Neuro-Cognitive Effects of Acute Tyrosine Administration on Reactive and Proactive Response Inhibition in Healthy Older Adults

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Abstract

The aging brain is characterized by altered dopamine signaling. The amino acid tyrosine, a catecholamine precursor, is known to improve cognitive performance in young adults, especially during high environmental demands. Tyrosine administration might also affect catecholamine transmission in the aging brain, thereby improving cognitive functioning. In healthy older adults, impairments have been demonstrated in two forms of response inhibition: reactive inhibition (outright stopping) and proactive inhibition (anticipatory response slowing) under high information load. However, no study has directly compared the effects of a catecholamine precursor on reactive and load-dependent proactive inhibition. In this study we explored the effects of tyrosine on reactive and proactive response inhibition and signal in dopaminergically innervated fronto-striatal regions. Depending on age, tyrosine might lead to beneficial or detrimental neurocognitive effects. We aimed to address these hypotheses in 24 healthy older human adults (aged 61–72 years) using fMRI in a double blind, counterbalanced, placebo-controlled, within-subject design. Across the group, tyrosine did not alter reactive or proactive inhibition behaviorally but did increase fronto-parietal proactive inhibition-related activation. When taking age into account, tyrosine affected proactive inhibition both behaviorally and neurally. Specifically, increasing age was associated with a greater detrimental effect of tyrosine compared with placebo on proactive slowing. Moreover, with increasing age, tyrosine decreased fronto-striatal and parietal proactive signal, which correlated positively with tyrosine's effects on proactive slowing. Concluding, tyrosine negatively affected proactive response slowing and associated fronto-striatal activation in an age-dependent manner, highlighting the importance of catecholamines, perhaps particularly dopamine, for proactive response inhibition in older adults.

Key words: dopamine; functional MRI; healthy aging; response inhibition

Significance Statement

Healthy aging comes with altered dopamine functioning and is associated with reduced performance on cognitive control tasks, such as response inhibition. However, it is yet unclear whether reactive or proactive response inhibition is modulated by dopamine. We addressed this question by administering the catecholamine precursor tyrosine in a double blind, placebo-controlled, randomized intervention study. Tyrosine decreased proactive response slowing, not reactive stopping, as a function of increasing age. Concurrently, proactive fronto-striatal and parietal blood oxygen level-dependent (BOLD) signal decreased after tyrosine with increasing age. These findings, especially in striatum, demonstrate that proactive, rather than reactive response inhibition, is dopamine dependent. Moreover, tyrosine’s effect on brain and cognition became detrimental with increasing age, questioning the cognitive enhancing potential of tyrosine in healthy aging.
Introduction

The aging brain is characterized by alterations in dopamine functioning (Kaasinen and Rinne, 2002; Braskie et al., 2008). Age-related decreases in dopamine receptor and transporter binding have been linked to impairments in cognitive functions such as attention, episodic and working memory (Bäckman et al., 2006) and age-related increases in dopamine synthesis capacity have been related to decreased neural reward processing (Dreher et al., 2008). In aged experimental animals, administration of a D1 receptor agonist improved working memory performance (Cai and Arnsten, 1997). Similarly, enhancing dopamine levels in humans with the drug L-Dopa (the direct dopamine precursor) improved age-related impairments in episodic memory performance and reinforcement learning (Chowdhury et al., 2012, 2013).

Tyrosine is a large non-essential neutral amino acid (LNAA), naturally present in food. Tyrosine is the precursor of the catecholamines, converted to dopamine via L-Dopa and the enzymes tyrosine hydroxylase (TH) and aromatic l-amino acid decarboxylase and to noradrenaline by dopamine β-hydroxylase (Molinoff and Axelrod, 1971). Research in rodents showed that orally administered tyrosine reaches the brain (Glaeser et al., 1983). Tyrosine administration increases dopamine metabolites in CSF, like homovanillic acid (HVA), in rats (Scally et al., 1977) and in patients with Parkinson’s disease (Growdon et al., 1982). In young adults, tyrosine administration improved cognitive control functions such as response inhibition, task switching, and working memory, especially in demanding circumstances (for review, see Deijen, 2005; Jongkees et al., 2015). In the aging brain, tyrosine may similarly improve cognitive functioning.

Aging is accompanied by deficits in inhibitory functions, both in terms of the inhibition of irrelevant information, e.g., sensory suppression (Gazzaley et al., 2008; Healey et al., 2008), as well as in terms of response inhibition, such as in stop-signal tasks (Kramer et al., 1994; Bedard et al., 2002; van de Laar et al., 2011). Two forms of response inhibition have been distinguished: reactive response inhibition is the process of canceling an ongoing response at the moment this is needed (i.e., outright stopping), whereas proactive response inhibition entails the preparation for stopping when this may become necessary, e.g., based on cues held in working memory. Age-related impairments have been shown in both reactive inhibition (measured with stop-signal reaction time; SSRT) and proactive inhibition (measured with anticipatory response slowing), particularly under high information load (i.e., high information processing demands for interpreting the stop-signal probability cues; Bloemendaal et al., 2016; Kleerekoper et al., 2016). It is unclear whether tyrosine-induced modulation of catecholaminergic signaling in older adults will affect reactive and/or proactive response inhibition. For reactive response inhibition, pharmacological animal and human genetic work shows both dopaminergic as well as noradrenergic involvement (Eagle et al., 2007; Congdon et al., 2009; Ghahremani et al., 2012; Rae et al., 2016; Schippers et al., 2016; see also Eagle and Baunez, 2010). Catecholaminergic modulation of proactive response inhibition has never been formally tested, but experimental animal work implicates dopamine in a variety of processes contributing to proactive response inhibition (Bari et al., 2009; Bari and Robbins, 2013). Further indirect evidence for a role of dopamine in proactive inhibition comes from neuroimaging studies. Specifically, midbrain signal was associated with stop-signal probability and RT adjustments (Boehler et al., 2011; Zandbelt et al., 2013).

In this neuro-imaging study, we investigated effects of acute, oral tyrosine administration on reactive and (load-dependent) proactive response inhibition and associated signal in dopamine-innervated fronto-striatal regions of the aging brain. We used a dose of 150 mg/kg body weight, in accordance with most previous studies in young volunteers (Shurtleff et al., 1994; Neri et al., 1995; Thomas et al., 1999; Magill et al., 2003; Mahoney et al., 2007), but see work by Colzato and colleagues for beneficial effects on cognition in young volunteers with much smaller doses (Colzato et al., 2013, 2014a, 2014b; Steenbergen et al., 2015). By investigating older adults, we could also assess the potentially beneficial effects of tyrosine in aging. We expected to find beneficial effects of tyrosine on brain and behavior across the group of older adults. Given our recent findings of age-related differences in the peripheral plasma response to oral tyrosine administration (van de Rest et al., 2017) and given differences in the effect of dopaminergic agents between younger and older adults (Turner et al., 2003), we also explored the possibility that tyrosine’s effects would vary as a function of age. Using a smaller age range, we do not expect generational differences to influence the effects of dopaminergic agents, such as differences in education or computer experience that would differ in a cross-sectional design with larger age differences. The oldest relative to the youngest older adults are presumably most dopamine deprived and, thus, may benefit most from administration of dopamine’s precursor tyrosine. However, whereas aging has been associated with reduced dopamine receptor and transporter binding,
it has also been shown to be accompanied by upregulation of (dorsal) striatal dopamine synthesis capacity (Braskie et al., 2008; Berry et al., 2016). This upregulation of synthesis capacity has been related to, if anything, worse rather than better neurocognitive functioning relative to young adults (Dreher et al., 2008; Berry et al., 2016). Moreover, a recent study with multiple oral tyrosine doses (100, 150, and 200 mg/kg, but no placebo condition) showed decreased cognitive performance with increased tyrosine dose in older adults (van de Rest et al., 2017). Therefore, in the current placebo-controlled study, administration of a high dose (150 mg/kg) of the dopamine precursor tyrosine might also impair instead of improve neuro-cognitive function in the oldest adults, with presumably the greatest upregulated dopamine synthesis capacity.

**Materials and Methods**

**Participants**

Participants met the following criteria: aged between 60 and 75, right handed, functioning within normal limits of general cognitive function [mini-mental state examination (MMSE); Folstein et al., 1975; cutoff ≥27 of 30], no depression or anxiety [hospital anxiety and depression scale (HADS) score <11; Bjelland et al., 2002], an estimated verbal IQ > 85 (Schmand et al., 1991), not suffering from neurologic or psychiatric disorders, no first degree relatives suffering from schizophrenia, bipolar disorder, or major depressive disorder, no history of alcohol or drug abuse, no habitual smoking defined as less than a pack of cigarettes a week for the last year, current or past (within last 12 months) participation in a specific cognitive training program, no contraindications for MRI, no daily use of β blockers, no use of medication interfering with tyrosine’s action (monoamine oxidase inhibitors and other antidepressants, sympathomimetic amines, and opioids), no thyroid problems and no low-protein diet, no endocrine or metabolic disorders such as hepatic or renal problems, no repetitive strain injury (RSI) or sensorimotor handicaps, blindness, or colorblindness.

Participants were recruited via adverts in local newspapers, websites, and associations for older adults. After informing potential participants about the inclusion criteria, 45 older adults were invited for a pre-screen session. After the pre-screen, we invited 33 participants for the test sessions (Fig. 1). Of these 33, 29 participants completed two test sessions. Of the four participants who did not complete all test sessions, three participants were excluded during test day 1 (due to panic on entering the scanner, high blood pressure, or vomiting) and one after...
test day 1 (due to headache after the test day). Of the 29 participants who completed both test days, a final sample of 24 healthy older adults were included in the analyses (mean age: 67.5, range 61–72, 15 men). Of the five participants who were not included in the analyses, two participants did not finish the stop-task on one of the sessions, and three participants were excluded before statistical data analysis: two due to excessive movement (>4-mm translation) and one due to signal intensity spikes.

The experiment was approved by the local ethics committee (CMO 2014-1172), and all participants gave written informed consent. The study was preregistered at the Dutch trial register (www.trialregister.nl) 159 under number NTR4938.

**Intervention**

Our participants received a dosage of 150 mg/kg tyrosine or placebo, adjusted to body weight as determined during the pre-screen session (Procedure). The European Food Safety Authority determined in July 2011 that tyrosine is proven to contribute to the normal synthesis of catecholamines (EFSA Panel on Dietetic Products Nutrition and Allergies, 2011). In accordance with most previous studies in young volunteers (Shurtleff et al., 1994; Neri et al., 1995; Thomas et al., 1999; Magill et al., 2003; Mahoney et al., 2007; but see Colzato et al., 2013, 2014a, 2014b; Steenbergen et al., 2015), we used a dosage of 150-mg tyrosine or placebo per kilogram of body weight.

For reference, a daily required intake of phenylalanine and tyrosine mixture have an equal energy value, similar structure and aftertaste. Equal taste experience for tyrosine and placebo was ensured in a formal sensory experiment by a specialized dietician from the Division of Human Nutrition of Wageningen University (E.Siebelink).

The placebo product was a mixture of 54 mg/kg dextrose-maltose (i.e., carbohydrates; product name Fantomalt by Nutricia) with maizena (110 mg/kg, ratio Fantomalt/cornstarch = −1/2). The ratio of Fantomalt to cornstarch was adjusted such that the placebo and tyrosine mixture have an equal energy value, similar structure and aftertaste. Equal taste experience for tyrosine and placebo was ensured in a formal sensory experiment by a specialized dietician from the Division of Human Nutrition of Wageningen University (E.Siebelink).

The tyrosine and placebo product were mixed with a carrier: banana-flavored yoghurt (Arla Food Nederland). Weighing of the doses and preparing and coding the samples was performed by a staff member who was not further involved in the study.

**Physiologic and mood measurements**

To monitor wellbeing, participants completed mood ratings, assessing calmness, contentness, and alertness (Bond and Lader, 1974). Moreover, we assessed levels of the catecholamine metabolites [HVA, vanillylmandelic acid (VMA), 3-methoxy-4-hydroxyphenylglycol (MOPEG), and 3,4-dihydroxyphenylacetic acid (DOPAC) in urine to measure peripheral effects.

**Procedure**

All participants were tested between November 2014 and August 2015 at the Donders Center for Cognitive Neuroimaging, Nijmegen, The Netherlands. Participants were pre-screened in a separate 4-h session and trained on the tasks. During this pre-screen session, participants signed informed consent, were screened on all the inclusion and exclusion criteria, and completed several neuropsychological measures: verbal IQ as measured with the Dutch version of the NART, MMSE, BIS-11, subscales and total score. Men and women are equally distributed across the whole group.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Mean (SEM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (years)</td>
<td>67.5 (0.6)</td>
</tr>
<tr>
<td>Sex (women/men)</td>
<td>W: 9; M: 15</td>
</tr>
<tr>
<td>Verbal IQ</td>
<td>114.5 (2.0)</td>
</tr>
<tr>
<td>HADS total</td>
<td>3.5 (0.5)</td>
</tr>
<tr>
<td>HADS anxiety</td>
<td>2.3 (0.4)</td>
</tr>
<tr>
<td>HADS depression</td>
<td>1.2 (0.3)</td>
</tr>
<tr>
<td>MMSE</td>
<td>29.1 (0.3)</td>
</tr>
<tr>
<td>BIS-11 motor</td>
<td>20.9 (0.6)</td>
</tr>
<tr>
<td>BIS-11 cognitive</td>
<td>14.8 (0.6)</td>
</tr>
<tr>
<td>BIS-11 non-planning</td>
<td>21.7 (0.7)</td>
</tr>
<tr>
<td>BIS-11 total</td>
<td>57.4 (1.1)</td>
</tr>
</tbody>
</table>

Data represent mean (SEM) except for the variable sex, for which data reflect frequencies. Verbal IQ is defined by scores on the Dutch version of the NART, MMSE, BIS-11, subscales and total score. Men and women are equally distributed across the whole group.

On both test sessions (of at least one week apart), the same procedure was followed except for the supplement taken: tyrosine or placebo, counterbalanced across participants.

An independent researcher randomized the order of tyrosine administration (tyrosine or placebo on the first test session) by means of a computer-generated order. The day before the test session from 10PM onwards, participants were asked to refrain from eating and drinking anything but water until arriving at the center the next morning, and to refrain from taking any medication that they would not take during both testing sessions, to avoid an unbalanced influence of this medication. The overnight fast prevents large variability in plasma LNAA levels between participants caused by the previous meal (Fernstrom et al., 1979). A similar fasting procedure has been adopted in other research using tyrosine supplementation (Lieberman and Wurtman, 1985; Banderet and Lieberman, 1989; Shurtleff et al., 1994; Mahoney et al., 2007; Colzato et al., 2013). Testing started from 8 or 10 A.M. at the latest and took ~4.5 h (Fig. 2). The time of testing was kept similar for the placebo and tyrosine session of each participant (i.e., maximal difference between test sessions was 1 h, except for one participant for who the difference was 1.5 h).

The test day started with assessing subjective feelings of wellbeing measured with Bond & Lader visual-analog ratings. Blood pressure and heart rate were assessed, and a urine sample was provided. Next, participants started re-familiarization with all three tasks (always in the same order). The yoghurt mixture (with tyrosine or pla-
cebo) was provided such that participants entered the scanner 90 min after ingestion. Maximal concentration of plasma elevation and cognitive effects are seen approximately after 1.5 h and are normalized after 6–8 h (Glaeser et al., 1979). The stop-signal task (described below) was the first task performed in the scanner, followed by a working memory task. Scanning took ~100 min. On exiting the scanner, a second urine sample was provided. After scanning, participants performed the third task measured effort discounting and completed a neuropsychological test battery assessing: immediate and delayed story recall (Wilson et al., 1985), digit span forward and backward (Wechsler, 1997), Stroop cards (Stroop, 1935), and verbal fluency (Tombaugh et al., 1999). Participants’ blood pressure, heart rate, and wellbeing were monitored three times during the test session.

**Experimental design: load-dependent stop-signal anticipation task**

Participants performed a stop-signal anticipation task consisting of three levels differing in information load, which were presented in alternating blocks (Bloemendaal et al., 2016; Fig. 3). The paradigm consisted of Go trials and Stop trials. On every trial, a bar moved at a constant speed from a lower horizontal line toward an upper horizontal line, reaching a middle line (flanked by two vertical lines) in 800 ms. The Go task was to bring the bar to a halt as close to the middle line as possible, by pressing a button with the right index finger. A minority of trials were Stop trials. On Stop trials, the bar stopped moving automatically before reaching the middle line (the stop signal). This stop signal instructed the participants to withhold the planned Go response. The middle horizontal line and the two vertical lines represented cues that indicated stop-signal probability context by varying in color. To manipulate information load, the task consisted of three levels that were alternated in short blocks (Last alinea of Experimental design section). Between levels, stop-signal probability cues were varied in amount as well as in complexity. The stop-signal probability could be anticipated on the basis of the color of the cues (i.e., horizontal and vertical lines, presented 500 ms before the onset of the moving bar). Level A was the level with the least information load, with only white cues (stop probability of 26%) and green cues (stop probability of 0%). In level B, there were five types of Go trials with varying stop-signal probability, using an intuitive color range for the cues (Zandbelt and Vink, 2010): green, 0%; yellow, 17%; amber, 22%; orange, 28%; and red, 35%, with a mean of 26% stop probability. The non-green trials are collectively called 0% trials. Level C consisted of the same types and numbers of stop-signal probability cues as level B. However, in level C only one of the vertical lines signaled the correct stop-signal probability context. The correct
side could be identified by the color of the middle line: a blue middle line indicated that the left vertical line color was valid, whereas a purple middle line indicated that the right vertical line color was valid.

We instructed participants that going and stopping were equally important and that it would not always be possible to suppress a response when a stop signal occurred. Participants were not informed of the exact stop-signal probabilities but were told that stop signals in all levels would not occur on trials with a green cue, and that stop signals in levels B and C were least likely in the context of a yellow cue and most likely in the context of a red cue, with the amber and orange cues coding intermediate, and, respectively, decreasing, stop-signal probabilities.

To ensure roughly equal numbers of successful and unsuccessful Stop trials, a staircase procedure adjusted stop-signal delay by 25 ms depending on stopping performance. Levels were presented in 34 blocks, each lasting 27 s and consisting of 10 trials, with an intertrial interval of 1000 ms. The sequence of trials and blocks were pseudo randomized (ensuring that the first three blocks of the task were always in order of levels A-B-C). Every level contained 70 trials with 0% (green) and 270 trials with >0% (white in level A and various colors in levels B and C) stop-signal probability. Of these 270 >0% trials, 70 were Stop trials, with a mean stop-signal probability of 26%. For levels B and C, each >0% trial type contained 50 Go trials, plus a varying amount of Stop trials per color resulting in varying stop-signal probabilities (in between brackets): 10 yellow (17%), 14 amber (22%), 19 orange (28%), and 27 red (35%). Two rest blocks of 20 s each were implemented at one-third and two-thirds of the task, respectively. The total task duration was ~45 min. During the pre-screening, each level was explained and practiced separately for 48 trials (level A) and 72 trials (levels B and C). Participants were asked to repeat task instructions to ensure sufficient understanding. Then they practiced the task (levels were presented in alternating blocks) for 10 min. On each test day, instructions were rehearsed and the 10-min practice was repeated.

Behavioral data analysis
Stop-signal task
All data were in accordance with the main assumptions of the race model (Logan and Cowan, 1984). Reactive response inhibition was measured by calculating the SSRT (stopping latency), according to the integration method (Verbruggen and Logan, 2009). Outliers on any outcome measure were determined using Grubbs’ test (i.e., they do not differ >2.8×SD from the mean; Grubbs, 1969). An ANOVA of SSRTs was used with the within-subjects factors level (A, B, C) and intervention (tyrosine, placebo).

RT slowing as a function of increasing stop-signal probability, indicated by the colored cues, indexed proactive response inhibition. Task non-compliance was determined by a negative difference on median RTs between 0% (green) and >0% (white) trials during the lowest cognitive load (level A). No participants were excluded based on this criterion. For each level, the slope of RTs was calculated as a function of stop-signal probability using a general linear model, resulting in a β value for the slowing slope. Hence, for level A, the slowing slope was calculated using the two proactive trial types [0% (green cues) and 26% (white)]. For levels B and C, the five proactive trial types were included in the slowing slope [0% (green cues), 17% (yellow), 22% (orange), 28% (amber) and 35% (red)]. Task instructions implied differential processing of 0% and >0% stop-signal probability trials, resulting in more slowing on >0% than 0% trials (i.e., a positive difference between these trial types). Level A consists of fewer cells than levels B and C and was therefore not compared with the other levels for the proactive inhibition analyses. An ANOVA was performed using the within-subjects factors level (levels B and C) and intervention (tyrosine, placebo). On lack of an interaction effect between level and intervention, the effect of intervention on slowing slope was assessed and reported on the average slowing slope irrespective of level. For effects of level independent of tyrosine manipulation, see Bloemendaal et al. (2016).

To assess the relation between age and tyrosine’s effects on reactive and proactive response inhibition, we added the covariate age in an ANOVA using factors intervention and level for proactive RT slowing and SSRT. On significant effects or interactions with intervention, we tested for a possible interaction between administration order and intervention. On lack of interaction effects with level, the effect of intervention and age was assessed irrespective of level.

Neuropsychological measures and additional measures: catecholamine metabolites in urine, physiologic measures, and wellbeing
Outliers on any outcome measure were determined using Grubbs’ test (Grubbs, 1969), resulting in exclusion of one outlier on the HVA and one on DOPAC urine levels. Performance on neuropsychological tasks (digit span, verbal fluency, story recall, Stroop, box completion, letter cancellation) was assessed using paired t-tests comparing scores on the tyrosine session with the placebo session. The effect of tyrosine administration on catecholamine metabolites HVA, VMA, MOPEG, and DOPAC in urine was determined using four ANOVAs with within-subjects factors time (T0, T2) and intervention (tyrosine, placebo). The effect of time on blood pressure, heart rate, and subjective wellbeing was assessed using ANOVAs with factor time (T20, T90, T240). The effect of tyrosine administration on catecholamine metabolites HVA, VMA, MOPEG, and DOPAC in urine was determined using four ANOVAs with within-subjects factors time (T0, T2) and intervention (tyrosine, placebo).
MRI data acquisition and pre-processing

Whole-brain imaging was conducted on a Siemens TIM Trio 3T scanner (Magnetrom Skyra Tim, Siemens Medical Systems), using a 32-channel head coil. Functional data were obtained using a multi-echo gradient T2*-weighted echo-planar scanning sequence (Poser et al., 2006) with blood oxygen level-dependent (BOLD) contrast (34 axial-oblique slices, repetition time, 2070 ms; echo-times, 9.0, 19.3, 30.0, and 40.0 ms; in plane resolution, 3.5 × 3.5 mm; slice thickness, 3 mm; distance factor, 0.17; field of view, 224 mm; flip angle, 90°). Visual stimuli were projected on a screen and were viewed through a mirror attached to the head coil. In addition, a high-resolution T1-weighted magnetization-prepared rapid-acquisition gradient echo anatomic scan was obtained from each participant (192 sagittal slices; repetition time, 2.3 s; echo time, 3.03 ms; voxel size 1.0 × 1.0 × 1.0 mm; field of view 256 mm).

Preprocessing and mass-univariate data analysis were performed using SPM8 software (Statistical Parametric Mapping; Wellcome Trust Center for Cognitive Neuroimaging). Realignment parameters were estimated for the images acquired at the first echo-time and subsequently applied to images resulting from the three other echoes. The echo images were combined by weighting with a parallel-acquired inhomogeneity-desensitized algorithm, assessing the signal-to-noise ratio as described by Poser et al. (2006). Thirty volumes, acquired before the task, were used as input for this algorithm. After data quality check (i.e., for signal intensity spikes), the echo combined and realigned images were slice time corrected to the middle slice. The functional images were coregistered to the T1 scan. A sample-specific template was created by segmenting each individual T1 and using diffeomorphic anatomic registration to place each participant’s gray and white matter images in a study-specific space (Ashburner, 2007). Deformation parameters were stored in a subject-specific flow field. The coregistered fMRI images and anatomic T1 scan were non-linearly normalized to the sample-specific anatomic template (using the subject-specific flow field), affine-aligned into a Montreal Neurologic Institute template, and finally smoothed using an 8.0-mm full width at half maximum Gaussian filter.

To exclude activation outside gray matter from second level analyses, GM normalized maps from all subjects in the sample were used to create an average gray matter mask, which was thresholded at a value of 0.4 (voxels with computed GM fractions >40% were selected, set after visual inspection) and applied as an explicit mask during second-level analysis.

fMRI task analysis

The general linear model was set up as in (Bloemendaal et al., 2016), including twelve regressors of interest. For each level, we included four regressors: one modeling all Go trials (i.e., containing 0% and >0% stop-signal probability trials) and a corresponding parametric (i.e., proactive) regressor modeling stop-signal probability (six regressors: Go level A, Proactive level A, Go level B, Proactive level B, Go level C, Proactive level C). In level A, the parametric regressor consisted of two trial types. In level B and C, the parametric regressors consisted of five trial types. An actual stop-signal appeared on a proportion of >0% trials. These Stop trials were separately modeled as StopSuccess trials and StopFailure trials, based on whether or not the behavioral response was inhibited (six regressors: StopSuccess level A, StopFailure level A, StopSuccess level B, StopFailure level B, StopSuccess level C, StopFailure level C). As regressors of non-interest, we included a regressor for missed trials across all levels (i.e., no button box response on a Go trial), as well as a regressor modeling task instructions at the beginning of each mini-block. Moreover, twenty four realignment parameters were modeled as regressors of non-interest (six rigid-body movement parameters, plus Volterra expansion of these: first derivatives and quadratic derivatives of the original as well as first derivatives; Lund et al., 2005). Finally, to prevent contribution of global signal changes, we included signal from segmented out-of-brain voxels in the model as regressor of non-interest. All regressors of interest were modeled as delta functions at the onset of the trial and were convolved with a canonical hemodynamic response function. Time series were high-pass filtered (128-s cutoff) and serial correlations were corrected using an autoregressive (AR1) model during classical (ReML) parameter estimation. Parameter estimates for the regressors of interest, derived from the mean least-squares fit of the model to the data, were used to estimate contrasts on the first level.

At the subject-specific, first level, we specified reactive and proactive contrasts within, across, and between levels. The first level contrast images were subsequently used in a second level random effects analysis to assess consistent effects across participants as well as the effects of intervention. Reactive response inhibition can be assessed using two different contrasts: StopSuccess > StopFailure or StopSuccess > Go. The contrast StopSuccess > StopFailure provides better control for stimulus-driven processing (i.e., presentation of the stop signal), and is orthogonal by design to the proactive inhibition contrast (which also involves the Go trials). The contrast StopSuccess > Go provides better control for the timing of the Go responses (i.e., Go and StopSuccess RTs are both slower than Stop Failure) and the outcome of the trial (i.e., both successful in Go and StopSuccess). We report effects on both types of contrasts. The parametric proactive regressors constituted the contrast for proactive response inhibition.

We assessed the main task effects of proactive and reactive response inhibition using the contrasts across levels and intervention. We explored level + intervention interactions within reactive and proactive response inhibition. On non-significant interactions, we assessed the effects of intervention across levels, comparing tyrosine and placebo sessions using a paired t test. On whole-brain corrected significant results, we assessed brain-behavior correlations in these clusters by extracting β weights using Marsbar (Tzourio-Mazoyer et al., 2002).

The sample-specific gray matter mask was applied as an explicit mask to the second-level statistics (see above). Statistical inference (pFWE < 0.05) was performed at the cluster level, correcting for multiple comparisons over the
whole brain. The intensity threshold necessary to determine the cluster-level threshold was set at $p = 0.001$, uncorrected. On significant cluster-level activation, we assessed simple effects using subsequent ANOVAs or paired $t$ tests.

Results

Behavioral results

Trait demographics and trait neuropsychological test scores are presented in Table 1. For a summary of statistical tests see Table 11.

Reactive response inhibition

Race model assumptions. Data were in compliance with the main assumptions of the race model (Logan and Cowan, 1984). For each level and age group separately, mean response times (RTs) were faster on StopFailure versus >0% trials (paired $t$ test, all $p < 0.05$) and mean RTs were faster for StopFailure RTs for short versus long SSDs (paired $t$ test, all $p < 0.05$). Inhibition functions represent the probability of successfully inhibiting a response for every SSD, where the probability to inhibit decreases as the stop-signal is presented more closely to the moment that the response is made (Logan and Cowan, 1984). For each level, individual inhibition functions were calculated and displayed decreasing inhibition probability as a function of SSD.

On non-significant tyrosine effects on SSRT between levels ($F_{(2,46)} = 1.19, p = 0.31$), we performed our analyses across levels. Tyrosine administration did not affect SSRT across levels ($F_{(1,23)} < 1$) and when assessing whether the effect of tyrosine administration depended on age, no effect was observed either ($F_{(1,22)} < 1$). Across intervention sessions, increasing age was associated with a slower SSRT (main effect of age: $F_{(1,22)} = 4.3, p = 0.05$, $\eta^2 = 0.16$).

Proactive response inhibition

On non-significant tyrosine effects on RT slowing between levels ($F_{(2,46)} < 1$), we performed our analyses...
across levels. Across levels and participants, tyrosine administration did not affect slowing values (reflecting increasing slowing with increasing stop chance; $F_{(1,23)} < 1$). However, when adding age as a covariate, intervention did modulate proactive RT slowing (intervention * covariate age interaction: $F_{(1,22)} = 5.7, p = 0.03$, $r^2 = 0.20$) (Fig. 4B). Specifically, increasing age was related to a greater detrimental effect of tyrosine administration on RT slowing. Administration order did not interact with tyrosine administration on proactive slowing ($F_{(1,21)} = 2.4, p = 0.14$) (Fig. 4D).

In sum, the effect of tyrosine administration on behavioral measures of reactive response inhibition (i.e., SSRT) was not significant. However, age negatively modulated the effect of tyrosine on proactive slowing: increasing age was associated with a greater detrimental effect of tyrosine on proactive slowing compared with placebo.

**fMRI results**

**Reactive and proactive inhibition activate frontoparietal networks and basal ganglia**

At our whole-brain corrected threshold of pFWE < 0.05 (cluster-level), main task effects revealed responses in a frontoparietal and striatal task network for reactive and proactive response inhibition (Figs. 5, 6; Tables 2–4) and deactivation of, e.g., motor cortex in the reactive response inhibition network, as shown previously for the current task in young and older adults (Bloemendaal et al., 2016) and for similar paradigms (Zandbelt and Vink, 2010; Kleerekooper et al., 2016).

**Tyrosine’s effects on reactive response inhibition (StopSuccess > StopFailure and StopSuccess > Go)**

In accordance with the behavioral results, no level * intervention interactions were observed during reactive response inhibition. Hence, effects of intervention are reported across level. Tyrosine did not affect neural signal during both contrasts of reactive inhibition (StopSuccess > StopFailure and StopSuccess > Go) (Fig. 4D). A positive correlation between age and the effect of tyrosine on reactive response inhibition (StopSuccess > StopFailure) was observed in the right angular gyrus (Fig. 7; Table 5). With increasing age, tyrosine increased angular gyrus responses compared with placebo. None of these clusters demonstrated a brain-behavior correlation between tyrosine’s effect on reactive inhibition β values and SSRT.

**Figure 5.** A, Main task effects across test sessions and level for reactive response inhibition – StopSuccess > fail. Images are thresholded at $p < 0.001$ uncorrected (for illustration purposes), cluster-level (pFWE < 0.05) significant clusters are listed in Table 2. B, Main task effects across test sessions and level for reactive response inhibition – StopSuccess > Go. Images are thresholded at $p < 0.001$ uncorrected (for illustration purposes), cluster-level (pFWE < 0.05) significant clusters are listed in Table 3.

**Figure 6.** Main task effects across test sessions and level for proactive response inhibition (parametric regressors of Go). Images thresholded at $p < 0.001$ uncorrected (for illustration purposes), cluster-level (pFWE < 0.05) significant clusters are listed in Table 4.
The reactive response inhibition contrast StopSuccess > Go did not yield a correlation with age.

Tyrosine’s effects on proactive response inhibition (parametric regressor of Go)

In accordance with the behavioral results, no level x intervention interactions were observed during proactive response inhibition. Hence, effects of intervention are reported across level.

Effect of intervention on neuropsychological measures and additional measures: catecholamine metabolites in urine, physiologic measures, and wellbeing

Tyrosine administration did not affect neuropsychological measures (Table 8), nor was there an interaction between age and the effect of tyrosine administration on these measures.

Catecholamine metabolites in urine

At the beginning of the test day (T0) as well as approximately 3 h after ingestion of tyrosine or placebo (T2), a urine sample was provided by the participants. As expected when consuming yoghurt after a night’s fast, HVA, MOPEG, and VMA concentrations in urine were increased −3 h after ingestion of the mixture (T2) relative to the start of the test day (T0); main effects of time on HVA: $F_{(1,22)} = 20.98, p < 0.001$; MOPEG: $F_{(1,22)} = 48.77, p < 0.001$; VMA: $F_{(1,23)} = 52.87, p < 0.001$; Table 8).

Tyrosine administration affected catecholamine metabolites. VMA concentrations were lower after tyrosine compared with placebo (time x intervention interaction, $F_{(1,22)} = 9.1, p = 0.006$, driven by intervention differences in concentrations on T2 (t(23) = 2.2, p = 0.036), not T0 (t(23) = 0.78, p = 0.94). However, DOPAC concentrations were higher after tyrosine compared with placebo (time x intervention interaction $F_{(1,22)} = 8.8, p = 0.007$, not driven
by intervention effects on either T2 ($t_{(22)} = -0.43, p = 0.67$) or T0 ($t_{(23)} = 1.3, p = 0.2$).

Independent of intervention and time of measurement, VMA and DOPAC levels were modulated by age (main effect of age, VMA: $F_{(1,22)} = 8.98, p = 0.007$; DOPAC: $F_{(1,21)} = 5.5, p = 0.024$); increasing age was associated with higher urine levels of VMA and DOPAC. MOPEG and VMA levels interacted between time of measurement and age; increasing age was associated with a larger increase from time point T0 to T2 independent of tyrosine or placebo administration (MOPEG: $F_{(1,22)} = 5.61, p = 0.027$; VMA: $F_{(1,22)} = 4.92, p = 0.037$).

**Wellbeing, blood pressure, and heart rate**

Wellbeing ratings were assessed at the beginning of the test day (T0), at the assumed peak of tyrosine plasma level (T1) and at the end of the test day, ~4 h after ingestion (T2). At the same time, systolic and diastolic blood pressure as well as heart rate were measured (Table 10).

Over the course of the test day, changes in wellbeing as measured on the Bond & Lader total and contentness subscales were observed (main effect of time, total: $F_{(2,44)} = 4.00, p = 0.025$; contentness: $F_{(2,44)} = 9.4, p < 0.001$, respectively). Wellbeing decreased on T1 compared with T0 (total: $F_{(1,23)} = 8.6, p = 0.008$; contentness: $F_{(1,23)} = 12.8, p = 0.002$), and increased to baseline again on T2 compared with T1 and T0 (T2 vs T1 total: $F_{(1,23)} = 4.5, p = 0.045$ and contentness: $F_{(1,23)} = 10.53, p = 0.004$, T2 vs T0 total: $F_{(1,23)} < 1$ and contentness: $F_{(1,23)} < 1$). The subscales calmness and alertness did not change over time. The effect of tyrosine on the Bond & Lader subscale scores during the baseline corrected therapeutic window (T1 – T0) did not interact with age.

Systolic and diastolic blood pressure changed during the course of the test day (main effect of time systolic: $F_{(2,44)} = 8.7, p = 0.001$ and diastolic: $F_{(2,44)} = 20.7, p < 0.001$). Diastolic pressure decreased on T1 compared with T0 ($F_{(1,23)} = 18.1, p < 0.001$), and both systolic and diastolic blood pressure increased on T2 compared with T1 (systolic: $F_{(1,22)} = 8.4, p = 0.008$ and diastolic: $F_{(1,23)} = 27.6, p < 0.001$). No effects of tyrosine administration or age were observed on systolic and diastolic blood pressure or heart rate during the baseline corrected therapeutic window (T1 – T0). Heart rate did not change over time.

The correlation between age and verbal IQ was non-significant ($r = -0.22, p = 0.323$). Verbal IQ was measured with the NART, which generally shows large cohort effects between age groups, with older adults performing better than younger adults (Uttl, 2002; see also unpublished observations from our lab (by M Bloemendaal, E Aarts, M van Holstein, R Cools) using the equivalent version in the native language of the participants). Therefore, this provides some confirmation of our hypothesis that there is minimal influence of generational differences on the current results with our small age range.

**Discussion**

The current study investigated the neuro-cognitive effects of acute tyrosine administration, a dopamine precursor, on reactive and proactive response inhibition in a healthy older sample (aged 61–72 years; mean age: 67.5). Behaviorally, across the group, no effects of tyrosine administration on measures of reactive (i.e., SSRT) and proactive response inhibition (i.e., response slowing) were observed, although, neurally, proactive neural signal in right middle cingulum, precentral and supramarginal gyrus was increased by tyrosine. When taking age into account, age was found to negatively modulate the effect of tyrosine on proactive behavioral response slowing independent of cognitive load (i.e., level): increasing age was associated with a greater detrimental effect of tyrosine on proactive slowing compared with placebo. Functional imaging results were concomitant with the behavioral results: with increasing age, tyrosine decreased fronto-striatal and parietal proactive signal, but increased reactive signal in the angular gyrus. Brain-behavior correlations underline the behavioral relevance of modulated signal in these areas involved in proactive inhibition: tyrosine’s effects on bilateral putamen (and left middle frontal gyrus) signal correlated positively with ty-

---

**Table 5. Whole-brain cluster-level significant regions yielding a positive correlation between age and effect of tyrosine during reactive response inhibition (StopSuccess > StopFailure)**

<table>
<thead>
<tr>
<th>Region</th>
<th>Peak MNI coordinates</th>
<th>$p$FWE value</th>
<th>Number of voxels in cluster</th>
</tr>
</thead>
<tbody>
<tr>
<td>Right angular gyrus</td>
<td>34 – 49 40</td>
<td>0.021</td>
<td>221</td>
</tr>
</tbody>
</table>
Rosine’s effect on proactive slowing. Such brain-behavior correlations were not observed for reactive inhibition. The age-dependent detrimental effects of tyrosine on proactive slowing are surprising given that prior work has shown almost exclusively beneficial effects of tyrosine administration on cognition (for review, see Jongkees et al., 2015). Critically, these prior studies have all assessed young, not older adults. A study in adult schizophrenia patients (mean age 37.8 years, SD 6.8) displayed increased errors in a smooth pursuit saccades task in eight patients during three weeks supplementation of 10-g tyrosine daily (Deutsch et al., 1994). In line with an overdose hypothesis, a recent study demonstrated decreased working memory (i.e., n-back) performance with tyrosine’s effect on proactive slowing. Such brain-behavior correlations were not observed for reactive inhibition.

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increasing tyrosine dose (from 100-150 to 200 mg/kg) in older adults (aged 60–75 years, mean age: 69.6; van de Rest et al., 2017). This cognitive overdose effect of tyrosine, which has so far been observed only in older adults, may be at least partly caused by a larger effective dose in older adults due to increased peripheral supply of tyrosine. Earlier research demonstrated increased plasma tyrosine levels in fasting older versus young women (Caballero et al., 1991) and increased plasma response in older versus young adults receiving the same dose (van de Rest et al., 2017). Critically, in this latter study, dose-dependent increases in plasma response correlated with dose-dependent decrements in working memory after tyrosine ingestion. Several peripheral processes can cause this presumed enhanced bioavailability; for example an age-related reduced first pass effect in the liver (Klotz, 2009), may result in higher amounts of tyrosine entering the blood stream in older adults. Furthermore, age-related insulin resistance (Caballero et al., 1991) may contribute to reduced peripheral amino acid uptake from the blood, resulting in higher amounts that reach the blood-brain barrier. The current results cannot identify the cause of enhanced bioavailability, which should be further studied. Results from the current study in urine metabolites support the idea of increased peripheral catecholamine precursor levels in older adults. Irrespective of intervention, we observed higher amounts of VMA and

Table 8. Effect of intervention on stop-signal task and neuropsychological tests

<table>
<thead>
<tr>
<th>Variable</th>
<th>Placebo</th>
<th>Tyrosine</th>
<th>p value</th>
<th>p value age</th>
</tr>
</thead>
<tbody>
<tr>
<td>Story immediate recall (points)</td>
<td>9.5 (0.6)</td>
<td>10.2 (0.7)</td>
<td>0.3</td>
<td>0.68</td>
</tr>
<tr>
<td>Story delayed recall (points)</td>
<td>9.0 (0.6)</td>
<td>9.2 (0.6)</td>
<td>0.8</td>
<td>0.47</td>
</tr>
<tr>
<td>Digit span forward (points)</td>
<td>7.5 (0.4)</td>
<td>7.21 (0.4)</td>
<td>0.14</td>
<td>0.75</td>
</tr>
<tr>
<td>Digit span backward (points)</td>
<td>6.1 (0.4)</td>
<td>6.25 (0.5)</td>
<td>0.84</td>
<td>0.93</td>
</tr>
<tr>
<td>Stroop effect (s)</td>
<td>81.4 (11.1)</td>
<td>95.88 (12.5)</td>
<td>0.18</td>
<td>0.38</td>
</tr>
<tr>
<td>Stroop effect (errors)</td>
<td>7.0 (0.2)</td>
<td>1.32 (0.3)</td>
<td>0.14</td>
<td>0.43</td>
</tr>
<tr>
<td>Verbal fluency DAT (items)</td>
<td>45.26 (2.3)</td>
<td>44.13 (2.3)</td>
<td>0.8</td>
<td>0.09</td>
</tr>
<tr>
<td>Block completion</td>
<td>92.23 (7.83)</td>
<td>83.42 (4.17)</td>
<td>0.36</td>
<td>0.84</td>
</tr>
<tr>
<td>Letter cancellation</td>
<td>249.29 (7.07)</td>
<td>250.17 (7.39)</td>
<td>0.61</td>
<td>0.35</td>
</tr>
</tbody>
</table>

Data represent mean (SEM). The first column with p values reflect the outcome of the paired t tests between tyrosine and placebo on the behavioral measure, the second column with p values reflects the interaction between tyrosine’s effect on the behavioral measure and age.

Table 9. Effect of tyrosine administration on catecholamine metabolites in urine

<table>
<thead>
<tr>
<th>Variable</th>
<th>Placebo</th>
<th>Tyrosine</th>
<th>p value</th>
</tr>
</thead>
<tbody>
<tr>
<td>HVA</td>
<td>2.80 (0.22)</td>
<td>2.74 (0.24)</td>
<td>0.43</td>
</tr>
<tr>
<td>MOPEG</td>
<td>1.08 (0.06)</td>
<td>1.14 (0.07)</td>
<td>0.097</td>
</tr>
<tr>
<td>VMA</td>
<td>1.75 (0.11)</td>
<td>1.75 (0.1)</td>
<td>0.006</td>
</tr>
<tr>
<td>DOPAC</td>
<td>1.21 (0.11)</td>
<td>1.10 (0.1)</td>
<td>0.013</td>
</tr>
</tbody>
</table>

Metabolite concentrations are expressed in mmol/mol creatinine, to correct for sample volume; p values reflect the interaction between drug and time from the ANOVA on tyrosine administration (tyrosine, placebo) and time (T0, T2). The effect of tyrosine on catecholamine metabolites did not interact with age.

Table 10. Wellbeing scales, blood pressure, and heart rate

<table>
<thead>
<tr>
<th>Variable</th>
<th>Placebo</th>
<th>Tyrosine</th>
<th>p value</th>
</tr>
</thead>
<tbody>
<tr>
<td>B&amp;L total</td>
<td>8.1 (0.3)</td>
<td>8.0 (0.2)</td>
<td>0.97</td>
</tr>
<tr>
<td>B&amp;L calmness</td>
<td>7.7 (0.4)</td>
<td>8.1 (0.3)</td>
<td>0.99</td>
</tr>
<tr>
<td>B&amp;L contentness</td>
<td>8.4 (0.25)</td>
<td>8.3 (0.2)</td>
<td>0.58</td>
</tr>
<tr>
<td>B&amp;L alertness</td>
<td>8.0 (0.3)</td>
<td>7.9 (0.3)</td>
<td>0.38</td>
</tr>
<tr>
<td>Systolic blood pressure</td>
<td>136.0 (3.8)</td>
<td>137.8 (3.8)</td>
<td>0.72</td>
</tr>
<tr>
<td>Diastolic blood pressure</td>
<td>84.0 (1.7)</td>
<td>82.8 (1.8)</td>
<td>0.06</td>
</tr>
<tr>
<td>Heart rate</td>
<td>63.4 (1.8)</td>
<td>63.4 (1.7)</td>
<td>0.72</td>
</tr>
</tbody>
</table>

B&L: Bond & Lader; p values reflect the outcome of the paired t tests between tyrosine and placebo on the baseline corrected physiologic measure (T1 – T0). No interactions between tyrosine administration and time or age were observed.
### Table 11. Summary of statistical analyses

<table>
<thead>
<tr>
<th>Data structure</th>
<th>Type of test</th>
<th>Statistic</th>
<th>CI for the difference of intervention effect (unless otherwise specified)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Behavioral results</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Reactive response inhibition</td>
<td>a1. RM ANOVA</td>
<td>$F_{(2,40)} = 1.19, p = 0.31, \eta^2_g = 0.05$</td>
<td></td>
</tr>
<tr>
<td>Interaction between intervention and level on reactive response inhibition (SSRT)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Effect of intervention on proactive slowing</td>
<td>d. RM ANOVA</td>
<td>$F_{(2,23)} = 1.55, p = 0.68, \eta^2_g = 0.01$</td>
<td></td>
</tr>
<tr>
<td>Interaction between intervention and covariate age on proactive slowing</td>
<td>e1. RM ANOVA with covariate age</td>
<td>$F_{(2,23)} = 5.7, p = 0.026, \eta^2_g = 0.21$</td>
<td></td>
</tr>
<tr>
<td>Main effect of time on diastolic blood pressure</td>
<td>a1. RM ANOVA</td>
<td>$F_{(2,23)} = 4.3, p = 0.05, \eta^2_g = 0.16$</td>
<td></td>
</tr>
<tr>
<td><strong>MRI results</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Reactive and proactive response inhibition task effects</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Task effects on reactive response inhibition across intervention and level (contrasts StopSuccess vs Failure and StopSuccess vs Go)</td>
<td>g. h. one sampled t test</td>
<td></td>
<td>Tables 2, 3</td>
</tr>
<tr>
<td>Task effects on proactive response inhibition across intervention and level</td>
<td>i. one sampled t test</td>
<td></td>
<td>Table 4</td>
</tr>
<tr>
<td>Intervention effects on reactive response inhibition (contrasts StopSuccess vs Failure and StopSuccess vs Go)</td>
<td>j. k. paired t test</td>
<td>pFWE &lt; 0.05, no significant clusters</td>
<td></td>
</tr>
<tr>
<td>Effect on intervention across levels on reactive response inhibition</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Interaction between intervention (difference between tyrosine and placebo) and covariate age on reactive response inhibition</td>
<td>l. m. one-way ANOVA</td>
<td>For contrast StopSuccess vs Failure, see Table 5; contrast StopSuccess vs Go, no significant results at pFWE &lt; 0.05</td>
<td></td>
</tr>
<tr>
<td>Intervention effects on proactive response inhibition</td>
<td>n. paired t test</td>
<td></td>
<td>Table 6</td>
</tr>
<tr>
<td>Effect on intervention on proactive slowing jls</td>
<td>w. one-way ANOVA with covariate age</td>
<td></td>
<td>Table 7</td>
</tr>
<tr>
<td>Brain-behavior correlations between functional jls showing detrimental effects of tyrosine administration with increasing age and behavioral intervention effect (on proactive slowing jls)</td>
<td>p1. correlation</td>
<td>Left middle frontal gyrus (r = 0.40, p = 0.054)</td>
<td></td>
</tr>
<tr>
<td>Additional measures</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Catecholamine metabolites in urine</td>
<td>q. RM ANOVA</td>
<td>$F_{(2,23)} = 20.98, p = 0.001$</td>
<td>T0 vs T2: $-0.52/0.20$</td>
</tr>
<tr>
<td>Main effect of time on HVA</td>
<td>r1. RM ANOVA</td>
<td>$F_{(2,23)} = 52.87, p = 0.000$</td>
<td>T0 vs T2: $-0.14/0.03$</td>
</tr>
<tr>
<td>Main effect of time on VMA</td>
<td>r2. RM ANOVA</td>
<td>$F_{(2,23)} = 9.1, p = 0.006$; T0: $t_{(23)} = -2.2, p = 0.038$; T2: $t_{(23)} = -2.07, p = 0.04$</td>
<td></td>
</tr>
<tr>
<td>Main effect of time on MOPEG</td>
<td>s. RM ANOVA</td>
<td>$F_{(2,23)} = 48.77, p = 0.000$</td>
<td>T0 vs T2: $-0.27/-.15$</td>
</tr>
<tr>
<td>Main effect of time on DOPAC</td>
<td>t. RM ANOVA</td>
<td>$F_{(2,23)} = 8.8, p = 0.007$; T0: $t_{(23)} = -4.3, p = 0.67$; T2: $t_{(23)} = 1.3, p = 0.2$</td>
<td></td>
</tr>
<tr>
<td><strong>Wellbeing, blood pressure, and heart rate</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Main effect of age on VMA</td>
<td>u1. RM ANOVA with covariate age</td>
<td>$F_{(2,23)} = 9.8, p = 0.007$</td>
<td>1.68/2.07</td>
</tr>
<tr>
<td>Main effect of age on DOPAC</td>
<td>v. RM ANOVA with covariate age</td>
<td>$F_{(2,23)} = 6.6, p = 0.027$</td>
<td></td>
</tr>
<tr>
<td>Main effect of age on Dopac</td>
<td>w. RM ANOVA with covariate age</td>
<td>$F_{(2,23)} = 5.5, p = 0.042$</td>
<td></td>
</tr>
<tr>
<td>Main effect of time on B&amp;L total</td>
<td>x. RM ANOVA</td>
<td>$F_{(2,46)} = 4.00, p = 0.03$; T1 vs T0: $F_{(2,23)} = 8.6, p = 0.01$; T2 vs T1: $F_{(2,23)} = 4.5, p = 0.05$; T2 vs T0 total: $F_{(2,23)} &lt; 1$</td>
<td>T1 vs T0: $-0.075/0.20$</td>
</tr>
<tr>
<td>Main effect of time on B&amp;L contentness</td>
<td>y. RM ANOVA</td>
<td>$F_{(2,46)} = 9.4, p &lt; 0.001$; T1 vs T0: $F_{(2,23)} = 12.8, p = 0.002$; T2 vs T1: $F_{(2,23)} = 10.53, p = 0.004$; T2 vs T0: $F_{(2,23)} &lt; 1$</td>
<td>T1 vs T0: $-0.08/0.82$</td>
</tr>
<tr>
<td>Main effect of time on systolic blood pressure</td>
<td>z. RM ANOVA</td>
<td>$F_{(2,46)} = 8.7, p &lt; 0.001$; T1 vs T0: $F_{(2,23)} = 18.1, p &lt; 0.001$; T2 vs T1: $F_{(2,23)} = 8.4, p = 0.008$</td>
<td>T1 vs T0: $-4.31/1.44$; T2 vs T1: $-10.77/-.62$</td>
</tr>
<tr>
<td>Main effect of time on diastolic blood pressure</td>
<td>aa. RM ANOVA</td>
<td>$F_{(2,46)} = 20.7, p &lt; 0.001$; T2 vs T1: $F_{(2,23)} = 27.6, p = 0.000$</td>
<td>T2 vs T1: $-8.56/-2.90$</td>
</tr>
</tbody>
</table>
DOPAC across time and a higher increase of MOPEG and VMA metabolites with time, as a function of age. Thus, increased peripheral supply of tyrosine to the blood-brain barrier might have resulted in the currently observed age-dependent overdose effects on proactive response inhibition, despite the same oral dose per kilogram of bodyweight in every participant and despite increased dopamine deficits with aging.

Mechanistically, the conversion of tyrosine to L-dopa by the rate-limiting enzyme TH is inhibited by its final end products, i.e., the catecholamines dopamine and noradrenaline, present in the cytoplasm (Daubner et al., 2011). Indeed, a very high dose of phenylalanine, the conditional precursor of tyrosine, reduced dopamine release in the rat striatum, whereas lower doses increased dopamine release (During et al., 1988). The authors speculated that TH inhibition resulted in net reduced dopamine synthesis due to sudden high amounts of cytoplasmic catecholamines. Reduction of dopamine synthesis by inhibiting TH may also occur further in the dopamine signaling cascade, when an excess of dopamine increases dopamine D2 autoreceptor binding (Lindgren et al., 2001). The aging brain might be more sensitive to overshoots in auto-regulation, for example due to increased inflammatory markers, such as cytokines, which increase with age (Michaud et al., 2013) and can alter TH availability and auto-regulatory dopamine transporter expression (Felger and Miller, 2012).

Detrimental effects of tyrosine may be less surprising when considering literature on increased dopamine synthesis capacity in older adults, which is consistently observed when using the PET tracer FMT (Dejesus et al., 2001; Braskie et al., 2008; Berry et al., 2016), although mixed results have been obtained with another aromatic amino acid decarboxylase substrate, FDOPA, with decreased signal-to-noise (Martin et al., 1989; Sawle et al., 1990; Bhatt, 1991; Dreher et al., 2008). Increased age-related dopamine synthesis was negatively correlated with reward-related BOLD signal (Dreher et al., 2008) and, similarly, the positive relation between dopamine synthesis and cognitive performance seen in young adults was absent in older adults (Berry et al., 2016). We speculate that administrating extra precursor to a system with already high dopamine synthesis capacity may result in its inhibition.

The current results provide a first indication of age-related effects of tyrosine administration on dopamine processing. We used age as a continuous measure to assess its relation to tyrosine’s effects on neurocognition. However, looking at the scatter plots, it seems that the detrimental effects of tyrosine administration on proactive inhibitory responses in brain and behavior were especially apparent in the middle-old, sometimes referred to as old-old, group (70–79 years) relative to the young-old (60–69 years) participants. This is in accordance with the sub-group definition by some authors (Forman et al., 1992; de Almondes et al., 2016), although others have defined young-old individuals as 65–74 years old (Zizza et al., 2009; Moon et al., 2018). In either definition, most of our participants were in the young-old group. However, age-related changes in the dopamine system have already been observed in similar age groups as in the current study, e.g., dopamine receptor binding differences in individuals up to 68 years old (Bäckman et al., 2000) and increased dopamine synthesis capacity in a group of older adults of an average 67 years old (Braskie et al., 2008). Nevertheless, previous dopamine findings were obtained by contrasting the effects between individuals with larger age differences than in the current study, although linear effects on striatal dopamine receptor binding with increasing age can be observed among the few participants that were in our age range of 61–72 years old (Wang et al., 1998). To strengthen our results, the peripheral and central mechanisms underlying age-dependent tyrosine effects on cognition should be investigated in future studies with larger age ranges and sample sizes, including measures of dopamine functioning. Moreover, the current results should be replicated using longitudinal designs, as cross-sectional designs cannot easily control for between-subject differences other than age, which could have contributed to the current results; even in this small age range.

Previous studies that have used 150 mg/kg, similar to the current study, have observed positive effects of tyrosine administration on cognition in young adults (Jongkees et al., 2015). However, it must be noted that these studies subjected participants to a stress intervention such as acoustic noise or a cold bath. One study did not use an external stress intervention other than the task at hand (Thomas et al., 1999) and only found positive effects of 150 mg/kg tyrosine administration on demanding multitasking. Stress or demanding circumstances increase neuronal firing and thereby catecholamine metabolism (Bliss et al., 1968), making these neurons more sensitive to precursor availability such as tyrosine (Scally et al., 1977). For this reason, a relatively high dose may be optimal in a high neuronal firing situation, but suboptimal during basal neuronal firing, even in young adults.

Effects of tyrosine administration were most prominent on behavioral and neural measures of proactive response inhibition. Behaviorally, with increasing age, tyrosine modulated only proactive response slowing, not SSRT. Neurally, tyrosine modulated signal in fronto-striatal and parietal regions during proactive inhibition, which was associated with its behavioral effects. Previous studies found evidence for dopaminergic modulation of proactive-like processes in response inhibition (such as post-error slowing and go accuracy; Bari et al., 2009; Bari and Robbins, 2013), without formally disentangling proactive and reactive response inhibition as in the current paradigm. Moreover, we show age-dependent effects of tyrosine on proactive response inhibition in, among others, bilateral putamen. The putamen was also found to be modulated by Go trial probability in a study by Dunovan et al. (2015), as well as in a network uniquely active during proactive response inhibition, as determined using independent component analysis (van Belle et al., 2014). These observations strengthen the interpretation that tyrosine has selectively modulated a functional network uniquely involved in processing proactive cues.
With noradrenaline being produced from dopamine, tyrosine administration could potentially also have contributed to increased noradrenaline synthesis. We cannot fully exclude this possibility, but given the presently observed tyrosine modulation of signal in the putamen, which is highly innervated by dopamine rather than noradrenaline (Nicola and Malenka, 1998), we hypothesize that the currently observed tyrosine effects are driven by dopaminergic neurons. Our hypothesis is strengthened by literature stating shortage of especially dopamine in the aging brain (Finch, 1973; Ota et al., 2006), rather than noradrenaline (Goldman-Rakic and Brown, 1981; Moretti et al., 1987).

During reactive response inhibition, tyrosine modulated angular gyrus signal with increasing age, which is only scarcely innervated by dopamine. Moreover, no behavioral effect of tyrosine on SSRT was observed, neither a correlation between tyrosine’s effect on this reactive region and SSRT. Therefore, the observed decrease in angular gyrus activation after tyrosine with increasing age might well reflect either an indirect result of tyrosine’s fronto-striatal effects on proactive inhibition or a noradrenergic effect in combination with a floor effect in SSRTs.

The majority of the catecholamine metabolites increased to a lesser extent after tyrosine compared with placebo administration (VMA significantly and MOPEG and HVA numerically). Only DOPAC levels increased after tyrosine compared with placebo. However, for unknown reasons, large baseline (T0) differences between intervention sessions were observed on this measure. This complicates the interpretation of the intervention effect on DOPAC levels. These mixed results should generally be interpreted with caution, as urine measures mostly reflect a systemic effect in combination with a floor effect in SSRTs.

In conclusion, we show age-related effects of tyrosine administration especially on proactive, not reactive, response inhibition, accompanied by signal changes in dopamine-rich fronto-striatal brain regions. Specifically, we observed that tyrosine’s effect on brain and cognition became detrimental with increasing age, questioning the cognitive enhancing potential of tyrosine in healthy aging.

Our results, particularly those in striatum, provide support for the hypothesis that proactive, but not reactive, response inhibition is modulated by dopamine.

References


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