Dose-Dependent Dissociation of Pro-cognitive Effects of Donepezil on Attention and Cognitive Flexibility in Rhesus Monkeys

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ABSTRACT

BACKGROUND: Donepezil exerts pro-cognitive effects by nonselectively enhancing acetylcholine (ACh) across multiple brain systems. Two brain systems that mediate pro-cognitive effects of attentional control and cognitive flexibility are the prefrontal cortex and the anterior striatum, which have different pharmacokinetic sensitivities to ACh modulation. We speculated that these area-specific ACh profiles lead to distinct optimal dose ranges for donepezil to enhance the cognitive domains of attention and flexible learning.

METHODS: To test for dose-specific effects of donepezil on different cognitive domains, we devised a multitask paradigm for nonhuman primates that assessed attention and cognitive flexibility. The nonhuman primates received either vehicle or variable doses of donepezil before task performance. We measured intracerebral donepezil and its strength in preventing the breakdown of ACh within the prefrontal cortex and anterior striatum using solid phase microextraction neurochemistry.

RESULTS: The highest administered donepezil dose improved attention and made the subjects more robust against distractor interference, but it did not improve flexible learning. In contrast, only a lower dose range of donepezil improved flexible learning and reduced perseveration, but without distractor-dependent attentional improvement. Neurochemical measurements confirmed a dose-dependent increase of extracellular donepezil and decreases in choline within the prefrontal cortex and the striatum.

CONCLUSIONS: The donepezil dose for maximally improving attention differed from the dose range that enhanced cognitive flexibility despite the availability of the drug in two major brain systems supporting these functions. These results suggest that in our cohort of adult monkeys, donepezil traded improvements in attention for improvements in cognitive flexibility at a given dose range.

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Muscarinic binding potential and higher AChE activity in the striatum than in other cortical regions (18). It is unclear how these differences affect ACh modulation of attention functions that depend on the PFC (19) and of flexible learning functions that are dependent on the striatum (20,21). One consequence of the brain area–specific sensitivity to ACh levels could be that a best dose for enhancing cognitive functions supported by the striatum might not sufficiently stimulate the PFC, and that a best dose for enhancing PFC functions might overstimulate the striatum.

To test for these possible implications of brain region–specific ACh action, we devised a drug testing paradigm for monkeys that assessed the effects of three different doses of donepezil across different domains of arousal, attention, and cognitive flexibility in a single testing session. We evaluated the attention domain with a visual search (VS) task that varied the number and perceptual similarity of distracting objects and quantified the domain of cognitive flexibility with a learning task asking monkeys to flexibly adapt to new feature-reward rules and avoid perseverative responding. This assessment paradigm goes beyond existing nonhuman primate studies of donepezil that so far have found enhanced short-term memory using delayed match-to-sample tasks (4,6,10,15,22–27), enhanced arousal and nonselective speed of processing (15,27), or no consistent effect (18) (surveyed in Table S1) and takes into account that studies in rodents report positive donepezil effects across a wider range of domains, including reversal learning (28), paired associate learning (29), object discrimination (30), and novelty detection (31), and variable results on serial choice tasks indexing attention functions (32) (surveyed in Table S2). With our design, we found that donepezil improves interference control over distractors at doses that caused an overall slower response (i.e., reduced speed of processing) and peripheral side effects. In contrast, a lower dose of donepezil caused no clear attentional effect but improved cognitive flexibility. These findings document domain-specific dose-response effects of donepezil for attention and cognitive flexibility.

METHODS AND MATERIALS

Nonhuman Primate Testing Protocol

Three adult male rhesus macaques were separately given access to a cage-mounted Kiosk Station that provided a touchscreen interface inside the animal’s housing unit to perform cognitive tasks (Figure 1A) (28) (see the Supplement). The behavioral tasks were controlled by the Unified Suite for Experiments (33).

Drugs and Procedures

We used donepezil (Sigma-Aldrich, catalog number D6821) in three doses, 0.06, 0.1, and 0.3 mg/kg, to operate within the dosing range of previous studies reporting pro-cognitive effects (surveyed in Tables S1 and S2). At this intramuscular (IM) range, plasma concentrations of donepezil are roughly the same when dosing with ~10x the concentration orally (15). Animals received saline as vehicle control or a dose of donepezil IM injection 30 minutes before starting task performance, taking into account its expected 1-hour half-life (34).

Administration was double blinded. Drug side effects were assessed 15 minutes after drug administration and after completion of the behavioral performance with a modified Irwin Scale (35–38) for rating autonomic nervous system (e.g., salivation) and somatomotor system functioning (e.g., posture, unrest). Monkey behavioral status was video-monitored throughout task performance (Figure 1A).

Behavioral Paradigms

In each experimental session, the monkeys performed a VS task to measure attentional performance metrics and a feature-reward learning (FL) task to measure cognitive flexibility metrics (39). Each performance day was made up of an initial set of 100 trials of VS, a set of 21 learning blocks with 35 to 60 trials each of the FL task, and a second set of 100 trials of the VS task (Figure 1Aii). Details of both tasks are provided in the Supplement. The VS task required the monkeys to find and touch a target object among 3, 6, 9, or 12 distracting objects to receive fluid reward (Figure 1B). The target was an object shown in 10 initial trials without distractors. Targets and distractors were multidimensional, 3D-rendered quadrille objects (33) that shared few or many features of different feature dimensions (colors, shapes, arms, body patterns), which rendered the search easier when there were no or few similarities among the features of targets and distractors or more difficult if the target-distractor (T-D) similarity was high (Figure 2A). The FL task required the monkeys to learn through trial and error which object feature was rewarded in blocks of ~35 to 60 trials (Figure 1C). The rewarded feature changed uncued and switched to a new feature of the same or different feature dimensions, which made the task similar to conceptual set-shifting tasks [e.g., (40,41)] but different by using a larger set of features that varied within and across sessions to vary task difficulty. In each trial, three objects were shown; they varied either in the features of one feature dimension (e.g., having different colors or body shapes) or in the features of two feature dimensions (e.g., having different colors and body shapes). Choosing the object with the correct feature was rewarded with a probability of 0.8. Blocks in which only one feature dimension varied (1D blocks) were easier because there was lower attentional load than in blocks with two varying feature dimensions (2D blocks).

Neurochemical Confirmation of Drug Effect

To evaluate the levels of donepezil in brain structures that are necessary for successful attention and learning performance, we measured the ACh metabolite choline and donepezil concentrations in the PFC and the anterior striatum (caudate nucleus) 15 minutes after administering low and high doses of donepezil (0.06 and 0.3 mg/kg, respectively, IM) in separate experiments. Measures of donepezil were made when we observed dose–limiting side effects at the 0.3-mg/kg dose, and the two tested doses were accompanied by pro-cognitive effects in our task (see results). We used microprobes that sampled the local neurochemical milieu with the principles of solid phase microextraction (SPME) (42) followed by quantification of the concentrations with liquid chromatography and mass spectrometry (42). The detailed procedures used are described in (43) and in the Supplement.
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Figure 1. Task design, metastructure, and VS performance as a function of distractor number. (A) Picture of one of the subjects working in the custom-built kiosk, interacting with the touchscreen and receiving fluid reward. (B) The metastructure of the multitask. Each experimental session consists of three superfblocks of VS, FL, and VS, in turn. Each VS block is preceded by 10 familiarization trials, which are identical to a VS trial but without any distractors. Each VS block contains trials with 3, 6, 9, or 12 distractors randomly selected and counterbalanced over the block. In contrast, each FL block contains 0 or 1 irrelevant feature dimensions in addition to the relevant feature dimension (the dimension with the rewarded feature value) counterbalanced over the session. (B) (i) From the grand pool of quaddles, which includes four feature dimensions and a variable number of feature values (9 shapes, 9 patterns, 8 colors, and 11 arms), three feature values from three feature dimensions are chosen. This 3 x 3 pool is then counterbalanced for dimension presentation and feature-reward association and is used for 2 weeks of data collection where all presented quaddles are selected from this 3 x 3 pool. (ii) Example trials. Two example VS trials (top) within the same block with 3 distractors (left) and 9 distractors (right). Each VS block will contain one of five backgrounds, with the VS blocks in the same day never having the same background. All distractors and target objects in VS blocks are three-dimensional objects, and distractors may be duplicated in each trial. Quaddles are spatially randomly presented at the intersections of a 5 x 4 virtual grid pattern on screen. The red box highlights the rewarded target object, which is invariant within the VS block, in these examples. Two example FL trials (bottom) within the same block containing 2D quaddles (1 distracting dimension plus the relevant dimension). The rewarded feature value in this block is the checkered pattern independent of what color feature value it is paired with. Quaddles may be presented in eight possible locations in a circle, each being 17° of visual radius away from the center of the screen. The red box signifies the rewarded target object, which is a variable combination of the rewarded feature value (the checkered pattern in this example) with a random feature value of the distractor dimension (color in this example). (C) The trial structure for both the FL (top) and VS (bottom) blocks of the task are very similar. A trial is initiated by a 0.3 to 0.5 s touch and hold of a blue square (3° visual radius wide), after which the blue square disappears for 0.3 to 0.5 s before task objects, which are 2.5° visual radius wide, are presented on screen. Once the task objects are on screen, the subject is given 5 s to visually explore and select an object via a 0.2 s touch and hold. A failure to make a choice within the allotted 5 s results in an aborted trial and does not count toward the trial count. Brief auditory feedback and visual feedback (a halo around the selected object) are provided on object selection, with any earned fluid reward being provided 0.2 s after object selection and lasting 0.5 s along with the visual feedback. Nonrewarded trials had a different auditory tone and a light blue halo around the selected, nonrewarded object. Rewarded objects had a higher pitch auditory tone, a light yellow halo around the selected rewarded object, and an accompanying fluid (water) reward. (D) Average VS performance by distractor number for the vehicle and all donepezil doses combined, both separated by the first vs. second VS block. VS performance was significantly different for block number (F1,1722 = 22.19, p < .001) and condition (F1,1722 = 19.0, p < .001). The inlay shows individual monkey average VS performance linear fits. (E) Average VS performance by distractor number between vehicle and 0.06, 0.1, and 0.3 mg/kg donepezil doses for the first VS block (F3,1722 = 10.77, p < .001). Both the 0.06- and 0.3-mg/kg doses were significantly different from vehicle (Tukey’s, p = .005 and p < .001, respectively). Error bars here reflect standard deviation in this panel. The set size effect of VS performance by distractor number for each condition. The 0.3-mg/kg dose set size effect was significantly shallower than the vehicle set size effect (H(3) = 11.46, p = .010; Tukey’s, p = .013). FL, feature-reward learning; n.s., not significant; Num, number; Perf, performance; Prop, proportion; VS, visual search.

Statistical Analysis
Data were analyzed with standard nonparametric and parametric tests as described in the Supplement.

RESULTS
Each monkey was assessed in 38 sessions in total, including 17 vehicle days and 7 days with each dose (0.06, 0.1, and 0.3 mg/kg). Drug side effects were noted after IM injections of the 0.3-mg/kg dose in the first 30 minutes after injection as changes in posture, sedation, vasovasodilation, and paleness of skin, but no adverse effects persisted beyond 30 minutes (Table S3). First, we confirmed that the monkeys performed the VS task with a high 84.4% (±0.54%) accuracy (monkeys Ig: 85.2% ± 0.81%; Wo: 88.3% ± 0.94%; St: 79.8% ± 0.97%) and showed the expected set size effect evident in decreased accuracy and slower reaction times with increasing numbers of distractors (Figure 1D and Figures S1 and S2). When the targets were more similar to distractors (high T-D similarity), VS performance decreased from 92.9% (±0.4%) to 85.5% (±0.3%) and 81.6% (±1.0%) for low, medium, and high T-D similarity, respectively (H[2] = 169.48, p < .001) (Figure 2B). In the feature-reward learning task, the monkeys reached the learning criterion faster in the easier 1D (low distractor load) condition (average trials to ≥80% criterion: 12.5 ± 0.2 SE) than in the 2D (high distractor load) condition (average trials to ≥80% criterion: 15.6 ± 0.2) (Figure 3A and Supplement).

Dose-Dependent Improvement of VS Accuracy and Slowing of Choice Reaction Times
Donepezil significantly improved the accuracy of the VS task (F1,1722 = 18.95, p < .001) (Figure 1D) but on average slowed search reaction times (F1,1722 = 4.83, p = .028) (Figure S1B).
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The slower choice reaction times were evident already for the single target object in the 10 target familiarization trials (Figure S1A). These main behavioral drug effects were evident prominently in the first VS block (Figure 1D and Figure S1A). We therefore focused our further analysis on the first search block.

The improved accuracy of VS was dose dependent ($F_{3,896} = 10.77, p < .001$). The 0.06-mg/kg dose enhanced performance by 2.5% ± 1.0%, 4.4% ± 1.3%, 6.1% ± 1.4%, and 6.3% ± 1.6% (mean ± SD) for 3, 6, 9, and 12 distractor trials, respectively (Tukey’s, $p = .005$). The 0.3-mg/kg dose enhanced performance by 2.7% ± 1.0%, 6.3% ± 1.2%, 8.5% ± 1.3%, and 11.0% ± 1.4% (mean ± SD) for 3, 6, 9, and 12 distractor trials, respectively (Tukey’s, $p < .001$) (Figure 1E). Thus, we found larger improvement when more distractors interfered with the target search. We confirmed this by fitting a regression line across performance at different numbers of distractors, which revealed overall, significantly shallower slopes with donepezil (slopes: $-0.013 ± 0.001$, $-0.009 ± 0.002$, $-0.015 ± 0.003$, and $-0.005 ± 0.002$ for vehicle, 0.06, 0.1, and 0.3 mg/kg of donepezil, respectively [$H_{3} = 11.46, p = .013$]). Pairwise comparison showed that the 0.3-mg/kg drug dose and the vehicle condition showed significantly different slopes (Tukey’s, $p = .013$) (Figure 1F).

In contrast to improving VS accuracy, donepezil slowed down reaction times across all distractor conditions at the 0.3-mg/kg dose relative to vehicle by on average 100 ± 40 ms, 238 ± 79 ms, 208 ± 99 ms, and 264 ± 102 ms (mean ± SD) for 3, 6, 9, and 12 distractors, respectively ($F_{3,896} = 15.15, p < .001$; Tukey’s, $p < .001$) (Figure S1C). The slope of the regression over different numbers of distractors did not differ between the 0.3-mg/kg dose and vehicle, which showed that reaction time effect was a nonselective effect that was independent of distractors (regression slope on reaction times: 0.061 ± 0.002, 0.065 ± 0.007, 0.067 ± 0.007, and 0.076 ± 0.009 [$H_{3} = 3.37$, not significant (n.s.)] for vehicle, 0.06, 0.1, and 0.3 mg/kg of donepezil, respectively) (Figure S1D).

Across sessions, VS accuracy was correlated with reaction times only for the vehicle (Pearson, $r = -.30, p < .001$) and 0.1-mg/kg donepezil dose condition (Pearson, $r = -.46, p = .034$) but not for the 0.06- and 0.3-mg/kg dose conditions in which monkeys showed improved accuracy, which suggests that the accuracy improvement is independent from a slowing of reaction speed (Figure S2A, B).

We next tested whether improved interference control over increasing number of distractor objects was likewise evident when increasing the similarity of distractor and target features (Figure 2A). First, we confirmed that higher T-D similarity overall reduced performance ($F_{2,672} = 16.17, p < .001$) (Supplement). Donepezil significantly counteracted this similarity effect and improved performance at the 0.06- and 0.3-mg/kg doses ($F_{2,672} = 7.75, p < .001$; Tukey’s, $p = .034$ and $p < .001$, respectively). This finding shows that the beneficial effect of donepezil significantly increased when there was higher demand to control perceptual interference from distracting objects (Figure 2B). This was also evident as a statistical trend of a shallower regression slope at 0.06- and 0.3-mg/kg doses of donepezil, which indicates less interference from distracting features when they were similar to the target (Figure 2C) [$H_{3} = 2.79$, n.s.; slope changes relative to vehicle for 0.06-, 0.1-, and 0.3-mg/kg doses were 0.0357 ± 0.0236, −0.0289 ± 0.0334, and −0.0656 ± 0.0197, respectively). The improved search performance with donepezil for VS with higher T-D similarity and with a higher number of distractors was evident in significant main effects, but there was no interaction, suggesting that they improved performance independent of each other ($F_{2,615} = 64.59, p < .001$; $F_{2,615} = 28.85, p < .001$; and $F_{2,615} = 0.69$, n.s., respectively) (Figure 2D). This independence was also suggested by the absence of a correlation of the T-D similarity effect and the number-of-distractor effect (Pearson, n.s.) (Figure S3).

Dose-Dependent Improvement of Flexible Learning Performance

Donepezil also improved feature-reward learning performance but only at the 0.06-mg/kg dose (Figure 3B), and it was most pronounced for the first third of the behavioral session ($F_{3,602} = 3.3, p = .020$) (Figure 3C). We therefore focused further analysis on the first of the learning blocks, which revealed that the learning improvement at the 0.06-mg/kg dose was significant for the low distractor load condition (significant interaction effect of drug condition and distractor load [condition × distractor load]: $F_{3,1052} = 3.59, p = .013$); for vehicle, 0.06-, 0.1-, and 0.3-mg/kg donepezil doses, the number of trials to criterion were 11.3 ± 4.7, 7.7 ± 0.9, 12.3 ± 1.3, and 11.0 ± 2.2, respectively, with the 0.06-mg/kg dose and the vehicle being significantly different ($p = .020, Bonferroni correction$) (Figure 3D). There was no change in the learning speed with other doses at low or high distractor load.

Beyond learning speed, we found overall slower choice reaction times at the 0.3 mg/kg donepezil dose (Figure 3E) (main effect of drug condition: $F_{3,1052} = 12.29, p < .001$). While reaction times were overall slower at the high distractor load ($F_{1,1052} = 7.18, p = .008$), there was no interaction with drug dose ($F_{3,1052} = 0.26$, n.s.) After visually inspecting the results, we separately tested the 0.3-mg/kg dose of donepezil and found that it led to significantly slower choice reaction time than the vehicle (Tukey’s, $p < .001$) (Figure 3E). The changes in choice reaction times did not correlate with changes in learning performance (number of trials to criterion) at any drug condition, indicating that they were independently modulated (Pearson, all n.s.) (Figure S2D).

We predicted that the faster learning at the 0.06 mg/kg donepezil dose could be due to a more efficient exploration of objects during learning, which would be reflected in reduced perseverative choices of unrewarded objects. Overall, perseverative errors (defined as consecutive unrewarded choices of objects with the same feature dimension) made up 20% of all errors. As expected, we found significantly shorter sequences of perseveration of choosing objects within distractor feature dimensions at the 0.06-mg/kg dose of donepezil (Figure 3F). For vehicle, 0.06-, 0.1-, and 0.3-mg/kg doses, the average length of perseverations in the distractor dimension was 2.1 ± 0.1, 1.8 ± 0.1, 1.9 ± 0.1, and 1.9 ± 0.1 trials, with the difference between the vehicle and the 0.06 dose being significant ($p = .021$). Perseverative choices in the target feature dimension were not different between conditions for 0.06-, 0.1-, and 0.3-mg/kg donepezil doses, the average
Dissociation of Attention and Learning Improvements, but Slowing Is Correlated

The effects of donepezil on feature-reward learning and VS might be related, but we found that learning speed and search accuracy were not correlated at the doses at which the drug improved learning and search (0.06-mg/kg dose) or improved only VS (0.3-mg/kg dose) (Pearson, all n.s.). A significant correlation was found only for the 0.1-mg/kg dose (Pearson, ρ: −0.54; p = .012) (Figure 4A). Learning at low or high distractor load and the set size (slope) effects in the VS task were uncorrelated (Pearson, all n.s.). However, at the 0.3-mg/kg donepezil dose, the T-D similarity effect (i.e., the search-slope change) in the VS task positively correlated with the difference of learning speed at high versus low distractor load in the learning task (Pearson, ρ: 0.60; p = .008). This effect signifies that better attentional search of a target among similar distractors is associated with poorer flexible learning of new targets when there are multiple object features to search through (high distractor load).

In contrast to accuracy, choice reaction times in the learning task and VS were significantly correlated for the 0.1-mg/kg donepezil dose (Pearson, ρ: 0.52; p = .016), the 0.3-mg/kg dose (Pearson, ρ: 0.66; p = .002), and the vehicle control condition (Pearson, ρ: 0.60; p < .001) (Figure 4B).

Determination of Extracellular Donepezil and Choline Levels in the PFC and Anterior Striatum

VS and flexible learning are realized by partly independent brain systems, including the PFC and anterior striatum (44). To determine whether extracellular levels of donepezil were increased to a similar magnitude in the PFC and anterior striatum, we measured its concentration after administering doses of either 0.06 or 0.3 mg/kg donepezil IM in the PFC, assumed to be necessary for efficient interference control during VS (19), and in the head of the caudate nucleus, which is necessary for flexible learning of object values (20,21). We used a recently developed microprobe that samples chemicals in the neural tissue based on the principles of SPME (42,43). We found that donepezil was available in both brain areas, and its extracellular concentration more than doubled after injecting 0.3 mg/kg compared with 0.06 mg/kg in both areas (F1,16 = 9.69, p = .007), with no significant difference between PFC and caudate (F1,16 = 1.44, n.s.) (Figure 5A). Donepezil should cause a depletion of the ACh metabolite choline (45). Using high-performance liquid chromatography–mass spectrometry analysis of the SPME samples, we found that 0.06 and 0.3 mg/kg donepezil reduced choline concentrations by 74.2% ± 14.9% (p = .005) and 85.7% ± 26.9% (p = .007) of their baseline concentrations in the PFC and, by 68.4% ± 13.8% (p = .022) and 81.0% ± 12.9% (p = .009) of respective baseline concentrations in the caudate (Figure 5B). The 11.5% and 12.6% stronger reduction of choline at the 0.3-versus 0.06-mg/kg dose in the PFC or caudate was not significant (n.s.).

To obtain an independent physiological marker of dose-dependent effects, we quantified how, during actual task performance, donepezil changed the heart rate before versus after drug administration (Supplement). The heart rate showed a transient peak ~20 minutes after the donepezil injection relative to baseline, which was significant for the 0.3-mg/kg dose (preinjection 102.3 ± 7.1 to postinjection 121.6 ± 2.6; p = .02) but not for the 0.06-mg/kg dose (preinjection: 90.3 ± 4.2 to postinjection: 94.8 ± 5.4; n.s.). The 0.3-mg/kg dose caused a significantly higher heart rate peak than the 0.06-mg/kg dose (p = .006) (Figure 5C).

DISCUSSION

Here, we dissociated donepezil’s improvement of attentional control of interference during VS from improvements of cognitive flexibility during FL. At the highest dose tested, donepezil reduced interference during VS, particularly when...
there were many distractors and a high similarity of distractors with the target, while concomitantly slowing down overall reaction times and inducing temporary peripheral side effects. In contrast, the lowest dose of donepezil did not affect target detection times during VS but improved adapting to new feature-reward rules and reduced perseverative responding. These findings document a dose-dependent dissociation of the best dose of donepezil for improving attention and cognitive flexibility.

**Different Donepezil Dose Ranges for Improving Interference Control and Flexible Learning**

Using a behavioral assessment paradigm with two tasks allowed us to discern the differences between the donepezil dose that maximally improved interference control (in the VS task) versus the dose that maximally improved flexible learning (in the reward learning task). In both tasks, donepezil modulated performance early within the session (first of two VS blocks and the first third of the learning blocks) consistent with its short half-life and rapid time to peak concentration with IM delivery (15,34). Our results focused therefore on these early time windows. We do not expect different conclusions if we had altered the task sequence (see Supplemental Discussion). At the 0.06-mg/kg dose, donepezil facilitated flexible learning of a new feature-reward rule and reduced the length of perseverative errors (Figure 3C, F). These behavioral effects are indicators of improved cognitive flexibility across reward learning and set-switching tasks (46–48). At the same 0.06-mg/kg dose, VS response times were unaffected (Figure S1) and donepezil doses that maximally improved interference control (in the VS task) and the dose that maximally improved flexible learning (in the reward learning task). In both tasks, donepezil modulated performance early within the session (first of two VS blocks and the first third of the learning blocks) consistent with its short half-life and rapid time to peak concentration with IM delivery (15,34). Our results focused therefore on these early time windows. We do not expect different conclusions if we had altered the task sequence (see Supplemental Discussion). At the 0.06-mg/kg dose, donepezil facilitated flexible learning of a new feature-reward rule and reduced the length of perseverative errors (Figure 3C, F). These behavioral effects are indicators of improved cognitive flexibility across reward learning and set-switching tasks (46–48). At the same 0.06-mg/kg dose, VS response times were unaffected (Figure S1) and
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Figure 5. In vivo extracellular measurements of choline and donepezil, as well as donepezil’s effect on heart rate. (A) Quantified concentration of extracellular unbound donepezil with 0.06- and 0.3-mg/kg donepezil administration in the PFC and CD. We were able to reliably detect higher donepezil concentrations with 0.3 mg/kg dosing relative to 0.06 mg/kg dosing (condition $F_{1,6} = 9.69, p = .007$) with solid phase microextraction. We also saw a trend toward higher detectable donepezil in the CD relative to the PFC at the 0.3-mg/kg dose tested; however, we found neither significant group nor interaction effects. (B) We used choline concentrations as a metric for donepezil bioactivity because it deactivates acetylcholinesterase and prevents acetylcholine’s degradation into choline. We extracted average sessionwise change in choline from baseline with 0.06 and 0.3 mg/kg donepezil doses within the PFC and CD. Although we found significant decreases in choline by up to $>80\%$ of baseline concentrations, we found no significant effect of dosing in either the PFC or CD. (C) The HR of our fourth monkey was monitored during the neurochemical experiments. This revealed a sharp and transient increase in HR after administration of donepezil at the 0.3-mg/kg dose (Supplement), which led to a higher average beats per minute (bpm). We found that we can significantly distinguish 0.06 and 0.3 donepezil administration via HR ($p = .006$). CD, caudate; HR, heart rate; n.s., not significant; PFC, prefrontal cortex.

VS accuracy was overall improved but independent of the number of distractors, i.e., independent of the degree of interference (Figure 1E, F). In contrast, at the higher donepezil doses, flexible learning behavior was indistinguishable from the no-drug vehicle control condition, showing that improving flexibility required donepezil at a lower dose.

This conclusion is opposite to that of the drug effects on VS performance, which was maximally improved at the 0.3-mg/kg dose. At this dose, subjects not only showed improved resistance to interference when there were more distracting objects (Figure 1E, F) but also improved resistance to distracting objects that were visually similar to the searched-for target (Figure 2B–D). These findings document that donepezil enhances robustness to distraction (49,50), which critically extends insights from existing primate studies with donepezil that mostly used simpler tasks to infer pro-cognitive effects on working memory or arousal (Table S1). The process of attentional control of interference also goes beyond a short-term memory effect measured with delayed match-to-sample tasks. In the VS paradigm we used, short-term memory of the target object is already necessary for performing the easier trials with 3 or 6 distractors, while an attention-specific effect can be inferred when there is greater improvement in performance with increased attentional demands in trials with 9 or 12 distractors. Thus, our study provides strong evidence that donepezil causes specific attentional improvement at higher doses, which supports the neurogenetic model of cholinergic modulation of attention (51) that recently has received functional support in studies reporting enhanced distractor suppression in nonhuman primates with nicotine receptor–specific ACh modulation (52–54) and improved suppression of perceptually distracting flankers in human subjects tested with a single dose (55). We should note, however, that at the high dose, donepezil already caused a nonselective slowing of reaction times indicative of peripheral side effects (see Supplemental Discussion).

The finding that different dose ranges improved flexible learning and VS distractor filtering suggests that these processes have partially independent Yerkes-Dodson style inverted-U dose-response curves (Figure 6). One reason supporting this suggestion is that flexible learning and distractor filtering are supported by partially different brain networks, which likely have differential sensitivity to cholinergic modulation. Lesion studies in nonhuman primates have shown that flexible reward learning is closely associated with the medial and orbitofrontal PFC and the striatum where lesions impair learning visual reward associations (46,56). In contrast, VS distractor filtering in primates depends on the dorsolateral PFC (dlPFC) and its connections with posterior parietal cortices, with bilateral dlPFC lesions impairing filtering distraction (57). Brain areas within these partly segregated networks for learning and distractor filtering might be differently sensitive to cholinergic modulation. For example, primate dlPFC has been documented to be uniquely sensitive to neuromodulation by catecholamines and ACh for spatial working memory and switching between distracting features (5,58), with ACh depletion in PFC causing deficits in attention but not learning.
During cognitive processes, ACh modulates synaptic efficacy postsynaptically in an inverted-U manner through both alpha 7 nicotinic receptors (60) and M1 muscarinic receptors (61). Such inverted-U curves for different receptors are not likely to be homogenous or fully overlapping when taking into consideration variable task demands within a cognitive domain or when considering different cognitive domains entirely (62). This is supported by studies showing disruption of rule-selective activity with iontophoretic M1 overstimulation of dIPFC neurons (63), while at lower doses, delay-cell firing and spatial tuning were enhanced (61). Our results may thus reflect different inverted-U curves along a construct of flexible attention shifting, required for optimal performance in our feature-reward learning task, and stable filtering of distractors required for optimal performance in our VS task (Figure 6).

Quantifying Extracellular Levels of Donepezil and Choline in PFC and Striatum

We confirmed the presence of extracellular donepezil in the PFC and anterior striatum at the doses tested (Figure 5A) and that it prevented ACH metabolism as evident in 68% to 86% reduced choline levels (Figure 5B). To our knowledge, this is the first quantification of donepezil action on the breakdown of ACh in two major brain regions in the primate. The observed reduction of choline is higher than reductions of AChE activity (of ~25%–70%) reported with positron emission tomography or in brain homogenate (64,65). Previous studies suggest that evaluating blood plasma levels or cerebrospinal concentrations may not predict how effectively AChE drugs influence behavioral outcomes (66). One likely reason is that intracerebral concentrations can be multifold higher than extracerebral concentration levels (64,67) and do not reflect the actual bioactive concentration available in target neural circuits. By confirming that donepezil prevented ACh breakdown in the PFC and striatum, we thus established a direct link between behavioral outcomes and local drug bioavailability in two brain structures that causally contributes to attention and learning (see above) (46,56,57,59,68). While our study showed that donepezil has a similar effect on ACh breakdown in both areas, it leaves unanswered whether or how choline concentrations in either brain area relate to finer performance variations across tasks because we measured choline only during one task and with insufficient statistical power to establish such a link at this stage.

The neurochemical measurements of donepezil in the PFC and striatum were achieved with a recently developed microprobe that samples neurochemicals through principles of SPME (42,43,69–71) and so far was used for testing the consequence of drugs only in rodents (70,72,73). We believe that leveraging this technique in primate drug studies will be important for clarifying whether systemically administered drugs reach the desired target brain systems in which they are supposed to exert their pro-cognitive effects.

In our study, confirming donepezil action in the PFC and striatum critically constrains the interpretation of the behavioral results, suggesting that different behavioral outcome profiles are not due an uneven drug availability. Rather, the different best doses for VS and flexible learning performance will likely be due to brain area–specific pharmacokinetic profiles of receptor densities, drug clearance profiles, or autoreceptor mechanisms that intrinsically downregulate local drug actions (74–76). One prediction from the specific distribution and kinetics of nicotinic or muscarinic receptors in the PFC and striatum is that donepezil might, at lower doses, act predominantly in the striatum via activation of muscarinic subreceptors because they have a particularly high binding potential (18) and respond stronger to muscarinic ACh receptor activation compared with the PFC (17) (see Supplemental Discussion). However, it is also possible that donepezil recruits nicotinic receptors, which are upregulated with chronic donepezil use (77). It is important to disentangle, in future studies, the role of nicotinic and muscarinic subreceptors in the PFC and striatum to optimize the clinical potential to improve learning and attention functions in conditions with cognitive impairment and particularly in dementia (see Supplemental Discussion).

In summary, our results provide rare quantitative evidence that a prominent Ach-enhancing drug exerts domain-specific cognitive improvements of attentional control and cognitive flexibility at a distinct dose range. A major implication of this finding is that for understanding the strength and limitations of pro-cognitive drug compounds, it will be essential to test their dose-response efficacy at multiple cognitive domains.

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