Thalamostriatal Interactions in Human Reversal Learning

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ABSTRACT

Cognitive flexibility, the ability to flexibly update thought and behaviour, is a crucial component of adaptation and survival, and it is disrupted in neuropsychiatric disorders and psychopathology. Reversal learning tasks are often used to investigate cognitive flexibility. Human studies of reversal learning typically focus on neural mechanisms required to identify changes in response contingencies, rather than on mechanisms required to learn and express a new response. However, animal studies of reversal learning have shown a key and specific role in this process for the dorsal striatum and its interactions with the centromedian parafascicular (CM-Pf) thalamus, but this system is not as well understood in humans. The aim of this study was to investigate the role of the human dorsal striatum and its interactions with the thalamus during probabilistic reversal learning, specifically with respect to learning a new response strategy. We used psychophysiological interaction (PPI) analysis of functional magnetic resonance imaging (fMRI) data, to measure task-dependent changes in connectivity between striatal and thalamic functional subdivisions. We show that connectivity between the dorsal striatum and the centromedian parafascicular (CM-Pf) thalamic complex was increased during reversal, but not initial, learning. The strength of this connectivity was associated with the ability to flexibly alter behaviour. This study helps to bridge the gap between animal studies of this system, and human studies of reversal learning and cognitive flexibility more generally.
SIGNIFICANCE STATEMENT

Cognitive flexibility is an important ability, enabling us to adapt to changes in the environment. Recent evidence from animal studies highlights the contribution of specific connections between the thalamus and the striatum in reversal learning, a key paradigm for the study of flexibility. This system remains understudied in humans. Using neuroimaging and computational modelling, we show that thalamostriatal connectivity increases specifically during reversal learning. The strength of this connectivity was associated with the ability to flexibly alter behaviour. In line with the animal literature, we show that this mechanism supports new learning, rather than the detection of the reversal *per se*. This highlights the separate contribution of thalamostriatal connectivity to flexibility compared to the better-understood role of the prefrontal cortex.
INTRODUCTION

Cognitive flexibility, the ability to alter behaviour in response to changes in environmental conditions, is crucial for adaptation and survival, and is disrupted in numerous neuropsychiatric disorders, such as Parkinson’s disease, Alzheimer’s disease, Huntington’s disease, autism, obsessive compulsive disorder and schizophrenia (Nilsson et al., 2015; Prado et al., 2017). Cognitive flexibility is typically measured using reversal learning paradigms, where a stimulus previously predictive of reward becomes irrelevant or starts to predict punishment, and another stimulus now becomes relevant for guiding behaviour (Nilsson et al., 2015). Evidence from the animal literature suggests a key specific role for thalamostriatal connectivity in reversal learning. For example, studies involving rodents have shown that disrupting connectivity between the centromedian-parafascicular nucleus (CM-Pf) of the thalamus and the dorsal striatum (DS) results in impaired reversal learning, whilst initial learning remains intact (Brown et al., 2010; Bradfield et al., 2013). The CM-Pf provides the main external input to the striatal cholinergic interneurons (CINs) (Ellender et al., 2013). Disruption of this connectivity has been shown to disrupt cholinergic signalling, which in turn results in impaired reversal learning. This impairment is specifically related to regressive errors after the reversal has been identified (rather than to identifying the occurrence of the reversal *per se*), suggesting interference between old and new learning (Bradfield et al., 2013). More specifically, evidence suggests that thalamostriatal connectivity modulates the gating of corticostriatal synapses by cholinergic activity (Smith et al., 2014). This system is of particular interest as it is implicated in pathological cognitive inflexibility, as observed, for example, in neurodegenerative disorders (Smith et al., 2014) and schizophrenia (Holt et al., 1999, 2005). For instance, patients with Parkinson’s disease, progressive supranuclear palsy and Huntington’s disease show significant neuronal loss in the CM-Pf (Henderson et al., 2000).

In humans, functional magnetic resonance imaging (fMRI) studies show activity related to reversal learning in frontal cortical areas, such as the orbitofrontal and anterior cingulate cortex (Cools et al., 2002; Remijnse et al., 2005; D’Cruz et al., 2011; Waegeman et al., 2014). These studies typically use two stimuli and multiple reversals, often focusing on activity during errors made after the contingency reversal has occurred, but before a change in behaviour has been established. Although this is relevant to the mechanism through which the reversal itself is identified, it provides less information on the mechanism required for learning and expressing a new behaviour. By contrast, multi-alternative tasks enable the study of processes relating to new, post-reversal learning. The use of such a task is an important element of this study, given recent evidence on how cholinergic-thalamostriatal interactions may contribute to flexibility as outlined above. Indeed, using proton magnetic resonance spectroscopy, we have recently shown task-related changes in cholinergic activity in the human DS during probabilistic reversal learning in such a task (Bell et al., 2017).

The aim of this study was to investigate the role of thalamostriatal interactions during human probabilistic reversal learning. We used high-resolution multiband fMRI in combination with psychophysiological interaction analysis (PPI) to test for changes in connectivity between the CM-Pf and the DS during reversal learning in a multi-alternative decision-making task. Based on prior evidence from animal models (Bradfield et al., 2013), we hypothesised that CM-Pf and DS activation would correlate during reversal, but not initial,
learning. To demonstrate the specificity of this result, we used the mediodorsal (MD) thalamus as a control region. The MD thalamus also projects to the striatum (Haber and Calzavara, 2009). However, lesions here result in an increase in perseverative behaviour during reversal learning (Chudasama et al., 2001), rather than an increase in regressive behaviour typically seen with CM-Pf lesions, suggesting that this system contributes to a different mechanism during reversal learning. Importantly, the MD thalamus does not project to the striatal cholinergic system (Gonzales and Smith, 2015), which is thought to be targeted by CM-Pf DS connectivity during reversal learning (Bradfield et al., 2013).
MATERIALS AND METHODS

Participants

The study was approved by the University of Reading Research Ethics Committee. 57 volunteers (20 female) between the ages of 18.5 and 30.6 (mean = 22.7, SD = 3.6) were recruited by opportunity sampling. All participants were healthy, right handed non-smokers and written informed consent was taken prior to participation.

One participant was excluded due to technical issues during data collection. 34 participants were excluded from the analysis reported here as they did not reach the task learning criteria specified below. 22 participants reached the specified learning criteria in both rounds and were included for analysis (12 female; mean age = 22.5, SD = 3.8).

Behavioural Data

Learning Task

The task was a probabilistic multi-alternative reinforcement learning task with a reversal components, described previously (Bell et al., 2017). It was programmed using MATLAB (2014a, The Mathworks, Inc., Natick, MA, United States) and Psychtoolbox (Brainard, 1997).

First, participants were presented with a fixation cross displayed in the centre of the visual display followed by four decks of cards. Each deck contained a mixture of winning and losing cards, corresponding respectively to a gain or loss of 50 points. The probability of getting a winning card differed for each deck (75%, 60%, 40%, and 25%) and the probabilities were randomly assigned across the four decks for each participant. Participants indicated their choice of deck by pressing the corresponding button on a button box. Outcomes were pseudo-randomised so that the assigned probability was true over every 20 times that deck was selected. Additionally, no more than 4 cards of the same result (win/loose) were presented consecutively in the 75% and 25% decks and no more than 3 cards of the same result in the 60% and 40% decks. A cumulative points total was displayed in the bottom right-hand corner throughout the session and in the centre of the visual display at the end of each trial (Figure 1A). Participants were instructed that some decks may be better than others, they are free to switch between decks as often as they wish, and they should aim to win as many points as possible.

The learning criterion was set as selection of either of the two highest decks on at least 80% of 20 consecutive trials. As the research question focused on the reversal, we wanted to encourage behavioural stability before the reversal to reduce intra-individual noise. Therefore, a “stability phase” was included at the end of the initial learning phase. The number of trials in this phase was equal to 60% of the number of trials taken to reach criterion. At the end of this phase the deck probabilities were reversed so that the high probability decks became low probability, and vice versa. Participants were not informed of the reversal. After reaching the learning criterion again, participants completed a second stability phase, again equal in
number to 60% of the number of trials taken to reach criterion after reversal. After this phase, the task ended (Figure 1B).

**Figure 1. Trial and task design**

![Figure 1](image)

**A.** Participants were instructed to choose between four decks of cards. Each deck had a different probability of generating winning cards (75%, 60%, 40%, and 25%). Once the predetermined learning criterion had been reached, the deck probabilities were reversed so that high probability decks became low probability decks and vice versa. Participants were not informed of this in advance and were simply instructed to gain as many points as possible. The time from initial deck presentation to deck choice is the decision making epoch referred to in the analysis. The length of time the feedback was displayed was the feedback epoch referred to in the analysis. RT = reaction time; SD = standard deviation. **B.** Schematic of the four task phases. Upon reaching criterion in the initial learning phase, participants completed a post criterion stability phase (lasting for 60% of the trials-to-criterion during initial learning). After this phase, the deck probabilities were reversed. Participants then completed a post-reversal learning phase and upon reaching criterion again, they completed another post criterion stability phase (lasting for 60% of the trials-to-criterion during reversal learning).

Participants were given 100 trials to reach criterion in both the initial learning and reversal phase. If participants did not reach criterion in the initial learning phase, they did not experience the reversal. The rationale was that participants who had not reached criterion during initial learning would likely not identify a change in contingencies during the reversal and therefore would not show a change in choice behaviour following a contingency reversal (at least not in a manner straightforwardly comparable with participants...
who had reached criterion). Following the reversal, participants were allowed 100 trials to reach criterion and enter the post-reversal stability phase, after which the task ended (Figure 1B).

The presentation timings were jittered. The stimuli were displayed for between 0.8 and 2.8s, with an average display time of 1.7s (standard deviation = 0.6s). Each trial lasted, on average, 8.3s (standard deviation = 1.3s). After the scanning session, participants were asked to rank the decks in order of preference from 1 to 4, with 4 being the best deck. Participants were instructed to give multiple answers if they thought the rankings changed. Participants were also asked to provide an estimate of the probability of winning on each deck using the numbers 1-100. As before, participants were instructed to give multiple answers if they thought the probabilities changed.

Performance was measured using the number of trials taken to reach criterion during the initial learning and reversal phases. Perseverative errors were defined as the trials after reversal until the probability of selecting the previously favoured deck reached chance level (0.25), i.e. the number of trials taken to identify the reversal and switch behaviour. Regressive errors were defined as selections of the previously favoured deck after the perseverative period had ended.

**Temporal Difference Reinforcement-Learning (TDRL) Model**

We modelled participants’ choice behaviour as a function of their previous choices and rewards using a TDRL algorithm (Sutton and Barto, 1998). This allows us to track trial-and-error learning for each participant, during each task phase, in terms of a subjective expected value for each deck. On each trial \( t \), the probability that deck \( c \) was chosen was given by a soft-max probability distribution,

\[
P(c_t = c) = \frac{e^{m_t(c)}}{\sum_j e^{m_t(j)}}
\]

where \( m_t(c) \) is the preference for the chosen deck and \( j \) indexes the four possible decks. The preference for the chosen deck was comprised of the participant’s expected value of that deck on that trial, \( V_t(c) \), multiplied by the participant’s individual value impact parameter \( \beta \) (equivalent to the inverse temperature):

\[
m_t(c) = \beta V_t(c)
\]

The parameter \( \beta \) describes the extent to which trial-by-trial choices follow the distribution of the expected values of the decks: a low \( \beta \) indicates choices are not strongly modulated by expected value, being effectively random with respect to this quantity (i.e. participants are not choosing based exclusively on value, indicating exploration of the available options); conversely, a high \( \beta \) indicates choices largely follow the expected value (i.e. participants choose the deck with the highest expected value; exploitation).

To update the subjective value of each deck, a prediction error was generated on each trial, \( pe_t \), based on whether participants experienced a reward or a loss (\( reward_t = +1 \) or \(-1 \) respectively). The expected value of the chosen deck was subtracted from the actual trial reward to give the prediction error,

\[
pe_t = reward_t - V_t(c)
\]

It has been shown that individuals differ in the degree to which they learn from better than expected outcomes (positive prediction errors) and worse than expected outcomes (negative prediction errors), (Gray,
1970; Niv et al., 2012; Christakou et al., 2013a; Bull et al., 2015). To account for this, two learning rate parameters were used to model sensitivity to prediction errors in updating the expected values: the weight of learning from better than expected outcomes (learning rate from positive prediction errors: $\eta^+$) and the weight of learning from worse than expected outcomes (learning rate from negative prediction errors: $\eta^-$). For example, individuals who are reward seeking will place a high weight on the former, whereas those who are loss-aversive will place a high weight on the latter. The prediction error on each trial was multiplied by either the positive ($\eta^+$) or negative ($\eta^-$) learning rate and used to update the value of the chosen deck.

\[
\delta_t = \eta^+ \times pe_t \quad \text{if } pe_t > 0
\]

\[
\delta_t = \eta^- \times pe_t \quad \text{if } pe_t < 0
\]

\[
V(\text{chosen}_t) = V(\text{chosen}_{t-1}) + \delta_t
\]

Thus, the model has three parameters of interest ($\beta$, $\eta^+$ and $\eta^-$). In psychological terms, $\beta$ captures the degree to which the subjective value of the chosen deck influenced decisions, while the learning rates capture the individual’s preference for learning from positive ($\eta^+$) or negative ($\eta^-$) prediction errors to guide choice behaviour during this task.

**Model Fitting**

As mentioned previously, individuals differ in the degree to which they learn from different prediction errors (described as learning rate asymmetry), which in turn can affect performance. Generally, it is assumed that the learning rate asymmetry is stable across the learning episode. Reinforcement learning relies on a trade-off between exploration of the available options and exploitation of the optimal choice, which in turn is likely driven by different learning rates. For example, during learning, participants must explore the decks to identify the optimal choice, therefore they should learn from positive and negative prediction errors equally. However, during periods of stability, participants must stick to the optimal deck and ignore minor losses. Therefore, they should place more weight on positive than negative prediction errors. Indeed, there is evidence that agents able to flexibly alter learning rate asymmetry based on reward history perform better on a probabilistic task (Cazé and van der Meer, 2013). To directly test this, the model was fit separately for each task phase (Figure 1B) per participant, providing parameters that maximised the likelihood of the observed choices given the model (individual maximum likelihood fit; Daw, 2011). The calculated deck values from the end of each phase were used as the initial deck values for the following phase, e.g. deck values at the end of the initial learning phase were used as the initial deck values in the first stability phase. There was no difference in the goodness of fit of the model between task phases when accounting for phase differences in trial number (likelihood repeated measures test: $F(3)=0.530$, $p=0.664$, partial eta squared= 0.030). To ensure the model produced consistent, interpretable parameter estimates, $\eta^+$ was limited to values between 0 and 1, and $\beta$ and $\eta^-$ were constrained by the following prior distributions (see Christakou et al., 2013b):

$\beta \sim \text{Gamma} (2,1)$

$\eta \sim \text{Beta} (1.2,1.2)$
Functional Magnetic Resonance Imaging

Data Acquisition

Data were collected using a Siemens Trio 3T MRI scanner with a 32-channel head array coil at the University of Reading. A multi-band echo planar imaging (EPI) sequence was used to acquire data during the learning task (voxel resolution = 1.8×1.8×1.8 mm; interleaved acquisition of 60 axial slices; no slice gaps; matrix size = 128×128, TE/TR=40/810 ms; flip angle 31°; multiband factor 6; partial Fourier factor = 1; bandwidth = 1502 Hz/Pixel). This was followed by the acquisition of a high-resolution whole brain T1 weighted structural image using an MPRAGE sequence parallel to the anterior-posterior commissure (voxel resolution=1x1x1 mm, field of view=250 mm, 192 sagittal slices, TE/TR=3.02/2020 ms, flip angle=9°).

Analysis of Functional Data

Analysis was performed using FSL version 5.0.8. (Stephen M. Smith, Mark Jenkinsona, Mark W. Woolricha, b, Christian F. Beckmannana, Timothy E.J. Behrens, Heidij Johansen-Berga, Peter R. Bannistera, Marilena De Lucaa, Ivana Drobnjaka, David E. Flitneya, Rami K. Niazya, James Saunders, John Vickersa, Yongy, 2004; Jenkinson et al., 2012). First, the brain was extracted from the T1 structural scan using the brain extraction tool (BET) (Smith, 2002). The following pre-statistics processing was used on the functional data: registration to the brain extracted structural scans, followed by registration to 1mm MNI space using FLIRT (Jenkinson and Smith, 2001; Jenkinson et al., 2002); motion correction using MCFLIRT (Jenkinson et al., 2002); brain extraction using BET (Smith, 2002); spatial smoothing (FWHM 3.0mm); high-pass temporal filtering.

Analysis of thalamostriatal interactions

For each participant, the activation time-series were extracted from each left and right thalamic ROI (CM-Pf and MD). Masks were generated based on the co-ordinates from Metzger et al., (2010). They used a behavioural task designed to separate thalamic activation in relation to emotional arousal (MD) versus attention and expectancy (CM-Pf). Based on this work, we created 6mm spherical ROIs surrounding the peak voxel in each thalamic region (MNI coordinates (x,y,z): MD: -9.00,-17.00, 14.00; CM-Pf: -3.78,-16.92, 0.22; Figure 2). The ROIs were then transformed to MNI space and the mean signal was extracted.
A general linear model (GLM) was generated which included four regressors based on the timings for each task epoch (Figure 1A). The regressors were created by convolving a box car function representing the onset and duration of an epoch with an ideal haemodynamic response function. Additionally, the feedback epoch was also modulated by prediction error by including the prediction error for each trial as generated by the TDRL model as a parametric modulator. Positive and negative prediction errors were included in separate regressors. The thalamic ROI timeseries were also included in the GLM, resulting in seven regressors. A separate GLM was created for each area, resulting in four GLMs in total (left and right CM-Pf and MD). FEAT was used to create interaction regressors between the ROI timeseries and the decision making and feedback epochs (Figure 1A; not modulated by prediction error). Higher level analysis was used to generate a group average and contrasts between initial learning and reversal learning. Age was included as a covariate as several brain regions are still maturing in the age range of the sample (Waegeman et al., 2014). Additionally, the number of trials in each phase was included as a covariate to control for different sized data sets. Region of interest analyses were carried out using cluster thresholding ($z=2.3$, $p < 0.05$).

Typically the striatum is divided into three sections based on morphology and cytoarchitecture: caudate, putamen and nucleus accumbens. However, studies of intrinsic functional connectivity have shown the striatum can be divided into more functional territories. For example, a resting state study divided the striatum into 5 functional subdivisions based on corticostriatal connectivity patterns (Choi et al., 2012). Three associative subdivisions were described in the caudate, extending into the anterior putamen; here we sum these three masks for ROI analysis of the associative dorsal striatum (DS). Similarly, a motor ROI was defined in the posterior putamen and a limbic ROI in the ventral striatum using the traditional three-region striatal model included in the FSL atlas (Tziortzi et al., 2014).
Figure 3. Striatal functional subdivisions

Location and extent of the masks for the left anterior caudate (AC; red; MNI coordinates: -12.37, 14.90, -0.27; number of voxels = 2067), dorsal caudate (DC; green; MNI coordinates: -13.51, 6.12, 14.61, number of voxels = 1607), ventral caudate (VC; blue; MNI coordinates: -22.01, 6.66, 2.07; number of voxels = 5208), putamen (Put; pink; MNI coordinates: -28.90, -8.89, 2.57; number of voxels = 2597), and ventral striatum (VS; yellow; MNI coordinates: -11.21, 11.08, -8.45; number of voxels = 1219).

Experimental Design and Statistical Analysis

Statistical analysis was performed using SPSS (IBM Corp. Released 2013. IBM SPSS Statistics for Windows, Version 22.0. Armonk, NY: IBM Corp). As the study used a repeated measures design, repeated-measures ANOVA tests were used to investigate changes in model parameters across task phases. Measures of connectivity were extracted and also compared across task phases. When assumptions of sphericity were violated the Greenhouse-Geisser correction was used. Correlation analyses were used to test the relationship between connectivity and performance. When assumptions of normality were not met we used a non-parametric correlation test. Kendall’s Tau ($\tau$) was used, as it provides a better estimate of the correlation strength in a small sample size compared to other non-parametric methods (Field, 2015).


RESULTS

Task Performance

Twenty two (22) participants reached criterion both during initial learning and after the reversal.

<table>
<thead>
<tr>
<th>Task Phase</th>
<th>Average Number of Trials</th>
<th>SD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Initial Learning (to criterion)</td>
<td>44</td>
<td>22</td>
</tr>
<tr>
<td>First Stability Phase</td>
<td>30</td>
<td>14</td>
</tr>
<tr>
<td>Reversal Learning</td>
<td>40</td>
<td>21</td>
</tr>
<tr>
<td>Second Stability Phase</td>
<td>27</td>
<td>14</td>
</tr>
<tr>
<td>Total</td>
<td>141</td>
<td>58</td>
</tr>
</tbody>
</table>

The behavioural model parameterises aspects of performance that potentially contribute differentially to initial learning compared to reversal. To test this, we compared model parameter estimates across the four task phases (Figure 1B). Figure 4 shows that the impact of the subjective value on decisions ($\beta$) average over the sample increased over time across the four phases. The figure also shows that the relative impact of positive and negative prediction errors on choices ($\eta^+/\eta^-$) changed across task phases: after initial learning, participants relied more on positive prediction errors (i.e. higher $\eta^+/\eta^-$). But when the contingencies reversed, they reverted to weighing positive and negative prediction errors more equally until they learned to criterion, after which the learning ratio increased again.

Specifically, there was a significant effect of task phase on the value impact parameter ($\beta$; $F(1.8,36.8) = 6.236$, $p = 0.006$, Greenhouse-Geisser corrected, partial eta squared = 0.229). Bonferroni-corrected post hoc tests showed that the $\beta$ value during initial learning was significantly lower than the $\beta$ values of all other task phases (first stability phase, $p = 0.004$; reversal learning phase, $p = 0.003$; second stability phase, $p = 0.013$; Figure 4A).

Regarding the learning rate parameters, a repeated measures ANOVA with a Greenhouse-Geisser correction showed a significant effect of task phase on the ratio of learning rates from positive and negative prediction errors ($\eta^+/\eta^-$; $F(1.2,24.7) = 7.043$, $p = 0.011$, partial eta squared = 0.251). Post hoc tests using the Bonferroni correction revealed the ratio of learning rates significantly increased from the initial learning phase to the first stability phase ($p = 0.003$), and again from the first stability phase to the reversal learning phase ($p = 0.015$). The learning rate ratio was also significantly higher in the second stability phase compared to the initial learning phase ($p = 0.026$) (Figure 4B).
There was a significant effect of task phase on both learning rates (learning rate from positive prediction errors, $\eta^+$; $F(3,63) = 7.021, p < 0.001$, partial eta squared = 0.251; learning rate from negative prediction errors, $\eta^-$; $F(3,63) = 7.091, p < 0.001$, partial eta squared = 0.252).

Post hoc tests using the Bonferroni correction revealed learning rates from positive prediction errors differed significantly between the initial learning phase and the second stability phase ($p = 0.001$) and between the reversal learning phase and the second stability phase ($p = 0.031$). Learning rates from positive prediction errors did not differ significantly between the other task phases (Figure 4B). Learning rates from negative prediction errors differed significantly between the initial learning phase and the first stability phase ($p = 0.003$) and between the initial learning phase and the second stability phase ($p = 0.004$). Learning rates from negative prediction errors did not differ significantly between the other task phases (Figure 4B).

Figure 4. Behavioural model parameter estimates across the task phases

A. Significant effect of task phase on value impact parameters ($\beta$; $F(1.8,36.8) = 6.236, p = 0.006$, partial eta squared = 0.229). *$p < 0.05$. B. Significant effect of task phase on the learning rate from positive prediction errors ($\eta^+$; $F(3,63) = 7.021, p < 0.001$, partial eta squared = 0.251) and negative prediction errors ($\eta^-$; $F(3,63) = 7.091, p < 0.001$, partial eta squared = 0.252). These changes are also expressed as the ratio of learning rates, which is used in the analysis.

Analysis of thalamostriatal interactions

There were no significant correlations between activity in either thalamic ROI (CM-Pf or MD) and activity in the DS during initial learning, or during the first stability phase.

During reversal learning, there was a significant correlation between activity in the left CM-Pf and activity in the left DS (corresponding to the ventral caudate (VC) subregion as defined in Choi et al. 2012, see Figure 3) during the feedback epoch (Figure 5A). This result was specific to the CM-Pf, with no significant correlations between activation in the MD and the DS. This striatal cluster was then used as a mask to extract
the average strength of CM-Pf DS connectivity for each task phase. A repeated-measures ANOVA showed a significant main effect of task phase on connectivity (F(3,63) = 5.510, p = 0.002, partial eta squared = 0.208). Connectivity was significantly higher during reversal learning compared to all task phases (initial learning phase, F(1,21) = 11.789, p = 0.002, partial eta squared = 0.360; first stability phase, F(1,21) = 7.015, p = 0.015, partial eta squared = 0.250; second stability phase, F(1,21) = 9.631, p = 0.005, partial eta squared = 0.314; Figure 5B). By contrast, there was no significant effect of task phase on connectivity between the left MD and DS (F(3,63) = 1.145, p = 0.338, partial eta squared = 0.052; Figure 5C).

Our computational model captures the mechanism of reversal learning in the observed reduction in the learning rate ratio from the stability phase to the reversal phase. To test the hypothesis that the strength of CM-Pf DS connectivity would facilitate this mechanism, we examined the correlation between the strength of the connectivity and the drop in learning rate ratio. A higher level of connectivity was weakly associated with a larger decrease in learning rate ratios ($t_{(22)} = -0.255, p = 0.048$ one tailed; Figure 5D). A higher level of connectivity was further associated with a smaller number of regressive errors ($t_{(22)} = -0.415, p = 0.010$; Figure 5E), in line with evidence from animal studies. There was no association between connectivity and perseveration ($t_{(22)} = 0.009, p = 0.954$).

These effects were specific to the CM-Pf and dorsal striatum connectivity. There were no significant associations between activity in either thalamic ROI (CM-Pf or MD) and activity in either the putamen or ventral striatum in any task phase.
Figure 5.

A. Group average map of the PPI analysis results showing a significant correlation during the feedback epoch in reversal learning between activation in the left DS (specifically ventral caudate) and the left CM-Pf (SVC p < 0.05; MNI coordinates centre: x = -22.7, y = 2.16, z = 5.59; number of voxels = 151; max Z = 3.01).

B. Significant effect of task phase on connectivity between the left CM-Pf and the left dorsal striatum (F(3,63) = 5.510, p = 0.002, partial eta squared = 0.208). Connectivity is significantly higher during the reversal learning phase compared to all other task phases.

C. There was no significant effect of task phase on connectivity between the left MD and the DS (F(3,63) = 1.145, p = 0.338, partial eta squared = 0.052).

D. The change in the ratio of learning rates (η+/ η-) from the first stability phase (following initial learning) to the reversal learning phase was associated with the strength of the connectivity between the left CM-Pf and the left dorsal striatum (t(22) = -0.255, p = 0.048, one tailed).

E. The number of regressive errors made during the reversal phase was associated with the strength of the connectivity between the left CM-Pf and the left DS (t(22) = -0.425, p = 0.010). CM-Pf: Centro-median parafascicular thalamic complex; DS: Dorsal Striatum; MD: Mediodorsal Thalamus; error bars represent the standard error; *p < 0.05.
DISCUSSION

We investigated the role of thalamostriatal connectivity in human reversal learning using a combination of connectivity analysis and computational modelling. We show enhanced connectivity between the CM-Pf and the DS that is specific to reversal learning. Moreover, the strength of the connectivity correlated with the ability to flexibly alter behaviour during probabilistic reversal.

Evidence from the animal literature has shown that disrupting connectivity between the CM-Pf and the dorsomedial (associative) striatum results in impaired reversal learning, whilst initial learning remains intact. The impact of such a manipulation on reversal learning is specific: animals are able to identify the reversal, but display interference from the initial learning when learning and expressing the new behaviour. This effect is thought to be driven by disruption of input specifically to the cholinergic system in the striatum (Brown et al., 2010; Bradfield et al., 2013). Our study provides evidence for this effect in humans. We have shown a significant correlation between activity in the CM-Pf and the dorsal striatum during reversal learning only, with no significant correlations during initial learning, or during the post-criterion stability phases. The strength of this connectivity was significantly higher during reversal learning compared to all task phases, in line with evidence from the animal literature. By contrast, there was no significant correlation between activity in the mediodorsal thalamus and the dorsal striatum, and no significant change in connectivity between these regions across task phases. The mediodorsal thalamus also projects to the dorsal striatum (Haber and Calzavara, 2009), but it does not project to the striatal cholinergic system (Gonzales and Smith, 2015), which is thought to interface the CM-Pf influence into corticostriatal function (Smith et al., 2014). Indeed, in a recent study using the same task we showed, for the first time in humans, evidence of cholinergic recruitment in the same dorsal striatal region during reversal learning (Bell et al., 2017). Together, these observations are in line with the notion that the change in CM-Pf-dorsal striatal connectivity we observed relates to a change in recruitment of the striatal cholinergic system.

We also show changes in learning rate asymmetry (measured here using the ratio of learning rates from positive and negative prediction errors) across the task. It has been previously shown that agents able to flexibly alter learning rate asymmetry based on reward history perform better on a probabilistic task (Cazé and van der Meer, 2013). However, the contribution of learning rate asymmetry to different stages of learning during a probabilistic reversal learning task has not been directly investigated. Using a reinforcement learning model, we looked at changes in the ratio between learning from positive and negative prediction errors throughout the task. During initial learning, participants placed equal weights on positive and negative prediction errors to identify which decks provide overall wins and which decks provide overall losses, making the ratio of the two learning rates small. By the first stability period, after criterion has been reached, participants have identified the optimal deck and are able to ignore any losses. Consequently, they increased the weight of learning from positive prediction errors and decreased the weight of learning from negative prediction errors, resulting in an increase in the ratio. During the reversal learning phase, participants start receiving more negative feedback. If they are to identify that this is no longer experienced as part of the learned probabilistic structure, but rather that the contingencies have changed, participants must
increase the weight of learning from negative prediction errors (thereby decreasing the ratio of the learning rates during reversal learning). This way, attending to worse than expected outcomes provides the opportunity to adaptively dismantle confidence in the previously learned response, making the change in learning rate ratio an important marker of reversal learning efficiency. During the last stability period, participants will have again identified the optimal deck and may once again ignore any losses, resulting in an increase in the learning rate ratio. Therefore, by continuously updating the learning rates based on feedback, participants are able to adapt to alterations in task structure. This is an important component of reversal learning, and cognitive flexibility more generally, and provides an insight into the more nuanced skills required for the ability to flexible alter behaviour.

Moreover, we found evidence for a relationship between the ability to flexibly alter behaviour and the strength of the CM-Pf-DS connectivity. Our results show a relationship between the drop in the learning rate ratio from the first stability phase to the reversal learning phase, and the strength of the CM-Pf-dorsal striatal connectivity, suggesting that connectivity between these two regions supports the ability to flexibly alter behaviour. Those with weaker connectivity demonstrated a smaller drop, which could potentially represent interference from the first stability phase, in which participants needed to place more weight on learning from positive prediction errors, and therefore had higher learning rate ratios. Relatedly, we also showed that those with weaker connectivity made more regressive errors, while there was no association between connectivity and perseveration. This is in line with evidence from the animal literature, which has shown that disrupting CM-Pf-dorsal striatal connectivity results in an increase in the number of regressive errors, whilst there is no effect on perseveration (Bradfield et al., 2013). This is thought to represent interference between new and existing learning, resulting from an inefficient partition of the conflicting contingencies into separate internal states or contexts (Bradfield and Balleine, 2017). It has been suggested that the initial and reversed contingencies are encoded within separate pools of neurons within the dorsal striatum. CM-Pf controlled cholinergic modulation may be used to select the appropriate pool of neurons for encoding and action selection based on the internal state (Stalnaker et al., 2016; Bradfield and Balleine, 2017).

In summary, we used connectivity analysis and computational modelling to investigate the role of thalamostriatal interactions in human reversal learning. We show increased connectivity between the CM-Pf and the dorsal striatum during reversal learning, the strength of which is associated with specific aspects of the ability to flexibly alter behaviour. This study helps to bridge the gap between animal studies of this system, and human studies of reversal learning and cognitive flexibility more generally, and highlights the contribution of thalamostriatal connectivity.
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