COGNITIVE EFFECTS OF DOPAMINE DEPLETION IN THE CONTEXT OF DIMINISHED ACETYLCHOLINE SIGNALING CAPACITY.

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Abbreviated title: Cognitive effects of dual dopamine and acetylcholine depletion

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ABSTRACT

A subset of patients with Parkinson’s disease acquires a debilitating dementia characterized by severe cognitive impairments (i.e. Parkinson’s Disease Dementia; PDD). PDD brains show extensive cholinergic loss as well as dopamine (DA)-depletion. We used a mutant mouse model to directly test whether combined cholinergic- and DA depletion leads to a cognitive profile resembling PDD. Mice carrying heterozygous deletion of the high-affinity, hemicholinium-3 sensitive choline transporter (CHT\textsuperscript{HET}) show reduced levels of acetylcholine throughout the brain. We achieved bilateral DA-depletion in CHT\textsuperscript{HET} and wild-type (WT) littermates via intra-striatal infusion of 6-hydroxydopamine (6-OHDA) or vehicle. Executive function and memory were evaluated using rodent versions of cognitive tasks commonly used with human subjects: the set-shifting task and spatial and novel-object recognition paradigms. Our studies revealed impaired acquisition of attentional set in the set-shifting paradigm in WT-6OHDA and CHT\textsuperscript{HET}-vehicle that was exacerbated in the CHT\textsuperscript{HET}-6OHDA mice. The object recognition test following a 24-hour delay was also impaired in CHT\textsuperscript{HET}-6OHDA compared to all other groups. Treatment with acetylcholinesterase (AChE) inhibitors physostigmine (0.05 or 0.1 mg/kg) and donepezil (0.1 and 0.3 mg/kg) reversed the impaired object recognition of the CHT\textsuperscript{HET}-6OHDA mice. Our data demonstrate an exacerbated cognitive phenotype with dual ACh-DA depletion as compared to either insult alone, with traits analogous to those observed in PDD patients. The results suggest that combined loss of DA and ACh may be sufficient for pathogenesis of specific cognitive deficits in PDD.

**Keywords:** Parkinson's disease dementia; mouse model; choline transporter; 6-OHDA; set-shifting; object recognition; acetylcholinesterase inhibitor; physostigmine; donepezil.
INTRODUCTION

Parkinson's disease (PD) has traditionally been viewed as a motor disorder caused by degeneration of dopaminergic neurons supplying dopamine (DA) to the striatum. From yearly stages, many PD patients show cognitive deficits known as mild cognitive impairment (Aarsland et al., 2011). Although its etiology is still unclear, the loss of DA in the basal ganglia circuitry appears important (Aarsland et al., 2011; Kehagia et al., 2010a; Sawamoto et al., 2008). In rodents (Braga et al., 2005; Da Cunha et al., 2006; De Leonibus et al., 2007; Moriguchi et al., 2011) and primates (Decamp et al., 2004; Lipina and Colombo, 2007), DA depletion impairs cognition. Most PD patients eventually develop dementia (Aarsland et al., 2003). Patients with PD and dementia (PDD) are impaired on tasks of executive function, defined as the ability to self-generate and alter plans and rules that guide behavior (Janvin et al., 2005; Noe et al., 2004; Parretta et al., 2005). Additionally, verbal and visual memory and visual-spatial abilities are impaired [(Higginson et al., 2005; Kuzis et al., 1999; Mosimann et al., 2004; Muslimovic et al., 2007; Noe et al., 2004), see also (Kehagia et al., 2010b) and references therein].

The pathogenesis of dementia in PD remains poorly understood. Since the loss of DA is comparable in non-demented PD and patients with PDD (Bychkov et al., 2008; Colloby et al., 2005; Hilker et al., 2005; Ito et al., 2002; Joyce et al., 2002), additional neural insults must contribute to dementia in PDD. Severe alterations in cholinergic pathways are found in PDD. PDD subjects show neuronal loss in the nucleus basalis (Nakano and Hirano, 1984; Whitehouse et al., 1983), the extent of which correlates with the severity of dementia (Perry et al., 1987). Choline acetyltransferase is reduced in the neocortex in PDD (Lange et al., 1993; Ruberg et al., 1982), and the reduction correlates with cognitive impairment (Dubois et al., 1983; Perry et al., 1985). The density of vesicular acetylcholine (ACh) transporters is reduced throughout the
neocortex in PDD but only in the parietal and occipital cortices in PD (Kuhl et al., 1996). PET markers for ACh pathways are reduced in the neocortex in PDD as compared to non-demented PD (Bohnen et al., 2003; Hilker et al., 2005; Klein et al., 2010). ACh esterase (AChE) inhibitors improve cognition in PDD patients (Aarsland et al., 2003; Giladi et al., 2003; Hutchinson and Fazzini, 1996). Benefits are modest, however, possibly due to dose-limiting side effects [reviewed in (Kurtz and Kaufer, 2011)]. Thus, both cholinergic deficit [reviewed in (Campbell et al., 2009; Hasselmo and Sarter, 2011; Klinkenberg and Blokland, 2010)] and loss of DA (Aarsland et al., 2011; Johnson and Galvin, 2011) impair cognition in humans and animals. We hypothesize that dementia in PD results from pronounced ACh deficit on the background of extensive DA depletion. Here we employ a novel mouse model to test the hypothesis that combined DA/ ACh depletion, but not the loss of ACh or DA alone, induces cognitive deficits characteristic for PDD, specifically, visual-spatial recognition and executive function.

To model widespread depletion of ACh, mutant mice heterozygous for a deletion of the high affinity, hemicholinium-3-sensitive choline transporter (CHT^{HET}) were used. CHT protein transports choline as a rate-limiting step in the synthesis of ACh (Ferguson et al., 2003). CHT^{HET} mice show roughly half the levels of ACh as compared to wild-type (WT) littermates (Bazalakova et al., 2007) but retain intact motor functions and basic cognitive abilities. Therefore, they appear ideal for assessing possible interactions between ACh- and DA-depletion on cognitive impairment.

**METHODS AND MATERIALS**

**Animals and Stereotaxic Surgery**

All animal procedures were approved by the Vanderbilt University Institutional Animal Care and Use Committee. Male mice with heterozygous deletion of the CHT protein (CHT^{HET})
have been previously described (Bazalakova et al., 2007; Ferguson et al., 2004; Ferguson et al., 2003). CHT\textsuperscript{HET} are congenic on a C57BL/6 background and were housed up to five per cage on a 12-hour light:dark cycle with food and water \textit{ad libitum}, except when food restriction was required for training. All experimental procedures were performed during the light phase of the diurnal cycle.

At 11-13 weeks of age, mice received bilateral intra-striatal injections of 6-hydroxydopamine (6-OHDA; 2 µg in 1µl of 0.05% ascorbic acid in phosphate-buffered saline) or vehicle at stereotaxic coordinates (from bregma) AP 0.65, ML ± 2.5, DV(skull) 3.2. Mice were anesthetized with ketamine/xylazine mixture (100 mg/kg/10 mg/kg, i.p.). Desipramine (20 mg/kg i.p; Sigma, St. Louis, MO) was given 30 min prior to 6-OHDA injection to protect noradrenergic terminals.

**Neurotransmitter measurements**

**Monoamines:** To assess the time course and extent of DA decline following intra-striatal 6-OHDA, unilaterally injected mice were decapitated under isoflurane anesthesia 7 (WT n = 5; CHT\textsuperscript{HET} n = 7), 14 (WT n = 6; CHT\textsuperscript{HET} n = 7), 21 (WT n = 4; CHT\textsuperscript{HET} n = 7), or 28 (WT n = 6; CHT\textsuperscript{HET} n = 6) days following surgery. The unilateral 6-OHDA lesion was employed in this particular case in order to determine the degree of DA loss at each post-surgical interval while conserving mice. Striatum was dissected on ice and rapidly frozen on dry ice prior to storage at -80°C until ready for monoamine analyses. HPLC determination of monoamine neurotransmitter and metabolites was performed by the Vanderbilt Center for Molecular Neuroscience Neurochemistry Core utilizing an Antec Decade II (oxidation: 0.4) electrochemical detector operated at 33° C and Phenomenex Kintex (2.6u, 100A) C18 HPLC column (100 x 4.60 mm).
Biogenic amines were eluted with a mobile phase consisting of 89.5% 0.1M TCA, \(10^{-2}\) M sodium acetate, \(10^{-4}\) M EDTA and 10.5 % methanol (pH 3.8) in the following order: noradrenaline, MHPG, Adrenaline, DOPAC, Dopamine, 5-HIAA, HVA, 5-HT, and 3-MT (Lindley et al., 1998). Solvent was delivered at 0.6 ml/min using a Waters 515 HPLC pump. HPLC control and data acquisition are managed by Empower software. Data from these analyses are expressed as ng/mg protein.

**Acetylcholine and choline:** To determine whether intra-striatal 6-OHDA altered ACh levels in the striatum or overlying cortex, bilaterally injected mice were rapidly decapitated 7 days following surgery (WT-vehicle \(n = 6\), WT-6OHDA \(n = 6\), CHT\(^{\text{HET}}\)-vehicle \(n = 6\), CHT\(^{\text{HET}}\)-6OHDA \(n = 5\)). Striatum and frontal cortex were dissected on ice following microwaving to inactivate AChE (Bertrand et al., 1994). Dissected tissue was rapidly frozen on dry ice and stored at -80°C until ready for use. ACh and choline levels were quantified using HPLC techniques by the Vanderbilt Center for Molecular Neuroscience Neurochemistry Core. The HPLC system was composed of a Waters 717+ autosampler, Water model 515 pump and Antec Decade electrochemical detector. Column employed was from Bioanalytical Systems (acetylcholine column) coupled with post column immobilized enzyme reactor (IMER) containing bound AChE and choline oxidase (ChO) that converted ACh to choline and further oxidized it to hydrogen peroxide. The final reaction product was detected amperometrically and quantified on a platinum working electrode (+400 mv) (Damsma et al., 1985).

Levels of ACh and choline from these analyses are expressed as nmol/mg protein.

**Quantitative Western Blot**
The expression of tyrosine hydroxylase (TH; an enzyme involved in the formation of DA) was measured in all animals tested for behavior. For the TH analysis, mice were decapitated under isoflurane anesthesia following completion of behavioral measures or, to create a timeline of TH decline following 6-OHDA lesion, at 7 (WT n = 5; CHT\textsuperscript{HET} n = 3), 14 (WT n = 3; CHT\textsuperscript{HET} n = 6), 21 (WT n = 3; CHT\textsuperscript{HET} n = 5), or 28 (WT n = 5; CHT\textsuperscript{HET} n = 5) days post-surgery. Brains were collected and rapidly frozen on dry ice. Both striata were outlined on precut 150\(\mu\text{m}\) thick coronal sections and scraped into 150\(\mu\text{l}\) of Lysis solution (Ambion, Austin, TX). Protein concentration in the samples was determined with the Bradford reagent (Bio-Rad, Hercules, CA). Samples were stored at -80°C until needed.

Protein sample preparation and Western blots were performed as previously described (Ahmed et al., 2007; Ahmed et al., 2010). Briefly, proteins were precipitated from Lysis buffer with nine volumes of methanol, pelleted by centrifugation, washed with 1mL of 90% methanol, dried, and dissolved in sodium dodecyl sulfate sample buffer at the final concentration of 0.1\(\mu\text{g}\) protein/\(\mu\text{l}\). For electrophoresis, 0.5\(\mu\text{g}\) protein from each sample were loaded on the gel. TH was detected with rabbit polyclonal primary antibody (Millipore, Temecula, CA) at 1:24,000. Secondary antibody was horseradish peroxidase-conjugated goat anti-rabbit at 1:25,000 (Jackson ImmunoResearch Laboratories, West Grove, PA). Blots were developed using SuperSignal enhanced chemiluminescence reagent WestPico (Pierce, Rockford, IL) and exposed to X-ray film for appropriate periods of time. Optical density of the bands was measured using Versadoc software (Bio-Rad, Hercules, CA). Standard curves were fitted to linear equations using Prism 4.0 (GraphPad Software, San Diego, CA).

**Immunohistochemistry**
TH immunostaining was performed as previously described (Ahmed et al., 2007; Ahmed et al., 2010). Briefly, mice were injected with ketamine/xylazine anesthesia 28 days following unilateral infusions of 6-OHDA and perfused intracardially with 4% paraformaldehyde. Following cryoprotection in 30% sucrose, brains were stored at -80°C until needed. Sections (30 μM) were obtained on a cryostat and incubated with rabbit polyclonal primary antibody for TH (1:500; Millipore, Temecula, CA) for 48 hours at 4°C. TH antibody was detected using fluorescently labeled goat anti-rabbit antisera (1:200; Vector Laboratories Inc., Burlingame, CA) and Streptavidine-Alexa 488 (1:200; Invitrogen, Eugene, OR), each incubated with TH antibody-labeled sections for 60 minutes at room temperature.

**Behavioral tasks**

**Rotarod:** Mice were tested for motor co-ordination and balance on the accelerating Rotarod apparatus (Model 7650, Ugo Basile, Napoli, Italy) following training in spatial/object recognition 19-21 days after surgery. Mice were placed on a cylinder, rotating at 5 rpm. Rotation speed increased to a maximum of 40 rpm gradually across 5 minutes. Latency (in seconds) to fall from the cylinder was recorded as the main dependent variable, with a maximum testing period of 500 seconds. Mice performed three trials per day for three consecutive days. The mean latency was calculated per day and used in statistical analyses.

**Set-shifting task (a.k.a. attentional set-shifting task):** A separate set of mice without previous training experience was used for these experiments. Procedures for the set-shifting task were adapted from previously used protocols for rats (Birrell and Brown, 2000; Wood et al., 1999) and mice (Bissonette et al., 2008; Colacicco et al., 2002). Procedures began following a one-week recovery period from intra-striatal vehicle or 6-OHDA infusion (WT-vehicle n = 7,
WT-6OHDA n = 8, CHT\textsuperscript{HET}-vehicle n = 7, CHT\textsuperscript{HET}-6OHDA n = 8). Data from two additional CHT\textsuperscript{HET}-vehicle mice were included in the reversal and striatal TH measures only, because these mice were trained in preliminary studies using a shorter protocol (i.e. reversal CHT\textsuperscript{HET}-vehicle n = 9). Mice were food restricted to 85% of their free-feeding weight and acclimated with the testing room and clear, polycarbonate chamber (30 x 30 x 30 cm with two 15 x 15 cm choice compartments at one end, Fig. 4A). Acclimation consisted of, on consecutive days: (1) exposure to the empty chamber for 10 minutes, (2) 4 trials with food cups holding uncovered food reward (1/2 piece of Honey-Nut Cheerios\textreg兵马), and (3) 4 trials with food reward hidden under unscented media. Media used during acclimation was not used again during the protocol. Testing continued for the next five days, with two sessions per day separated by two hours to prevent satiation during afternoon testing. Mice were trained to find food hidden under media scented by dried spices (i.e. odors). The location of the food reward (i.e. a correct response) was determined either by one of the odors (in which case the media was irrelevant) or one of the medias (in which case the odor was irrelevant). Mice were randomly assigned to approach an odor or media. Table 1 lists the odor-media combinations used per session. At each trial, mice were allowed to explore both bowls before making a choice by digging. During the first four trials of each session, mice were allowed to retrieve the food reward from the correct bowl after initially making an incorrect response. Following the fourth trial, no digging could be done following an incorrect choice.

Mice were initially trained on a simple discrimination (SD) where two odors were each paired with only one media type for odor-media set-shifting, or vise versa for media-odor set-shifting. Referred to as direction of shift, mice in each group were counterbalanced to ensure no bias towards a particular dimension. Following the example in Table 1, mice learned that clove-
scented dishes held food while basil-scented did not, with both clove and basil mixed into the same media. Session 2, a compound discrimination (CD), added a second media, for which clove and basil were each mixed with the previously used media plus an additional, novel media. In the CD, mice learned that clove-scented dishes held food, regardless of the media, and basil-scented never held food, regardless of the media. An intra-dimensional shift (IDS) modeled the compound discrimination paradigm with, importantly, novel odors and medias for each IDS. Reversal of the third intra-dimensional shift (session six) rewarded an odor that was incorrect in the immediately preceding session. An extra-dimensional shift (EDS) during the last session rewarded a media instead of an odor or the reverse. Poor performance during the EDS indicates successful formation of the attentional set. Mice were trained in each session until achieving the learning criterion of eight consecutively correct trials or for a maximum of 32 trials. The session maximum of 32 trials was chosen to avoid lack of performance due to satiation. A trial maximum of two minutes was imposed. For each session, the correct dish was counterbalanced to the right or left chamber compartment. Also, the direction of extra-dimensional shift (i.e. odor-media or media-odor) was counterbalanced for each treatment group. For each session, trials to criterion (i.e. eight consecutively correct trials) and quantity of errors were recorded. Performance on the EDS was further analyzed as the ratio of trials to criterion for the EDS versus the immediately preceding IDS (i.e. IDS VI).

**Spatial and Object Recognition:** A separate set of mice without previous training experience was used for these experiments. Following a one-week post-lesion recovery period, mice were transferred to the Center for Molecular Neuroscience's Laboratory for Neurobehavior Core holding facilities and given one week to acclimate. On training day, mice were placed in a clean holding cage and placed in the training anteroom for a minimum of 30 minutes prior to
initiation of the task. Training occurred across four sessions, initially in an empty chamber (clear polycarbonate, 50 x 50 x 40 cm) followed by three sessions with five objects (wiffle ball, plastic orange, clay leaf, plastic funnel, rubber duck) in a specific configuration (Figure 5A). A retention delay of either 5 minutes (WT-vehicle n = 10, WT-6OHDA n = 7, CHT\textsuperscript{HET}-vehicle n = 7, CHT\textsuperscript{HET}-6OHDA n = 7) or 24 hours (WT-vehicle n = 11, WT-6OHDA n = 11, CHT\textsuperscript{HET}-vehicle n = 9, CHT\textsuperscript{HET}-6OHDA n = 10) was imposed, followed by a spatial recognition test. Two previously encountered objects were moved to novel locations, resulting in a novel configuration of the same five objects. Mice were exposed to the novel configuration a second time, followed by an object recognition test in the final session. The object recognition test consisted of introducing a novel object (wheel caster) in place of a familiar, non-displaced object.

Mouse behavior was captured using an overhead camera and analyzed by AnyMaze tracking software (Stoelting, Wood Dale, IL) in real-time. Exploration was quantified as the amount of time spent within one centimeter of each object. Tracking was centered on the nose of the mouse and time spent on top of an object was subtracted. For the spatial recognition test, data were expressed as the percent of total exploration time spent with the two displaced objects. Similarly for the object recognition test, data were expressed as the percent of total exploration time spent with the one novel object. Animals were eliminated if exploration at any of the three acquisition sessions totaled less than five seconds. Sessions lasted six minutes with five-minute inter-session intervals. Preliminary studies were performed to verify a lack of preference amongst the six objects used.

The effect of physostigmine (eserine hemisulphate salt; Sigma, St. Louis, MO) and donepezil (donepezil hydrochloride, Sigma) on visual-spatial recognition was determined in a
separate group of WT-vehicle and CHT\textsuperscript{HET}-6OHDA mice. AChE inhibitors were chosen because they are predominantly used in the treatment and shown to effectively ameliorate cognitive impairments in PDD (Aarsland et al., 2002; Fuchs et al., 2004; Leroi et al., 2004). Immediately post-training (i.e. following the final acquisition session), mice were injected with physostigmine (i.p. 0.05 [n=10] or 0.1 [n=9] mg/kg), donepezil (i.p. 0.1 [n=7] or 0.3 [n=9] mg/kg or saline-vehicle (n = 9). As a control, vehicle-treated WT mice (n = 12) were also tested. To minimize stress from the injection, mice were handled for five out of seven days prior to training. Handling occurred for five minutes per day outside the vivarium, in a room distinctly different from the training room.

**Data Analysis**

All animals used in behavioral testing were analyzed for the striatal expression of TH upon completion of behavioral testing. Mice with 75\% or more sparing of TH-positive innervation in the striatum (as compared to the mean value of the respective vehicle group, WT-vehicle or CHT-vehicle) were eliminated from final analysis. StatView software (SAS Institute, Cary, NC) was used for statistical analysis. Most statistical analyses were performed using two-way ANOVA’s with Treatment (vehicle, 6-OHDA) and Genotype (WT, CHT\textsuperscript{HET}) as main factors. Timelines for changes to TH, DA, DOPAC, 3-MT, HVA, norepinephrine, and serotonin were analyzed using a three-way ANOVA with the additional independent variable of Duration of recovery (7, 14, 21, 28 days). Two behavioral tests were analyzed using repeated measure two-way ANOVA with Treatment and Genotype as between group factors. The repeated measure was Day of testing for rotarod analysis and Session number for analysis of the set-shifting task. Effects of physostigmine on performance of the recognition task were analyzed
using a one-way ANOVA with full treatment (WT-vehicle-saline, CHT$^{\text{HET}}$-6OHDA-saline, CHT$^{\text{HET}}$-6OHDA-0.05, CHT$^{\text{HET}}$-6OHDA-0.1) as the independent variable. When relevant, post-hoc analysis was performed using the Games-Howell test (Games and Howell, 1976). For all tests, the value of $p < 0.05$ was considered significant.

**RESULTS**

**Characterization of dopaminergic and cholinergic depletion**

The parameters of the 6-OHDA lesion were optimized to achieve partial DA depletion centered on the motor region of the caudate-putamen. This lesion paradigm was chosen because deep bilateral DA depletion compromises mouse viability and may result in unwanted motor defects. Intra-striatal infusions of 6-OHDA led to a partial loss of dopaminergic fibers in the striatum, with apparent sparing of dopaminergic neurons in the substantia nigra (Fig. 1A). The lesion was similar in WT and CHT$^{\text{HET}}$ mice and remained stable for 4 weeks following surgery as evidenced by the level of TH (Fig. 1B,D). The amount of TH was reduced by 6-OHDA treatment [$F(1,27) = 157.2, p < 0.0001$]. Similar declines were found in WT (~58%) and CHT$^{\text{HET}}$ (~54%) mice (no Genotype effect; $F[1,27] = 0.03, p =0.87$) and remained constant across 7, 14, 21 and 28 days following surgery ($F[3,27] = 1.14, p=0.35$).

DA was ~33% lower in 6-OHDA than vehicle-treated hemispheres (Fig. 1C; $F[1,40] = 55.9, p < 0.0001$), with no effect of Genotype and number of recovery days. The catecholamine metabolites DOPAC, HVA, and 3-MT were also significantly lower in the 6-OHDA-treated hemispheres ($p < 0.001$), with no effect of Genotype or duration of recovery (Fig. 1E). Norepinephrine (NE) content was also slightly, albeit significantly ($p<0.01$) decreased across
genotypes and recovery days, although no significant differences were seen in either genotype on individual days (Fig. 1F). Serotonin level (Fig. 1F) remained unaffected by the 6-OHDA lesion.

TH significantly declined in mice treated with 6-OHDA as compared to vehicle (Fig. 1B, D, C). Similar declines were found in WT (~58%) and CHT\textsuperscript{HET} (~54%) mice that remained constant across 4 weeks following surgery. The degree of TH loss was less than that seen in the brains of human PD and PDD patients at post-mortem (Bychkov et al., 2008), although postmortem material represents end-stage disease with likely more extensive loss of TH than in living PD patients. Taken together, the data show that the partial lesion was stable and reproduced the pattern of TH and DA loss seen in PD patients. However, the magnitude of the DA depletion in the mice underrepresented that seen in PDD patients.

As found previously (Bazalakova et al., 2007), ACh levels were 52% lower in striatal and cortical tissue in CHT\textsuperscript{HET} mice than in WT littermates (Fig. 2A, B; p<0.001). Intra-striatal infusions of 6-OHDA did not affect ACh levels regardless of Genotype (Fig 2 A, B). Striatal ACh levels in the CHT\textsuperscript{HET}-vehicle and CHT\textsuperscript{HET}-6OHDA groups were similar and significantly lower than in both vehicle and 6-OHDA-treated WT groups. The ratio of ACh to choline displayed the same pattern of effects as seen for ACh alone (Fig 2C, D). The ratio of ACh to choline in the frontal cortex and striatum was significantly lower in CHT\textsuperscript{HET} mice than WT, regardless of 6-OHDA treatment (p = 0.001 for the effect of Genotype in both regions). No genotype or treatment differences were found for the ACh precursor, choline, in either brain region (Fig 2E, F).

**Behavioral Evaluation**

**Rotarod.** To ascertain whether CHT\textsuperscript{HET} with loss of DA show motor deficits, we compared the performance of all groups on rotarod (WT-vehicle n = 7, WT-6OHDA n = 6,
As previously shown (Bazalakova et al., 2007), we observed no main effects of Genotype or 6-OHDA Treatment. There was a significant improvement across the three days of testing \[F(2,50) = 29.26, p < 0.0001\] with mice remaining on the rotarod for a shorter period of time on day one compared to days two \(p < 0.01\) and three \(p < 0.01\). Latencies to fall from the rotarod were higher than previously published (Bazalakova et al., 2007), potentially due to the extensive handling received by mice during preceding behavioral testing. To verify proper rotarod procedures, naïve wild type and CHT\(^{HEt}\) mice were tested with no prior experimental procedures (e.g. surgery, handling). Latency to fall in naïve mice was similar to previously published reports \(n = 6; \text{average seconds} \pm \text{S.E.M}: \text{day 1} = 123.5 \pm 14.39, \text{day 2} = 211.17 \pm 15.45, \text{day 3} = 263.5 \pm 29.07\). The results demonstrate intact motor performance in mice with loss of ACh, DA or combined ACh-DA depletion, which is the prerequisite for cognitive assessments of these animals.

**Set-Shifting Task.** The impairment of executive functions is believed to be the core component of cognitive impairment in PDD (Emre, 2004; Robottom and Weiner, 2009). We evaluated the ability of DA- and ACh-depleted mice to form and shift attentional sets. WT-6OHDA mice showed 49±10.8\% of TH relative to vehicle control and CHT\(^{HEt}\)-6OHDA showed 56.6±9.3\% relative to CHT\(^{HEt}\)-vehicle group (mean±S.E.M.). Figure 4A schematically shows the experimental chamber and setup, and Table 1 lists the stimuli used. Two-way ANOVA analysis of SD, CD, and IDS I through III individually showed no significant effects of Genotype (Fig. 4B), Treatment or Genotype x Treatment interaction. During the reversal session, all groups showed markedly worsened performance as compared to IDS III \[F(1,28)=50.9, p<0.0001, \text{across genotypes and treatments by three-way repeated measure ANOVA}\] with no group differences, demonstrating that all groups were capable of forming affective sets. In IDS
IV that followed the reversal session, both 6-OHDA-treated groups performed significantly, albeit slightly, worse than vehicle-treated groups [F(1,28)=5.1, p=0.032 for Treatment] (Fig. 4B).

The experimental groups demonstrated different dynamics of performance across sessions. The analysis of trials to criterion across all IDS for each experimental group by repeated-measure ANOVA with Session as within group factor yielded significant effects of Session for all groups (p=0.019, 0.03, 0.017, for WT-vehicle, WT-6OHDA, and CHT\textsuperscript{HET}-vehicle, respectively) except CHT\textsuperscript{HET}-6OHDA (p=0.25). These data suggest that there was a distinct behavioral pattern from session to session in all groups except CHT\textsuperscript{HET}-6OHDA. A more detailed analysis of the Session effects revealed that prior to the reversal, WT-vehicle mice, but no other group, needed progressively fewer trials in IDS I-III to reach criterion (p = 0.01) indicating consistently improved performance across sessions. After the reversal, in IDS IV, both vehicle-treated groups performed significantly worse than in IDS III preceding the reversal (p<0.05). WT-vehicle significantly improved performance in subsequent IDS as compared to the reversal (p<0.05). WT-vehicle was the only group to show signs of improvements immediately after the reversal in IDS IV (p=0.09 Reversal-IDS IV). Other experimental groups in IDS IV performed similarly to that in the reversal (p>0.5). WT-6OHDA and CHT\textsuperscript{HET}-vehicle groups (p<0.05) improved from the reversal through IDS VI as well as from IDS IV through IDS VI (p<0.01 and 0.05, respectively). CHT\textsuperscript{HET}-6OHDA did not differ in performance across IDS IV-VI or in comparison to the reversal (p>0.5). Thus, CHT\textsuperscript{HET}-6OHDA was the only experimental group that failed to improve across a series of training sessions.

In EDS, there were several effects. Genotype [Fig. 4B; F(1,26) = 17.1, p < 0.0001] and Treatment [F(1,26) = 7.44, p = 0.01] significantly affected performance. The WT-vehicle group required a greater number of trials to reach the criterion as compared to the CHT\textsuperscript{HET}-vehicle and
CHT^{HET}-6OHDA (p < 0.01) groups. Furthermore, the CHT^{HET}-6OHDA was the only group that committed significantly fewer errors in EDS (p < 0.05) and showed significantly lower EDS/IDS VI trial-to-criterion ratio than WT control (Fig. 4C). These data are consistent with the failure of CHT^{HET}-6OHDA mice to improve performance across training sessions and together suggest a failure to form an attentional set.

The WT-vehicle group required approximately twice as many trials to reach criterion in the EDS as compared to the preceding IDS VI session (p<0.001 according to repeated measure ANOVA). Both WT-6OHDA and CHT^{HET}-vehicle mice demonstrated impaired performance in EDS as compared to IDS VI requiring roughly 50% more trials to criterion (p<0.05). CHT^{HET}-6OHDA showed no relative increase in trials to criterion between IDS VI and EDS, and post hoc analysis showed CHT^{HET}-6OHDA performance as significantly different from all other groups (Fig. 4D). The ratio of trials to criterion during the EDS versus IDS VI was not affected by the direction of shift (i.e. odor to medium or medium to odor). The data suggest that mice with combined DA/ACH depletion were unable to form an attentional set.

**Spatial and Object Recognition.** Memory is another cognitive domain affected in PDD. To assess this domain, we used object and spatial recognition tests (Fig. 5A). Following a 5-minute retention delay, recognition scores in spatial or object test were not affected by Genotype or Treatment. These data indicate that dual DA/ACh depletion did not impair performance in relatively simple short-term visual-spatial memory tasks.

A more difficult version of the same tasks, with a 24-hour delay, revealed memory impairments in both the spatial and object tests. On the spatial memory test, a main effect of Genotype (Fig. 5C; p = 0.02) indicated impaired performance in CHT^{HET} mice, either vehicle- or 6-OHDA-treated, as compared to WT groups, but post hoc comparison revealed no significant
group differences. The performance in the object memory test was significantly affected by Genotype [F(1,36)=4.25, p=0.046], with the effect of Treatment and Genotype x Treatment interaction approaching significance (p=0.055 and 0.067, respectively). CHT$^{\text{HET}}$-6OHDA group was the only one that performed significantly worse than WT-vehicle (Fig. 5C). Both 6OHDA-treated groups had similar levels of TH depletion: WT-6OHDA mice showed 45.8±5.2% level of TH relative to the vehicle control and CHT$^{\text{HET}}$-6OHDA - 53.2±6.2% relative to CHT$^{\text{HET}}$-vehicle group (mean±S.E.M.).

A number of confounding factors might have affected the mouse performance in the recognitions tasks. One obvious possibility would be simply an overall lower activity or reduced motivation for novelty of CHT$^{\text{HET}}$-6OHDA mice, resulting in shorter exploration time. This explanation is refuted by the fact that there were no group differences in total distance traveled or total exploration time of all objects during any of the session of the recognition protocol (Fig 6A,B). Our finding that a deficit is revealed following a 24-hour but not 5-minute delay suggests that lower exploration of the novel object in CHT$^{\text{HET}}$-6OHDA results from a memory problem rather than a defect in motivation.

A deficit in 24-hour delayed object recognition unique to CHT$^{\text{HET}}$-6OHDA mice presented an opportunity to test whether increasing ACh via inhibition of AChE in the brain would rescue the phenotype. The previously found impaired spatial memory impairment was not replicated (Fig. 5D; p > 0.7). However, significant group differences were found, again, on the object memory test [Fig. 5D; F(3,36) = 4.99, p < 0.01]. The performance was impaired in the CHT$^{\text{HET}}$-6OHDA mice compared to WT-vehicle (p < 0.01), both given peripheral saline-control. The object recognition deficit in CHT$^{\text{HET}}$-6OHDA mice was reversed by systemic administration of physostigmine at 0.05 mg/kg (p < 0.02) and 0.1 mg/kg (p < 0.01). Physostigmine-treated
CHT^{HET}-6OHDA mice demonstrated the same level of novel object exploration as WT-vehicle mice (Fig. 5D). CHT^{HET}-6OHDA mice treated with another AChE inhibitor, donepezil, were not significantly different in the object recognition performance from WT-vehicle mice (Fig. 5D). Physostigmine or donepezil treatment did not affect locomotion or exploration behavior, since there were no group differences in distance traveled (Fig. 7A) or total exploration time (Fig. 7B). CHT^{HET}-6OHDA treated with vehicle demonstrated the TH level of 49.4±6.0% (mean±S.E.M.) of the respective vehicle control. Drug-treated groups were also similar: 0.5 mg/kg physostigmine showed 35.6±5.6% relative to CHT^{HET}-vehicle group, 1 mg/kg showed 36.5±4.3%, 0.1 mg/kg dopenesil showed 34.3±3.1% and 0.3 mg/kg showed 45.4±6.8% of the CHT^{HET}-vehicle control.

DISCUSSION

To define the role of dopaminergic and cholinergic deficits in the pathogenesis of dementia in PDD, we studied the effect of DA/ ACh depletion on cognition in a mouse model. To that end, we used the CHT^{HET} mouse, with a global ~50% loss of ACh, that underwent bilateral 6-OHDA lesion into the motor caudate-putamen to achieve partial, bilateral striatal DA depletion. The procedure resulted in a stable depletion of both neurotransmitters (ACh and DA) in the mouse brain without causing gross motor deficits. Therefore, the double-depleted mouse is a suitable model to investigate the impact of the combined loss of DA and ACh on specific cognitive functions affected in PDD.

Executive functions are one of the cognitive domains most impaired in PDD. To evaluate executive functions, we tested mice on the set-shifting task. The test is highly relevant because it shares important features with a test commonly used in human studies of executive
functioning, the Wisconsin Card Sorting Test. Non-demented PD patients display selective deficits is set-shifting performance (Cools et al., 2001; Monchi et al., 2004), possibly due to a combination of nigrostriatal and mesocortical DA deficiency. Patients with PDD demonstrate perseveration and difficulty acquiring the task (Beatty and Monson, 1990; Beatty et al., 1989; Litvan et al., 1991; Tröster et al., 1998). The set-shifting task depends on the functioning of the prefrontal cortex (Birrell and Brown, 2000; Bissonette et al., 2008; McAlonan and Brown, 2003). Dysfunction of the cholinergic system may also play a role in the pathogenesis of executive disturbances (Chen et al., 2004; Tzavos et al., 2004).

Set formation is evidenced by poorer performance during EDS trial as compared to preceding IDS in normal humans, primates and rodents (Birrell and Brown, 2000; Bissonette et al.; Dias et al., 1996; Owen et al., 1993a). Since each IDS used novel odors and medias with the rewarded dimension remaining constant, mice had the opportunity to learn that one dimension consistently led to reward while the other dimension did not (i.e. to form an attention set). By assigning the previously correct dimension as incorrect, EDS requires an extra effort, in a process known as set-shifting, to abandon an attentional set learned in preceding IDS sessions and to learn the new rules. EDS performance is affected in the opposite directions by both set formation during IDS and set-shifting during EDS. Mice, in comparison to rats, have been reported to have difficulty in forming attentional sets (Bissonette et al., 2008; Garner et al., 2006). In our behavioral paradigm, WT-vehicle mice displayed a strong impairment in EDS, indicative of a successful formation of attentive set. Mice with DA or ACh depletion demonstrated significant, albeit less pronounced, deficit in EDS performance. Mice with double depletion showed no deficit in the EDS performance thus presenting no evidence of attention set formation. An alternative interpretation is that better EDS learning reflects greater cognitive flexibility allowing for faster
set-shifting, but this explanation is unlikely. Animals that learn an attentional set would be expected to improve performance across IDS sessions and show poor performance on the EDS, as was seen in WT-vehicle and, to some extent, in WT-6OHDA and CHT$^{\text{HET}}$-vehicle mice. In contrast, CHT$^{\text{HET}}$-6OHDA mice did not benefit from the repetition of IDS sessions, learning at the same rate in each subsequent IDS session. This further argues that DA-depleted CHT$^{\text{HET}}$ animals approached each IDS trial as a separate event to which no previous experience applied. Our data supports the idea that double DA/ACh depletion is detrimental for the ability to form attentional set. Previous data indicated that the loss of NE affected set-shifting performance in rats (McGaughy et al., 2008; Tait et al., 2007). However, the degree of the NE depletion (70%) far exceeded the minimal loss detected in this study. Furthermore, the NE depletion impaired set-shifting, but not set formation. Therefore, it is unlikely that NE contributed to the deficit in set-shifting performance in CHT$^{\text{HET}}$-6OHDA mice. Loss of ACh alone induced by the lesion of the basal forebrain did not alter the set-shifting performance in rats (McGaughy et al., 2008; Tait and Brown, 2008), which underscores a unique phenotype in the set-shifting paradigm detected in double DA/ACh depleted mice. Thus, the evidence suggests that loss of DA in the cortico-striatal circuit associated with PD and further loss of ACh in a proportion of PD patients might play an essential role in the pathogenesis of executive deficits in PDD.

Studies have demonstrated memory deficits in PDD (Higginson et al., 2005; Kuzis et al., 1999; Noe et al., 2004; Owen et al., 1993b; Varanese et al., 2010). The impairment is most evident on visual-spatial memory tasks (Noe et al., 2004; Owen et al., 1993b). A rapid decline in visual-spatial performance is characteristic for PDD patients, compared with non-demented PD (Johnson and Galvin, 2011; Mosimann et al., 2004) and may be a contributing factor in memory deficits. Spatial and object recognition is a classic test used to evaluate visuo-spatial memory in
animals. We have employed the test to examine the effect of double DA/ ACh depletion in mice on their ability to recognize a new object or a new position of familiar objects. In object recognition with 24-hour delay, the CHT\textsuperscript{HET}-6OHDA was the only group that showed a deficit in performance. This result differs from data reported with mice hemizygous for vesicular acetylcholine transporter that were impaired on object recognition with both short and long delays (Prado et al., 2006). The reason for this discrepancy is unclear, although differences in the method might have played a role. Importantly, our approach allowed for the selective detection of a deficit due to combined DA/ ACh loss. The fact that in our model deficits in object recognition were unique to the double-depleted group, whereas both single DA- and ACh-depleted groups showed intact performance, suggests that combined DA/ ACh depletion is required to induce cognitive deficits. It is possible that more profound ACh or DA depletion alone would lead to similar cognitive impairment. For example, Wisman et al. (2008) (Wisman et al., 2008) showed that double DA/ ACh depletion with 6-OHDA and 192 IgG-saporin led to severe impairments in working and reference memory in Morris water maze, with loss of DA alone causing similar cognitive impairments only mildly exacerbated with additional ACh lesioning. However, at the level of depletion in our model, which importantly does not cause gross motor or cognitive deficits, only double DA/ ACh loss resulted in impairment in cognitive domains affected in PDD. Interestingly, the deficit in object recognition seen in DA-depleted CHT\textsuperscript{HET} mice was completely reversible by inhibition of ACh breakdown suggesting that an increase in available ACh is beneficial for cognition even when the deficit is not caused exclusively by the loss of ACh.

Although PD has traditionally been viewed as a motor disorder, there is evidence that PD patients from early stages show cognitive and memory deficits collectively referred to as mild
cognitive impairment (Aarsland et al., 2011; Johnson and Galvin, 2011; Sawamoto et al., 2008).
Cognitive problems associated with mild cognitive impairment, although quite heterogeneous,
embrace deficits in executive functions, flexibility, planning and working memory and are
believed to be driven by subcortical DA deficiency (Aarsland et al., 2011; Kehagia et al., 2010b).

In nonhuman primates (Decamp et al., 2004; Lipina and Colombo, 2007), bilateral DA depletion
impaired performance on a variety of cognitive tasks. Mice with bilateral striatal 6-OHDA
lesions displayed deficiency in spatial recognition with short, but not long, delay (De Leonibus et
al., 2007). Bilateral lesioning of the substantia nigra caused defects in working, cued spatial and
long-term memory in rodents [(Ardayfio et al., 2008; Braga et al., 2005; Da Cunha et al., 2006;
Ferro et al., 2005; Hefco et al., 2003; Lindnera et al., 1999; Miyoshi et al., 2002; Tadaiesky et al.,
2008), see also (Dunnett and Lejos, 2010) and references therein]. In some cases, motor
alterations (Ardayfio et al., 2008; Ferro et al., 2005; Lindnera et al., 1999) have also been
described. Some studies found no cognitive detriments in specific memory tasks such as novel
object recognition or passive avoidance in bilaterally DA-depleted rodents (Branchi et al., 2008;
De Leonibus et al., 2007). Therefore, loss of DA seems an important contributing factor to
cognitive deficits in PDD but by itself may not result in cognitive impairment. The loss of ACh
alone may be sufficient to impair cognition as evidenced by detrimental effects of anticholinergic
drugs on cognitive performance both in humans and animals (Campbell et al., 2009; Klinkenberg
and Blokland, 2010). Obviously, the degree of the neurotransmitter loss is an important factor.
However, our data suggest that a combination of a moderate ACh deficit on the background of
modest DA depletion may produce a specific cognitive state not seen in either condition alone.

In spite of a high prevalence of dementia in PD (Aarsland and Kurz, 2010), its
pathophysiology remains obscure. In treatment, the PDD clinic appropriates strategies from the
Alzheimer’s disease field capitalizing on drugs that enhance ACh signaling, AChE inhibitors. AChE agents provide only modest improvement in cognitive performance [reviewed in (Aarsland et al., 2011; Kehagia et al., 2010a; Maidment et al., 2006; Williams-Gray et al., 2006)]. The mechanistic and translational studies of dementia in PDD lag far behind that of motor deficits in PD or dementia in Alzheimer’s disease. So far, the understanding of PDD pathophysiology has relied on human postmortem studies and imaging studies of PDD patients. Both of these approaches, although providing invaluable information, are not amenable to experimental manipulation and thus afford limited opportunities to probe for causal links or discover new therapeutic targets. Our studies with the DA-depleted CHT\textsuperscript{HET} mouse demonstrate that combined modest loss of DA and ACh, but not the depletion of each neurotransmitter alone, causes deficits in the cognitive domains specifically affected in PDD, executive functions and visual-spatial memory. Since this behavioral phenotype reminiscent of PDD has been reproduced in the DA-depleted CHT\textsuperscript{HET} mouse based on known neuropathological features of PDD, e.g. loss of DA in the basal ganglia combined with loss of ACh, our data suggest that both loss of DA and ACh contribute to dementia in PDD. The ability of AChE inhibitors, known to improve cognition in PDD, in reversing visual-spatial memory deficits in DA-depleted CHT\textsuperscript{HET} mice serves as further confirmation of essential similarities between the neuropathology in the DA-depleted CHT\textsuperscript{HET} mouse and PDD.

It is clear that the neuropathology of the DA-depleted CHT\textsuperscript{HET} mouse does not cover the whole spectrum of the PDD neuropathology. Post-mortem analysis of brain tissue has revealed several neuropathological features of PDD brains. The cortical Lewy bodies (Halliday et al., 2008; Mattila et al., 2000) and Alzheimer’s-related pathologies (Aho et al., 2008; Jellinger, 2009; Jellinger et al., 2002) seen post-mortem frequently correlate to ante-mortem cognition, although
recent imaging studies failed to detect high amyloid load in PDD patients (Edison et al., 2008). The role of Lewy bodies and Alzheimer’s type pathologies in PDD is a hotly debated area, with some authors favoring the accumulation of Lewy bodies and others favoring Alzheimer’s type pathologies as the prime contributor to the PDD pathogenesis (Aarsland et al., 2005; Hurtig et al., 2000; Jellinger, 2009). Importantly, ACh depletion is the most consistently found neural difference between non-demented PD and PDD and consistently correlates with cognitive score (Bohnen et al., 2006; Choi et al., 2012; Kuhl et al., 1996; Perry et al., 1985; Perry et al., 1987). Our studies in the CHT\^HET\textsuperscript{-}6OHDA mouse suggest that downstream consequences of combined DA-ACh depletion may be sufficient to bring about essential features of dementia in PDD, which is important for the evaluation of treatments and interventions for therapeutic potential in patients.

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The authors report no biomedical financial interests or potential conflict of interests.

AUTHOR CONTRIBUTIONS

E.V.G. and L.Z. conceived and designed the experiments. L.Z., E.B., E.L.T. and C.S. performed the experiments. L.Z., C.S. and E.V.G. analyzed the data. R.D.B. contributed CHT mice. L.Z., E.V.G. and R.D.B. wrote the paper.

TRANSLATIONAL IMPACT

Clinical issue. Cognitive deficits and dementia are prevalent in Parkinson's disease (PD), although the disease has been traditionally viewed as a motor disorder. Dementia has a strong impact on the quality of life of PD patients and even on their life expectancy. Currently, dementia in PD is recognized as one of the most pressing and challenging problems in PD clinical management. However, our understanding of the pathophysiology of dementia in PD and the ability to treat the condition in patients lags behind that for motor symptoms. The present work translates the findings from clinical studies in human patients into mechanistic animal experiments. To model the dementia in PD in non-human animals, factors known to contribute to the pathophysiology of dementia in PD, such as loss of subcortical dopamine (DA) and loss of acetylcholine (ACh), were integrated. Results. We found that mice with combined depletion of DA and ACh demonstrate cognitive abnormalities that are specific for PD-associated dementia such as diminished executive functions and impaired visual-spatial memory. Importantly, mice that only have the loss of one brain neurotransmitter, DA or ACh, did not
show these defects. Furthermore, visual-spatial memory deficit in these mice was reversible by acetylcholinesterase inhibitors physostigmine and dopenesil, the only drugs currently available to manage dementia in PD. **Implications and Future Directions.** The study provides first mechanistic support for the loss of ACh and DA as possible causal factors in PD-associated dementia as opposed to previous research that only established correlative association. Furthermore, the study draws the attention to the importance of combined neurotransmitter deficits as opposed to the loss of each neurotransmitter alone in PD dementia. The work provides the foundation for further mechanistic studies of PD dementia utilizing the double-depleted mouse.
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Figure 1: Partial dopaminergic lesion in the striatum was stable and independent of genotype. Mice received a unilateral partial 6-OHDA lesion (control hemisphere was injected...
with vehicle) as described in Methods and Materials. (A) Low power photomicrograph of representative striatal (upper panel) and nigral (lower panel) sections immunostained for TH. Photographs illustrate partial depletion of TH-positive fibers in the dorso-lateral caudato-putamen (white arrow) in the hemispheres injected intra-striatally with 6-OHDA as compared with vehicle. Note minimal loss of dopaminergic cells in the substantia nigra. (B) Representative Western blot for TH showing reduced amount of TH in the hemisphere lesioned with 6-OHDA. Standards correspond to the amount of TH found in indicated amount of the total striatal proteins from the rat brain. DA measured by HPLC (C) and TH measured by Western blot (D) showed significant declines in 6-OHDA groups. Data (mean±S.E.M.) are shown as ng/mg protein for DA (C) and arbitrary units for TH (C). * - p <0.05, ** - p<0.01 to corresponding vehicle group according to one-way repeated measure ANOVA with Treatment as a repeated measure factor performed separately for each genotype and each post-lesion day. (E) The DA metabolites DOPAC (left), HVA (middle), and 3-MT (right) (expressed as the percentage of vehicle-treated hemispheres) were also reduced following 6-OHDA treatment. * - p <0.05, ** - p<0.01 to corresponding vehicle group according to one-way repeated measure ANOVA with Treatment as a repeated measure factor performed separately for each genotype and each post-lesion day. (F) Norepinephrine (left) and serotonin (right) in the 6-OHDA-lesioned hemispheres expressed as percents of values in the vehicle-treated hemispheres.
Figure 2: The level of acetylcholine was not altered by dopaminergic lesion in either genotype. (A, B) ACh, measured by HPCL, in sham-operated or 6-OHDA-treated WT or CHT^{HET} in the prefrontal cortex (A) or striatum (B). Tissue was collected 7 days following surgery. * - p < 0.001 to corresponding vehicle group across genotypes. (C, D) The ratio of ACh to choline in the frontal cortex (C) and striatum (D). * - p = 0.001 across genotypes. (E, F) The cortical (E) and striatal (F) levels of choline were not affected by 6-OHDA or Genotype.
Figure 3: The dopaminergic lesion or genotype did not compromise rotarod behavior. Rotarod performance was measured by latency to fall, in seconds. * indicates significant difference in performance between Day 1 and Day 2 and between Day 1 and Day 3 across genotypes and treatments at p < 0.01. # indicates a significant difference between WT-vehicle and WT-6OHDA on day 2.

Figure 4: Double depletion of acetylcholine and dopamine impaired attentional set formation. (A) Schematic representation of the training apparatus. Food cups filled with media...
(light or dark fill) scented with two different odors (black or white circles) were placed in choice compartments, one hiding a food reward (star). Trials were initiated by removing a PVC door, symbolized by the dashed line. (B) Set-shifting training procedure by sessions shown as the number of trials to reach learning criterion (8 consecutively correct responses) for each session (means±S.E.M.). ^ - p < 0.05, ^^ - p<0.01 according to repeated measure ANOVA with Session as a repeated measure factor applied separately for each genotype-treatment combination across sessions indicated by the brackets. a - p <0.05, b - p<0.01, c - p<0.001 by repeated measure ANOVA with Session as factor separate for each group. ). * - p < 0.05, ** - p<0.01 as compared to WT-vehicle according to Gamed-Howell post hoc test following one-way ANOVA with Group (combinations of genotype-treatment) as the main factor. (C) The number of errors committed during the EDS trial (means±S.E.M.). (D) The ratio of trials to criterion against the immediately preceding IDS session. * - p < 0.05, ** - p<0.01 as compared to WT-vehicle according to Gamed-Howell post hoc test following one-way ANOVA with Group (combinations of genotype-treatment) as the main factor.
Figure 5: Mice with double depletion of acetylcholine and dopamine demonstrate a deficit in delayed object recognition. (A) Schematic representation of the spatial and object recognition task. An acclimation session with no objects was followed by three sessions with five objects in a specific configuration. Memory was challenged by a 5-minute or 24-hour retention delay. Two sessions with an altered spatial configuration (two object misplaces,
circled) were given, the first being quantified for spatial recognition memory. One session with a novel object (circled) was scored for object recognition memory. The arrow points to the time of administration of physostigmine for the experiment depicted in panel D. (B) Spatial and object recognition performance with a retention delay of 5 minutes. (C) Spatial and object recognition performance with a retention delay of 24 hours. A modest impairment in performance in the spatial memory test was seen in CHT\textsuperscript{HET} mice regardless of the treatment (p<0.05). * - p<0.05 to all other groups according to Game-Howell post hoc test following one-way ANOVA with Group (combinations of genotype-treatment) as main factor (D) Physostigmine and donepezil had no effect on the spatial recognition test. Both doses of physostigmine reversed impaired object recognition memory in CHT\textsuperscript{HET}-6OHDA mice. Similarly, both dozes of donepezil reversed the object recognition deficit in CHT\textsuperscript{HET}-6OHDA mice * - p < 0.01 to WT-vehicle-saline group; # - p < 0.05, ## - p<0.01 to CHT\textsuperscript{HET}-6OHDA-saline group.
Figure 6: No differences were detected in total exploration and distance traveled during the delayed visual-spatial recognition task. (A) Total distance traveled during visual-spatial recognition task with 24 h delay in all sessions including acclimation, in which no objects were present, is expressed in centimeters (cm) (mean±S.E.M). Significance levels are given for the effect of Session across all sessions based on one-way repeated measure ANOVA with Session as repeated measure factor and apply to all experimental groups (brackets). (B) Total exploration time of the five objects available during each session, is expressed as total time in seconds (mean±S.E.M). There was an overall reduction across groups in the total exploration time across the acclimation through Acquisition III session (repeated measure ANOVA with Session as the repeated measure factor, p=0.0002), but individually only the CHT^{HET}-vehicle...
group showed a significant decrease (p<0.05). Similarly, the total exploration time significantly decreased across the spatial recognition through the object recognition session (p=0.0004), with the WT-6OHDA group showing a significant effect (p<0.001).

Figure 7: Treatment with AChE inhibitors did not affect total exploration and distance traveled during the visual-spatial recognition task. (A) Total distance traveled is expressed in centimeters (cm) (m±S.E.M.) during each session of training, including acclimation, in which no objects were present. Significance levels are given for the effect of Session across all sessions based on one-way repeated measure ANOVA with Session as the repeated measure factor and apply to experimental groups under the brackets. (B) Total time, in seconds, spent in exploration of all five objects present during a given session. Significance levels are given for the effect of
Session across all sessions based on one-way repeated measure ANOVA with Session as the repeated measure and apply to experimental groups under brackets.

| Table 1. Summary of the set-shifting experiment, with specific odors and media used¹. |
|----------------------------------|----------------|----------------|----------------|----------------|
|        | Odor 1     | Odor 2     | Medium 1     | Medium 2     |
| SD     | Cloves*    | Basil      | Aspen Bedding | n/a           |
| CD     | Cloves*    | Basil      | Aspen Bedding | Sponge       |
| IDS I  | Thyme*     | Cumin      | Yarn          | Plastic pellets |
| IDS II | Oregano*   | Nutmeg     | Paper clips   | Cotton T-shirt |
| IDS III| Sage*      | Black pepper | Bed fluff | Bermuda grass |
| Reversal| Sage      | Black pepper* | Bed fluff* | Bermuda grass |
| IDS IV | Cinnamon*  | Turmeric   | Eco-bedding  | Moss         |
| IDS V  | Garlic*    | Ginger     | Seashells    | Felt fabric  |
| IDS VI | Rosemary*  | Chili powder | Nitrile | Aluminum foil |
| EDS    | Orange     | Onion      | Styrofoam*   | Shredded newspaper |

SD – simple discrimination; CD – compound discrimination; IDS – intra-dimensional shift; EDS – extra-dimensional shift; stimuli indicating reward are bold underlined with asterisk.

¹- The exact experimental setup shown was used for odor rewarded (in IDS sessions) animals. For medium rewarded mice, the same exemplars were used, only one of the media was indicative of the reward (in the SD session, both media were present but only one of the odors). The direction of extra-dimensional shift (odor to medium or medium to odor) was counterbalanced within treatment groups.