

Brain Areas Consistently Linked to Individual Differences in Perceptual Decision-making in Younger as well as Older Adults before and after Training

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Abstract

■ Perceptual decision-making performance depends on several cognitive and neural processes. Here, we fit Ratcliff's diffusion model to accuracy data and reaction-time distributions from one numerical and one verbal two-choice perceptual-decision task to deconstruct these performance measures into the rate of evidence accumulation (i.e., drift rate), response criterion setting (i.e., boundary separation), and peripheral aspects of performance (i.e., nondesideration time). These theoretical processes are then related to individual differences in brain activation by means of multiple regression. The sample consisted of 24 younger

and 15 older adults performing the task in fMRI before and after 100 daily 1-hr behavioral training sessions in a multitude of cognitive tasks. Results showed that individual differences in boundary separation were related to striatal activity, whereas differences in drift rate were related to activity in the inferior parietal lobe. These associations were not significantly modified by adult age or perceptual expertise. We conclude that the striatum is involved in regulating response thresholds, whereas the inferior parietal lobe might represent decision-making evidence related to letters and numbers. ■

INTRODUCTION

It has been a fundamental goal of cognitive neuroscience to link cognitive and neural processes. One of the many problems in this endeavor is that global cognitive performance measures are influenced by many underlying processes. For example, the performance measures most commonly used in experimental psychology, mean reaction times for correct decisions (RT) and accuracy, are not only a function of the efficiency of the targeted cognitive processes but also of the subjects' response criterion, as speed and accuracy are in a tradeoff relationship: Hasty decisions are more prone to errors, whereas careful and accurate judgments take more time (Wickelgren, 1977). To deconstruct global cognitive performance measures and, for example, disentangle criterion setting from process efficiency, mathematical models can however be applied. Here, we use Ratcliff's (1978) diffusion model (Ratcliff & Tuerlinckx, 2002) to disentangle the speed-

accuracy tradeoff (Ratcliff, 2002; Ratcliff & Rouder, 2000) and to relate individual differences in perceptual decision-making processes to underlying brain regions (Bogacz, Wagenmakers, Forstmann, & Nieuwenhuis, 2010).

The diffusion model aims to explain data from two-choice RT tasks in a comprehensive way, utilizing accuracy information as well as the shape of the RT distribution for correct and erroneous responses. This is achieved by assuming that perceptual-decision behavior involves several processes that come with associated parameters (Figure 1). First, the quality of evidence accumulation during the decision process, the drift rate, is a central parameter. It describes how quickly information is accumulated in a random walk-like diffusion process that progresses from a starting point toward one of two response boundaries, one for correct and one for wrong responses. Higher drift rates indicate faster accumulation of evidence; that is, a more efficient decision process. A second central parameter of the model characterizes response criteria. This more strategic aspect of decision behavior is implemented by differences in the distance between the response boundaries, the boundary separation. Wider boundary separation means more conservative responding because more evidence needs to be accumulated before a boundary is reached and a response is initiated. Still, another

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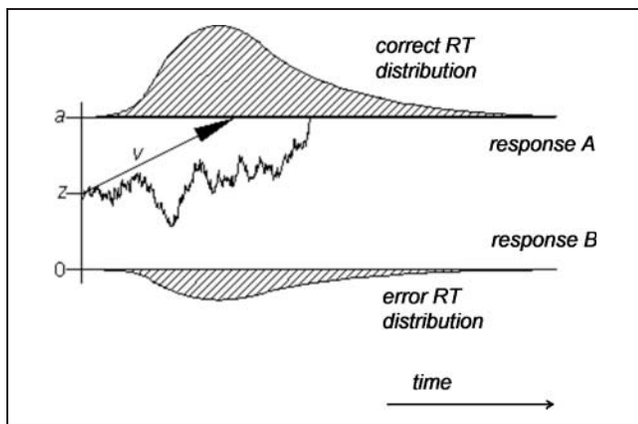


Figure 1. An illustration of the diffusion model with one simulated paths with drift rate v , boundary separation a , and starting point z .

parameter is nondecision time, combining the efficiency of peripheral sensory and motor aspects of the decision process.

In this study, we applied the diffusion model to data from a two-choice decision task with numerical or verbal content. In the numerical task, participants had to judge whether briefly presented numbers were odd or even. In the letter task, participants decided whether a selection of letters were consonants or vowels. Interindividual differences in the estimates of the diffusion-model parameters were then linked to differences in brain activation as observed with functional magnetic resonance imaging (fMRI) during performance of these tasks. In line with the assumptions of the diffusion model and its theoretical application to neuroscience (Heekeren, Marrett, & Ungerleider, 2008), we hypothesized that the drift rate, indicating perceptual evidence accumulation, should be associated with activation in stimuli-specific brain regions, that is, in regions involved in processing of numbers and letters (Joseph, Cerullo, Farley, Steinmetz, & Mier, 2006; James, James, Jobard, Wong, & Gauthier, 2005; Dehaene, Piazza, Pinel, & Cohen, 2003; Eger, Sterzer, Russ, Giraud, & Kleinschmidt, 2003; Joseph, Gathers, & Piper, 2003), such as the left inferior parietal lobe. This prediction is, for example, supported by studies on monkeys that have modeled neurophysiological data from intracranial recordings as diffusion processes (Ratcliff, Hasegawa, Hasegawa, Smith, & Segraves, 2007; Ratcliff, Cherian, & Segraves, 2003; Kim & Shadlen, 1999). In Kim and Shadlen's task, monkeys had to perform a direction of motion visual discrimination task while data were recorded in cells downstream of area MT (lateral intraparietal area, frontal eye fields, superior colliculus, and dorsolateral prefrontal cortex). Their results suggest that decisions are formed by computing the difference between the activities of populations of neurons in area MT that code for opposite directions of motion, and this decision variable is represented in the downstream cells. The results from a large number of studies support the view that decision-making is based on the integration of noisy sensory evidence that is repre-

sented by sensory neurons (e.g., Smith & Ratcliff, 2004; Gold & Shadlen, 2001). For the boundary parameter, we predicted, based on previous studies focussing on the speed-accuracy tradeoff, an association with activity in the striatum and the presupplementary motor area (Forstmann et al., 2008; Ivanoff, Branning, & Marois, 2008; Van Veen, Krug, & Carter, 2008). The basal ganglia (striatum and pallidum) may implement a generic action-selection mechanism that releases those actions from inhibition that are desirable and maintains inhibitory control over others (Mink, 1996), thereby acting as a gate-keeper (see also Frank, 2006). Thus, it is plausible to assume that activity in the striatum is related to criterion setting.

To address these hypotheses, we utilized data from a large-scale study (COGITO Study; Schmiedek, Bauer, Lövdén, Brose, & Lindenberger, 2010; Schmiedek, Lövdén, & Lindenberger, 2010) including younger and older participants performing six tests of perceptual speed (including the two tasks reported in the present article), three tests of episodic memory, and three tests of working memory in approximately 100 daily 1-hr sessions. Before and after the 100-day-long longitudinal phase, extensive data on two-choice RT performance were collected, allowing for estimation of the diffusion parameters as well as for investigating age differences and practice-related changes in these parameters. In this article, we report data for a subsample of participants that additionally performed the two-choice RT tasks in the MRI scanner at pretest and posttest.

Previous studies on age differences in diffusion-model parameters have primarily reported that more conservative response criteria in old age as well as age-related slowing of nondecision time contribute to age-related slowing of perceptual decision making (Starns & Ratcliff, 2010; Ratcliff, Thapar, & McKoon, 2001, 2006; Ratcliff, Thapar, Gomez, & McKoon, 2004; Thapar, Ratcliff, & McKoon, 2003). Practice has been reported to result in faster evidence accumulation and less conservative response criteria, with older adults changing more in these parameter estimates (Ratcliff et al., 2006). fMRI studies of activation changes as a function of practice on lower-level sensory/motor tasks, in contrast to higher-level cognitive tasks such as working memory, have demonstrated decrease of activity in task-related areas after practice (Kelly & Garavan, 2005). In addition, activity in brain areas that have been associated with the so-called resting-state or default-mode (Raichle et al., 2001) network increase during task performance after practice (Mason et al., 2007). This pattern likely reflects more automatic, less attention-demanding, execution of the task posttraining (Kelly & Garavan, 2005). Thus, the data reported in this study allow for examining the generality of the predicted link between regional brain activation and aspects of perceptual decision-making behavior at different average levels of global performance and of the underlying processing components, across younger and older adults, and for nonexpert and highly skilled, less effortful, execution of the task.

chosen stimuli out of the two response categories. In the figural task, participants decided whether the stimuli were either symmetric or asymmetric to stimuli that were either the upper or lower two lines to the left and right of the “calculator 8” (symmetric condition), or the two possible combinations of one upper and one lower line at the left and right (asymmetric condition). In the numerical task, participants decided whether the displayed number was odd or even. Stimuli were “3,” “5,” and “7,” for the odd condition and “2,” “4,” and “6” for the even condition. In the verbal task, participants decided whether letters were either consonants or vowels. Stimuli were “F,” “H,” and “P” for the consonant condition and “A,” “E,” and “U” for the vowel condition. No performance feedback was provided during the task.

Pretest and Posttest

Participants completed behavioral pretests and posttests during 10 sessions that consisted of 2 to 2.5 hr of cognitive test batteries and self-report questionnaires. The pretest brain imaging session was conducted after the behavioral pretest and immediately before the longitudinal practice phase. The posttest imaging session was completed shortly after the completion of the behavioral posttest. The sessions were separated by an average of 179 days ($M_{\text{young}} = 180$, $SD_{\text{young}} = 21.0$; $M_{\text{old}} = 179$, $SD_{\text{old}} = 31.1$). Below we describe the assessment of the two-choice RT performance in more detail.

Behavioral assessment of two-choice RT data. Performance on the two-choice RT tasks was assessed within one behavioral session of about 2.0 to 2.5 hr at pretest and again at posttest. In these sessions, participants worked on the verbal, numerical, and figural versions of the task. Tasks and stimuli were identical for pretest and posttest. In each session, participants worked on 25 blocks consisting of 40 items each for each task, with breaks after sets of 5 blocks (length of break determined by participants). Each block consisted of 20 odd/consonant/symmetric and 20 even/vowel/asymmetric stimuli. Masking times were distributed randomly, but with equal frequencies (10 items per masking time) across blocks, resulting in a total of 250 items per masking time and task. At pretest, participants additionally did first work on two blocks of each task without masking and then on another three practice blocks. These blocks were not used in the current analyses. Participants responded on a customized button box connected to the parallel port for high resolution of RT measurements.

The diffusion model was fit to the data for each task and each subject by minimizing a Chi-square value with a general SIMPLEX minimization routine that adjusts the parameters of the model until it finds the parameter estimates that minimize the Chi-square value (see Ratcliff & Tuerlinckx, 2002, for a full description of the method). For each experimental condition, the 0.1, 0.3, 0.5, 0.7, 0.9

quantile RTs for correct and error responses and the corresponding accuracy values were entered into the minimization routine. The quantile RTs and the diffusion model were used to generate the predicted cumulative probability of a response as a function of quantile RTs. By subtracting the cumulative probabilities for each successive quantile from the next higher quantile, one gets the proportion of responses between adjacent quantiles. These are the expected values to be compared to the observed proportions of responses between the quantiles (i.e., the proportions between 0, 0.1, 0.3, 0.5, 0.7, 0.9, and 1.0, which are 0.1, 0.2, 0.2, 0.2, 0.2, and 0.1) multiplied by the number of observations for the Chi-square computation. Summing over $(\text{Observed} - \text{Expected})^2 / \text{Expected}$ for all conditions results in a single Chi-square value to be minimized. Estimated parameters were drift rate, boundary separation, nondecision time, across-trial variability in drift rate (normally distributed), across-trial variability in starting point (uniformly distributed), across-trial variability in nondecision time (uniformly distributed), and the probability of contaminant responses stemming from a uniform distribution. Drift rates were allowed to vary across masking time conditions. For the analyses reported here, only the drift rate estimates for the conditions with masking time of four screen cycles were used, as those matched the masking time in the scanner session. All other parameters were assumed to be invariant across masking time conditions. As shown in Figure 3, plotting characteristics of observed data versus theoretical expectations based on estimated parameters indicated that the model did fit the data quite well with very few individuals showing deviations. No alternative models were tested, so there was no model selection.

Behavioral task in MRI scanner. The task consisted of 53-sec-long task blocks alternating between the verbal and the numerical tasks described above, intermixed with 16-sec fixation blocks. Each task block included 16 trials (8 odd/8 even; 8 consonants/8 vowels) presented with jittered interstimulus intervals between 2000 and 8000 msec. In total, eight blocks per task were presented, distributed over four runs. Thus, in total, 128 trials per task were presented, with randomly chosen stimuli out of the response categories. Masking times were 50 msec for all participants and assessments.

MRI procedures. Images were collected with a 3-T Magnetom Trio MRI scanner system (Siemens Medical Systems, Erlangen, Germany). Whole-brain functional images were collected using a T2*-weighted EPI sequence sensitive to BOLD contrast (TR = 2000 msec, TE = 30 msec, image matrix = 64×64 , FOV = 224 mm, flip angle = 80° , slice thickness = 3.5 mm, distance factor = 0%, voxel size $3.5 \times 3.5 \times 3.5 \text{ mm}^3$, 32 axial slices). One hundred forty-seven image volumes were acquired per run all aligned to AC–PC. Anatomical images were acquired using a T1-weighted sagittal 3-D spoiled gradient-echo (SPGR) image

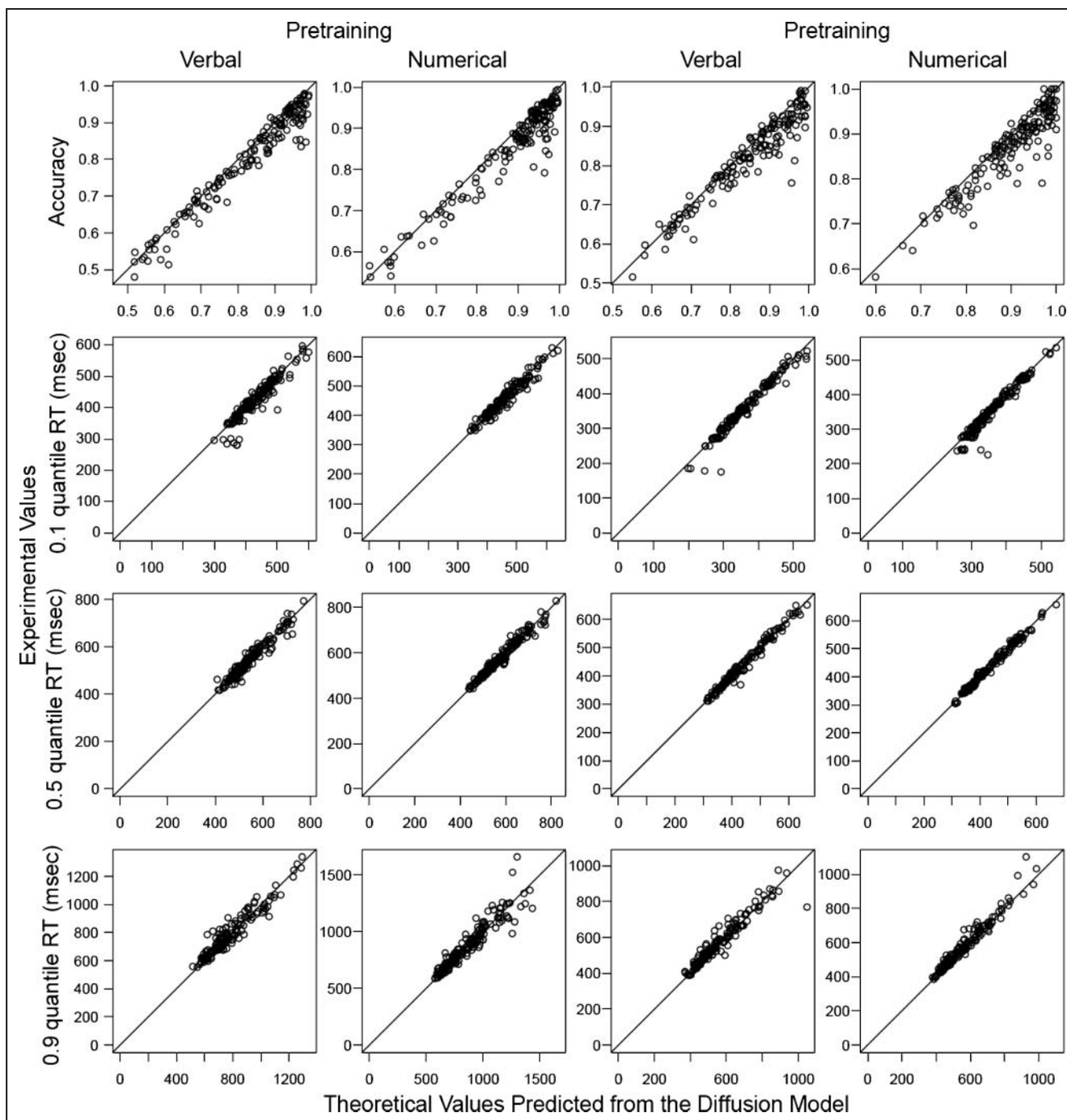


Figure 3. Plots of accuracy and RT quantiles for data (y -axis) and predicted values from fits of the diffusion model (x -axis).

(TR = 24 msec, TE = 8 msec, acquisition matrix = $256 \times 256 \times 124$, FOV = 250×250 mm², flip angle = 30°, slice thickness = 1.5 mm) on a GE Signa system (General Electric, Milwaukee, WI).

fMRI Data Analysis

Preprocessing. The fMRI data were analyzed using the SPM5 software (Wellcome Department of Cognitive Neurology, London, UK). The first three volumes of all EPI series

were excluded from the analysis to allow the magnetization to approach a dynamic equilibrium. Data processing started with slice time correction and realignment of the EPI datasets. A mean image for all EPI volumes was created, to which individual volumes were spatially realigned by means of rigid-body transformations. The high-resolution structural image was coregistered with the mean image of the EPI series. Then the structural image was normalized to the Montreal Neurological Institute (MNI) template, and the normalization parameters were applied to the EPI images

Table 1. Mean and Standard Deviation of RT and Accuracy from Task Performance in the Scanner

	Young Numerical Task		Young Verbal Task		Old Numerical Task		Old Verbal Task	
	Pretest	Posttest	Pretest	Posttest	Pretest	Posttest	Pretest	Posttest
RTs (msec)	742 (123)	584 (107)	793 (130)	593 (99)	824 (169)	697 (94)	814 (208)	710 (95)
Accuracy	0.89 (0.116)	0.94 (0.068)	0.74 (0.133)	0.91 (0.062)	0.73 (0.148)	0.93 (0.060)	0.55 (0.110)	0.83 (0.141)
Drift rate	0.351 (0.114)	0.406 (0.133)	0.230 (0.116)	0.317 (0.121)	0.430 (0.132)	0.578 (0.171)	0.465 (0.124)	0.497 (0.159)
Boundary	0.159 (0.030)	0.083 (0.022)	0.116 (0.029)	0.075 (0.015)	0.124 (0.031)	0.116 (0.080)	0.121 (0.032)	0.097 (0.024)
Nondecision	0.401 (0.039)	0.310 (0.031)	0.386 (0.035)	0.316 (0.033)	0.461 (0.050)	0.403 (0.062)	0.466 (0.048)	0.423 (0.051)

Diffusion-model parameters from behavioral assessment outside the scanner.

The boundary parameter revealed a significant main effect of Task [$F(1, 37) = 15.35, p < .001$], with a stricter decision criterion for the numerical compared to the verbal task, and a significant effect of Time point [$F(1, 37) = 36.62, p < .001$], reflecting a general reduction in strictness of the boundary over time. Moreover, we found a significant Time point \times Age interaction [$F(1, 37) = 11.80, p < .05$], with a stronger decline in the boundary parameter for the younger compared to the older adults. The threefold interaction of Task \times Time point \times Age was again significant [$F(1, 37) = 8.01, p < .05$], indicating larger decreases over time for younger adults in the numerical task, which contrast with the larger decreases in the verbal task for the older adults. No significant main effect of Age group was found.

The nondecision parameter revealed a significant main effect of Time point [$F(1, 37) = 154.40, p < .001$], reflecting a general reduction in nondecision time, and a significant main effect of Age [$F(1, 37) = 55.70, p < .001$], indicating shorter nondecision times in the young. Moreover, we found a significant Time point \times Age interaction [$F(1, 37) = 8.21, p < .01$], with a stronger reduction of nondecision time for the younger compared to the older adults.

fMRI Data

Full-factorial ANOVA

A random effects analysis of the factor Time point from the full-factorial ANOVA containing the factors Task (numerical vs. verbal) and Time point (pretraining vs. posttraining) and the factor Age (young vs. old) revealed significantly more activity at pretest compared to posttest in brain areas that have been associated with the so-called cognitive control network. In detail, we found higher pretest compared to posttest activity in dorsal anterior cingulate cortex (dACC), bilateral insular cortex, left inferior frontal junction (IFJ), left inferior frontal gyrus triangularis (IFG), and right fusiform gyrus (Figure 4A and Table 2A). Increased activity at posttest compared to pretest was found in brain areas that have been associated with the so-called default mode network, namely, ventromedial prefrontal cortex

(vmPFC) and the precuneus, as well as the right superior temporal gyrus (STG) (Figure 4B and Table 2B).

Multiple Regression Analysis with Diffusion-model Parameters

Next, we examined the predictive effects of the diffusion-model parameters on the whole-brain activation maps in a multiple regression model also including the factors Task, Age, and Time point as predictors.

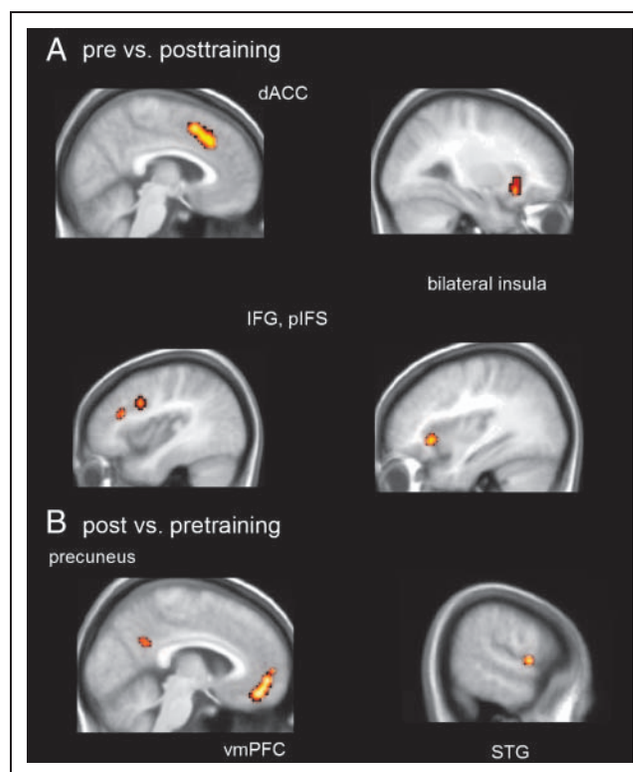


Figure 4. Activation map averaged over 39 subjects ($p < .05$, family-wise error corrected, $k > 5$) mapped onto an averaged image of the participants' normalized (MNI) T1-weighted images. (A) Significantly higher brain activity in the pretraining compared to the posttraining scanning session; (B) Significantly higher brain activity in the posttraining compared to the pretraining scanning session.

Table 2. Brain Areas Showing Significant Differences between Pre- and Posttest

Area	BA	Peak Coordinates (MNI)	Z-score	Extent
<i>(A) Pretraining vs. Posttraining</i>				
Rostral cingulate zone	32	-6, 18, 45	6.85	188
Left insular cortex, inferior frontal gyrus	47	-30, 21, -9	5.53	26
Right fusiform gyrus	19	27, -66, -6	5.41	14
Left inferior frontal junction	44	-42, 9, 33	5.39	13
Right insular cortex, inferior frontal gyrus	47	33, 24, -3	4.94	20
Left inferior frontal gyrus triangularis	45	-39, 27, 21	4.93	7
<i>(B) Posttraining vs. Pretraining</i>				
Ventromedial prefrontal cortex	10/11	-3, 51, -15	5.44	52
Right superior temporal gyrus	22	60, 3, 3	5.21	9
Precuneus	23	-9, -54, 27	4.83	7

The absolute level of drift rate was associated, over and above the other parameters in the model, with interindividual differences in activity in the left inferior parietal lobule (IPL, BA 40; MNI coordinates: -39 -42 54), indicating that participants with higher drift rates show higher activity in IPL. Interindividual differences in boundary separation, on the other hand, were found to uniquely predict activation in the left striatum (putamen; -18 -6 15), indicating that participants with higher boundary parameters showed higher activity in the striatum. These correlations were not influenced by outliers, which the scatterplots in Figure 5 confirm. The nondecision parameter predicted individual differences in the left supramarginal gyrus (BA 40; -45 -33 27).

Even with lowered threshold ($p < .005$, uncorrected), the interaction terms between Age and Time point on the one hand and the diffusion-model parameters on the other hand were not significant in the brain regions reported, supporting the generality of the observed effects. ROI analyses further demonstrate the similarity of these associations across age groups: When computing the correlation coefficients separately for the age groups, we find a significant correlation of percent signal changes in the striatum ROI and boundary parameter for both young [$r(24) = .44, p < .05$] and old [$r(15) = .73, p < .01$] participants. Similarly, we found a correlation of percent signal changes in the left IPL and drift parameter for both young [$r(24) = .51, p < .05$] and old [$r(15) = .51, p = .054$] participants, although the correlation in the older participants was only marginally significant.

Because it has been hypothesized that the drift rate should be associated with brain areas that are responsible for stimulus-specific information accumulation (Heekeren et al., 2008), we additionally computed two separate models, one for the numerical task and one for the verbal task. When applying a more lenient threshold ($p < .001$, clus-

ter of voxels > 10), the resulting brain areas that correlated with the drift rate were the left inferior parietal lobe (BA 40; -45 -39 -51), the left middle occipital gyrus (BA 19; -33 -81 30), and the left superior temporal gyrus (BA 22; -66 -42 15) for the numerical task. For the verbal task, interindividual differences in drift rate predicted activity in the bilateral inferior parietal lobe (BA 40; -36 -42 54; 51 -27 51) and left ventral premotor cortex (BA 6; -57 -3 36).

Finally, it can be predicted that the associations between individual differences in regional activity and individual differences in the boundary and drift rate estimates should generalize to individual differences in practice-related changes. To probe this prediction, we correlated changes (posttest - pretest) in the diffusion parameters to changes in the brain areas identified from the multiple regression analyses. In line with the positive level-level associations, these change-change associations had a positive direction (IPL drift rate: $r = .19, p = .248$; striatum boundary: $r = .53, p < .001$).

DISCUSSION

Perceptual decision-making performance (i.e., RT and accuracy) is a function of several underlying cognitive processes (e.g., Heekeren et al., 2008). In this study, we applied Ratcliff's diffusion model to deconstruct RT and accuracy data from a verbal and a numerical two-choice RT task into three separate parameters: the drift rate, indicating the accumulation rate of perceptual evidence (i.e., the efficiency of the decision process); the boundary separation, reflecting the more strategically influenced response criteria; and the nondecision time, combining the efficiency of sensory and motor aspect of performance. Interindividual differences in these parameters were then related to interindividual differences in brain activation as observed with

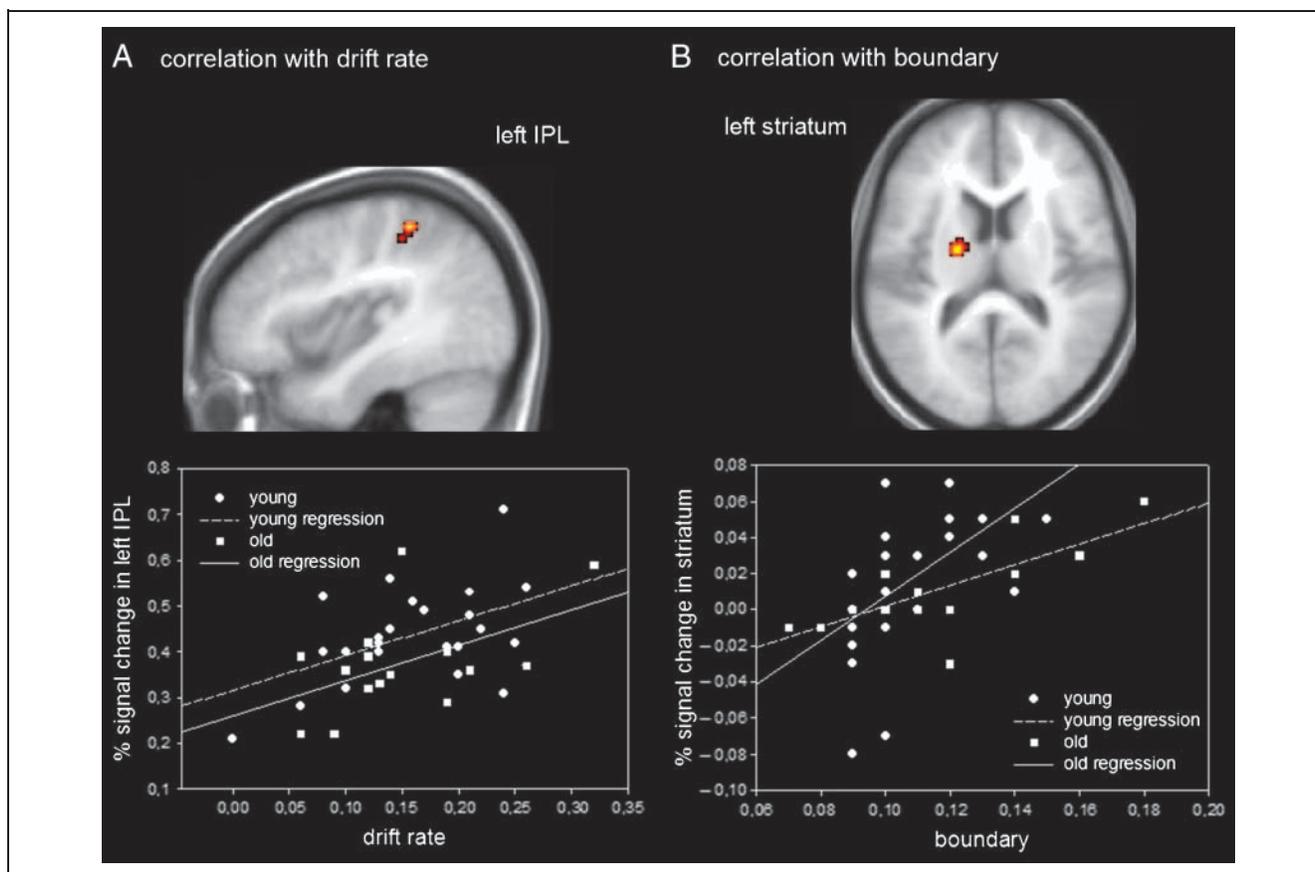


Figure 5. Activation map averaged over 39 subjects ($p < .05$, family-wise error corrected, $k > 5$) mapped onto an averaged image of the participants' normalized (MNI) T1-weighted images. (A) Brain activity (% signal change averaged over pre- and posttraining sessions and over numerical and verbal tasks) correlated with the drift rate and (B) brain activity (% signal change averaged over pre- and posttraining sessions and over numerical and verbal tasks) correlated with the boundary parameter. Note that the regression lines are plotted separately for old and young participants although there is no significant effect of age on the associations.

fMRI during performance of these tasks. In line with our hypotheses, we found that individuals with higher values of drift rate show stronger brain activity in areas that have been reported in processing of numbers and letters, namely, the left IPL, and that individuals with higher values in the boundary parameter showed a higher magnitude of striatal activity.

The parietal lobe, especially the horizontal segment of the intraparietal sulcus (HIPS), has been associated with number processing, and in particular, with the manipulation of numbers, such as in comparison or approximation (Dehaene et al., 2003). For example, patients with parietal lesions have been reported to suffer from difficulties in number processing (Dehaene, Dehaene-Lambertz, & Cohen, 1998). Moreover, the parietal lobe contains an abstract representation of numerical magnitude independent of number notation (Eger et al., 2003). However, the left IPL has also been reported as being specific for single letter processing (Joseph et al., 2003, 2006; James et al., 2005). The fact that the left IPL is involved in both accumulation of numerical as well as letter information is supported by the consistent relation between the drift rate and the magnitude of activation in this region across

separate analyses for the numerical and verbal task. Moreover, several studies have shown that activation in parietal regions correlates with successful memory, suggesting that the strength of activation might contribute to the eventual decision-making (Wheeler & Buckner, 2003). Hence, parietal brain regions have been suggested to function as a mnemonic accumulator, accumulating signal until a criterion is reached and judgments are made, as described by signal-detection theory (Cabeza, Ciaramelli, Olson, & Moscovitch, 2008; Wagner, Shannon, Kahn, & Buckner, 2005). Our findings are in line with this evidence and suggest that the IPL might be involved in the accumulation of perceptual evidence-related letters and numbers.

The association between interindividual differences in the boundary parameter and the strength of activity in the striatum is consistent with at least two lines of previous research. First, it has been speculated that the basal ganglia are critical in speed-accuracy tradeoff (Bogacz & Gurney, 2007; Lo & Wang, 2006) by directly regulating the chain of neuronal responses leading to motor acts (Aron et al., 2003; Band & van Boxtel, 1999; Mink, 1996). For example, the striatum has been linked to the suppression of movements during antisaccade tasks (Raemaekers et al., 2002) as well as

to initiating movements in monkeys (Lebedev & Nelson, 1999), therefore suggesting that the striatum is involved in both initiation and inhibition of motor responses. A second line of research has explored the neural correlates of speed–accuracy tradeoff by instructing participants to either respond as quickly as possible or as accurately as possible (Bogacz et al., 2010; Forstmann et al., 2008; Ivanoff et al., 2008; Van Veen et al., 2008). In terms of the diffusion model, such manipulations affect the boundary parameter of the decision process. When instructing participants to respond accurately, the boundaries are set wider. More information must thus be accumulated, assuming an unaltered drift rate, before a decision can be made, leading to more accurate and slower responses. In contrast, instructing participants to respond as fast as possible should lead to narrower boundaries. In accordance with our findings, Forstmann et al. (2008) report an association between the striatum and a caution parameter derived from a linear ballistic accumulator (LBA) model when comparing performance after speed and accuracy instruction. Likewise, Van Veen et al. (2008) report an association of basal ganglia activity with speed–accuracy tradeoff. But in both of these studies, higher activity in the striatum goes along with the speed compared to the accuracy instruction, whereas our results suggest enhanced striatal activity in relation to higher response caution. However, the activation reported by Forstmann et al. involved the caudate nucleus rather than the putamen (as in this study), whereas van Veen et al. report activity in both the caudate nucleus and the putamen. The present findings are, however, in line with findings of higher striatal activation during trials with a higher probability of response inhibition (Vink et al., 2005) when considering the boundary parameter as an indicator of cautious responding. In summary, the current evidence suggests that the striatum is involved in the regulation of response thresholds rather than, or at least in addition to, the initiation or suppression of responses. This is in line with a model of decision-making and response selection by Frank (2006) and Frank and Claus (2006) that conceptualizes the basal ganglia as a system that gates information flow from frontal cortex to the motor system. The circuit is assumed to facilitate or suppress action representations in frontal cortex. Two cell populations in the striatum are assumed to have opposing effects on the selection of an action. Activation in so-called “go” neurons facilitates the execution of an action considered, whereas activation in “no-go” neurons suppresses the competing responses, thereby constituting a gating mechanism in the striatum. Moreover, our findings are in line with the suggestion that failures in the fronto-striatal dopaminergic neural circuits underpin impulsive and compulsive acts (Fineberg et al., 2010).

As indicated by the absence of interactions between the observed associations (i.e., drift rate with the IPL and boundary separation with the striatum) and the factors Time point and Age, these associations show a fair amount of generalization over the adult lifespan and over different

levels of task-specific expertise, despite the major effects these factors had on performance and the parameters of the diffusion model. Specifically, we observed higher drift rates and lower boundary parameters at posttest than at pretest, which replicates the findings of Dutilh, Vandekerckhove, Tuerlinckx, and Wagenmakers (2009, see also Ratcliff et al., 2006). Our data extend those previous findings by showing larger changes over time in these parameters for younger as compared to older adults. Consistent with the observed positive associations between individual differences in level of activity and level of the diffusion-model estimates, the corresponding associations between individual differences in changes also tended to go in the positive direction. In particular, the significant change–change correlation (post–pretest) between brain activity in the striatum and the boundary parameter suggests that the striatum may also play a role in practice-related changes in boundary separation. Our finding that older adults have considerably longer nondesideration times compared to young adults, on the other hand, replicate previous studies (Ratcliff et al., 2001, 2006; Ratcliff, Thapar, Gomez, et al., 2004; Ratcliff, Thapar, & McKoon, 2004; Thapar, Ratcliff, & McKoon, 2003). Finally, brain activation during performance of these tasks displayed changes over time: We found a decrease in task-related areas (left IFG, left IFJ, bilateral insula, dACC, right fusiform gyrus) and an increase in the so-called default-mode areas (vmPFC, precuneus, right STG). This pattern of changes is likely to reflect more automatic (i.e., less effortful) and perceptual expertise-like performance of the tasks at posttest (Kelly & Garavan, 2005), but the mean level changes over time point are difficult to decisively interpret without an appropriate control group. We thus conclude that individual differences in striatal activity are related to criterion setting and that the rate of evidence accumulation for perceptual decisions of numerical and letter nature is related to stimulus-specific brain regions in the parietal lobe.

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