Habitual versus goal-directed action control in Parkinson disease

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Habitual versus Goal-directed Action Control in Parkinson Disease

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Abstract

This study presents the first direct investigation of the hypothesis that dopamine depletion of the dorsal striatum in mild Parkinson disease leads to impaired stimulus–response habit formation, thereby rendering behavior slow and effortful. However, using an instrumental conflict task, we show that patients are able to rely on direct stimulus–response associations when a goal-directed strategy causes response conflict, suggesting that habit formation is not impaired. If anything our results suggest a disease severity–dependent deficit in goal-directed behavior. These results are discussed in the context of Parkinson disease and the neurobiology of habitual and goal-directed behavior.

INTRODUCTION

If an act became no easier after being done several times, if the careful direction of consciousness were necessary to its accomplishment on each occasion, it is evident that the whole activity of a lifetime might be confined to one or two deeds (...). A man might be occupied all day in dressing and undressing himself (Maudsley, 1876, p. 155).

Since Maudsley (1876) wrote these words on the importance of habit formation, it has been proposed by many psychologists that instrumental behavior becomes habitual with extensive practice (Bolles, 1972; Kimble & Perlmutter, 1970; Tolman, 1932; James, 1890). The underlying associative mechanism was first described by Thorndike (1911) as a gradual stamping in of associations between contextual stimuli (S) and responses (R) that lead to rewarding outcomes (O). Via these direct S→R associations, instrumental actions can be activated with minimal cognitive effort, thereby freeing up cognitive resources. The ability to form habits allows for fast selection of appropriate responses in stable contexts and therefore plays a crucial role in much of our everyday decision-making.

Although the notion of habit formation has been around for a long time, a gradual shift from internal to external control over behavior with practice was not demonstrated experimentally until the 1980s. Adams (1982) showed that after extensive training, instrumental behavior of rats loses its direct sensitivity to the incentive value of the outcome, suggesting a transition from goal-directed behavior mediated by O→R associations1 to S→R habits directly driven by external cues (de Wit, Corlett, Aitken, Dickinson, & Fletcher, 2009). This overtraining paradigm was used only very recently to demonstrate that in humans, as in animals, practice leads to the development of behavioral autonomy (Tricomi, Balleine, & O’Doherty, 2009). Moreover, humans as well as other animals will revert to a habitual strategy even early in training if a goal-directed strategy causes response conflict (de Wit, Corlett, et al., 2009; de Wit, Niry, Wariyar, Aitken, & Dickinson, 2007).

Recently, the neurobiology of these distinct habitual and goal-directed control mechanisms has received an increasing amount of attention (Daw, Niv, & Dayan, 2005; Yin, Knowlton, & Balleine, 2004; Joel & Weiner, 2000). Although the BG has long been implicated in habit memory (Packard & Knowlton, 2002; Mishkin, Malamut, & Bachevalier, 1984), behavioral neuroscience studies with rodents have only recently begun to elucidate the specific neural mechanisms of experimentally defined behavioral control processes. This work has shown that the dorsomedial striatum and the prefrontal cortex subserve goal-directed actions (Yin, Knowlton, & Balleine, 2005; Corbit & Balleine, 2003; Killcross & Coutureau, 2003; Balleine & Dickinson, 1998), whereas habit formation is reflected in a shift in control toward the dorsolateral striatum (DLS) (Yin & Knowlton, 2006; Yin et al., 2004, 2005). Dopamine is thought to be crucially involved in this process (Wise, 2004), and the dopaminergic projection from the substantia nigra to the dorsal striatum has been implicated in the reinforcement of habits (Faure, Haberland, Conde, & El Massioui, 2005; see also Reynolds, Hyland, & Wickens, 2001).

Homologue areas (Joel & Weiner, 2000) are thought to be involved in human instrumental behavior. Several fMRI...
studies implicate the ventromedial pFC (vmPFC; de Wit, Corlett, et al., 2009; Tanaka, Balleine, & O’Doherty, 2008; Valentin, Dickinson, & O’Doherty, 2007) and the anterior caudate nucleus (O’Doherty et al., 2004) in goal-directed control. On the other hand, Tricomi et al. (2009) recently provided the first evidence for progressive recruitment of the human homologue area of the rodent DLS, namely, the dorsal putamen, with prolonged instrumental training. However, this evidence is merely correlational in that Tricomi et al. observed increased fMRI activations in this area as a function of practice. So far, there is no direct evidence that in humans the dorsal striatum plays a critical supporting role in S→R habit formation.

One approach to studying the importance of mesocorticolimbic circuits and dopamine for habitual control is by investigating the effects of Parkinson disease (PD). PD is associated with progressive nigrostriatal and mesocorticolimbic dopamine depletion and is accompanied by subtle cognitive impairments even in the early stages, resembling those seen in frontal lobe patients (Owen et al., 1992, 1995; Taylor, Saint-Cyr, & Lang, 1986). Mild PD is a particularly good model for assessing the hypothesized distinct roles of different parts of the striatum because studies have shown that, in early PD, dopamine depletion is most severe in the dorsal striatum, only later progressing to areas associated with goal-directed action control, including more ventral and medial parts of the striatum and pFC (Agid et al., 1993; Kish, Shannak, & Hornykiewicz, 1988). In keeping with this neurobiological pattern of dopamine depletion, it has long been hypothesized that, already at an early stage of the disease, PD is accompanied by a disruption of habit formation (Knowlton, Mangels, & Squire, 1996), which would render even simple everyday activities or performing more than one action at once effortful for these patients (Brown & Marsden, 1990, 1991). For instance, on the basis of evidence for impaired implicit, incremental associative learning of relationships between stimuli and outcomes in a probabilistic classification task but intact acquisition of declarative knowledge, Knowlton et al. (1996) have argued that PD patients exhibit a habit memory deficit (but see, Witt et al., 2006). Since then, similar feedback-based (probabilistic) learning impairments have been observed in a variety of instrumental learning paradigms (Frank, Samanta, Moustafa, & Sherman, 2007; Frank, Seeberger, & O’Reilly, 2004; Shohamy et al., 2004; Shohamy, Myers, O’Nialor, & Gluck, 2004). Accordingly, it is now well accepted that mild PD can be accompanied by instrumental learning impairments (but see Swainson et al., 2006). However, a major problem with many of these instrumental learning studies is that paradigms were used that cannot distinguish between the deployment of habitual versus goal-directed associative structures in instrumental behavior as defined experimentally in the animal literature.

The primary aim of the present study is to address this confound between habit-based and goal-directed behavior in the PD literature. To this end, we employed a behavioral procedure that has been used successfully in previous studies to establish habits in both animals and humans. Specifically, we assessed the concurrent learning of multiple biconditional instrumental discriminations in which fruit pictures and points functioned both as discriminative stimuli and as outcomes for left and right key-presses (de Wit, Corlett, et al., 2009; de Wit et al., 2007; for animal studies with an equivalent task, see de Wit, Ostlund, Balleine, & Dickinson, 2009; de Wit, Kosaki, Balleine, & Dickinson, 2006; Dickinson & de Wit, 2003). The three types of discriminations are illustrated in Figure 1 (for a more elaborate explanation, see Methods). Whereas four different fruits functioned as stimuli and as outcomes in the standard biconditional discrimination, the other two discriminations each involved only two fruit pictures that were either the same in each component of the discrimination (congruent) or opposites (incongruent). In the latter incongruent discrimination, fruit pictures should become associated with opposite responses via S→R versus O→R associations. Critically, whereas performance on congruent and standard discriminations can be supported by both goal-directed associative structures as well as stimulus–response (S–R) habit formation, the incongruent discrimination requires predominant reliance on S→R associations to prevent response conflict due to O→R associations. In previous studies, this reliance on the habit system was reflected in overall poor performance on this incongruent discrimination relative to the congruent and standard discriminations that receive additional support from the goal-directed system (de Wit, Corlett, et al., 2009; de Wit et al., 2007). To assess directly the degree to which subjects adopted habit or goal-directed learning strategies to solve the different discriminations, we employed an “instructed” outcome devaluation test at the end of training (for detailed description, see Methods). If subjects formed O→R associations during training and were successful in using the instructed value to guide their behavior during the test, they should direct their actions toward the still-valuable fruit outcomes at the expense of devalued goals. In line with our theoretical account of incongruent performance, previous studies have shown that outcome devaluation test performance of young healthy volunteers is indeed impaired for the incongruent relative to the other discriminations (de Wit, Corlett, et al., 2009; de Wit et al., 2007).2

To summarize, with this study we aimed to extend the existing correlational evidence for the role of the striatum in the ability to form habits in humans (Tricomi et al., 2009) by assessing whether mild PD patients, characterized by relatively severe dopamine depletion in the DLS, exhibit a significant S→R habit formation deficit. In keeping with the above-reviewed literature, we predicted that mild PD patients would exhibit disproportionate difficulty with the learning of the incongruent discrimination, which requires the use of S→R habits. Conversely, performance on the subsequent outcome devaluation test should not be negatively affected early in the disease as it relies on
goal-directed associative structures, which should at that stage be relatively intact. Importantly, we also investigated whether there was a negative relationship between disease severity and outcome devaluation test performance in our sample because of progressive dopaminergic depletion of areas that support goal-directed action control. Finally, to investigate the role of dopamine in the hypothesized habit formation deficit, we compared performance of groups of patients on versus off their normal regimen of dopaminergic medication.

METHODS

This study was approved by the Peterborough and Fenland Local Research Ethics Committee. All subjects gave written consent.

Patients

Thirty PD patients were recruited from the Brain Repair Centre at Addenbrooke’s Hospital, Cambridge, UK. All patients were diagnosed by a neurologist, and all were receiving dopaminergic medication. Fifteen patients were tested taking their medication as usual (On group; 11 men/4 women), whereas the other half was asked to abstain from their medication 18 hr before the test session (Off group; 11 men/4 women). This procedure allowed us to investigate the effect of medication withdrawal using a between-subjects design (to prevent practice effects associated with a within-subject design). Average number of hours since taking the last dose was approximately 4.5 hr for patients in the On group and approximately 20 hr for the Off group. We endeavored to match the patient groups in terms of the type of medication as much as was feasible. As can be seen in Table 1, 23 of the 30 patients tested were receiving L-dopa. The remaining seven patients all received the D3 (and to lesser extent D2/D4) receptor agonist ropinirole. Demographics and clinical characteristics of the PD patients are detailed in Table 2. None of the patients had a significant neurological history unrelated to PD, and all patients were nondemented (Mini-Mental State Examination [MMSE] > 24) and nondepressed (Beck Depression Inventory [BDI] < 30, with a range of 0–25 in the patients and 4–19 in the controls).

Table 1. Medications

<table>
<thead>
<tr>
<th></th>
<th>PD On group</th>
<th>PD Off group</th>
</tr>
</thead>
<tbody>
<tr>
<td>L-Dopa</td>
<td>10</td>
<td>13</td>
</tr>
<tr>
<td>Perigolide (D1/D2)</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Amantadine</td>
<td>5</td>
<td>6</td>
</tr>
<tr>
<td>COMT inhibitor</td>
<td>0</td>
<td>3</td>
</tr>
<tr>
<td>Antidepressants</td>
<td>2</td>
<td>4</td>
</tr>
<tr>
<td>MAO-B inhibitor</td>
<td>0</td>
<td>3</td>
</tr>
<tr>
<td>Benzodiazepine</td>
<td>1</td>
<td>2</td>
</tr>
</tbody>
</table>
Table 2. Background Details

<table>
<thead>
<tr>
<th></th>
<th>Age</th>
<th>Edu</th>
<th>NART</th>
<th>BDI</th>
<th>MMSE</th>
<th>UPDRS</th>
<th>H&amp;Y</th>
<th>Hours Since Last Dose</th>
</tr>
</thead>
<tbody>
<tr>
<td>On (n = 15)</td>
<td>64.8 (6.5)</td>
<td>12.4 (3.2)</td>
<td>33.6 (9.1)</td>
<td>7.7 (4.0)</td>
<td>28.9 (1.4)</td>
<td>36.7 (13.2)</td>
<td>1.7 (0.6)</td>
<td>4.3 (4.9)</td>
</tr>
<tr>
<td>Off (n = 15)</td>
<td>61.1 (2.1)</td>
<td>12.9 (2.7)</td>
<td>38.9 (9.0)</td>
<td>7.8 (6.9)</td>
<td>29.6 (0.8)</td>
<td>41.4 (18.9)</td>
<td>1.8 (0.8)</td>
<td>19.8 (2.8)</td>
</tr>
<tr>
<td>CS (n = 14)</td>
<td>63.0 (8.1)</td>
<td>13.1 (3.1)</td>
<td>40.0 (5.3)</td>
<td>6.3 (4.8)</td>
<td>29.3 (1.0)</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
</tr>
</tbody>
</table>

Values are presented as mean (SD).

Edu = education; NART = National Adult Reading Test; BDI = Beck Depression Inventory; MMSE = Mini-Mental State Examination; UPDRS = Unified Parkinson Disease Rating Scale; H&Y = Hoehn & Yahr; N/A = not applicable.

Table 3. Results of the Background Neuropsychological Tests

<table>
<thead>
<tr>
<th></th>
<th>FAS</th>
<th>Sem Flu</th>
<th>Str Words</th>
<th>Str Colors</th>
<th>Str Interference</th>
<th>PRM</th>
<th>SRM</th>
</tr>
</thead>
<tbody>
<tr>
<td>On (n = 15)</td>
<td>42.7 (13.8)</td>
<td>33.7 (6.2)</td>
<td>86.0 (15.7)</td>
<td>62.1 (12.3)</td>
<td>33.0 (10.4)</td>
<td>88.7 (10.2)</td>
<td>77.9 (12.8)</td>
</tr>
<tr>
<td>Off (n = 15)</td>
<td>43.7 (13.7)</td>
<td>31.8 (11.9)</td>
<td>93.0 (24.2)</td>
<td>64.5 (13.0)</td>
<td>32.6 (12.0)</td>
<td>92.5 (8.6)</td>
<td>80.6 (7.1)</td>
</tr>
<tr>
<td>CS (n = 14)</td>
<td>41.1 (15.4)</td>
<td>33.2 (6.2)</td>
<td>101.6 (17.6)</td>
<td>69.9 (14.9)</td>
<td>39.7 (12.5)</td>
<td>93.8 (6.7)</td>
<td>83.6 (7.2)</td>
</tr>
</tbody>
</table>

Values are presented as mean (SD).

FAS = letter fluency; Sem Flu = semantic fluency; Str = Stroop; PRM = pattern recognition memory; SRM = spatial recognition memory.

(Beck, Ward, Mendelson, Mock, & Erbaugh, 1961). The average disease duration was 6.2 years for the On group ($SEM = 0.7$) and 8.5 years for the Off group ($SEM = 1.4$). None of the patients showed evidence for a dopamine dysregulation syndrome. The severity of PD symptoms was assessed during the testing session with the Hoehn and Yahr (1967) rating scale and the 44-item Unified Parkinson Disease Rating Scale (UPDRS) (Fahn, Elton, & Committee, 1987). Hoehn and Yahr ratings ranged between I and III. We expected higher UPDRS scores for the Off group than for the On group. However, the two groups did not differ significantly in terms of disease severity (as reflected by their UPDRS scores), $F < 1$, suggesting that had medication status been matched, disease severity was likely more severe for the patients in the Off group than that in the On group. Therefore, any effects of medication might reflect effects of disease severity.

Controls

Fifteen healthy age- and IQ-matched control volunteers were recruited through local advertisement in the Cambridge community. The data of one subject had to be excluded because of a technical error, leaving 14 subjects in the control (CS) group (9 men and 5 women). The background details of the control subjects are presented in Table 2. Separate one-way ANOVAs established that there were no significant differences between the On, Off, and CS groups in terms of: age ($F < 1$); education ($F < 1$); and premorbid IQ (as assessed with the National Adult Reading Test [NART]; Nelson, 1982), with estimated verbal IQ scores of 114, 119, and 120 for the On, Off, and CS groups, respectively, $F(2, 41) = 2.67$, $MSE = 64.89$; MMSE, $F(2, 41) = 1.77$, $MSE = 1.16$; BDI ($F < 1$). BDI did differ considerably between subjects, perhaps partly because positive scores can reflect the motor symptoms of PD rather than true depression and could be expected to affect instrumental performance. However, we failed to find evidence for correlations between the BDI score and the discriminative performance during training and accuracy of test performance (with Pearson correlations of $-0.12$ and $-0.14$).

Background Neuropsychological Tests

In addition to the instrumental conflict task, all volunteers received several background neuropsychological tests: letter and semantic fluency tasks (Benton, 1968), Stroop (1935) task, pattern recognition memory (PRM), and spatial recognition memory (SRM) (Sahakian et al., 1988). The results are presented in Table 3. Separate one-way ANOVAs showed that performance of the three groups on these tasks was statistically indistinguishable: letter fluency, $F < 1$; semantic fluency, $F < 1$; Stroop (in terms of Stroop interference divided by Stroop words), $F < 1$; PRM, $F(2, 41) = 1.30$, $MSE = 75.07$; SRM, $F(2, 41) = 1.28$, $MSE = 89.84$. Finally, in PD patients, disease severity in terms of UPDRS score did not correlate with performance on these neuropsychological tests, letter fluency ($r = -0.18$), semantic fluency ($r = -0.18$), Stroop ($r = -0.22$), PRM ($r = -0.08$), SRM ($r = 0.09$), nor with age ($r = 0.21$), education ($r = 0.05$), NART ($r = -0.29$), MMSE ($r = -0.16$), and BDI ($r = 0.25$).

Procedure

The full experiment took approximately 2 hr. The background and experimental tasks were always administered...
in the following order: NART, FAS, MMSE, instrumental conflict task, PRM and SRM, Stroop task, BDI, UPDRS, and Hoehn and Yahr (1967) rating scale. The computerized experimental task was adapted from the version used by de Wit et al. (2007). The main changes were that subjects received a demonstration of the task, as also in a previous fMRI study with this paradigm (de Wit, Corlett, et al., 2009; de Wit, Ostlund, et al., 2009), and that the instrumental training phase was longer than that in previous studies to ensure that subjects acquired the instrumental discriminations.

Stimuli

The stimuli consisted of colored icons representing the eight different fruits: orange, pineapple, pear, apple, banana, cherry, grape, and coconut (see also de Wit et al., 2007). For the demonstration of the task, we used three colored icons, representing beer, wine, and coffee. All pictures were presented on a standard PC monitor, and responses on a left (m) and right (z) key were recorded on a standard keyboard using a program written in Visual Basic 6.0.

Demonstration of Conflict Task and Instructions

All subjects received a demonstration of the conflict task, using the following instructions on the computer screen:

In this game, you will get the chance to earn points by collecting items from inside a box on the screen by opening the box by pressing either the right or the left key. If you press the correct key, the box will open to reveal a drink inside and points will be added to your total score. However, if you press the incorrect key, the box will be empty and no points will be added to your total. Your task is to learn which is the correct key to press. Sometimes it will be the left-hand key and sometimes the right-hand key. The picture on the front of the door should give you a clue about which is the correct response. To give you an impression of the game you will be asked to play later on, we will first give you some demonstration trials. Just follow the instructions on the screen.

Having read these instructions, subjects were shown a picture of a closed box with a picture of a glass of beer on the front door. At the bottom of the screen, we showed them the instructions “Press Left.” Pressing the left key led to a picture of an open empty box. On the following screen, subjects were again shown a picture of a glass of beer on the front door of a box, but this time with the instruction “Press Right.” Pressing the right key was rewarded with another glass of beer and 1 point. Subjects were then shown in the same fashion that a cup of coffee signaled that pressing the right key would not be rewarded, whereas pressing the left key was rewarded with a glass of wine and 1 point. Subjects were then given the following instructions:

You have had a chance to learn which was the correct key to press for two different pictures. In the following demonstration, you will no longer be told which response to make, and your task is to press the correct key. Only the first keypress on each trial will count and the quicker a correct response is made the more points will be added to your total, so try to respond as quickly as possible!

Subsequently, subjects received four practice trials with the beer stimulus and four trials with the coffee stimulus, randomly intermixed. Pressing the correct key for the beer and the coffee was rewarded with points and with either beer or a glass of wine inside the box, respectively. Pressing the incorrect key was always followed by an empty box. As in the real experiment, the faster a response was made, the more points were earned. The number of points awarded for correct responses within the following RT ranges was as follows: 0–1 sec, 5; 1–1.5 sec, 4; 1.5–2 sec, 3; >2–2.5 sec, 2; >2.5 sec, 1. The outcome display showed a picture of the drink outcome and the number of points earned. This display remained present for 1 sec before being replaced by the stimulus display of the next trial after a 1.5-sec intertrial interval. The total score was always displayed at the top of the screen. At the end of the discrimination training phase, subjects received instructions for the (outcome-cued) outcome devaluation test that was designed to assess the strength of O→R associations:

In the next phase, two open boxes will appear on the screen with different drinks inside them. One drink was earned by a left response in the first stage and the other by a right response. Although both drinks were valuable previously, one of them is now devalued and earns no points, whereas the other is still valuable and gains points. The devalued drink will have a cross on it. You should respond by pressing the key that earns a valued drink. The points you earn now will not be shown on the screen but you will see your final total at the end of the game. As in the training phase, only your first response will count.

The subjects were then shown two open boxes on the screen (one above the other), one containing a beer and one containing a glass of wine. On the first trial, the wine had a red cross superimposed on it, signifying that the left response associated with it no longer earned any points, whereas on the second trial the beer was shown with a cross, signifying that the right response was no longer rewarded. Each keypress marked the end of that trial and was immediately followed by the next test trial. Subjects therefore did not receive feedback about their performance during the test to ensure that their choices were
Discrimination Training

Once the experimenter had ensured that the instructions and demonstration had been understood (and if necessary, had rerun the demonstration until the instructions were clear), each participant was presented with the first trial. As with the demonstration phase, participants were shown boxes bearing a fruit and were required to use this information to select the left or right keypress. A correct response led to another fruit picture and a gain of a minimum of 1 point and a maximum of 5, depending on RT. Incorrect responses led to an empty box on the screen and 0 points. Three discriminations were trained together: cue-outcome incongruent, cue-outcome congruent, and standard (see Figure 1). Two fruit icons were assigned to the congruent discrimination, two to the incongruent discrimination, and four to the standard discrimination.

Performing the correct response to a fruit stimulus yielded the same fruit icon as the outcome in the congruent discrimination but the other fruit icon in the incongruent discrimination. In the example of a congruent discrimination in Figure 1, a banana signals that pressing the left key will be rewarded with another banana, whereas grapes signal that right keypresses will be rewarded with grapes. In contrast, in the incongruent example, each fruit functions as a stimulus and outcome for opposing responses: An apple signals that pressing the left key will be rewarded with a pineapple, whereas in the other component of the discrimination, the pineapple signals that pressing the opposite, right, key will be rewarded with the apple. Finally, in the standard discrimination, two fruit icons acted as the stimuli and the other two as the outcomes with the assignment of stimulus–outcome pairs remaining constant across training. Hence, in the example, the correct left response to a coconut stimulus consistently yields a cherry outcome, whereas a correct right response to the orange stimulus yields a pear outcome. As can be seen in the right panel of Figure 1, performance on the congruent and standard discriminations can be supported by an S→O→R associative structure. In contrast, O→R associations can cause response conflict in the case of the incongruent discrimination (for a detailed description and theoretical background, see de Wit, Corlett, et al., 2009; de Wit et al., 2007).

The icons were paired arbitrarily for the three discriminations, and the left response was correct for one icon and the right response for the other icon of a pair. To ensure that the identity of the icons was not confounded with discrimination type, the assignments of these icons and response pairings to the different discriminations were permuted across participants, in such a way that there were eight possible combinations, with each fruit icon functioning overall twice as stimulus and twice as outcome for each discrimination, once with a right and once with a left response.

Discrimination training consisted of eight 12-trial blocks. Within each block, there were two trials with each of the component contingencies from each of the three discriminations, which were presented in a random order that varied across participants. Therefore, every participant received a total of 16 trials with each component of the three discriminations (twice as much as in the original experiment of de Wit et al., 2007).

Outcome Devaluation Test

Following discrimination training, the participants were reminded of the instructions for the outcome devaluation test. This test consisted of four trials from each of the three discriminations, two with one of the outcomes devalued and two with the other outcome devalued. These 12 trials were presented in a different random order for each participant.

Questionnaires

Subjects were asked to indicate on a printed questionnaire for each fruit that had functioned as a discriminative stimulus, whether the right or the left response had been correct, and which fruit was presented inside the box following a correct response for that discriminative stimulus.

Data Analysis

Statistical analysis was performed using SPSS 15.0. We employed repeated measures ANOVA (RM-ANOVA), complemented with two-tailed t tests, to investigate whether PD patients and controls differed in accuracy (percentage of correct responses per block of training) and RT (sec) during discrimination training, in accuracy (average percentage of correct responses) during the outcome devaluation test, and in accuracy (number of correct answers out of two per discrimination) on the questionnaires. In
addition, we compared patients on versus off medication on all of the abovementioned dependent measures. We included each patient’s sum UPDRS score as a covariate in these RM-ANCOVAS because disease severity varied considerably within our sample (with UPDRS scores ranging from 19 to 82.5). All p values involving repeated measures factors are based on Greenhouse–Geisser sphericity corrections, and all significant (p < .05) first-order interactions involving the factor of interest (discrimination type) are reported.

RESULTS

Discrimination Training—Accuracy

To investigate the acquisition of instrumental discriminations, we conducted an RM-ANOVA on the percentage of correct responses, with the between-subjects factor Group (controls/PD patients) and within-subject factors Block and Discrimination. As can be seen in Figure 2, the instrumental discriminations were acquired gradually, as supported by a significant effect of Block, F(7, 294) = 17.14, MSE = 541.9, p < .0005. There was no Group × Block interaction, F = 1.02, MSE = 552.2, but discriminative performance was negatively affected overall in the PD patients relative to the CS group, F(1, 42) = 3.99, MSE = 6311, p = .05. We failed, however, to find evidence for a Group × Discrimination interaction, F(2, 84) = 1.70, MSE = 1479, indicating that the congruence effect did not differ between PD patients and controls (see Table 4 for average performance on the three discriminations). A main effect of Discrimination, F(2, 42) = 11.01, MSE = 872.1, p < .0005, prompted pairwise comparisons. Congruent performance was significantly better than both incongruent and standard performance (ps < .01), but, unlike in previous studies, the difference between standard and incongruent performance was only marginally significant (p = .06). Importantly, two-tailed t tests on the percentage of correct responses on the final block of training show that both patients and controls performed significantly above chance on all discriminations (PD, ts ≥ 2.60; CS, ts ≥ 6.01).

To investigate whether medication affected instrumental learning, we conducted a separate RM-ANCOVA on the patients on/off medication. To take into account disease severity, we included each patient’s UPDRS score as a covariate. This analysis only yielded a significant effect of Block, F(7, 189) = 2.96, MSE = 630.7, p < .05. There was no main effect of Group (on/off), F < 1, nor a Discrimination × Group interaction, F(2, 54) = 1.23, MSE = 1006. Finally, we failed to find an effect of the UPDRS covariate, F < 1. In the present analysis, there was no significant effect of Discrimination, F < 1, but we do not wish to attribute significance to this null effect in the patient group as...
we did not find evidence for a Group × Discrimination effect in the prior analysis.

**Discrimination Training—Reaction Time**

Fast responding was encouraged during discrimination training. In an RM-ANOVA of the RTs (sec) of PD patients versus controls, we only found a significant effect of Block, $F(7, 294) = 44.94, MSE = 0.15, p < .0005$, reflecting that subjects gradually learned to respond faster, as is depicted in the right panel of Figure 2. PD patients and control subjects responded equally fast overall, $F(1, 42) = 1.62, MSE = 6.47$, and there were no differences in RT depending on discrimination type, $F = 1.03, MSE = 0.237$. We also conducted a separate RM-ANCOVA on the patient data to investigate whether RT was affected by medication status and disease severity. There were no significant main effects of medication status, nor of disease severity, $Fs < 1$, nor any first-order interactions.

**Outcome Devaluation Test**

As can be seen in Table 4, performance was best on the congruent test trials and worst on the incongruent trials. This was confirmed with an RM-ANOVA with the between-subjects factor group (PD/CS) and the within-subject factor discrimination. A significant effect of Discrimination, $F(2, 84) = 30.89, MSE = 787.6, p < .0005$, was further investigated with pairwise comparisons, which showed that congruent performance was significantly superior to standard performance, which in turn was better than incongruent performance ($ps < 0.05$). Separate $t$ tests for the PD/CS groups showed that performance was above chance level only for the congruent and standard discriminations ($ps < 0.05$). Therefore, we replicated the congruence effect during test observed in previous studies (de Wit, Corlett, et al., 2009; de Wit et al., 2007). We failed, however, to find an effect of group on the acquisition/deployment of goal-directed R-O knowledge. The patients and controls did not differ in their level of performance overall, $F < 1$, nor did discrimination interact with group, $F = 1.02, MSE = 803.2$.

Again, we conducted a separate RM-ANCOVA on test performance of the PD patients, with medication status as a between-subjects variable and UPDRS score as a covariate. In line with our hypothesis, a significant main effect of UPDRS, $p < .05$, indicated that disease severity negatively affected test performance. A two-tailed Pearson correlational analysis on the average test performance and UPDRS score yielded a significant negative correlation ($r = -.37, p < .05$), as shown in Figure 3. We also found that patients on medication tended to perform worse overall than patients off medication, but the effect of medication failed to reach significance, $F(1, 27) = 3.38, MSE = 519.1, p = .08$. Given that the medicated patients were likely to be clinically more severely affected, this apparent effect of medication might reflect an effect of disease severity, in line with the effect of UPDRS reported above.

**Outcome Devaluation Test—Reaction Time**

An RM-ANOVA with the between-subjects factor group (PD/CS) and the within-subject factor discrimination established that RTs during test did not differ between the three discriminations, $F(2, 84) = 1.13, MSE = 1.395$, with average RTs of 2.5 sec on both congruent and standard trials and 2.8 sec on incongruent trials. RTs of patients and controls were also statistically indistinguishable, $F < 1$, with average RTs of 2.8, 2.5, and 2.5 sec for the On, Off, and CS groups, respectively. The separate RM-ANCOVA on the PD patients showed that RT also did not depend on medication status, $F < 1$, and did not correlate with disease severity, $F(1, 27) = 1.61, MSE = 10.06$.

**Questionnaires**

Participants were asked to indicate the correct response and outcome for both discriminative stimuli of each discrimination and were given a point for each correct answer. The scores for response and outcome for each discrimination ranged therefore from a minimum of 0 to a maximum score of 2. These scores (see Table 4) were analyzed separately. One subject in the control group failed to fill out the questionnaire, leaving 13 subjects in that group.

As can be seen in Table 4, there was a high level of explicit S-R knowledge in all three groups, with average scores of 1.5, 1.6, and 1.6 for the On, Off, and CS groups, respectively. An RM-ANOVA comparing scores of PD patients and controls did not yield any significant effects. Explicit knowledge of the S-R relationships was the same for the different discriminations, $F(2, 82) = 1.65, MSE = 0.39$, and did...
Explicit memory of the instrumental S→R associations. Difficulties with the internal generation of actions and cognitive plans as well as enhanced performance on the outcome devaluation test (de Wit, Corlett, et al., 2009). However, goal-directed support may similarly rely on the gradual building up of associations, and impaired goal-directed control may therefore also contribute to the present learning deficit. In line with the latter possibility, we found evidence for a disease severity–dependent impairment of performance on the subsequent outcome devaluation test. Performance on this test should not be negatively affected by impaired habit formation as it is mediated by goal-directed knowledge, so this result suggests that progressive PD leads to impaired goal-directed control. In the remainder of this article, we will discuss the implications of as well as potential issues with the presented evidence.

Our evidence for disrupted goal-directed control is consistent with other lines of evidence suggesting that progressive PD leads to a cognitive profile characteristic of more ventral corticostriatal circuits (Agid et al., 1993; Kish et al., 1988). In a previous fMRI study with the conflict task, we showed that the vmPFC was engaged during performance on the outcome devaluation test (de Wit, Corlett, et al., 2009). Moreover, there is evidence to suggest that activations in ventral corticostriatal circuits are modulated by contingency, with vmPFC tracking local changes in correlations between action and outcome rates (Tanaka et al., 2008). Therefore, pFC and anterior caudate nucleus may work together to support goal-directed learning. Interestingly, a goal-directed deficit may concur with a separate literature that highlights a shift from internal to external control in PD (van Spaendonck, Berger, Horstink, Born, & Cools, 1995; Brown & Marsden, 1988; Cools, van den Bercken, Horstink, van Spaendonck, & Berger, 1984). Observations that PD patients have no problem initiating actions when presented with an unambiguous external stimulus (e.g., Rahman, Griffin, Quinn, & Jahanshahi, 2008; Praamstra, Stegeman, Cools, & Horstink, 1998) suggest that if anything remains intact in PD, it is the ability to act on direct S→R associations. Difficulties with the internal generation of actions and cognitive plans as well as enhanced...
cue-reliance and stimulus-driven behavior are more in line with a goal-directed impairment.

On the basis of work by, for example, Frank et al. (2004, 2007), showing that instrumental learning is dopamine dependent, we would expect a goal-directed deficit during learning and test to be remediated by dopaminergic medication. However, in the present study, we did not find evidence for superior goal-directed action in patients on medication. In fact, performance of the On group was marginally worse than that of the Off group.

It is important to point out that a caveat of this study is that the On and Off groups were not matched well in terms of disease severity, with patients on medication being clinically more strongly affected. The absence of an effect of medication status should be replicated in future studies with more carefully matched patient groups.

One might argue that the instrumental conflict task is not sufficiently sensitive to detect habit formation deficits in PD. However, our results strongly suggest that incongruent performance relies on habit formation. When subjects were asked to select responses on the basis of the instructed value of the instrumental outcomes, they were able to do this only on congruent and standard trials. In contrast, performance on incongruent trials of the outcome devaluation test did not differ from chance level (see also de Wit, Corlett, et al., 2009; de Wit et al., 2007), indicating a lack of goal-directed control over incongruent performance. Further support for this possibility comes from the recent fMRI study with the conflict task, which showed not only that vmPFC is recruited during the outcome devaluation test but also that this area is preferentially engaged during congruent and standard training relative to incongruent (de Wit, Corlett, et al., 2009; de Wit, Ostlund, et al., 2009). We argue that in the absence of goal-directed control over incongruent performance, response selection was guided by direct S→R associations (see also de Wit, Corlett, et al., 2009; de Wit, Ostlund, et al., 2009; de Wit et al., 2007). Thus, the intact acquisition of the incongruent discrimination by PD patients suggests that the ability to form habits in the presence of conflicting O→R associations is not affected.

It could still be argued that a relative S→R habit deficit in PD patients was masked by their reliance on a more declarative rule formation strategy to solve the incongruent discrimination. Moody, Bookheimer, Vanek, and Knowlton (2004) showed that mild PD patients are able to perform normally on a probabilistic task, but at the same time these patients showed activations in temporal brain areas rather than the striatal areas that were activated in the control participants, raising the possibility that the PD patients adopted an alternative, more declarative strategy. Although we have so far not directly investigated the possibility of propositional encoding of the incongruent discrimination, we cannot exclude the possibility that this represents an alternative viable approach to the instrumental conflict task. However, our current study provides two lines of evidence against this possibility. First of all, PD patients performed at chance level during the incongruent trials of the outcome devaluation test. Successful encoding of the incongruent rule should have allowed them to select the appropriate response for each valuable outcome. Second, a questionnaire at the end of training failed to produce evidence for superior declarative knowledge of the instrumental contingencies in PD patients relative to controls.

However, it remains possible that conflict-induced habit formation relies on processes that are different from those underlying habit formation in the context of extensive training, and our data, therefore, do not exclude the possibility that PD is accompanied by deficits in training-induced habit formation. Nevertheless, we should stress that according to dual-system accounts of instrumental action, S→R associations are strengthened even in the early stages of acquisition, and indeed Tricomi et al. (2009) showed that the dorsal putamen was engaged from the outset of instrumental training. Of course goal-directed associations should usually dominate behavior control early on, but conflict due to R→O associations can cause a reliance on S→R associations from the outset (de Wit, Corlett, et al., 2009; de Wit, Ostlund, et al., 2009; de Wit et al., 2007). The possibility that the formation of strong S→R associations with extensive practice is affected in PD disease could be investigated directly with the kind of paradigm recently employed by Tricomi et al. If S→R habit formation through extensive practice is impaired in PD patients, they should paradoxically outperform control subjects on a subsequent outcome devaluation test.

Finally, it seems important to note that although it is often assumed that the dorsal striatum is crucially involved in S→R habit formation in humans, direct evidence is so far not overwhelming. Human studies have shown that this area is involved in procedural learning, but caution is warranted in equating this with the acquisition of S→R habits as it has been studied in the animal studies that implicate the dorsal striatum. An exception is the recent study of Tricomi et al. (2009), which produced correlational evidence for a role of this area in habit learning. Moreover, in the present study, we addressed for the first time the question whether intact functioning of the human dorsal striatum is a prerequisite for S→R habit formation. Although we failed to produce favorable evidence, this question clearly deserves further scrutiny in human studies employing experimental tasks that are analogous to the carefully constructed paradigms used in animal studies.

In summary, we investigated in PD patients and age-matched controls the ability to form habits by assessing trial-and-error learning of S-R mappings using an instrumental discrimination task but failed to find evidence for a relative impairment in the formation of S→R associations in PD patients. In fact, impaired performance with progressive disease severity on a subsequent outcome devaluation test suggests there may be a deficit in goal-directed control. This goal-directed deficit may be due to progressive depletion of ventral corticostriatal circuits.

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Therefore, our research does not lend support for the hypothesis that habit formation is disrupted in mild PD patients and consequently highlights the need for caution in accepting the habit account of effortful action in PD. The present findings represent an important initial step toward understanding the effects of PD on goal-directed versus habitual behavior and will hopefully inspire further investigations of instrumental dysfunction in PD with behavioral models that capture this crucial distinction.

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Notes
1. Competing associative accounts of goal-directed action stress the importance of either the forward $R\to O$ association or the backward $O\to R$ association (for a review, see de Wit & Dickinson, 2009). As our research does not aim to distinguish between these accounts, we will for simplicity’s sake refer to $O\to R$ associations. 2. It could be argued that the ultimate goal in the conflict task was to earn points (rather than the more specific fruit picture outcomes) and that subjects could rely on response-“correct” outcome learning to the same degree in all discrimination learning conditions. However, such a general goal should not allow for the activation of appropriate goal-directed actions via $S\to O\to R$ associations as the general outcome (of points) should become associated both with right and left responses (we refer the interested reader to the “differential outcomes” literature; see, e.g., Urcuioli, 2005). Furthermore, the outcome devaluation effect provides evidence for reduced reliance on the fruit picture outcome in the incongruent condition.

REFERENCES


