Age-Related Differences in Ventral Striatal and Default Mode Network Function During Reciprocated Trust

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Manuscript information
Number of pages: 31
Number of figures (5), tables (1)
Number of words for abstract (229), introduction (639), and discussion (1035)

Acknowledgments
This work was supported, in part, by grants from the National Institutes of Health (R21-MH113917, R03-DA046733, and RF1-AG067011 to DVS and R15-MH122927 to DSF) and a Pilot Grant from the Scientific Research Network on Decision Neuroscience and Aging [to DVS; Subaward of NIH R24-AG054355 (PI Gregory Samanez-Larkin)]. We thank Elizabeth Beard for assistance with task coding, Dennis Desalme, Ben Muzekari, Isaac Levy, Gemma Goldstein, and Srikar Katta for assistance with participant recruitment and data collection, and Jeffrey Dennison for assistance with data processing. DVS was a Research Fellow of the Public Policy Lab at Temple University during the preparation of this manuscript (2019-2020 academic year).

Conflict of interest statement
The authors have no conflicts to disclose.

Data and code availability
Data can be found on \url{https://openneuro.org/datasets/ds003745}. Analysis code can be found on \url{https://github.com/DVS-Lab/srndna-trustgame}. Thresholded and unthresholded statistical maps are located on \url{https://identifiers.org/neurovault.collection:10447}.

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Abstract

Social relationships change across the lifespan as social networks narrow and motivational priorities shift to the present. Interestingly, aging is also associated with changes in executive function, including decision-making abilities, but it remains unclear how age-related changes in both domains interact to impact financial decisions involving other people. To study this problem, we recruited 50 human participants (N_{younger} = 26, ages 18-34; N_{older} = 24, ages 63-80) to play an economic trust game as the investor with three partners (friend, stranger, and computer) who played the role of investee. Investors underwent functional magnetic resonance imaging (fMRI) during the trust game while investees were seated outside of the scanner. Building on our previous work with younger adults showing both enhanced striatal responses and altered default-mode network (DMN) connectivity as a function of social closeness during reciprocated trust, we predicted that these relations would exhibit age-related differences. We found that striatal responses to reciprocated trust from friends relative to strangers and computers were blunted in older adults relative to younger adults, thus supporting our primary pre-registered hypothesis regarding social closeness. We also found that older adults exhibited enhanced DMN connectivity with the temporoparietal junction (TPJ) during reciprocated trust from friends compared to computers while younger adults exhibited the opposite pattern. Taken together, these results advance our understanding of age-related differences in sensitivity to social closeness in the context of trusting others.
Significance

Our social relationships change as we grow older. Despite the ubiquity of these changes, little is known about how they impact our neural and behavioral responses to financial decisions involving other people. We examined this problem in the context of a trust game played with three partners varying in social closeness (friend, stranger, computer) while participants underwent fMRI. We found both blunted striatal activation and enhanced DMN-TPJ connectivity to reciprocated trust from friends relative to strangers and computers in older compared to younger adults. These results suggest that older adults demonstrate altered representations of social closeness within financial exchanges involving trust, which may expose older adults to increased risk for financial exploitation perpetrated by strangers.
Introduction

The transition to older adulthood is dynamic, characterized by shrinking social networks, prioritization of social relationships and socially-centered goals, and an emphasis on positive relative to negative social experiences (Carstensen, 1992, 1995; Charles, 2010). These age-related socioemotional changes are coupled with changes in neural networks supporting executive and social function (Andrews-Hanna et al., 2007; Devitt & Schacter, 2020; Hughes et al., 2020; Laurita et al., 2020; Persson et al., 2006; Spreng et al., 2020), and taken together, such patterns have implications for older adults’ abilities to successfully engage in social interactions. While an emerging body of literature has highlighted changes in decision-making in older relative to younger adults (Burr et al., 2021; Lighthall et al., 2018; Seaman et al., 2018) we know surprisingly little about how this translates to social decisions (i.e., trusting others) and processing outcomes (e.g., reciprocity, betrayal) involving people of varying degrees of social closeness (e.g., friends, strangers).

Building and maintaining social relationships across the lifespan requires the ability to appraise someone as (un)trustworthy and effectively interpret their behavior within social interactions (Fareri, 2019). Research in young adults indicates that learning to trust draws on initial impressions of others that are dynamically updated with experienced patterns of reciprocity, a social reward signal (Chang et al., 2010); this process consistently recruits reward-related neural circuits (e.g., ventral striatum, medial/lateral prefrontal cortex; Bellucci et al., 2016; Fareri et al., 2012; Fouragnan et al., 2013). Moreover, our prior work (Fareri et al., 2015) demonstrates that these circuits differentially encode the value of reciprocity as a function of social closeness: reciprocity from friends relative to strangers elicits enhanced activation of the ventral striatum. This increased value signaling may in turn reinforce already close bonds, and in fact, decisions to trust are strongly related to beliefs about safety (Chen et al., 2021). Yet, how these social learning processes shift as we transition to older adulthood remains an outstanding question.

Critically, trust-based interactions are also predicated on an ability to create and adapt models of others and their intentions. Such theory-of-mind processes engage a network of brain regions including the temporoparietal junction (TPJ), posterior cingulate cortex (PCC), and medial prefrontal cortex (mPFC). Together, these regions comprise the “social brain,” which shares substantial overlap with the default-mode network (DMN; Mars et al., 2012). The DMN shows enhanced reactivity to social relative to non-social outcomes (Fareri et al., 2020) and may serve to prime us for engagement with the social world (Meyer, 2019). One possibility is that functional
alterations within or between the DMN and reward-related networks may underlie the increased positivity and focus on beneficial social engagement in older adults, in turn contributing to an increased risk for exploitation (Castle et al., 2012; Harlé & Sanfey, 2012; Spreng, Shoemaker, et al., 2017).

Here, we sought to systematically investigate age-related differences in the effects of social closeness on trust behavior and the neural representation of reciprocity. In a pre-registered study (https://aspredicted.org/MVZ_ODI), we implemented a variant of a trust game task used in a previous study (Fareri et al., 2015): participants played with a computer, a stranger, and a close friend who accompanied them to the experiment. While older adults may prioritize interactions with close others to meet emotional goals (e.g., maintaining a relationship), evidence also suggests that knowledge-based goals (e.g., social learning) can shift priorities toward interactions with less close/unknown others (Carstensen et al., 2003). Thus, we hypothesized lower rates of trusting trusting friends relative to strangers in older (ages 63-80) relative to younger adults (ages 18-34). We additionally expected blunted striatal responses to reciprocity from friends relative to strangers in older adults. We also hypothesized that age-related differences in striatal responses to reciprocity would be tied to increased DMN-striatal connectivity, which in turn would mediate expected age-related differences in trust behavior; we expected this mediation to be moderated by self-reported social closeness.
Methods

Participants

Fifty participants (26 young adults, ages 18-34; 24 older adults, ages 63-80) were recruited to participate in this study. This sample size was pre-registered and established before data collection. Young adult participants were recruited primarily through the Temple University Psychology Department participant pool and received course credit (along with a task bonus in gift cards) for participation. Older adult participants were recruited using a range of efforts, including at local community and senior centers, newspaper advertisements, and local flyers. All participants were screened before data collection to rule out current major psychiatric or neurologic illness, as well as MRI contraindications. Older adults were screened to rule out dementia using the Telephone Interview for Cognitive Status, with a score under 30 meeting exclusion criteria (Brandt and Folstein, 2012). All participants included in analyses had at least 3 usable runs of the task. These exclusions left a final sample of 48 total participants, with 26 younger adults (mean age: 23.2 years, SD: 4.07 years; 35% male) and 22 older adults (mean age: 69.3 years; SD: 4.38 years; 50% male). All participants gave written informed consent as part of a protocol approved by the Institutional Review Board of Temple University.

Procedures

Participants completed two appointments. During the first appointment, participants underwent a mock MRI scan to help control for motion and acclimate to the scanner. They also completed a brief neuropsychological test battery including measures of estimated premorbid intelligence, specific cognitive domains (e.g., attention, executive functioning, episodic memory, language), and self-reported everyday functioning (older adults only). Participants completed a second appointment with a self-selected friend of the same identified sex and age group (within 5-10 years, who was not a family member or spouse). During the second appointment, participants completed a 45-minute survey that included measures of mood, media usage, and emotion regulation (not presented here) as well as questions about perceived social closeness with their friend and the stranger (Inclusion of Other in Self Scale (IOS); Aron et al., 1992)), and basic demographics.
**Experimental Paradigm**

After completing the survey, participants and their friends were introduced to a sex- and age-matched (within 10 years) stranger (confederate) and were then trained on the Trust Game task (adapted from [Fareri et al. 2015]). MRI participants were told that they would be playing a game called the investment game in real time with their friend, the stranger and a computer partner. On a given trial of the task, the MRI participant would play with one of their three partners as indicated by a photo and name presented on the screen. Participants were instructed that they would start each trial with $8 and that they would have a choice between sending (investing) different proportions of that $8 to their partner on a given trial. The amounts that could be sent varied on a trial to trial basis, ranging from $0-$8. Participants would have up to 3 seconds to indicate via a button press on an MRI compatible response box which of the two investment options they preferred. Participants were instructed that whatever amount they chose to invest would be multiplied by a factor of 3 (i.e., an investment of $6 would become $18 for the partner), and that their partner could decide to split the multiplied amount evenly with them (reciprocate) or keep it all for themselves (defect). Upon entering their response, participants would see a screen that said ‘waiting’, (1.5s) during which time they believed that their decision was being presented to their partner in another room in the research suite. After the waiting screen, a variable ISI was presented (mean = 1.42s), and participants were then notified (2s) whether their partner decided to split that amount evenly with them (reciprocate) or keep (defect) all of the money. Unbeknownst to participants, all outcomes were predetermined, and all partners were preprogrammed to reciprocate 50% of the time as per our previous work (Fareri et al., 2015).

After task training, the MRI participant, the stranger and close friend were split up into different rooms before beginning the actual experimental session. Participants underwent a 90-minute MRI scan that included up to 5 runs of the trust game task, as well as two additional, but separate tasks (not discussed here). Each run of the task consisted of 36 trials, with 12 trials per partner. See Figure 1A and our previous work (Fareri et al., 2015) for task details.

**Behavioral Analyses**

We examined participants’ decisions to trust (i.e., invest) as a function of partner and age by conducting a 3 (Partner) x 2 (Age Group) repeated measures ANCOVA. We also controlled for individual differences in reaction time by z-scoring participants’ mean reaction time during the
decision phase and including these values as a covariate in this analysis. We similarly performed a 3 (Partner) x 2 (Age Group) repeated measures ANOVA on participants’ reaction time during the decision phase.

**Neuroimaging Data Acquisition**

Neuroimaging data were collected at the Temple University Brain Research and Imaging Center (TUBRIC) using a 3.0 Tesla Siemens Prisma scanner equipped with a 20-channel phased array head coil. Functional images sensitive to blood-oxygenation-level-dependent (BOLD) contrast were acquired using a single-shot T2*-weighted echo-planar imaging sequence with slices roughly parallel to the axial plane collected in descending order [repetition time (TR): 2.02 s; echo time (TE): 23 ms; matrix 74 x 74; voxel size: 2.97 x 2.97 x 2.80 mm; 36 slices (15% gap); flip angle: 76°]. To facilitate co-registration and normalization of functional data, we also collected high-resolution T1-weighted structural scans (TR: 2.4 s; TE: 2.2 ms; matrix 192 x 192; voxel size: 1.0 mm³; 192 slices; flip angle: 8°) and B₀ field maps (TR: 645 ms; TE₁: 4.92 ms; TE₂: 7.38 ms; matrix 74 x 74; voxel size: 2.97 x 2.97 x 2.80 mm; 36 slices, with 15% gap; flip angle: 60°). In addition, we also collected T2-weighted structural images (TR: 3.2 s; TE: 567 ms; matrix 192 x 192; voxel size: 1.0 mm³; 192 slices; flip angle: 120°); these images are included with our data on OpenNeuro.org, but we did not use them in our preprocessing or analyses.

**Preprocessing of Neuroimaging Data**

Neuroimaging data were converted to the Brain Imaging Data Structure (BIDS) using HeuDiConv version 0.5.4 (Halchenko et al., 2019). Results included in this manuscript come from preprocessing performed using fMRIPrep 20.1.0 (Esteban, Blair, et al., 2018; Esteban, Markiewicz, et al., 2018), which is based on Nipype 1.4.2 (K. Gorgolewski et al., 2011; K. J. Gorgolewski et al., 2018). The details described below are adapted from the fMRIPrep preprocessing details; extraneous details were omitted for clarity.

**Anatomical data preprocessing**

The T1-weighted image was corrected for intensity non-uniformity (INU) with N4BiasFieldCorrection (Tustison et al., 2010), distributed with ANTs 2.2.0 (Avants et al., 2008),
and used as T1w-reference throughout the workflow. The T1w-reference was then skull-stripped with a Nipype implementation of the antsBrainExtraction.sh workflow (from ANTs), using OASIS30ANTs as target template. Brain tissue segmentation of cerebrospinal fluid (CSF), white-matter (WM) and gray-matter (GM) was performed on the brain-extracted T1w using FAST (FSL 5.0.9, (Zhang et al., 2001)). Volume-based spatial normalization to MNI152NLin2009cAsym standard space was performed through nonlinear registration with antsRegistration (ANTs 2.2.0), using brain-extracted versions of both T1w-reference and the T1w template. To this end, the *ICBM 152 Nonlinear Asymmetrical template version 2009c* (Fonov et al., 2009) template was selected for spatial normalization.

Functional data preprocessing

For each of the BOLD runs contained per subject, the following preprocessing steps were performed. First, a reference volume and its skull-stripped version were generated using a custom methodology of *fmriprep* (Esteban, Markiewicz, et al., 2018). Head-motion parameters with respect to the BOLD reference (transformation matrices, and six corresponding rotation and translation parameters) are estimated before any spatiotemporal filtering using mcflirt (FSL 5.0.9, (Jenkinson et al., 2002)). BOLD runs were slice-time corrected using 3dTshift from AFNI 20160207 (Cox & Hyde, 1997). A B0-nonuniformity map (or fieldmap) was estimated based on a phase-difference map calculated with a dual-echo GRE (gradient-recall echo) sequence, processed with a custom workflow of *SDCflows* inspired by the *epidewarp.fsl* script and further improvements in HCP Pipelines (Glasser et al., 2013). The fieldmap was then co-registered to the target EPI (echo-planar imaging) reference run and converted to a displacements field map (amenable to registration tools such as ANTs) with FSL’s fugue and other *SDCflows* tools. Based on the estimated susceptibility distortion, a corrected EPI (echo-planar imaging) reference was calculated for a more accurate co-registration with the anatomical reference. The BOLD reference was then co-registered to the T1w-reference using FLIRT (FSL 5.0.9, (Jenkinson & Smith, 2001)) with the boundary-based registration (Greve & Fischl, 2009) cost-function. Co-registration was configured with nine degrees of freedom to account for distortions remaining in the BOLD reference. The BOLD time-series (including slice-timing correction when applied) were resampled onto their original, native space by applying a single, composite transform to correct for head-motion and susceptibility distortions. These resampled BOLD time-series will be referred to as *preprocessed BOLD in original space*, or just *preprocessed BOLD*.
The BOLD time-series were resampled into standard space, generating a preprocessed BOLD run in MNI152NLin2009cAsym space.

Additionally, a set of physiological regressors were extracted to allow for component-based noise correction (CompCor, (Behzadi et al., 2007)). Principal components are estimated after high-pass filtering the preprocessed BOLD time-series (using a discrete cosine filter with 128s cut-off) for the two CompCor variants: temporal (tCompCor) and anatomical (aCompCor). tCompCor components are then calculated from the top 5% variable voxels within a mask covering the subcortical regions. This subcortical mask is obtained by heavily eroding the brain mask, which ensures it does not include cortical GM regions. For aCompCor, components are calculated within the intersection of the aforementioned mask and the union of CSF and WM masks calculated in T1w space, after their projection to the native space of each functional run (using the inverse BOLD-to-T1w transformation). Components are also calculated separately within the WM and CSF masks. For each CompCor decomposition, the k components with the largest singular values are retained, such that the retained components' time series are sufficient to explain 50 percent of variance across the nuisance mask (CSF, WM, combined, or temporal). The remaining components are dropped from consideration. As an additional confound, we also estimated framewise displacement (FD). FD was computed using the relative root mean square displacement between affines (Jenkinson et al., 2002).

All resamplings can be performed with a single interpolation step by composing all the pertinent transformations (i.e., head-motion transform matrices, susceptibility distortion correction when available, and co-registrations to anatomical and output spaces). Gridded (volumetric) resamplings were performed using antsApplyTransforms (ANTs), configured with Lanczos interpolation to minimize the smoothing effects of other kernels (Lanczos, 1964).

**Neuroimaging Analyses**

Neuroimaging analyses used FSL version 6.03 (Jenkinson et al., 2012; S. M. Smith et al., 2004). We specifically focused on two types of models (activation and connectivity) to quantify how reciprocated trust and social closeness were associated with BOLD responses. Both types of models were based on a general linear model with local autocorrelation (Woolrich et al., 2001). Our first model focused on the brain activation evoked during the trust task and used a total of nine regressors of interest. We used three regressors to model the decision phase (duration = response time) associated with each of the partners (computer, stranger, friend). To
model brain activation associated with outcomes (reciprocate and defect) by partner, we used six additional regressors (duration = 1 second). Each task-related regressor was convolved with the canonical hemodynamic response function.

Our second type of model focused on the task-dependent connectivity associated with the trust task. To estimate these changes in connectivity, we used psychophysiological interaction (PPI) analysis (Friston et al., 1997; O'Reilly et al., 2012). Notably, recent meta-analytic work has shown that PPI reveals consistent and specific patterns of connectivity across multiple seed regions and psychological contexts (D. V. Smith et al., 2016; D. V. Smith & Delgado, 2017). We first estimated a network PPI model that focused on task-dependent changes in connectivity with the DMN (Fareri et al., 2020; Utevsky et al., 2017). The DMN and nine additional networks were based on prior work (S. M. Smith et al., 2009). Network time courses were extracted with a spatial regression component of the dual regression approach (Filippini et al., 2009; Nickerson et al., 2017) and entered into a model with the nine task regressors from the activation model described above. PPI regressors were formed by multiplying each of the nine task regressors by the DMN regressor, yielding a total of 28 regressors.

We also conducted exploratory seed-based analyses using regions defined from our analyses (see Results). For each participant in these analyses, the average time course of a seed was extracted and entered as a regressor into a model with the nine task regressors from the activation model described above. PPI regressors were formed by multiplying each of the nine task regressors by the seed regressor, yielding a total of 19 regressors in each seed-based PPI model.

Both activation and connectivity models included a common set of confound regressors. We first modeled out missed responses by including an additional task-related regressor to account for the full duration of the choice screen. We also included additional regressors for the six motion parameters (rotations and translations), the first six aCompCor components explaining the most variance, non-steady state volumes, and the framewise displacement (FD) across time. Finally, high-pass filtering (128s cut-off) was achieved using a set of discrete cosine basis functions.

We combined data across runs, for each participant, using a fixed-effects model. Group-level analysis was carried out using FLAME (FMRIB's Local Analysis of Mixed Effects) Stage 1 and Stage 2 (Beckmann et al., 2003; Woolrich et al., 2004). Our group-level model focused on comparisons between older and younger groups; these comparisons included covariates to
account for gender, temporal signal to noise ratio (tSNR), mean framewise displacement, and mean response time. All z-statistic images were thresholded and corrected for multiple comparisons using an initial cluster-forming threshold of \( z > 3.1 \) followed by a whole-brain corrected cluster-extent threshold of \( p < 0.05 \), as determined by Gaussian Random Field Theory (Worsley, 2001).

**Deviations from Pre-registration**

We note that our analyses deviated slightly from those defined in our pre-registration in the following ways. To test the hypothesis that older adults would demonstrate reduced levels of trust relative to younger adults (H1.1), we conducted a repeated measures ANOVA instead of a mixed effects regression in order to be consistent with our previous work (Fareri et al., 2015). In order to examine the relation between self-reported relationship closeness and reward-related neural responses (H2.1), we also included social closeness as a covariate in an offline analysis with extracted parameter estimates, as opposed to at the whole brain level. This was done in order to maximize statistical power and preserve degrees of freedom at the whole-brain level. Given that we did not observe evidence for age-related DMN-striatal connectivity (H2.2; see below), we did not test H2.3.

Additionally, we conducted a number of exploratory analyses based on both previous literature as well as patterns that emerged in the data which complement our pre-registered questions. We conducted a task-based generalized network PPI analysis to examine connectivity of the Executive Control Network (ECN) during the processing of trust game outcomes, based on our previous work (Fareri et al., 2020; Utevsky et al., 2017). Last, we conducted whole-brain seed-based PPI analyses using the fusiform face area (FFA), ventromedial prefrontal cortex (vmPFC) and ventral striatum, TPJ and PCC as seed regions in order to investigate age-related differences in responses to trust game outcomes as a function of social closeness.
Results

Social closeness shapes trust decisions

As a manipulation check, we investigated whether participants exhibited differences in self-reported social closeness to friends and strangers. A 2x2 repeated-measures ANOVA on IOS scores revealed a significant main effect of partner (F(1,41) = 70.40, p < .001), such that participants reported feeling closer to friends (M = 4.50, SD = 1.70) relative to strangers (M = 2.14, SD = 1.52). Contrary to our hypotheses, we observed neither a significant interaction of partner and age group (F(1,41) = 0.10, p = 0.76), nor a main effect of age group (F(1,41) = 2.53, p = .12), though the group difference is trending in the hypothesized direction of blunted effects in older adults (M_{old} = 4.09, SD = 1.82, M_{young} = 4.85, SD = 1.54). We note that due to technical error, a number of participants (n = 5) lacked closeness ratings for the stranger and were unable to be included in this analysis.

Figure 1: Task and Behavior

(A) We used an economic trust game task to measure trust. Participants played the role of the investor with three distinct investees of varying social closeness (friend, stranger, and computer as a nonsocial control). On each trial of the trust task, participants were first shown an image of their partner with a neutral expression (note that images shown here are for illustrative purposes only; the experiment used real pictures of friends and strangers). While viewing their partner, participants were given a choice to invest a greater or lesser amount of money with their partner. Money invested into the partner triples. When money was invested into the investee, the participant was then shown a screen indicating that the investee is contemplating the investment (“Waiting…”).
the outcome of the investee’s decision (i.e., reciprocate and share the money evenly, or defect and take all of the money). (B) We examined whether partner identity impacted the amount invested, which serves as a metric of trust. We found that both older and younger participants invested money as a function of partner.

Our first goal was to examine differences in trust behavior as a function of age and partner. To test pre-registered hypothesis 1.1, we conducted a 3 (partner) x 2 (age group) repeated measures ANOVA on the average amount of money invested during the trust game. Controlling for individual differences in overall reaction time, we observed partial support for pre-registered hypothesis 1.1: a significant main effect of partner on amount of money invested emerged (F(2,90) = 18.49, p < .001; see Figure 1B), such that participants sent more money to their friends relative to both strangers (t(88) = 2.86, p < .01) and computers (t(88) = 5.14, p < .001). This result replicates prior work from our group (Fareri et al., 2015). However, inconsistent with hypothesis 1.1, we did not observe an interaction between partner and age (F(2,90) = 2.16, p = 0.12), though we did note a trending difference between older and younger adults in amount of money invested overall (F(1,45) = 2.77, p = .10), with older adults investing marginally more money in their partners than younger adults. We also conducted an exploratory 3 (partner) x 2 (age group) repeated measures ANOVA on reaction time. This analysis revealed a significant effect of age (F(1,46) = 7.60, p < .01), with older adults exhibiting significantly longer reaction times when making their decisions. A trending main effect of partner also emerged (F(2,92) = 2.75, p = .069), such that participants responded more quickly when interacting with their friends relative to the computer (t(92) = 2.34, p = .064).

**fMRI Results: Trust Game Outcomes**

**Blunted striatal responses to social closeness in older adults**

Based on our expected behavioral results and our prior work, we were also interested in characterizing differences in task-based neural activation and connectivity specifically during experienced outcomes (i.e., reciprocity vs. defection of trust) in the trust game. We first hypothesized that participants would exhibit enhanced reward-related responses within the striatum when experiencing reciprocity from a close friend relative to other partners, and that this effect would be stronger in younger adults and associated with relationship closeness (pre-registered hypothesis 2.1). To test this hypothesis, we conducted a confirmatory whole-brain contrast of reciprocate >
defect. This analysis revealed robust activation of the bilateral ventral striatum (Rt. ventral striatum: x, y, z = 7.5, 13.2, -4.4; Lt. ventral striatum: x, y, z = -10.3, 16.2, -1.2; see Fig. 2A).

We next conducted a 2 (age group) x 3 (partner) repeated-measures ANOVA on extracted parameter estimates from the bilateral ventral striatum to investigate age differences in processing of trust game outcomes. We observed a significant partner x group interaction (F(2,92) = 5.94, p = .004; see Figure 2B): older adults exhibited diminished striatal activation when experiencing reciprocate relative to defect outcomes from friends as compared to strangers (t(46) = -2.68, p = .01), whereas young adults demonstrated a trend toward the opposite pattern (t(46) = 1.89, p = .065), consistent with patterns demonstrated in prior work from our group (Fareri et al., 2015). We note that re-running these analyses including tSNR, mean framewise displacement (FD) and sex as covariates reduced the partner x group interaction effect to a trend (F(2,92) = 2.30, p = .107).

![Figure 2: Blunted Effects of Social Closeness on Striatal Social Reward Responses in Older Adults.](https://identifiers.org/neurovault.image:512088; Unthresholded: CC-BY 4.0 International license available under a Creative Commons Attribution 4.0 International license).

(A) We identified voxels in the striatum whose activation increased for reciprocate outcomes relative to defect outcomes, irrespective of age. The overlaying Z statistic image was thresholded parametrically (Gaussian Random Field Theory) using GRF-theory-based maximum height thresholding (i.e., voxel level threshold) with a (corrected) significance threshold of P=0.05 (Thresholded: https://identifiers.org/neurovault.image:512088; Unthresholded: https://identifiers.org/neurovault.image:512088).
(B) We interrogated this striatal region further to probe for group differences in the effects of partner on reward processing. Although younger adults exhibited an enhanced striatal response to the contrast of reciprocate > defect when the partner was a friend relative to a stranger, older adults exhibited the opposite effect, suggesting that the effects of social closeness on reward related neural responses are blunted in older adults.

Last, we examined whether differences in self-reported social closeness with friends and strangers was associated with differences in the striatal response to trust game outcomes with friends and strangers. A linear regression regressing the difference in striatal BOLD for reciprocate > defect for friends > strangers on the difference in reported social closeness with those same partners revealed a trend of an effect \( (b = 13, SE = 7.86, t = 1.66, p = .104) \), such that the closer participants felt to their friends relative to strangers, the greater the difference in striatal BOLD responses to reciprocity experienced with friends. We note that this is a weak but encouraging effect, and that including tSNR, FD and sex as covariates diminishes the strength of this effect \( (p = .18) \). Taken together these results provide partial support for pre-registered hypothesis 2.1.

**Altered cortical responses to social closeness in older adults**

We also conducted an additional exploratory whole-brain interaction contrast to highlight regions demonstrating age-related differences in responses to trust game outcomes. This analysis revealed significant clusters of activation in the anterior insula, posterior cingulate and both dorsal and ventral medial prefrontal cortex. Extracting parameter estimates from the insula \( (x, y, z = 37, 25, 8) \) revealed a reduced response to defect relative to reciprocate outcomes specifically in the stranger condition in older adults (Fig 3A, B). Older adults also demonstrated a reduced response to reciprocate relative to defect outcomes experienced with friends in the PCC \( (x, y, z = 4, -58, 28; \) Fig 3C, D).
Figure 3: Age-related Differences in Neural Responses to Social Outcomes

(A) We found that the anterior insula exhibited a larger response to defect outcomes relative to reciprocate outcomes in younger adults (plotted here as defect > reciprocate), consistent with the idea that older adults are less sensitive to outcomes associated with losses (Thresholded: https://identifiers.org/neurovault.image:512048; Unthresholded: https://identifiers.org/neurovault.image:512076). (B) For descriptive purposes, we extracted beta values within this anterior insula region and confirmed that older adults showed a blunted response to defect outcomes. (C) In the opposite contrast (reciprocate > defect), we found several regions whose outcome-related activation was enhanced in younger adults (Thresholded: https://identifiers.org/neurovault.image:512037; Unthresholded: https://identifiers.org/neurovault.image:512065). (D) For descriptive purposes, we extracted the response within the posterior cingulate cortex (PCC) and confirmed that the response to reciprocation in this region was enhanced in younger adults. We note that Z statistic images were thresholded parametrically (Gaussian Random Field Theory) using clusters determined by $Z>3.1$ and a (corrected) cluster significance threshold of $P=0.05$. 

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Older adults exhibit enhanced network connectivity during trust game outcomes with friends

Our task-based activation analyses revealed engagement of both reward related circuits and regions comprising the default mode network (e.g., PCC, mPFC) during experiences of reciprocity relative to violations of trust. As outlined in our pre-registration, we predicted that we would observe age-related differences in default mode connectivity, specifically with the striatum, when experiencing reciprocity relative to defection from different partners (H2.2). To test this prediction, we conducted a generalized network psychophysiological interaction (nPPI) analysis during the outcome phase of the task using the DMN as our seed network (cf. Fareri et al., 2020). A contrast of reciprocate > defect as a function of social closeness (friend > computer) and age (young > old) revealed enhanced connectivity between the DMN and TPJ (x, y, z = 64, -40, 24.5; see Fig. 4) when experiencing reciprocity from friends relative to the computer in older, compared to younger adults, though we did not observe enhanced connectivity between the DMN and striatum as predicted in pre-registered hypothesis 2.2. Because of the lack of support for H2.2, we did not pursue our final pre-registered hypothesis (H2.3), which posited DMN-striatal connectivity would mediate partner related effects in trust behavior.

Figure 4: Older Adults Show Enhanced DMN-TPJ Connectivity as a Function of Social Closeness
(A) We used a network psychophysiological interaction (nPPI) analysis with the default mode network (DMN) as a seed. We examined connectivity with DMN as a function of age differences and social closeness (friend > computer) during the outcome phase (reciprocate > defect). This double subtraction analysis [(friend reciprocate > friend defect) > (computer reciprocate > computer defect)] indicated that...
DMN connectivity with the right temporal-parietal junction (TPJ) was enhanced in older adults (Thresholded: https://identifiers.org/neurovault.image:512032; Unthresholded: https://identifiers.org/neurovault.image:512035). (B) For illustrative purposes, we extracted the parameter estimates within this TPJ region. We note that Z statistic images were thresholded parametrically (Gaussian Random Field Theory) using clusters determined by Z>3.1 and a (corrected) cluster significance threshold of P=0.05.

We also explored task-based connectivity of the Executive Control Network (ECN) during the processing of trust game outcomes. This exploratory analysis was motivated by previous work implicating the ECN in reward-processing and goal directed behavior more generally (Fareri et al., 2020; Grill et al., 2021; Waltz et al., 2013). We conducted a network PPI contrast of reciprocate > defect for friends > strangers as a function of age group. Here, we found that relative to younger adults, older adults demonstrated increased connectivity between the ECN and a region encompassing mid/posterior insula (x, y, z = 34, 4, -4) when experiencing defect outcomes with one’s friend, but not with the stranger or computer (see Table 1).

*Young adults demonstrate enhanced vmPFC-hippocampus connectivity during reciprocity*

Finally, we also conducted additional exploratory seed-based PPI analyses investigating group differences in responses to trust game outcomes as a function of social closeness using the following seeds: FFA, vmPFC and ventral striatum, TPJ and PCC. Robust group differences emerged (see the NeuroVault associated with this study (https://identifiers.org/neurovault.collection:10447) for relevant maps), but here we highlight one result of interest from an analysis with the vmPFC as a seed region. This analysis interestingly revealed increased vmPFC-hippocampus (HPC; Right HPC: x, y, z = 22, -14, -20.5; Left HPC: -28, -8, -24) connectivity in younger relative to older adults during reciprocate relative to defect outcomes (see Figure 5).
**Figure 5: Reduced vmPFC-Hippocampal Connectivity in Older Adults.**

(A) In an exploratory seed-based PPI analysis, we examined age-related differences in connectivity with vmPFC. We found that older adults exhibited reduced vmPFC connectivity with a hippocampal (HPC) cluster extending into the amygdala during reciprocate relative to defect outcomes (Thresholded: https://identifiers.org/neurovault.image:512046; Unthresholded: https://identifiers.org/neurovault.image:512074). (B) Interrogation of this hippocampal cluster revealed that the pattern was stable across all partner types. We note that Z statistic images were thresholded parametrically (Gaussian Random Field Theory) using clusters determined by Z>3.1 and a (corrected) cluster significance threshold of P=0.05.
Discussion

Older adulthood is associated with a heightened focus on experiences with close others in conjunction with shrinking social networks, particularly when emotional goals are prioritized (Carstensen, 1992; Isaacowitz et al., 2021). This changing social world thus colors many daily decisions, including trusting a range of individuals with assets, potentially creating opportunities for exploitation and abuse (Spreng et al., 2016). Although decision neuroscience has made significant progress in characterizing how age-related differences in affect and motivation contribute to decision making (Samanez-Larkin & Knutson, 2015), relatively less is known about age-related differences in social decision-making and social outcome processing. Using an iterated trust game in which participants played with three partners who varied in social closeness with the participant (computer, stranger, friend), we show that while younger and older adults demonstrated similar rates of trust with friends and strangers, neural processing of reciprocity and defection varied with respect to age and social closeness. Striatal responses to reciprocated trust from friends relative to strangers and computers were blunted in older adults relative to younger adults, and older adults also showed enhanced DMN connectivity with the rTPJ during reciprocated trust from friends compared to computers. Taken together, these results suggest that older adults demonstrate altered representations of social closeness within financial exchanges involving trust, which may have downstream effects for their ability to adapt their behavior in the long-term.

Altered recruitment of corticostriatal circuits in older adults when processing reciprocity and defection is partially consistent with earlier work looking at social outcome processing in aging. Early investigations noted that relative to younger adults, older adults demonstrated altered recruitment of corticostriatal circuits (e.g., dorsolateral PFC, insula) when faced with unfair offers in an ultimatum game relative to younger adults (Harlé & Sanfey, 2012), and an enhanced anticipatory response in the ventral striatum to social rewards in older relative to younger adults (Rademacher et al., 2014). Findings have also shown reduced striatal activation in older adults when exposed to social outcomes that are inconsistent with initial impressions of others (Suzuki et al., 2019). Here, differences in social value-related signaling in the ventral striatum and the insula in older adults may suggest an inappropriate weighting of social outcomes with different social partners that may have implications for the increased susceptibility to financial exploitation in this population.
Our study also importantly builds on evidence (Hughes et al., 2020; Spreng, Cassidy, et al., 2017; Weissberger et al., 2020) suggesting that older adults at heightened risk for financial exploitation show altered activation and connectivity in regions supporting social cognition. For example, decreased mPFC activation and weaker DMN connectivity has been related to diminished mentalizing about unknown others in older relative to younger adults (Cassidy et al., 2021; Hughes et al., 2019). Here, we report enhanced DMN-TPJ connectivity in older adults when experiencing reciprocity from friends relative to strangers. Coupled with reported stronger ingroup (vs. outgroup) trust bias in older adults (Cassidy et al., 2020), the enhanced DMN-TPJ connectivity may suggest a heightened ability to incorporate social outcomes into models of known close others. We also found reduced vmPFC-HPC connectivity in older adults during experiences of reciprocity, which, when considered in light of evidence of reduced hippocampal involvement in feedback-based learning and memory in older adults (Lighthall et al., 2018), may suggest significant changes in the ways in which social and value-related information during trust-based interactions are integrated to inform social learning in aging samples. Future work may examine how age-related connectivity changes during social outcome processing may relate to neural differences during trust-related decision-making as a function of social closeness.

We note that our results merit further consideration of a few important caveats. First, we observed minimal age-related differences in willingness to trust others as a function of social closeness. Though this could be partly attributed to our small sample size (we were powered to detect moderate to large effects), it is also possible that age-related differences in behavior are more subtle, potentially requiring advanced computational methods to estimate latent factors in the decision-making process (Miletić et al., 2020). An alternative possibility is that because the choices in this task were designed to assess levels of trust in one given individual at a time, there were benefits to placing trust in a stranger, so as to learn about them (Isaacowitz et al., 2021). Future work may implement alternative two-stage designs (Daw et al., 2011) in which older adults have the option to choose to interact on a given trial with a close or unknown other before deciding how much to invest. We also note that other work has reported inconsistencies in age-related differences in trust decisions, and it appears that experimental context may be an important moderator of these effects (Bailey et al., 2015; Frazier et al., 2021). In addition, the effects of social closeness on behavior could be influenced by other aspects of the relationships that we did not assess, including social network size (Kwak et al., 2018) and relationship quality (Santini et al., 2015). Last, age-related differences in brain activation and connectivity could be
linked to confounding variables which we did not assess. We attempted to control for this possibility by including covariates relating to gender, head motion, data quality, and response time. Nevertheless, it is possible that other variables, such as white matter integrity and vascular health (Prins & Scheltens, 2015), may have contributed to the age-related neural differences we observed.

Despite these limitations, our results support the conclusion that older adults exhibit altered neural responses to social closeness during trust-based social interactions. We believe that efforts to characterize age-related differences in these types of decisions may have implications for vulnerability to social victimization and abuse among the elderly, particularly financial exploitation (Lichtenberg, 2016; Lichtenberg et al., 2013; Spreng et al., 2016). Although financial exploitation is a multifaceted issue that can occur in a wide range of scenarios (Beals et al., 2015; Nguyen et al., 2021), many instances of financial exploitation may stem from placing too much trust in others (Nguyen et al., 2021; Shao et al., 2019). We speculate that future efforts to understand how older adults modulate decision making and reward processing as a function of social closeness will lay the groundwork for identifying novel risk factors for financial exploitation.
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