Reconciling psychological and neuroscientific accounts of reduced motivation in aging

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Abstract

Motivation is a hallmark of healthy aging, but the motivation to engage in effortful behavior diminishes with increasing age. Most neurobiological accounts of altered motivation in older adults assume that these deficits are caused by a gradual decline in brain tissue, while some psychological theories posit a switch from gain orientation to loss avoidance in motivational goals. Here, we contribute to reconcile the psychological and neural perspectives by providing evidence that frontopolar cortex (FPC), a brain region involved in cost-benefit weighting, increasingly underpins effort avoidance rather than engagement with age. Using anodal transcranial direct current stimulation together with effort-reward trade-offs, we find that the FPC’s function in effort-based decisions remains focused on cost-benefit calculations but appears to switch from reward seeking to cost avoidance with increasing age. This is further evidenced by exploratory, independent analysis of structural brain changes, showing that the relationship between the density of frontopolar neural tissue and willingness to exert effort differs in young versus older adults. Our results inform aging-related models of decision making by providing preliminary evidence that, in addition to cortical thinning, changes in goal orientation need to be considered in order to understand alterations in decision making over the lifespan.
Introduction

Reduced motivation to engage in demanding activities is a widespread phenomenon in aging, even in otherwise healthy individuals (Depping & Freund, 2011). Previous studies observed reduced willingness to engage in effortful behavior with increasing age (Hess & Ennis, 2012; Westbrook, Kester, & Braver, 2013). However, these deficits might be specific for self-regarding rather than for other-regarding benefits (Beadle, Sheehan, Dahlben, & Gutchess, 2015; Lockwood et al., 2020) and, moreover, some studies reported no difference in effort aversion with increasing age (Seaman et al., 2018; Seaman et al., 2016). Diminished motivation has been related to frailty in older adults (Semprini, Lubrano, Misaggi, & Martorana, 2012) and is a hallmark of several age-related conditions, including dementia and Alzheimer disease (Mortby, Maercker, & Forstmeier, 2011). Conversely, higher levels of motivation are indicative of mental and physical health as well as general well-being at older ages (Mortby et al., 2011). Because motivation is important for healthy aging, and because the population of older adults is expected to strongly increase in the future (United Nations, 2017), it is crucial to understand the brain mechanisms underlying age-related changes in motivation. Knowledge about the neurobiological basis of motivational deficits in older adults may foster the development of interventions for prevention or rehabilitation of these deficits and the associated health problems.

Research on younger adults identified a core network of brain regions involved in motivating goal-directed behavior (for reviews, see Le Heron, Apps, & Husain, 2018; Soutschek & Tobler, 2018; Studer & Knecht, 2016). Within this network, frontopolar cortex (FPC) plays a key role for the motivation to engage in effortful behavior (Soutschek, Kang, Ruff, Hare, & Tobler, 2018). Increasing FPC excitability with non-invasive brain stimulation enhanced the willingness to engage in rewarded mental and physical effort in younger adults, consistent with the hypothesized role of FPC for weighting costs against benefits of competing action alternatives (Mansouri, Koechlin, Rosa, & Buckley, 2017) and for healthy...
motivation more generally (Canu et al., 2015; Jorge, Starkstein, & Robinson, 2010; O'Callaghan, Shine, Lewis, & Hornberger, 2014). However, it is unknown whether changes in FPC structure contribute to the increased effort aversion in older adults. Age-related impairments in cognition are often thought to result from structural brain changes (Dennis & Cabeza, 2012; S.-C. Li, Biele, Mohr, & Heekeren, 2007; Lighthall, 2020; Samanez-Larkin & Knutson, 2015; West, 1996). From this perspective, we expect the stronger effort aversion in older than in younger adults to be mediated by age-related loss of FPC grey matter volume (GMV). Note that this presupposes that the function of the FPC for motivating effort engagement remains constant over lifespan but gradually declines in efficacy as result of structural neural changes.

In addition to the assumption of gradual declines in structure and function in aging neuroscience, psychological theories posit that goals in decision making change over the lifespan. In particular, motivational goals in older adults may switch from reward maximization to loss minimization (Depping & Freund, 2011; Hennecke & Freund, 2017). Under the assumption that resources for mental and physical effort are limited (Boksem & Tops, 2008), the exertion of effort can be considered as the loss of limited resources that needs to be weighed against the value of the reward at stake. With diminishing available resources for effort exertion, older adults are hypothesized to increasingly focus on preventing loss of resources for less valuable goals. From this perspective, lower motivation for effortful behavior in older adults stems from changes in goal structure in addition to gradual loss of neural tissue. That is, instead of promoting the engagement in rewarded effort as in younger adults, FPC might strengthen the goal to avoid costly expenditure of resources in older adults.

To determine the role of the FPC for motivation in older adults, the current study combines correlative analyses of structural brain changes with transcranial direct current stimulation (tDCS). tDCS is a non-invasive brain stimulation technique that facilitates the determination of the functional roles of a brain region by assessing how changing its
excitability causally alters behavior. If the FPC preserves its computational role in motivating engagement in rewarded effort in older adults despite reduced gray matter volume (as we originally hypothesized in line with the assumptions of aging neuroscience), increasing FPC excitability should enhance the willingness to engage in rewarded effort in older adults, similar to previous findings in younger adults (Soutschek et al., 2018). However, contrary to our original hypothesis and to our findings for younger adults, the current data provide evidence that increasing FPC excitability leads to lower willingness to exert effort in older adults. This is further supported by exploratory analyses suggesting an age-specific relationship between FPC gray matter volume and willingness to exert effort. By combining correlative structural analyses with causal brain stimulation methods, our study joins neural with psychological accounts of decision making in aging, substantiating the notion that changes in goal setting crucially influence altered decision making in older adults in addition to cortical thinning (Samanez-Larkin et al., 2007).

Materials and Methods

Participants

Older adults. The sample of healthy older participants included a total of 26 volunteers ($M_{\text{age}} = 69.2$ years, $SD_{\text{age}} = 4.2$, range = 65-80, 12 female). Participants were recruited via the University of the Third Age (UZH3) at the University of Zurich. According to an apriori power analysis, a sample size of 30 participants allows detecting a significant tDCS effect with a power of 80% (alpha = 5%, one-tailed), assuming the effect size (Cohen’s $d = 0.46$) for the main effect of anodal versus sham tDCS in our study on younger adults (Soutschek et al., 2018). It is worth noting though that effect sizes from single previous studies may overestimate the true effect size in the population. However, due to the outbreak of the COVID-19 pandemic, we were able to collect the data of only 26 volunteers before the pre-registered study termination date in June 2020, which reduced the statistical power to 74%. Because also
after the termination date the ongoing pandemic impeded assessing further older adults as members of a high-risk group for COVID-19, we decided to terminate the study and analyze the data. For study enrollment, volunteers had to be aged between 65-80 years, have a score ≥ 27 in the Montreal cognitive assessment test (MoCA; Nasreddine et al., 2005), and no history of psychological or neurological disorders. We note that a cutoff score of 27 in the MoCA has routinely been used in past studies, though more recent evidence suggests that it might be relatively high (Elkana, Tal, Oren, Soffer, & Ash, 2020). Participants furthermore had to fulfill all inclusion criteria for tDCS and imaging experiments. These criteria included: no history of psychiatric or neurological disorders, stroke, heart attack, or head injuries, and no metal in the body or head. The study was approved by the Cantonal Ethics Committee of Zurich and the design and the hypotheses of the tDCS study were preregistered on clinicaltrials.gov (identifier: NCT03197181). We obtained written informed consent from all participants before participation. Participants were reimbursed with 30 Swiss francs per hour plus a performance-dependent monetary bonus (see below).

Younger adults. We compared effort preferences in the older adults group with two separate samples of younger adults. The first sample of younger adults (control group 1) performed the mental and physical effort tasks under FPC tDCS, and the results for this dataset have already been reported (Soutschek et al., 2018). To compare tDCS effects between younger and older adults, we selected the younger adults in the anodal tDCS (N = 43, M_{age} = 22.5, SD_{age} = 3.0, range = 18-30, 22 female) and sham tDCS (N = 48, M_{age} = 23.2, SD_{age} = 2.3, range = 20-30, 24 female) groups but discarded the cathodal stimulation group because the older adults cohort comprised only anodal and sham tDCS conditions.

Because in the first sample of younger adult structural scans were available only for a minority of participants, we decided, post-hoc, to collect data from a second control group of younger adults in order to explore whether differences in GMV explain age-related
differences in effort preferences. Control group 2 included 28 younger adults (M\text{age} = 24.1\ years, SD\text{age} = 3.0, range = 19-33, 14 female). A power analysis based on a previous study on age differences in effort-based decision making (Westbrook et al., 2013) suggested that 23 participants per age group should be sufficient to detect a significant difference with a power of 80% (alpha = 5%, one-tailed), assuming a Cohen’s d of 0.76 observed for the Age × Effort level interaction in this study. All younger adults in control group 2 performed the physical effort decision task (see below) and underwent MRI to collect a structural brain scan, but they did not receive tDCS during task performance. For study enrollment, participants in control group 2 had to be aged between 18-35 years and fulfill the inclusion criteria for MRI.

**Stimuli and task design**

In the *effort-based decision task* (Figure 1), participants decided whether or not they were willing to exert mental or physical effort for different monetary rewards (Soutschek et al., 2018). For physical effort exertion (Figure 1A), participants had to squeeze a handgrip dynamometer for 20 s with 20%-100% of their individually determined maximum grip force (Soutschek et al., 2020; Soutschek et al., 2018). For mental effort exertion (Figure 1B), participants had to cross all instances of the letter “e” in a text composed of groups of random letters (i.e. pseudo-words) according to a demanding rule (the two letters before and the two letters after an “e” must not comprise vowels). In analogy to calibrating physical effort levels to the individual maximum strength, the mental effort levels were calibrated to the individual performance level as follows: Participants had to work on the text task for 2 minutes, and the number of lines completed within these 2 minutes was defined as 20% mental effort level. For example, if a participant completed 4 lines within 2 min, 20% mental effort in the decision task required the performance of 4 lines of this task, whereas 100% mental effort required 20 lines.
To make the task as intuitive as possible, we symbolized the monetary rewards that participants could obtain in a given trial by the number of red apples on a tree. We used five reward magnitudes (1, 3, 6, 9, and 12 apples), and informed participants that one apple would be exchanged for 0.1 Swiss francs after the experiment. The five possible effort levels were illustrated with white bars at the tree trunk (20%, 40%, 60%, 80%, and 100% effort), and the effort level required in a given trial was indicated by a red bar at the corresponding height of the trunk (Figure 1C). Participants had to accept or reject the presented offer by pressing the left or right arrow key on a standard keyboard (key-choice assignment was counterbalanced across participants) within 10 s. The chosen option turned red for 1 s and then the next trial started. We administered separate blocks for mental and physical effort, with the current type of effort displayed on top of the tree.

To disentangle motivation to exert effort from exhaustion, participants did not exert the effort immediately after accepting an offer (Soutschek et al., 2020; Soutschek et al., 2018; Westbrook et al., 2020). Instead, one trial of the decision task was randomly selected at the end of the experiment and implemented. If participants had accepted the offer, they had to exert the corresponding amount of effort to obtain the monetary bonus. If they had rejected the offer, they did not have to exert effort after the experiment and received no bonus. We informed participants that each choice had an equal chance of being selected after the experiment. To avoid influences of risk aversion of effort-based choices (e.g., participants might fear to be unable to exert 100% physical effort for 20s), participants could try to exert the required effort until they succeeded.

**Procedure**

Participants from the older adult cohort took part in three experimental sessions: In the first session, we screened participants for exclusion criteria and collected a structural T1 scan. In sessions 2 and 3, participants performed the effort-based decision task while undergoing
anodal or sham tDCS over FPC (see below). At the start of these sessions, we familiarized participants with the required mental and physical work and calibrated task difficulty to individual performance levels. To remind them of the mental effort, they had to complete as many lines of text as possible during 2 min, whereas to illustrate physical effort, they had to squeeze the dynamometer for 20 s with 50% of their maximum grip force. Next, participants were instructed about the effort-based decision task, and they performed 5 trials for each effort type as practice.

During tDCS, participants performed 4 blocks of the mental and physical effort decision tasks (2 blocks per effort type). Each block contained 25 trials, with all combinations of reward magnitudes and effort levels being presented once within a block. At the end of the experiment, participants rated their current mood and the perceived aversiveness of the stimulation, but these measures showed no significant differences between anodal and sham tDCS, mood: $t < 1, p = 0.38$, perceived aversiveness of stimulation: $t < 1, p = 0.60$.

The group of younger adults participated in one experimental session only where they performed the physical effort-based decision task (without tDCS). Younger adults performed only the physical effort-based decision task because this task allows for a more stringent calibration of effort levels to the individual maximum performance (i.e., maximum grip strength) than the mental effort task, which is a crucial precondition for comparing choice behavior across age groups.

tDCS protocol

In two separate sessions, older adults received anodal or sham tDCS (in counterbalanced order) using a 16-channel tDCS stimulator (neuroConn, Ilmenau, Germany). A smaller, 5×5 cm electrode was placed over the right FPC via neuronavigation using the same MNI coordinates (x=32, y=54, z=21) as in our previous study (Soutschek et al., 2018). A larger, 10×10 cm electrode was placed over the vertex, defined by the middle of the nasion-
inion distance (see Figure 1D for electrode positioning and the modeled current density for anodal stimulation using the Comets2 toolbox (Jung, Kim, & Im, 2013)). We used a larger vertex than FPC electrode to minimize the stimulation effect at the “control” vertex site relative to the FPC site (Nitsche et al., 2007). Both electrodes were fixed by rubber straps.

During performance of the effort-based decision task, we stimulated with 1mA current strength in the anodal tDCS condition, whereas for sham tDCS we turned the current off after 30s. Participants started task performance 4min after start of the stimulation to account for possible delays in the onset of tDCS effects.

**Magnetic Resonance Imaging**

We collected high-resolution whole-brain T1-weighted images with a fast field echo (FFE) sequence (number of slices = 170 slices, repetition time = 3.7ms, echo time = 8ms, voxel size = $1\times1\times1\text{mm}^3$, flip angle = $8^\circ$, field of view = $256\times256\text{mm}^2$). These scans were acquired on a Philips Achieva 3T whole-body scanner with a 32-channel head coil (Philips Medical Systems, Best, The Netherlands) at the Laboratory of Social and Neural Systems Research (University of Zurich, Switzerland).

**Behavioral analyses**

Behavioral data in the effort-based decision tasks were analyzed with mixed generalized linear models (MGLMs) using the lme4 package (Bates, Mächler, Bolker, & Walker, 2014) in R version 3.6.1. The alpha threshold was set to 5%. For all MGLMs, we used the optimizer “nloptwrap” and increased the number of iterations to $2e^6$ in order to ensure model convergence. In addition, we computed Bayes factors as indicators of how strongly the data favor the alternative over the null hypothesis ($BF_{10}$) with the brms package (function “hypothesis”). As prior for the null hypothesis, we assumed Cauchy-distributed priors centered around zero. To assess whether FPC tDCS modulates effort preferences in
older adults, we regressed binary choices in the decision tasks on fixed-effect predictors for tDCS (0=sham, 1=anodal), Effort type (-1=mental effort, 1=physical effort), Effort level (z-standardized), Reward magnitude (z-standardized), and the interaction terms assessing how tDCS and Effort type modulate Effort level and Reward magnitude (MGLM-1). All main effects and interaction terms were also modelled as random slopes in addition to participant-specific random intercepts. Due to technical issues with the tDCS stimulator in the anodal session, both the anodal and sham data of two participants were excluded from this analysis.

We also compared the effects of tDCS on effort preferences between our current cohort of older adults and our previously published dataset involving younger adults (control group 1). To assess whether tDCS effects depend on age, we conducted an MGLM on the combined data sets (MGLM-2) that included the same predictors as MGLM-1 and additionally predictors for Age (z-standardized) as well as the interactions between Age and all predictors of MGLM-1.

We note that MGLM-2 does not allow comparing the influences of reward magnitude and required effort between younger and older adults due to differences in task procedures. We therefore compared effort preferences in the older adult cohort with a second control group of younger adults which had performed the physical effort task in the same way as the older adults. To test the hypothesis that willingness to engage in effort decreases with increasing age, we regressed younger and older adults’ binary choices in the effort-based decision task on z-transformed fixed-effects predictors for Age, Effort level, Reward magnitude, and all interaction terms (MGLM-3). As random effects, we modelled participant-specific random intercepts and random slopes for Effort level, Reward magnitude, and the interaction term. In the cohort of older adults, we restricted this analysis to the data from the sham session. As post-hoc tests, we computed MGLMs that assessed the impact of age on willingness to exert effort separately for each effort level (MGLM-4). For these post-hoc
tests, we adjusted the p-values using Bonferroni correction ($p_{\text{corrected}} = p_{\text{uncorrected}} \times 5$, which is equivalent to a corrected alpha threshold of $\frac{5\%}{5} = 1\%$).

**Voxel-based morphometry (VBM) analysis**

Preprocessing of the structural brain scans was performed using the CAT12 toolbox (Gaser & Dahnke, 2016) and comprised three steps. Each structural image first was normalized to Montreal Neurological Institute (MNI) space and, second, segmented into grey matter (GM), white matter, and cerebrospinal fluid (CSF). Third, the normalized segmented images were spatially smoothed with a Gaussian kernel (FWHM=8mm). We also determined each participant’s total intracranial volume (TIV) using the routines implemented in the CAT12 toolbox.

To test whether differences in motivation between older and younger adults are related to changes in brain structure, we conducted a whole-brain VBM analysis (using SPM12) regressing preprocessed GM voxels on the following predictors: acceptance rates for 20% effort (based on the individual coefficients from MGLM-3 for 20% effort), age, and the interaction between age and acceptance rates. We used the 20% level as for this effort level we observed significant age difference in motivation. As covariate of no interest, we entered the participant-specific TIV. In order to assess GMV changes in the FPC region stimulated in the tDCS experiment, we performed small-volume correction with a spherical ROI (diameter=16mm, twice the smoothing kernel) centered at the coordinates used for the placement of the FPC tDCS electrode ($x=32$, $y=54$, $z=21$). For exploratory whole-brain analyses, we controlled for multiple comparisons using family-wise error (FWE) correction at the peak or cluster level, employing a cluster-inducing threshold of $p < 0.001$ (Eklund, Nichols, & Knutsson, 2016).

**Results**
Anodal stimulation over FPC lowers willingness to engage in effort in older adults

We tested the computational role of FPC for motivation by assessing whether excitatory anodal FPC tDCS increases the willingness to exert effort. MGLM-1 suggested that participants were less willing to engage in effort with increasing effort levels, $\beta = -2.94$, $z = 3.97$, $p < 0.001$, $BF_{10} = 2e28$, as well as with decreasing rewards available, $\beta = 3.05$, $z = 4.21$, $p < 0.001$, $BF_{10} = 1e18$, and effort discounting was steeper the higher the rewards at stake, $\beta = -0.89$, $z = 2.21$, $p = 0.04$, $BF_{10} = 52.6$ (Figure 2A/B). Contrary to our findings for younger adults and our original hypothesis (Soutschek et al., 2018), a main effect of tDCS indicated that older adults were less, rather than more, willing to exert effort under anodal compared with sham tDCS, $\beta = -1.99$, $z = 2.43$, $p = 0.02$, $BF_{10} = 3.9$ (Figure 2C and Table 1). Note that if as prior for the Bayes factor we assumed the effect size from our previous study on younger adults (instead of a prior centered around zero), the Bayes factor of 4.4 would even more strongly favor the alternative over the null hypothesis (the likelihood of the null hypothesis, i.e. that the magnitude of the tDCS effect is equal to the tDCS effect in the younger adults study, relative to the alternative hypothesis is only 0.23). While there was a non-significant trend for higher acceptance rates for physical than for mental effort, $\beta = 0.61$, $z = 1.83$, $p = 0.07$, $BF_{10} = 4.3$, tDCS effects did not significantly differ between mental and physical effort, $\beta = -0.26$, $z = 0.64$, $p = 0.52$, $BF_{10} = 2.1$. No further effect including the factor tDCS was significant, all $z < 0.39$, all $p > 0.39$, and we also observed no significant differences between mental and physical effort (besides the trend-level main effect of Effort type), all $z < 0.52$, all $p > 0.60$. Taken together, increased FPC excitability lowered the motivation to engage in effort in older adults, in contrast to the role of the FPC in younger adults (Soutschek et al., 2018).

To directly assess whether FPC stimulation has opposite effects on effort-based decisions in younger versus older adults, we compared the current data in older adults with our previous tDCS data set for younger adults (for details, see Soutschek et al. (2018)). The
sample of younger adults included 43 participants in the anodal group and 48 participants in the sham group (we note that the sample of younger adults was considerably larger than the sample of older adults). The cathodal stimulation group was discarded from this analysis, because we had used only anodal and sham tDCS in the study on older adults. MGLM-2 on the combined data set revealed a significant interaction between Age and tDCS (anodal versus sham), \( \beta = -0.28, z = 2.19, p = 0.03, \text{BF}_{10} = 3.8 \), providing further evidence that FPC tDCS has dissociable effects on effort-based decisions in younger and older adults. We also observed a significant Age × Effort level interaction, \( \beta = 0.63, z = 3.81, p < 0.001, \text{BF}_{10} = 96.4 \), suggesting that effort preferences under sham differed between younger and older adults as a function of the required effort level (Table 2). We note though that the tasks used in the older adults and the younger adults samples were not fully comparable, because we had used different calibration procedures for mental effort and because the task for younger adults had included additional reward and effort levels. As these differences might affect effort preferences, the comparison between effort preferences in older and younger adults based on these data should be interpreted with caution. Nevertheless, the findings provide preliminary and consistent evidence that FPC might have dissociable roles for the motivation of effortful behavior in older and younger adults.

Changes in FPC structure explain individual differences in cost-benefit weighting in an age-specific manner

The finding that increased FPC excitability in older adults reduced motivation raises the question as to why FPC stimulation shows opposite effects on motivation in older versus younger adults. Given the hypothesized role of FPC for integrating costs and benefits for action alternatives (Mansouri et al., 2017), one possibility is that this result is due to a shift in goals from gain maximization to effort minimization, consistent with psychological theories of aging (Depping & Freund, 2011; Hennecke & Freund, 2017). FPC activity may serve to
enhance or boost the current goal over its costs, instead of simply promoting reward over effort in a fixed, goal-independent manner. This sort of goal-dependent role could explain why FPC activity would result in reward-seeking overriding effort costs more often in younger adults (resulting in a higher willingness to engage in effort), whereas in older adults, it seems to promote effort avoidance.

To test for goal-dependent influences of FPC in another way, we explored the relation between FPC gray matter volume and behavior in older versus younger adults. Studies linking structural brain changes to age-differences in decision making or cognition typically assume that a region’s computational role remains constant over age but gradually declines (Dennis & Cabeza, 2012; S.-C. Li et al., 2007; Lighthall, 2020; Samanez-Larkin & Knutson, 2015; West, 1996). Contrary to this assumption, our tDCS findings suggest that correlations between FPC structure and willingness to exert effort over age may not only reflect age-related decreases in functional efficacy in performing cost-benefit computations, but also a shift from gain to loss orientation. Together, the theories that goals switch from gain maximization to loss minimization (Depping & Freund, 2011; Hennecke & Freund, 2017) and that the FPC is goal-dependent predict that the relationship between FPC GMV and willingness to exert effort changes with age. Specifically, in younger adults FPC GMV should be associated with a higher willingness to exert effort, whereas in older adults higher FPC GMV should correlate with a lower willingness to engage in effort.

To test this assumption, we collected data from a new group of younger adults, who – just as our cohort of older adults did – performed a physical effort-based decision task and underwent MRI for a structural brain scan. We first aimed to replicate previous findings that older adults are more effort-averse than younger adults (Hess & Ennis, 2012; Westbrook et al., 2013) by regressing binary choices to accept versus reject effortful offers in the physical effort-based decision task on continuous predictors for Age, Effort level, Reward magnitude, and the interaction terms (MGLM-3; Table 3). In the sample of older adults, we restricted this
analysis to the physical effort-based task in the sham condition, because control group 2 of younger adults had performed only the decision task for physical effort without receiving tDCS (but undergoing MRI to collect individual structural scans; see Materials and Methods). A significant Age × Effort level interaction, \( \beta = 0.69, z = 2.01, p = 0.04, BF_{10} = 5.5 \), suggested that younger and older adults differed in their motivation to engage in effort depending on the required demands (Figure 3A). Post-hoc MGLMs (MGLM-4; separately for each effort level) revealed that older adults had reduced acceptance rates for the lowest, 20% effort level, \( \beta = -0.57, z = 3.38, p = 0.004 \), Bonferroni-corrected, \( BF_{10} = 57.3 \). The 40% effort level showed a non-significant trend for higher acceptance rates in the younger than in the older adult cohort, \( \beta = -0.37, z = 2.39, p = 0.07 \), Bonferroni-corrected, \( BF_{10} = 2.9 \). For all other effort levels, we observed no significant age differences, all \( z < 1.37 \), all \( p > 0.8 \), Bonferroni-corrected. There was no evidence that age modulated the sensitivity to monetary rewards, \( \beta = -0.29, z = 0.89, p = 0.37 \), \( BF_{10} = 1.3 \). Taken together, these findings suggest that aging is associated with increased sensitivity to low levels of effort, a hallmark of apathy (Le Heron, Plant, et al., 2018).

Next, we asked whether these age-related differences in behavior can be explained by variation in brain structure. To test whether the link between FPC GMV and motivation changes with age, we regressed FPC GMV on predictors for age, willingness to exert effort (given by the individual coefficients from the MGLM for the 20% effort level, as this model had shown significant age differences in behavior), and the interaction term (controlling for total intracranial volume). We observed a significant interaction effect between age and willingness to exert effort in FPC (\( x = 24, y = 62, z = 27; t = 3.87; p = 0.03 \), small-volume corrected with 16 mm spherical mask centered at coordinates for the FPC tDCS electrode placement; note that an exploratory whole-brain analysis revealed no further significant clusters, see Table 4). Plotting this interaction effect by extracting individual GMV from the significant FPC cluster suggests that in younger adults, as expected based on our previous
findings, greater FPC GMV tended to predict higher willingness to exert effort, Spearman’s rho = 0.30, p = 0.06, one-tailed, BF$_{10}$ = 1.9, whereas in older adults FPC GMV was negatively correlated with acceptance rates, rho = -0.39, p = 0.046, two-tailed, BF$_{10}$ = 5.8 (Figure 3B). Although these effects are relatively weak, they are nevertheless consistent with the observed opposite effects of FPC tDCS on motivation in younger and older adults.

Discussion

Reduced motivation is a prevalent problem in aging, but its neurobiological roots remain unclear so far. Here, we provide preliminary evidence that changes in both brain structure and the goal orientations underlying cost-benefit tradeoffs need to be considered in order to explain the lower willingness to engage in goal-directed effort in older adults. First, we replicated previous findings of increased effort aversion in older compared with younger adults (Hess & Ennis, 2012; Westbrook et al., 2013), in line with increased apathy (Le Heron, Plant, et al., 2018). We note that two previous studies reported no evidence for stronger physical effort aversion in older compared with younger adults (Seaman et al., 2018; Seaman et al., 2016), though this might partially be explained by a floor effect because effort aversion was rather low in all age groups in these studies. Interestingly, older adults showed stronger effort aversion than younger adults particularly to low effort requirements, which may potentially reflect a floor effect (as acceptances rates for high effort in younger adults were already rather low). Noteworthy, there was no evidence that the influence of reward magnitude on choices differed between age groups, speaking against the possibility that older adults valued monetary rewards less than younger adults. Many neuroscientific accounts of decision making in older adults suggest that such behavioral deficits result from neural changes (Lighthall, 2020; Samanez-Larkin & Knutson, 2015), implicitly assuming that goals and neural computations remain constant throughout adulthood, and only few studies considered psychological accounts of aging to explain age-related changes in neural
functioning (Samanez-Larkin et al., 2007). Psychological theories posit that increased effort aversion in older adults reflects a switch from gain maximization to loss minimization goals (Depping & Freund, 2011; Hennecke & Freund, 2017).

Our study provides a further step towards bridging this gap by suggesting that changes in both brain structure and goal orientation contribute to the increased effort sensitivity in aging. First, increasing FPC excitability lowered the willingness to engage in mental or physical effort in older adults. This is in contrast to our own previous findings for younger adults, in which FPC stimulation increased motivation (Soutschek et al., 2018). We note that previous studies on motor function, language processing, and memory have reported similar effect sizes for stimulation effects in younger and older adults (or, if anything, even stronger stimulation effects in older adults) despite GMV loss in older adults (for a review, see Perceval, Floel, & Meinzer, 2016). Therefore, the opposite stimulation effects are unlikely to be caused by age-related changes in FPC GMV per se. Second, the tDCS findings are further supported by the VBM analysis providing evidence that the relationship between FPC GMV and willingness to exert effort changes with age, such that younger versus older adults show positive versus negative correlations between FPC GMV and motivation for rewarded effort (though we note that the correlations between GMV and motivation within each age group are relatively weak and should be interpreted with caution).

These findings are inconsistent with the assumption that the FPC plays a functional role in promoting reward obtainment that remains constant with age. Instead, a goal-dependent role for FPC appears to be the best explanation for the age-dependent link between motivation and FPC. We speculate that FPC may bias the evaluation of reward benefits versus effort costs in different directions due to differences in internal goals in younger and older adults. This is consistent with psychological theories claiming that goals change from gain to loss orientation over the lifespan (Depping & Freund, 2011), though as caveat we note that we did not directly measure goal orientation in the current study. We also note that our study
followed a cross-sectional instead of a longitudinal design, such that one needs to be careful with conclusions regarding changes in the FPC’s functional role on an individual level. Neurally, this shift in the FPC’s computational role may be linked age-related changes in the input and output between FPC and brain regions representing rewards and effort costs. For example, aging appears to be associated with impairments in the dopaminergic reward system (S. C. Li & Rieckmann, 2014) and with increased functional connectivity strength between FPC and supplementary motor cortex (Q. Li et al., 2020), a region that encodes physical effort costs and sends effort cost information to FPC (Bonnelle, Manohar, Behrens, & Husain, 2016; Burke, Brunger, Kahnt, Park, & Tobler, 2013). Thus, increased aversion to low levels of effort in older adults may also be a consequence of re-organization within motivation networks.

These conclusions have implications for neural accounts of decision making across the lifespan. Such accounts often aim to relate quantitative properties of neural tissue (like GMV, cortical thickness, or fiber connections) with the degree of cognitive functioning, but neglect that goals, perhaps instantiated by a brain region’s relative weighting of inputs, may also change with increasing age (but see Samanez-Larkin et al., 2007). We note that evidence for similar dissociations between younger and older adults was reported in other stimulation studies as well. For example, cathodal stimulation of left and anodal stimulation of right lateral prefrontal cortex reduces risk-taking in younger adults (Fecteau et al., 2007), while older adults show a stronger preference for risky prospects under the same stimulation protocol (Boggio et al., 2010). These findings too can be reconciled with the hypothesized change from gain maximization to loss avoidance over the lifespan. Thus, an implication of our results is that future neural investigations of decision making in older adults should consider both changes in neural tissue and in goal orientation as mediators of altered behavior in aging.
It is worth mentioning some limitations of the current investigation. First, we emphasize the exploratory nature of the comparisons between younger and older adults, which we conducted to explain the unexpected tDCS effects in older adults. Second, while some of the p-values for the tDCS effects and the group comparisons are relatively close to the statistical threshold, it is worth pointing out that the tDCS and the GMV findings provide converging evidence for age-specific roles of FPC for motivation. Nevertheless, given the limited sample size and given that we could assess four older adults less than planned apriori due to the COVID-19 pandemic, both the tDCS and the GMV findings should be considered preliminary. Third, the mental effort task confounded effort with delay because higher effort levels required participants to work on more lines of text, which increased the delay until payment by a few minutes. However, the fact that we observed no significant differences between mental and physical effort suggests that time considerations had virtually no impact on participants’ choices.

Fourth, previous studies observed that the strength of tDCS effects depends on cortical thickness of the stimulated region and/or on neurochemical concentrations of GABA or glutamate (Filmer et al., 2020; Filmer, Ehrhardt, Bollmann, Mattingley, & Dux, 2019). Because in the current study we did not measure neurochemical concentrations and had acquired a structural scan only for a minority of the younger adults in the tDCS experiment, it is possible that differences in neurochemical concentrations or cortical thickness between older and younger adults explain the dissociable effects of anodal tDCS in these groups. In fact, concentrations of neurotransmitters such as GABA or dopamine differ between younger and older adults (Huang et al., 2017; Mora, Segovia, & Del Arco, 2008) and might contribute to age-related changes in decision making and reward sensitivity (Samanez-Larkin & Knutson, 2015). From this view, neurochemical changes constitute a plausible neurobiological mechanism underlying the FPC’s age-related change in functioning observed in our study.
A further limitation of the current findings is that the age difference in effort preferences was significant only for the lowest (20%) effort level. While we suggested that this result might hint to a potential floor effect (for example, 43% of all younger adults had an acceptance rate of 0% for the 100% effort level), there are other plausible explanations as well. According to the selective engagement theory (Hess, 2014), when deciding to mobilize effort, older adults, compared to younger adults, are more strongly guided by the personal relevance of goals or by self-presentation concerns. As it has been suggested that individuals occasionally engage in high effort demands for self-presentation purposes (Inzlicht, Shenhav, & Olivola, 2018), a potential (albeit speculative) hypothesis for the lack of age differences at higher effort levels is that high effort demands more strongly activated self-presentation motives in older adults than in younger adults. In any case, we acknowledge that the current data do not provide a conclusive answer to the question as to why the age difference in effort avoidance was strongest for the lowest effort level.

Taken together, our findings provide insights into the psychological and neurobiological roots of altered motivation in healthy aging. One source of the increased avoidance of low effort in older adults might be a goal-dependent functioning of FPC. Given the apparent goal-dependent nature of FPC function, and most likely other brain regions as well, neural interventions aimed at preventing or counteracting loss of motivation in older adults should consider how goals are set and pursued as well.
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Declaration of interest

The authors declare to have no conflict of interest.

Open practices statement

The raw behavioral data, a script for the behavioral data analysis, and the results of the VBM analysis are available on Open Science Framework (https://osf.io/ejs47/). The tDCS experiment was preregistered on clinicaltrials.gov (identifier: NCT03197181).
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Figure captions

**Figure 1.** Experimental tasks. Older adult participants performed both a physical effort and a mental effort task, younger adults the physical effort task only. (A) For physical effort exertion, participants had to squeeze a handgrip with variable levels of their maximum grip force for 20 s. (B) The mental effort exertion required crossing all letters “e” according to a given rule in a text with random letter sequence. Levels of mental effort were manipulated via the number of lines to be completed (with 20% defined as the number of lines a participant completed within 2 min). (C) In the effort-based decision task, participants made a series of choices between accepting or rejecting offers to engage in rewarded mental or physical effort. The magnitude of the available reward was illustrated by the number of apples on the tree, while required effort demands were indicated at the tree trunk. (D) Current density modelling suggests that tDCS effects were most pronounced in right frontopolar cortex.
Figure 2. Brain stimulation results for older adults. (A, B) Illustration of the effects of frontopolar stimulation on willingness to exert effort as a function of effort level and reward magnitude, separately for (A) sham and (B) anodal stimulation (collapsed across mental and physical effort because tDCS effects did not significantly differ between mental and physical effort). Error bars indicate standard error of the mean. (C) Contrary to previous findings for younger adults (Soutschek et al., 2018), anodal (relative to sham) tDCS over frontopolar cortex lowers the motivation to engage in rewarded mental or physical effort in older adults. Black dots indicate individual coefficients for the main effect of tDCS in MGLM-1.
Figure 3. Relation of willingness to exert physical effort, age and frontopolar cortex volume. (A) Boxplots showing the willingness to exert physical effort as function of effort level (20-100%), separately for younger and older adults. Older adults show a significantly lower willingness to exert low levels (i.e., 20% of maximum effort) of physical effort for rewards than younger adults. Red lines indicate the median, black dots represent individual data points. (B) The relationship between the willingness to exert low levels of effort and frontopolar (FPC) gray matter volume (GMV) differed between younger and older adults. For display purposes, we show the younger and older adults as separate groups in different colors. While in younger adults higher FPC GMV tends to predict increased willingness to exert effort (green dots and regression line), in older adults greater FPC GMV is associated with a lower motivation to engage in rewarded effort (purple dots and regression line). Asterisks indicate significant effects (* p < 0.05, ** p < 0.01).
Table 1. Results of MGLM-1 assessing the impact of frontopolar tDCS (anodal versus sham) on willingness to exert effort in older adults. Standard errors of the mean are in brackets. BF\textsubscript{10} is the Bayes factor indicating how strongly the data favor the alternative over the null hypothesis (note that no Bayes factor can be computed for fixed-effect intercepts in mixed models).

|                  | Beta (SE) | Z  | P     | BF\textsubscript{10} |
|------------------|-----------|----|-------|----------------------|
| Intercept        | 1.62 (0.56) | 2.89 | 0.004 |                      |
| Reward           | 3.05 (0.72) | 4.21 | <0.001 | 1e18                 |
| Effort           | -2.94 (0.74) | 3.97 | <0.001 | 2e28                 |
| Effort type      | 0.61 (0.33) | 1.83 | 0.07  | 4.3                  |
| tDCS             | -1.99 (0.89) | 2.24 | 0.02  | 3.9                  |
| tDCS × Reward    | -0.60 (0.94) | 0.64 | 0.52  | 2.2                  |
| tDCS × Effort    | 0.94 (1.17) | 0.81 | 0.42  | 1.3                  |
| tDCS × Effort type| -0.26 (0.41) | 0.64 | 0.52  | 2.1                  |
| Reward × Effort  | -0.89 (0.42) | 2.11 | 0.04  | 52.6                 |
| Effort type × Reward | 0.04 (0.35) | 0.13 | 0.90  | 0.5                  |
| Effort type × Effort | -0.05 (0.43) | 0.12 | 0.91  | 1.8                  |
| tDCS × Effort type × Reward | -0.11 (0.41) | 0.27 | 0.79  | 1.2                  |
| tDCS × Effort type × Effort | 0.06 (0.50) | 0.12 | 0.90  | 3.6                  |
| tDCS × Reward × Effort | 0.55 (0.64) | 0.85 | 0.39  | 0.8                  |
| Effort type × Reward × Effort | -0.17 (0.34) | 0.51 | 0.61  | 0.4                  |
| tDCS × Effort type × Reward × Effort | 0.19 (0.40) | 0.47 | 0.64  | 0.5                  |
Table 2. Results of MGLM-2 comparing the impact of frontopolar tDCS (anodal versus sham) on willingness to exert effort between younger and older adults. BF$_{10}$ is the Bayes factor indicating how strongly the data favor the alternative over the null hypothesis (note that no Bayes factor can be computed for fixed-effect intercepts in mixed models).

|                  | Beta (SE) | Z     | P       | BF$_{10}$ |
|------------------|-----------|-------|---------|-----------|
| Intercept        | -0.84 (0.28) | 2.19  | 0.03    |           |
| Reward           | 3.73 (0.22)  | 16.61 | <0.001  | 2.0e17    |
| Effort           | -3.68 (0.18) | -20.28| <0.001  | 4.4e16    |
| Effort type      | 0.76 (0.12)  | 6.11  | <0.001  | 3.4e17    |
| tDCS             | 0.35 (0.26)  | 1.34  | 0.18    |           |
| Age              | 0.48 (0.37)  | 1.30  | 0.19    | 3.7       |
| tDCS × Reward    | 0.16 (0.17)  | 0.95  | 0.34    |           |
| tDCS × Effort    | 0.08 (0.14)  | 0.55  | 0.59    | 0.9       |
| tDCS × Effort type| -0.06 (0.10) | 0.61  | 0.54    | 0.7       |
| tDCS × Age       | -0.28 (0.13) | 2.19  | 0.03    | 3.8       |
| Reward × Effort  | -0.58 (0.10) | 6.00  | <0.001  | 781.3     |
| Effort type × Reward | 0.14 (0.09) | 1.61  | 0.11    | 0.6       |
| Effort type × Effort | 0.19 (0.11) | 1.66  | 0.10    | 1.8       |
| Reward × Age     | -0.57 (0.21) | 2.74  | 0.006   | 41.7      |
| Effort × Age     | 0.63 (0.16)  | 3.81  | <0.001  | 96.4      |
| Effort type × Age| -0.17 (0.11) | 1.58  | 0.11    | 10.3      |
| tDCS × Effort type × Reward | 0.07 (0.06) | 1.10  | 0.27    | 0.9       |
| tDCS × Effort type × Effort | -0.08 (0.09) | 0.91  | 0.36    | 0.7       |
| tDCS × Reward × Effort | -0.01 (0.08) | 0.15  | 0.88    | 0.6       |
| Effort type × Reward × Effort | -0.18 (0.07) | 2.59  | 0.01    | 1.8       |
| tDCS × Reward × Age | -0.03 (0.09) | 0.33  | 0.74    | 0.5       |
| tDCS × Effort × Age | -0.03 (0.07) | 0.42  | 0.67    | 0.5       |
| tDCS × Effort type × Age | 0.03 (0.05)  | 0.55  | 0.58    | 0.4       |
| Reward × Effort × Age | -0.01 (0.09) | 0.16  | 0.88    | 0.5       |
| Effort type × Reward × Age | -0.08 (0.06) | 1.37  | 0.17    | 1.1       |
| Effort type × Effort × Age | 0.02 (0.09)  | 0.19  | 0.85    | 0.6       |
| tDCS × Effort type × Reward × Effort | -0.01 (0.06) | 0.25  | 0.80    | 0.5       |
| tDCS × Effort type × Reward × Age | 0.03 (0.04)  | 0.77  | 0.44    | 0.4       |
| tDCS × Effort type × Effort × Age | -0.05 (0.05) | 1.02  | 0.31    | 0.9       |
| tDCS × Reward × Effort × Age | 0.03 (0.04)  | 0.66  | 0.51    | 0.4       |
| Effort type × Reward × Effort × Age | 0.08 (0.05)  | 1.38  | 0.17    | 0.4       |
| tDCS × Effort type × Reward × Effort × Age | -0.00 (0.03) | 0.02  | 0.98    | 0.3       |
Table 3. Results of MGLM-3 assessing the impact of age on willingness to exert physical effort. Standard errors of the mean are in brackets. BF_{10} is the Bayes factor indicating how strongly the data favor the alternative over the null hypothesis (note that no Bayes factor can be computed for fixed-effect intercepts in mixed models).

| Effect             | Beta (SE)   | Z     | P      | BF_{10} |
|--------------------|-------------|-------|--------|---------|
| Intercept          | 1.82 (0.34) | 5.39  | <0.001 |         |
| Reward             | 3.51 (0.36) | 9.78  | <0.001 | 2e17    |
| Effort             | -3.39 (0.38)| 8.95  | <0.001 | 4e20    |
| Age                | -0.36 (0.32)| 1.14  | 0.25   | 1.6     |
| Age × Effort       | 0.69 (0.34) | 2.01  | 0.04   | 5.5     |
| Age × Reward       | -0.29 (0.32)| 0.89  | 0.37   | 1.3     |
| Reward × Effort    | -0.79 (0.25)| 3.13  | 0.002  | 23.8    |
| Age × Reward × Effort | -0.12 (0.24)| 0.49  | 0.62   | 1.1     |
Table 4. Anatomical locations and MNI coordinates of the peak activations showing an interaction effect between age and willingness to engage in 20% physical effort

| Region                        | Hem | BA | X   | Y   | Z   | k   | t    |
|-------------------------------|-----|----|-----|-----|-----|-----|------|
| Frontopolar cortex            | R   | 10 | 24  | 62  | 27  | 20  | 3.87 |
| Insula                        | R   | 13 | 36  | -6  | 2   | 29  | 3.55 |
| Supplementary motor area      | L   | 6  | -9  | -14 | 63  | 21  | 3.55 |

The frontopolar cortex cluster is significant after applying small-volume correction within a spherical mask centered on the coordinates for the FPC tDCS electrode placement (see main text). The results in the insula and supplementary motor area do not survive corrections for multiple comparisons at the whole brain level (p < 0.001 uncorrected, minimum cluster size k ≥ 20 voxels), but are reported here for completeness and to facilitate the formation of future hypotheses and replication studies. We do not make any inferences on these insula and supplementary motor area results in the current manuscript. Hem = Hemisphere (L = left, R = right); BA = Brodmann area.