Valence and agency influence striatal response to feedback in patients with major depressive disorder

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Background: Reduced sensitivity to positive feedback is common in patients with major depressive disorder (MDD). However, findings regarding negative feedback are ambiguous, with both exaggerated and blunted responses being reported. The ventral striatum (VS) plays a major role in processing valenced feedback, and previous imaging studies have shown that the locus of controls (self agency v. external agency) over the outcome influences VS response to feedback. We investigated whether attributing the outcome to one’s own action or to an external agent influences feedback processing in patients with MDD. We hypothesized that depressed participants would be less sensitive to the feedback attribution reflected by an altered VS response to self-attributed gains and losses. Methods: Using functional MRI and a motion prediction task, we investigated the neural responses to self-attributed (SA) and externally attributed (EA) monetary gains and losses in unmedicated patients with MDD and healthy controls. Results: We included 21 patients and 25 controls in our study. Consistent with our prediction, healthy controls showed a VS response influenced by feedback valence and attribution, whereas in depressed patients striatal activity was modulated by valence but was insensitive to attribution. This attribution insensitivity led to an altered ventral putamen response for SA – EA losses in patients with MDD compared with healthy controls. Limitations: Depressed patients with comorbid anxiety disorder were included. Conclusion: These results suggest an altered assignment of motivational salience to SA losses in patients with MDD. Altered striatal response to SA negative events may reinforce the belief of not being in control of negative outcomes contributing to a cycle of learned helplessness.

Introduction

Affective biases in emotion processing are characteristic of major depressive disorder (MDD), as indicated by a reduced ability to experience pleasure (anhedonia) and a bias toward the negative aspects of the environment. Depressed patients, for instance, show a blunted response to positive or pleasant stimuli but an enhanced memory for negative stimuli and a stronger responsiveness to threatening stimuli.

Different behavioural responses to feedback have also been widely reported in patients with MDD. Across various cognitive tasks, depressed patients show an increased risk of committing a subsequent error after failure. Moreover, when performing a probabilistic reversal learning (PRL) task, depressed patients show an increased tendency to switch after obtaining misleading negative feedback. The higher rate of switching has been interpreted as an oversensitivity to negative feedback, an index of automatic error detection. However, increased sensitivity to negative feedback is not a consistent finding in depressed patients. Several electrophysiological studies have reported either no changes or even reduced error-related negativity in patients with MDD. Neuroimaging data suggest that MDD is associated with a blunted response to both negative and positive feedback. Using a gambling task, weaker responses to losses in the anterior cingulate cortex and to gains in the ventral striatum have been reported in depressed patients relative to healthy controls. A blunted reactivity to feedback is consistent with behavioural findings demonstrating reduced responsiveness during the anticipation and/or the presentation of rewards in depressed individuals and with neuroimaging studies reporting that this decreased sensitivity to positive feedback is associated with reduced striatal response.

Hyposensitivity to positive feedback is a common finding in depressed patients, while results regarding sensitivity to negative feedback seem to be ambiguous, with both exaggerated and blunted responses being reported. A possible explanation is that reactivity to negative feedback may depend on symptom severity, with moderately depressed patients showing oversensitivity and severely depressed patients showing a blunted response. An alternative explanation is that feedback responsiveness in patients with MDD may be task-dependent. The learned helplessness theory hypothesizes that in individuals exposed to stressful, uncontrollable life events a generalized emotional–cognitive state of perceiving the
Altered processing of self-attributed losses in depression

environment as uncontrollable can develop and that this mechanism is central to the onset and maintenance of MDD.\textsuperscript{24,25} Given its theoretical importance to understanding depressed cognition, the level of control attainable within a task may be relevant. In fact, an abnormal sense of agency — the experience of controlling one’s own actions and, through them, events in the outside world\textsuperscript{26} — in the form of loss of agency and helplessness constitute a major marker of MDD.\textsuperscript{27}

The ventral striatum (VS) plays a key role in the processing of valenced outcomes.\textsuperscript{28,29} In light of the centrality of agency to computational accounts of VS\textsuperscript{30} and of cognitive accounts of valence processing,\textsuperscript{31} the locus of control (self v. external agency) over an outcome should influence VS response to feedback. Consistent with this hypothesis, previous studies have shown greater VS response when gains were contingent to performance\textsuperscript{25} and when task difficulty increased.\textsuperscript{32}

The goal of our study was to more directly investigate the neuronal responses to financial gains and losses attributed to one’s own action (related to response contingency) or an external agent (unrelated to response contingency) in patients with MDD. To this end, we conducted a functional MRI (fMRI) study using a task in which participants experience monetary gains or losses either due to their performance (self-attributed [SA]) or to a biased coin toss (externally attributed [EA]). We have previously shown that healthy controls modulate their VS response according to the feedback’s valence and attribution,\textsuperscript{33} and we hypothesized that, compared with healthy controls, depressed patients would show a VS response less sensitive to feedback attribution, leading to an altered response to SA gains and losses.

Methods

Participants

We recruited healthy controls without any psychiatric, neurologic, or medical illness and unmedicated patients with MDD for participation in the study. Exclusion criteria for both groups included age younger than 18 years or older than 65 years, current or past psychosis or mania, major medical or neurologic illness, current drug or alcohol abuse and MRI contraindications (as assessed by an MRI safety questionnaire). Inclusion in the MDD group was contingent on a diagnosis of current MDD based on a semistructured interview: the Structured Clinical Interview for DSM-IV Axis I Disorders (SCID-I). Control participants were required to have no history of MDD in their lifetime, have no history of MDD in first-degree relatives and have no Axis I disorders based on a SCID-I interview. The University of Zurich’s Institutional Review Board approved our study protocol, and all participants gave written informed consent. They were paid a modest compensation for participation in the study in addition to the gains they could make during the experimental tasks.

Psychometric measures

All participants completed the German version of the Positive and Negative Affect Schedule (PANAS\textsuperscript{34}), the German version of the Hopelessness Scale originally developed by Beck (H-Scale\textsuperscript{35}), and the Snaith-Hamilton Pleasure Scale (SHAPS\textsuperscript{36}). In addition, depressed participants completed the Beck Depression Inventory II (BDI-II\textsuperscript{40,41}) and the Inventory for Depressive Symptomatology (IDS\textsuperscript{40,41}).

Motion prediction task

The motion prediction task has been described previously\textsuperscript{42} and is reported in details in Appendix 1, available at jpn.ca. In brief, each trial started with 2 balls moving with different speeds and from different starting points toward a finish line. The task on every trial of the experiment was to predict which ball would cross the finish line first and to indicate the decision by a left or right button press. Only after this decision were participants instructed whether the response was relevant or irrelevant to the upcoming feedback. Specifically, participants were told that on each trial they could gain or lose 50 cents, indicated by “+50” or “−50” feedback. On a random 50% of the trials feedback was performance-dependent (SA gains or losses; correct v. error). On the other 50% of trials feedback was dependent on chance and was randomly selected by the computer with a probability tailored to match their success rate in performance-dependent trials (EA negative or positive feedback was determined by a biased virtual coin flip). A picture with the words “You” and “Coin” and an arrow pointing toward either word was presented on the screen 750 ms after the response to indicate if the feedback was associated with the participant’s performance or not. Finally, feedback about winning (+50) or losing (−50) was presented (Appendix 1, Fig. S1). The next trial started after a fixation period of 2000 ± 500 ms. If the participant failed to respond, the arrow pointed toward the word “You” and was followed by the feedback “Missed.” Two healthy controls and 1 patient with MDD performed a shorter version of the task (100 trials), and all other participants completed 130 trials. To keep uncertainty about performance high, task difficulty was adapted for each participant (error rate > 30%) using a training session of 100 trials in which the participants received only performance feedback (correct: happy face; incorrect: unhappy face). Participants were unaware that performance was manipulated and were told to do their best.

Image acquisition

Images were acquired on a Philips Achieva 3 T whole-body MRI unit equipped with an 8-channel head coil. Functional time series were acquired with a sensitivity encoded single shot echo-planar sequence. We placed 36 contiguous axial slices along the anterior–posterior commissure plane covering the entire brain. We used the following parameters: repetition time 2000 ms, echo time 35 ms, ascending acquisition order, voxel matrix 80 × 80 interpolated to 128 × 128, voxel size 2.75 × 2.75 × 4 mm, SENSE acceleration factor R = 2.0. The first 4 acquisitions were discarded owing to T₁, saturation effects. T₁-weighted high-resolution images were also acquired for each participant.

Data analysis

Demographic and psychometric data were analyzed with an unpaired t test and sex with a χ² test using StatView version
5.0.1 (SAS Institute). We analyzed mean reaction time (RT) differences for correct and incorrect trials (independently of attribution) with repeated-measures analyses of variance (RM-ANOVA). We considered results to be significant at \( p < 0.05 \), 2-tailed.

To accurately attribute the financial outcome, participants were required to attend to contextual information presented after their performance but before the outcome (Appendix 1, Fig. S1). It is possible that depressed patients had a reduced ability to discriminate the 2 attribution contexts. However, we have previously shown that in healthy controls mean RT in trials following an SA feedback is longer than in trials following an EA feedback (independent of valence).34 This post-SA feedback RT slowing is a behavioural measure indicating that participants were able to discriminate feedback attribution. Therefore, RTs for post-SA and post-EA gains and losses were also analyzed using RM-ANOVA, with valence and attribution as within-subjects factors and diagnosis as an independent factor.

Image processing was carried out using Statistical Parametric Mapping software version 8 (Wellcome Department of Imaging Neuroscience). The preprocessing of functional images included motion correction, coregistration to a standard template, alignment to the first volume for each participant, spatial normalization to the Montreal Neurological Institute (MNI) template (voxel size \( 2 \times 2 \times 2 \) mm), and smoothing using an 8 mm full-width at half-maximum Gaussian kernel. Statistical analysis was performed by modeling the different conditions convolved with a hemodynamic response function and its temporal derivative as explanatory variables within the context of the general linear model on a voxel-by-voxel basis. Our \( 2 \times 2 \) factorial design independently manipulated the agency and valence of feedback. Four regressors corresponding to the 4 feedback conditions of interest (SA losses, EA losses, SA gains, EA gains) were modelled as well as several covariates of no interest (missed feedback, realignment parameters, time of the motor response; Appendix 1). Feedback was modelled as an event (duration = 0) and occurred 4.13 s after the onset of the trial. These regressors of no interest explained additional variance in the blood oxygen level–dependent (BOLD) signal.35

A fixed-effect model at a single-participant level was specified, giving images of parameter estimates for each contrast that were then used for a second-level random effects analysis (Appendix 1). We entered individual participant contrasts into a second-level analysis to examine group differences in activation using 2-sample \( t \) tests with age, sex and years of education as covariates. Activation differences were identified with a family-wise error (FWE)–corrected at peak level for multiple parameter contrasts over space (i.e., the chance of finding an activation with this amplitude or a greater amplitude in a manner that controls for multiple comparisons over space; Appendix 1). For the small volume–corrected analyses, we created a functional mask of the left and right VS using the gains – losses contrast across all participants (Appendix 1, Fig. S2). We calculated the mean percent signal change using MarsBar from the SPM toolbox.

Results

Participants

We included 25 healthy controls and 21 unmedicated patients with MDD in our study. All participants were right-handed and had normal or corrected-to-normal vision. Groups did not differ in sex distribution, age, years of education or performance in the motion prediction task (Table 1). Fourteen patients with MDD were medication-naive, and 3 stopped treatment with antidepressant medications 6 weeks or more before the study. All other patients had been free from medications for at least 2 years. Six patients with MDD had comorbid anxiety disorders: 2 had social phobia (1 of whom also had an eating disorder), 1 had generalized anxiety disorder (GAD) and 3 had panic disorder (1 of whom also had GAD).

Behavioural data

Demographic, psychometric and behavioural data, including mean RT for correct and incorrect trials (across SA and EA trials) and mean RT for trials following SA or EA feedback are presented in Table 1.

The mean RT for incorrect trials was significantly longer than for correct trials \( (F_{1,44} = 23.1, p < 0.001) \), but we found no effect of diagnosis \( (p = 0.45) \) or diagnosis \( \times \) correctness interaction \( (p = 0.74) \). Moreover, on trials following SA feedback (post-SA losses/gains), mean RT was longer than for trials following EA feedback (post-EA losses/gains) \( ; F_{1,44} = 11.81, p = 0.002 \). However, we found no effect of diagnosis \( (p = 0.44) \) or diagnosis \( \times \) attribution \( (p = 0.17) \), diagnosis \( \times \) valence \( (p = 0.54) \) or diagnosis \( \times \) attribution \( \times \) valence interactions \( (p = 0.38) \).

Imaging data

Whole-brain analyses investigating the agency × valence interaction effects

In healthy controls, the 1-sample \( t \) test for the contrast \( (SA gains – EA gains) – (SA losses – EA losses) \) showed a significant activation in the right ventral putamen \( (p = 0.012_{\text{FWE}} \) at peak level, \( t = 7.59 \), MNI space \( x, y, z = 20, 12, -14 \) \). The reported coordinates index the location of the peak (local maxima). In contrast, we found no significant activation in depressed patients even when a lower statistical threshold (uncorrected \( p < 0.005 \) ) was investigated.

When comparing depressed patients and healthy controls, the 2-sample \( t \) test revealed a trend toward a significant whole-brain difference in the right ventral putamen \( (p = 0.05_{\text{FWE}} \) at peak level, \( t = 5.51 \), MNI space \( x, y, z = 28, 8, -8 \) \). When we conducted a small volume–corrected analysis we found a significant difference in the right VS \( (p = 0.002_{\text{FWE}} \) at peak level, \( t = 4.58 \), MNI space \( x, y, z = 26, 6, -10 \) ) but no significant difference in the left VS. Figure 1 displays the significant 15 voxels \( (p < 0.05_{\text{FWE}} \) at peak level) in the right ventral putamen along with the mean percent signal change for each feedback condition.

When we repeated the analysis excluding all the depressed patients with comorbid anxiety disorders, a significant effect in the right ventral putamen was still present (Appendix 1).
No correlations emerged between mean percent signal change in any of the 4 feedback conditions and depression severity (BDI, IDS) or anhedonia and hopelessness scores (SHAPS, H-Scale).

Small volume–corrected analyses investigating the simple effects of attribution in the VS
To further investigate the interaction effect reported previously, we conducted post hoc between-groups analyses of the simple effects of attribution. The 2-sample t test showed a significant effect for the contrast SA losses – EA losses (p = 0.049,ν at peak level, t = 3.33, MNI space x, y, z = 22, 4, –8). We found no effect for the contrast SA gains – EA gains.

Whole-brain analyses investigating the main effect of valence and attribution
We found no significant between-group differences, and the results for each diagnostic group separately are reported in Appendix 1, Tables S1 and S2.

Model-based prediction error analyses
As the VS is considered to signal reward prediction error and altered prediction error signal in the striatum has been reported in patients with MDD, we also considered a model-based prediction error analysis to further investigate whether the diagnosis × agency × valence interaction effect found in the ventral putamen was due to prediction error differences in patients with MDD (Appendix 1). For each feedback condition, we included a parametric modulator that coded trial-by-trial prediction errors (as a first-level covariate). However, the diagnosis × agency × valence interaction effect described previously remained significant when the prediction errors were included as covariates. Conversely we found no significant differences in prediction error magnitude between conditions. This suggests that all interesting variance was captured by a qualitative comparison of our factorial conditions.

Discussion
The present study investigated how feedback attribution influences neuronal feedback processing in patients with MDD. We hypothesized that healthy controls would show a VS response modulated by valence and attribution, whereas depressed patients would have a VS response less sensitive to feedback attribution, leading to an altered processing of SA gains and losses. Consistent with our hypothesis, we found altered striatal response in depressed patients that seems to reflect a failure to modulate the putamen activity according to the feedback’s attribution, leading to an altered putamen activity during the processing of SA losses – EA losses in patients compared with controls. In contrast, we found no evidence that MDD was associated with an altered VS response to gains.

The VS is involved in several processes that are relevant when evaluating feedback, including signalling reward prediction error, incentive motivation and motivational salience. Disentangling the contribution of these different processes was not the aim of our study. However, we conducted a supplementary

Table 1: Demographic and clinical characteristics of the study sample

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>MDD, n = 21</th>
<th>Control, n = 25</th>
<th>Statistic</th>
<th>p value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Male sex</td>
<td>14 (67)</td>
<td>12 (52)</td>
<td>χ² = 1.0</td>
<td>0.37</td>
</tr>
<tr>
<td>Age, yr</td>
<td>39.3 ± 13.1</td>
<td>33.5 ± 10.4</td>
<td>t = 1.7</td>
<td>&gt; 0.99</td>
</tr>
<tr>
<td>Education, yr</td>
<td>16.4 ± 2.6</td>
<td>15.9 ± 2.6</td>
<td>t = 0.6</td>
<td>0.58</td>
</tr>
<tr>
<td>BDI-II score</td>
<td>25.4 ± 8.4</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>IDS score</td>
<td>33.4 ± 8.3</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Single MDD episode</td>
<td>5 (24)</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Never medicated</td>
<td>14 (67)</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Comorbid anxiety disorder</td>
<td>6 (29)</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>STAI-Trait score</td>
<td>56.3 ± 10.7</td>
<td>33.0 ± 9.0</td>
<td>t = 7.9</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>SHAPS score</td>
<td>3.4 ± 3.0</td>
<td>0.5 ± 1.1</td>
<td>t = 4.5</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>H-Scale score</td>
<td>73.9 ± 12.9</td>
<td>44.3 ± 10.2</td>
<td>t = 8.7</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>PANAS positive score</td>
<td>10.5 ± 3.2</td>
<td>13.6 ± 3.9</td>
<td>t = 2.9</td>
<td>0.006</td>
</tr>
<tr>
<td>PANAS negative score</td>
<td>7.7 ± 2.2</td>
<td>5.6 ± 0.9</td>
<td>t = 4.7</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>%SA gains</td>
<td>61.2 ± 7.8</td>
<td>59.4 ± 6.7</td>
<td>t = 0.8</td>
<td>0.41</td>
</tr>
<tr>
<td>%EA gains</td>
<td>59.3 ± 6.1</td>
<td>58.3 ± 6.9</td>
<td>t = 0.5</td>
<td>0.60</td>
</tr>
<tr>
<td>%Miss</td>
<td>2.1 ± 3.3</td>
<td>1.5 ± 3.3</td>
<td>t = 0.6</td>
<td>0.53</td>
</tr>
<tr>
<td>RT correct</td>
<td>542.3 ± 162.7</td>
<td>516.2 ± 97.2</td>
<td>t = 0.6</td>
<td>0.50</td>
</tr>
<tr>
<td>RT incorrect</td>
<td>578.5 ± 139.8</td>
<td>547.8 ± 112.8</td>
<td>t = 0.8</td>
<td>0.41</td>
</tr>
<tr>
<td>RT post-SA gain</td>
<td>566.6 ± 149.2</td>
<td>529.7 ± 111.8</td>
<td>t = 1.0</td>
<td>0.34</td>
</tr>
<tr>
<td>RT post-SA loss</td>
<td>560.8 ± 174.5</td>
<td>525.1 ± 101.2</td>
<td>t = 0.9</td>
<td>0.39</td>
</tr>
<tr>
<td>RT post-EA gain</td>
<td>531.5 ± 139.4</td>
<td>519.8 ± 109.3</td>
<td>t = 0.3</td>
<td>0.75</td>
</tr>
<tr>
<td>RT post-EA loss</td>
<td>542.2 ± 155.5</td>
<td>512.2 ± 100.0</td>
<td>t = 0.8</td>
<td>0.43</td>
</tr>
</tbody>
</table>

BDI-II = Beck Depression Inventory II; EA = externally attributed; H-Scale = Hopelessness Scale; IDS = Inventory of Depressive Symptomatology; MDD = major depressive disorder; PANAS = Positive and Negative Affect Scale; RT = reaction time; SA = self-attributed; SD = standard deviation; SHAPS = Snaith–Hamilton Pleasure Scale; STAI = Spielberger Trait Anxiety Inventory.

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analysis that strongly suggests prediction error signals did not contribute to our VS results (Appendix 1). Consistent with our results in healthy controls, personal responsibility for positive outcomes has been associated with greater VS response.\textsuperscript{32,33} Moreover, in our tasks only SA feedback provided information about performance, thus it is likely that controls processed SA feedback as more salient than EA feedback, leading to a VS response modulated by feedback valence and attribution. In contrast, the internal bias of having no control over the environment may have led depressed patients to an altered salience attribution and consequently a VS response insensitive to feedback attribution. This interpretation is consistent with recent findings showing altered functional connectivity between the salience network and the striatum in unmedicated depressed patients.\textsuperscript{50} More importantly, we have recently investigated functional alterations present at rest and during performance of the motion prediction task in a subgroup of individuals involved in the present study using independent component analysis. We found that 1 intrinsic network showed greater amplitude of low-frequency fluctuations specifically during task performance in depressed patients compared with healthy controls (unpublished data, 2014). This network included the anterior insula and the ventrolateral prefrontal cortex. Considering the role of these brain regions in processing salient events,\textsuperscript{51,52} these results strongly suggest that differences in motivational salience may contribute to the altered VS response in patients with MDD in the present study.

Interestingly, we have recently reported an altered striatal response to feedback attribution in healthy controls homozygous for the Val allele of the brain-derived neurotrophic factor (BDNF) Val66Met polymorphism.\textsuperscript{53} Whereas Met carriers showed a VS response modulated by feedback valence and attribution, Val/Val carriers showed a VS response insensitive to feedback attribution that led to an altered VS response to SA losses – EA losses, which is similar to our findings in depressed patients. Moreover, increased striatal response to performance-dependent monetary losses has been reported in healthy adolescents characterized as behaviourally inhibited,\textsuperscript{54} a temperamental trait associated with an increased risk for anxiety disorder early in life.\textsuperscript{55} As trait anxiety and neuroticism, known risk factors for MDD, are reportedly higher in Val/Val than in Met carriers,\textsuperscript{56–59} these results may lead to the speculation that altered motivational salience processing of SA losses is present in populations at risk for MDD. Future studies are warranted to investigate this hypothesis.

Reduced striatal response to positive feedback has previously been reported in depressed patients.\textsuperscript{15,18,19} In contrast, we found no evidence of a blunted striatal response to gains (but rather a potential increase for EA gains). However, greater activity in the right putamen has been also found by Smoski and colleagues\textsuperscript{22} during the anticipation of monetary rewards compared with pleasant images in patients with mild depression. Since financial rewards have a higher incentive than pleasant images or than simply being correct, it is possible that differences in incentive motivation in the present study compared with previous studies contributed to our VS findings. Reduced reward responsiveness has also been correlated with anhedonia scores in depressed and healthy controls.
individuals and with anhedonia symptoms 1 month later. Moreover, reduced striatal reward prediction error signals have been found in patients with MDD and have been shown to correlate with anhedonia scores. However, anhedonia scores in our study were not particularly high on average but were highly variable among patients (Table 1). It remains possible that a blunted response to positive feedback, by which reward prediction errors or not, is more prevalent in depressed individuals with high anhedonia scores.

**Limitations**

Several limitations need to be mentioned. First, we included patients with comorbid anxiety disorders, thus it is possible our findings are not representative of functional abnormalities occurring in the entire MDD population. However, when all the depressed patients with comorbid anxiety disorders were excluded from the analysis, a significant valence × attribution effect in the VS was still present (Appendix 1), indicating that comorbid anxiety disorder cannot account for our results. Second, to accurately attribute the financial outcome, participants were required to attend to contextual information presented after their performance but before the outcome (Appendix 1, Fig S1). Although our study was conducted in mildly depressed patients, it is still possible that patients had a reduced ability to discriminate the 2 attribution contexts. We have previously shown in healthy individuals that RTs in trials following SA feedback are longer than in trials following EA feedback. This post-SA feedback slowing is a behavioural measure indicating that participants are able to differentiate feedback attribution. In the present study, we observed post-SA feedback slowing in patients and controls, strongly indicating that patients were able to accurately discriminate feedback attribution. Third, although the target of learning (i.e., 60% expected reward rate) was identical for both agency conditions (SA and EA were matched), the nature of learning was different. During the processing of SA feedback participants learned whether their actions led to a reward (instrumental learning), whereas during EA feedback participants learned the probability of receiving a reward independent of their actions (noninstrumental learning; Appendix 1). Fourth, imaging studies have also demonstrated a clear dissociation between regions involved in the processing of feedback anticipation and outcome, and differences in the hemodynamic response between depressed patients and healthy controls have been reported during both phases of feedback processing. Although this dissociation could not be investigated with our task’s design, a better understanding of the neural network involved in anticipation of controllable and uncontrollable feedback is important to understand altered feedback processing in patients with MDD. Finally, we used an 8 mm smoothing kernel because this is more commonly used in fMRI studies and thus leads to coordinates being more easily comparable with those of previous reports. However, this is likely to bias the spatial localization of ventral striatal responses posteriorly.

**Conclusion**

Our results show that in depressed patients the striatal response to feedback was insensitive to feedback attribution, leading to an altered VS response during SA losses – EA losses compared with healthy controls. Our findings suggest an altered assignment of motivational salience to SA losses in patients with MDD. This altered sensitivity to feedback attribution may reinforce the belief of not being in control of negative events. This cycle is compatible with mechanisms leading to learned helplessness and, translated into clinical terms, appears to be relevant for psychotherapy. A combination of mindfulness and cognitive interventions may raise the awareness for self-determined gains and help correct the depressed individual’s bias in processing of feedback.

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**Competing interests:** None declared.

**Contributors:** J. Chumbley and S. Spinelli designed the study. J. Spätii, N. Doerig and J. Brakovski acquired the data, which J. Chumbley, M. Grosse Holtforth, E. Seifritz and S. Spinelli analyzed. S. Spinelli wrote the article. All authors reviewed and approved it for publication.

**References**


With the exception of the study by Ruchsow et al. (2013) mentioned earlier, the research on the role of the habenular complex and the reward system in clinical depression has been rather limited compared to other brain regions involved in reward processing.

However, recent studies suggest that the habenular complex might play a specific role in the processing of negative affect. For instance, a study by Ruchsow et al. (2013) using a computational model of the habenular complex found that the structure of the model could influence the symptoms of depression and anxiety. The authors suggested that this may be because the habenula is involved in the processing of negative affect and that its dysfunction could lead to changes in the way the brain processes negative information.

Additionally, a study by Jiang et al. (2014) found that the habenula is involved in the processing of negative affect in healthy controls, but not in patients with depression. This suggests that the habenula may be more involved in the processing of negative affect in patients with depression, which could explain why the habenula is often involved in the control of negative affect in psychiatric disorders such as depression.

In conclusion, the role of the habenula in reward processing and its potential involvement in the processing of negative affect in depression warrants further investigation. The results of these studies highlight the importance of considering the habenula in the understanding and treatment of psychiatric disorders, particularly in the context of reward processing and negative affect.

References: