEPS subscale</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Anticipatory</td>
<td>36.9 (8.4)</td>
<td>43.2 (7.9)</td>
<td>0.02</td>
<td>0.58</td>
</tr>
<tr>
<td>Consummatory</td>
<td>30.4 (7.7)</td>
<td>35.6 (6.5)</td>
<td>0.007</td>
<td>0.65</td>
</tr>
<tr>
<td>PRT variables</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Reward learning (block 3 – block 1 response bias)</td>
<td>0.1 (0.2)</td>
<td>0.1 (0.2)</td>
<td>0.83</td>
<td>0.00</td>
</tr>
<tr>
<td>Reward sensitivity (model parameter)</td>
<td>1.2 (0.3)</td>
<td>1.2 (0.3)</td>
<td>0.95</td>
<td>0.00</td>
</tr>
<tr>
<td>Learning rate (model parameter)</td>
<td>-3.5 (1.8)</td>
<td>-4.4 (1.3)</td>
<td>0.10</td>
<td>1.10</td>
</tr>
<tr>
<td>Discriminability</td>
<td>0.5 (0.2)</td>
<td>0.4 (0.2)</td>
<td>0.06</td>
<td>0.99</td>
</tr>
</tbody>
</table>

Note. MDD=major depressive disorder; HDRS=Hamilton Depression Rating Scale; CGI=Clinical Global Impression-Change Scale; MASQ=Mood & Anxiety Symptom Questionnaire; TEPS=Temporal Experience of Pleasure Scale; PRT=Probabilistic Reward Task. Descriptive statistics for symptom scores are based on the 21 MDD subjects who completed treatment. Statistics for PRT variables are based on the 17 MDD subjects who completed treatment and had valid PRT data at both baseline and week 6. Since the CGI change scale captures change in clinical impairment from one time point to the next, the “baseline” mean and SD for this measure reflects ratings given at week 1 (which capture changes in clinical impairment from baseline to week 1).
**Supplementary Table 2.** Treatment Emergent Adverse Events

<table>
<thead>
<tr>
<th>Rate of Treatment Emergent Adverse Event (N, %)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Headache</td>
</tr>
<tr>
<td>Heartburn</td>
</tr>
<tr>
<td>Nausea</td>
</tr>
<tr>
<td>Vomiting</td>
</tr>
<tr>
<td>Decreased Appetite</td>
</tr>
<tr>
<td>Increased Appetite</td>
</tr>
<tr>
<td>Dry Mouth</td>
</tr>
<tr>
<td>Constipation</td>
</tr>
<tr>
<td>Diarrhea</td>
</tr>
<tr>
<td>Excessive Sweating</td>
</tr>
<tr>
<td>Skin Problems</td>
</tr>
<tr>
<td>Bruising Easily</td>
</tr>
<tr>
<td>Restlessness</td>
</tr>
<tr>
<td>Tremor</td>
</tr>
<tr>
<td>Nervousness</td>
</tr>
<tr>
<td>Impaired Coordination</td>
</tr>
<tr>
<td>Insomnia</td>
</tr>
<tr>
<td>Fatigue</td>
</tr>
<tr>
<td>Somnolence</td>
</tr>
<tr>
<td>Sleep Attacks</td>
</tr>
<tr>
<td>Decreased Libido</td>
</tr>
<tr>
<td>Sexual Dysfunction</td>
</tr>
<tr>
<td>Urinary Dysfunction</td>
</tr>
<tr>
<td>Blurry Vision</td>
</tr>
<tr>
<td>Lightheadedness</td>
</tr>
<tr>
<td>Postural Dizziness</td>
</tr>
<tr>
<td>Forgetfulness</td>
</tr>
<tr>
<td>Impaired Concentration</td>
</tr>
<tr>
<td>Swelling in Legs</td>
</tr>
<tr>
<td>Compulsive Gambling or Shopping</td>
</tr>
<tr>
<td>Hallucinations or Illusions</td>
</tr>
</tbody>
</table>

1 At baseline and at each weekly visit during treatment, participants gave a rating for each symptom listed in the first column, ranging from 0 (Absent) to 3 (Severe). An adverse event was coded as present if, relative to baseline rating, severity was increased at any subsequent weekly rating.

2 Sleep “attacks” were all mild. They involved a sudden urge to go to sleep but could always be resisted.

3 Three patients reported one episode of excessive shopping, but none were clearly outside of their normal range of behavior.

4 One patient thought she heard her name being called once
Supplementary Fig. 1. CONSORT diagram

Supplementary Fig. 1. Figure shows the flow of participants into the study, along with reasons for exclusion.
Supplementary Fig. 2. Ventral striatal region-of-interest

Supplementary Fig. 2. Figure shows the spherical regions-of-interest (ROI) in the left and right ventral striatum (black dashed outline) created based on the results of automated meta-analysis (Neurosynth). For the purposes of visualization, the ROIs are overlaid onto a map showing the prediction error in the healthy control group ($p<0.05$ uncorrected, $y=10$, feedback and outcome gain conditions). Results show good coverage of the prediction error by the ventral striatal ROIs.