Discrepancies between fluid and crystallized ability in healthy adults: a behavioral marker of preclinical Alzheimer’s disease

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ABSTRACT

Measures of core cognitive processes (fluid abilities) are highly correlated with measures of knowledge (crystallized abilities) in healthy adults. In early stages of Alzheimer’s disease (AD), fluid abilities, however, decline more rapidly than crystallized abilities. We hypothesized that cognitively normal older adults who evidenced lower fluid ability compared with crystallized ability (an ability discrepancy) would show evidence of early AD neuropathology indexed via in vivo measures of amyloid-beta (Aβ) deposition and cortical thickness in AD-vulnerable regions. A sample of older adults (n = 112) aged 65 to 89 underwent a cognitive battery, structural magnetic resonance imaging, and a subset (n = 75) also completed positron emission tomography scanning to measure Aβ deposition using F-18 Florbetapir. Of this sample, 60 older adults (43 with available positron emission tomography scans) evidenced a discrepancy where fluid ability was lower than crystallized ability. The magnitude of the ability discrepancy was independently associated with a greater Aβ deposition and thinner cortex in AD-vulnerable regions, as well as age. The data suggest that such a discrepancy may be a marker of preclinical AD.

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1. Introduction

Detecting Alzheimer’s disease (AD) early is of great importance, especially because potential interventions are likely to be most effective at the earliest stages of the disease (Sperling et al., 2011b). AD is characterized by marked declines in episodic memory and aggregation of plaques composed of amyloid-beta (Aβ), neuronal tangles (composed of tau protein filaments), and cortical atrophy. Both Aβ imaging and autopsy results have yielded evidence that a significant proportion of cognitively normal older adults (approximately 25%–45%) harbor Aβ deposits (e.g., Bennett et al., 2006; Mintun et al., 2006; Pike et al., 2007; Price and Morris, 1999)—a phase characterized as preclinical AD (e.g., Jack et al., 2012, 2013; Sperling et al., 2011a). Studies have increasingly reported that cognitively normal older adults with elevated levels of Aβ show subtle behavioral deficits on cognitively challenging tasks, particularly episodic memory (e.g., Hedden et al., 2013; Resnick et al., 2010). A number of researchers also have reported effects in other domains, including processing speed, executive function, and reasoning (e.g., Rodrigue et al., 2012; Snitz et al., 2013; Wirth et al., 2013a,b). Collectively, these deficits are all in cognitive domains that comprise fluid abilities and, like episodic memory, evidence decline in early stages of AD (e.g., Albert et al., 2011; Bastin and Salmon, 2014; Kaszniak, 1986; Rosen, 1983). In the present study, we used behavioral theories and measures commonly used in studies of cognitive aging as an important predictor of preclinical AD. Specifically, we used a “discrepancy” score based on the subtraction of fluid from crystallized ability and correlated this score with 2 AD brain markers: Aβ deposition and cortical thinning (Bakker et al., 2008; Dickerson et al., 2009; Wong et al., 2010).

Fluid ability represents a set of core processes underlying intelligence that includes processing speed, working memory, and reasoning (e.g., Wechsler, 1944, 1997). These core abilities are drawn on to engage in complex thought, solve problems, and function in everyday life (e.g., Johnson et al., 2004; Kaszniak, 1986; Lezak, 1995; Wechsler, 1944). Fluid ability tasks are typically unfamiliar and do not rely on past experience to perform the required mental operations. In contrast, people do bring accrued knowledge and experience to many situations, and this experiential component of intelligence has been termed “crystallized ability.” Crystalized ability is usually assessed by vocabulary scores (e.g., Ekstrom et al., 1976; Wechsler, 1944; Zachary and Shipley, 1986) or word
pronunciation scores (e.g., the National Adult Reading Test; Blair and Spreen, 1989). (Because vocabulary and word pronunciation measures are highly correlated with one another \( r = 0.78; \) e.g., Blair and Spreen, 1989; O’Carroll and Gilcrease, 1986), we consider them both under the term “crystallized ability.”).) In cognitively normal adults, fluid and crystallized abilities are highly correlated with one another \( r = 0.70; \) Cattell, 1965, see also Kaufman et al., 1996), suggesting that individuals with higher fluid ability also have higher crystallized ability. At the same time, it is well-known that individuals afflicted with AD show declines in fluid ability faster than declines in crystallized ability (e.g., O’Carroll and Gilcrease, 1986; Wechsler, 1944) and that fluid ability decline is predictive of the rate of symptom progression in AD (e.g., Albert et al., 2011; Bastin and Salmon, 2014; McKhann et al., 2011; Schmid et al., 2013; Tabert et al., 2006). Because crystallized ability is spared initially in AD (e.g., O’Carroll and Gilcrease, 1986; Wechsler, 1944), it is considered to be a good measure of premorbid cognitive ability (e.g., Lehrl et al., 1995; Wechsler, 1944; Yates, 1956).

These differential rates of AD-related cognitive decline led us to reason that older adults who had a higher crystallized ability compared with their fluid ability (a positive ability discrepancy score) might harbor AD neuropathology. Consistent with this idea, a greater ability discrepancy has been shown not only to differentiate healthy adults from those with probable AD, but these discrepancies also have been shown to increase as disease severity worsens (e.g., Dierckx et al., 2008; Lezak, 1995; McCarthy et al., 2005). Hence, we hypothesized that larger discrepancy scores between crystallized versus fluid abilities in healthy adults would be associated with individuals in a preclinical stage of AD as evidenced by elevated levels of Aβ and more cortical thinning in AD-sensitive brain regions.

Our primary aim was to use ability discrepancy scores to distinguish between normal and pathologic aging. It is important to note, however, that ability discrepancies also have been observed in older relative to younger adults in cross-sectional analyses (e.g., Eisdrerfer et al., 1959; Rabbit, 1993). These age discrepancies are small in magnitude and could occur because the samples used included adults at the preclinical AD stage. In a similar way, consider that normal aging is characterized by a relatively modest memory deficit compared with young adults (e.g., Hertzog et al., 2003; Park et al., 1996), but AD is characterized by a very large memory deficit (e.g., McKhann et al., 2011). Perhaps differences in ability discrepancy parallel memory deficits in that a small discrepancy characterizes normal older adults, but a large discrepancy is associated with elevated levels of Aβ and more cortical thinning—both signs of preclinical AD. Thus, the modest ability discrepancy effect that occurs in normal older adults does not preclude that large differences in this marker might be diagnostic of AD pathology.

We hypothesized that ability discrepancy might covary with the aforementioned brain markers of preclinical AD. To address this issue, (1) the analyses between ability discrepancy and AD brain markers controlled for age, and (2) we focused the analyses on older adults that had positive ability discrepancy scores while also conducting supplemental analyses to test whether individuals with an exaggerated ability discrepancy were more likely to be associated with AD brain markers relative to other older adults in the sample.

2. Material and methods

2.1. Participants

We drew our sample from 112 adults aged 65—89 who underwent magnetic resonance imaging (MRI) scanning, genotyping, and cognitive testing to measure fluid and crystallized ability as a part of the Dallas Lifespan Brain Study. From these 112, we selected only subjects who had an ability discrepancy score (z-score crystallized minus z-score fluid ability) greater than 0 (n = 60). Subjects with negative scores were excluded in the primary analyses. In fact, they might be associated with nondementia-related impairments due to early deficits in vocabulary ability (e.g., Copet et al., 2010; Satz, 1976). All 60 subjects selected had structural MRI data, and those that had behavioral data collected within 12 months also were invited to undergo positron emission tomography (PET) scanning (43 of the 60; for more PET recruitment details, see Rodrigue et al., 2012).

All subjects were community-dwelling volunteers who were free from neurological conditions, major cardiovascular and psychiatric disorders, and no reported head injury with loss of consciousness >10 minutes. Subjects also were excluded for reported drug/alcohol abuse, and major heart surgery or chemotherapy within 5 years of testing. All subjects were fluent English speakers, right-handed, and provided written informed consent as approved by the university human investigations committee guidelines. A summary of the sample characteristics is provided in Table 1.

2.2. Overall protocol

All subjects had completed 2 neuropsychological assessment visits and 1 visit for MRI scanning. For those subjects with Aβ data, a fourth visit for a PET scan was completed.

2.2.1. Neuropsychological assessment

To assess fluid and crystallized abilities, we selected measures available from the Dallas Lifespan Brain Study that were highly reliable, represented multiple domains of cognition, and are commonly used across many studies (e.g., Johnson et al., 2004; Wechsler, 1997). The 6 fluid measures spanned the cognitive domains of processing speed, working memory, and reasoning. Processing speed was measured using the number of correct items on the Digit Symbol task (Wechsler, 1997) and the Digit Comparison task (Hedden et al., 2002; adapted from Salthouse and Babcock, 1991). Working memory was measured using the number of correct items on the Letter–Number Sequencing task (Wechsler, 1997) and the sum of the number of words in each set on perfectly recalled trials on the Operation Span task (Turner and Engle, 1989). Reasoning was measured using the total score on the ETS Letter Sets test (Ekstrom et al., 1976) and accuracy on the Raven’s Progressive Matrices task (Raven, 1938). Crystallized ability was measured using the number of correct items for both the ETS Advanced Vocabulary Test I–IV (Ekstrom et al., 1976) and the Shipley Vocabulary (Zachary and Shipley, 1986). We submitted all these measures to a factor analysis, confirming that the 6 fluid measures and 2 crystallized measures would generate separate factor loadings (see Section 2.3.1).

2.2.2. Structural MRI acquisition and processing

Subjects underwent MRI scanning on a 3T Philips Achieva scanner equipped with an 8-channel head coil. High-resolution anatomical images were collected using a T1-weighted magnetization prepared rapid gradient echo sequence with 160 sagittal slices, \( 1 \times 1 \times 1 \text{ mm}^3; \) \( 256 \times 256 \times 160 \) matrix, repetition time = 8.18 ms, echo time = 3.76 ms, flip angle = 12°, field of view = 220 mm. To estimate cortical thickness, T1-weighted images were processed using a surface-based processing stream provided by FreeSurfer v5.1 (http://surfer.nmr.mgh.harvard.edu), including bias-field correction, intensity normalization, skull-stripping (Dale et al., 1999; Segonne et al., 2004). Following these steps, FreeSurfer segments of gray/white matter and pial surfaces were used to estimate distance between boundaries,
which were checked for accuracy and edited, if necessary, by extending white matter boundaries or removing nonbrain tissue. The resulting surface maps were then transformed to a common space and smoothed using a 10-mm full width at half maximum (FWHM) Gaussian filter. From these maps, 9 bilateral regions of interest (ROIs) were hand-traced on the cortical surface and averaged across hemispheres and included anterior medial temporal lobe, inferior temporal gyrus, temporal pole, angular gyrus, superior frontal gyrus, superior parietal lobule, supramarginal gyrus, precuneus, and inferior frontal sulcus (see Fig. 1). These ROIs were selected based on prior research identifying them as regions vulnerable to AD, also known as AD signature regions (Bakkour et al., 2009; Dickerson et al., 2009). An additional bilateral region in the occipital cortex served as a control ROI.

2.2.3. PET imaging acquisition and processing
To measure Aβ deposition, subjects underwent PET scanning with 18F-Flurbetapir (Avid Radiopharmaceuticals). Subjects were injected with a 370 MBq (10 mCi) bolus of 18F-Flurbetapir for a 10 minutes emission and a 10 minutes transmission scan. A 2-frame by 5-minute each dynamic emission acquisition protocol was started 50 minutes postinjection. An internal rod source transmission scan was acquired for 7 minutes. The transmission image was reconstructed using back projection and a 6-mm FWHM Gaussian filter. The emission images were processed by iterative reconstruction, 4 iterations and 16 subsets with a 3-mm FWHM ramp filter. Each subject’s PET scan was spatially normalized to a Florbetapir uptake template (2 × 2 × 2 mm³ voxels) using SPM8 and in-house MATLAB scripts and visually inspected for registration quality. Aβ counts were assessed from 8 bilateral ROIs in Montreal Neurological Institute (MNI) space in regions where amyloid typically aggregates in AD patients (Wong et al., 2010). These ROIs included subregions in orbital frontal, dorsolateral prefrontal, anterior cingulate, posterior cingulate, lateral temporal, lateral parietal, and occipital cortex, as well as precuneus. To minimize inclusion of nonspecific white matter binding in these ROIs, we identified the gray/white matter boundary threshold in the PET images in a separate group of young adults (age < 35, n = 9), and eroded each ROI by the resulting binarized white matter mask. Within these final ROIs, counts were extracted, averaged across voxels within each ROI, and then normalized by counts in the gray matter of the cerebellum to produce standardized uptake value ratios.

2.3. Statistical analysis
All statistical analyses were performed using SPSS version 20 (IBM Corp, Armonk, NY, USA). Bias-corrected and accelerated Cs were determined using 1000 bootstrap samples, and significance was set at p < 0.05.

2.3.1. Deriving measures of fluid and crystallized ability
Fluid and crystallized measures were first calculated using factor analysis. Factor analysis was chosen because it is a widely used technique that captures similar variance across measures to help reduce data to a smaller set of measures. Specifically, the 8 neuropsychological measures (2 each for processing speed, working memory, reasoning, and vocabulary) were z-transformed and then entered into a factor analysis using direct oblimin rotation (Kline, 1994). This rotation was chosen because it allows the resulting factors to be correlated with one another, thus better representing the true underlying nature of the data. We intentionally constrained the model to 2 factors because of our a priori interest in fluid and crystallized abilities, but we also inspected the resulting eigenvalues to ensure that 2 factors were the best fit for the data set.
2.3.2. Calculating the ability discrepancy score

Using the results from the factor analysis, we subtracted the fluid score from the crystallized score to create an ability discrepancy score for each subject. Increasingly positive values represented an increasingly greater discrepancy in ability (e.g., Kaufman, 1990; Matarazzo and Herman, 1985; Schretlen et al., 1994). Our primary analyses focused on individuals who had an ability discrepancy score greater than 0.

2.3.3. The effects of ability discrepancy across regions on cortical thickness

Here, we tested the hypothesis that older adults with higher positive discrepancy scores would have lower cortical thickness across the 9 AD signature regions. We conducted a mixed analysis of covariance (ANCOVA) with ability discrepancy score treated as a continuous independent variable, ROIs as a within-subjects independent variable, and their interaction. The dependent measure was cortical thickness. This analysis allowed us to test whether ability discrepancy exerted a consistent effect across all 9 ROIs or whether there was an interaction, suggesting that ability discrepancy exerted a different relationship across the ROIs. Testing this interaction enabled a closer inspection of the robustness of this relationship and could potentially reveal particularly sensitive regions to preclinical AD. In addition, we controlled for age and sex in the analysis to generalize the results to all older age groups (65–90 year olds) and both sexes. These analyses provided input for which ROIs to include as summary scores to simplify the data when testing for independent associations of ability discrepancy and preclinical AD brain markers (see Section 2.3.4).

2.3.4. The effects of ability discrepancy across regions on cortical Aβ deposition

In this analysis, we tested the hypothesis that ability discrepancy scores would exert a main effect on Aβ deposition. We also assessed whether the effects exerted a consistent effect across the 8 regions used from which Aβ standardized uptake value ratios were extracted. We conducted an ANCOVA with ability discrepancy as a continuous variable, ROI as a within-subjects variable, and their interaction. The dependent measure was a continuous measure of Aβ deposition. We also controlled for age and sex. In addition to treating Aβ deposition as a continuous variable, we also used a threshold approach, whereby Aβ was treated as a categorical value and subjects were classified as Aβ+ or Aβ−. Using the threshold value of 1.22 (see Rodrigue et al., 2012), 17 (40%) were classified as Aβ+ and 26 (60%) as Aβ− (out of 43 subjects) for older adults in the positive ability discrepancy group. In the negative discrepancy group, 14 (44%) were classified as Aβ+ and 18 (56%) as Aβ− (out of 32 subjects). These proportions are consistent with previous research suggesting that approximately 25%–45% of cognitively normal older adults harbor Aβ deposits (e.g., Bennett et al., 2006; Mintun et al., 2006; Pike et al., 2007; Price and Morris, 1999). This categorical approach is commonly used (e.g., Dickerson et al., 2009; Landau et al., 2012; Okello et al., 2009; Resnick et al., 2010; Snitz et al., 2013; Wirth et al., 2013a,b) and provided additional reliability of the results.

2.3.5. Independent contributions of cortical thickness and Aβ deposition on ability discrepancy

Because cortical thickness and Aβ deposition are both markers of preclinical AD, they might not provide independent information regarding the status of preclinical AD. On the other hand, models of preclinical AD (e.g., Jack et al., 2012, 2013) propose that Aβ deposition precedes neurodegeneration (e.g., cortical thinning), potentially leading to independent contributions of each measure. To assess whether the 2 types of AD brain markers independently affected ability discrepancy, we also conducted a multiple regression analysis. We treated cortical thickness and Aβ deposition as predictor variables and ability discrepancy as the outcome variable. ROIs that were significant in the ANCOVAs described previously were averaged separately to derive a summary measure of cortical thickness and Aβ deposition. As in the previous analyses, age and sex were included as covariates. To maximize degrees of freedom, backward regression was used to eliminate insignificant higher-order factors.

In addition to the primary regression analysis, a supplemental regression analysis was conducted on the full sample with group (positive and negative discrepancy) as a between-subjects factor. An interaction with group and either of the AD brain markers would provide evidence that any of the relationships with the AD brain markers depended on which end of the ability discrepancy scale older adults fell.

3. Results

3.1. Factor analysis of neuropsychological assessment

The factor analysis yielded 2 factors that, as expected, clearly differentiated crystallized from fluid abilities. For the first factor, the zero-order factor loadings (i.e., structure coefficients) were largest for Shipley vocabulary (0.91) and ETS advanced vocabulary (0.84), thus representing crystallized ability. This factor explained 32.20% of the variance in the data. For the second factor, the factor loadings, in order of magnitude, were Digit Symbol (0.74), Digit Comparison (0.69), ETS Letter Sets (0.65), Letter–Number Sequencing (0.46), Raven’s (0.45), and Operation Span (0.45). The second factor was designated as the fluid ability factor and explained an additional 16.57% of the variance. The other eigenvalues were less than 1.0 (the third eigenvalue had a value of 0.88) and explained little variance beyond the first 2 factors. Consistent with prior work (e.g., Cattell, 1965; Kaufman et al., 1996), the crystallized and fluid ability factors were positively correlated with one another, r(112) = 0.47, p < 0.001, CI (0.32, 0.61).

3.2. Calculating ability discrepancy scores and partitioning groups

Subtracting fluid from crystallized ability resulted in an ability discrepancy score for each subject that ranged from −2.97 to 2.32 (M = −0.01). Sixty of the 112 subjects had an ability discrepancy score greater than 0 as well as MRI scans and hence were retained for the primary analyses pertaining to cortical thickness. Of these 60 subjects, 43 subjects were retained for the Aβ analysis as they had completed both MRI and PET scans. A comparison of the positive and negative discrepancy groups indicated that the 2 groups did not differ in chronological age, mini mental state exam (MMSE), cortical thickness averaged across the 9 ROIs, or Aβ averaged across the 8 ROIs (all p's > 0.26). However, the positive group had higher levels of education (M = 16.82 vs. 15.57; t[110] = 2.75, p = 0.007) and relatively more men than women (48% male/52% females vs. 23% male/77% female; χ²(1) = 7.66, p = 0.006) compared with the negative group. In the negative group, mean cortical thickness was not related to mean cortical Aβ (r[28] = 0.02, p = 0.94). We next assessed the same associations within the positive group. No relationship was observed between mean cortical thickness and mean cortical Aβ (r[39] = −0.23, p = 0.15). Among age, sex, education, and MMSE, only age significantly correlated with ability discrepancy (r [60] = 0.26, p = 0.05; for the other variables all p’s > 0.49).

3.3. Effects of ability discrepancy on cortical thickness

We conducted an ability discrepancy × ROI ANCOVA on cortical thickness. This ANCOVA yielded a main effect of ability discrepancy and...
of ROI, but no interaction. The ability discrepancy effect was significant $\left( F[1, 56] = 7.54, p = 0.008, \eta^2_p = 0.12 \right)$; subjects with a greater ability discrepancy had lower cortical thickness. The accompanying parameter estimates for the main effect of ROI $\left( F[8, 448] = 5.49, p < 0.001, \eta^2_p = 0.09 \right)$ indicated that temporal regions were thicker than frontal and parietal regions $\left( t[59] = 16.42, p < 0.001, d = 2.36 \right)$ and $\left( t[59] = 26.64, p < 0.001, d = 3.77 \right)$, respectively, and frontal regions were thicker than parietal regions $\left( t[59] = 15.67, p < 0.001, d = 1.63 \right)$. However, the absence of a significant interaction $\left( F[8, 448] = 1.15, p = 0.33, \eta^2_p = 0.02 \right)$ suggested that the ability discrepancy effect generalized across regions. (Controlling for APOE status and education did not change the significance of the results nor was APOE status or education a significant factor $\left[ \text{all } p's > 0.05 \right]$.) Because we had a modest age range $\left[ 65-89 \right]$, we also tested whether age interacted with ability discrepancy, but no interaction with age was significant $\left[ \text{all } p's > 0.46 \right]$.

Thus, the cortical thickness values in each of the 9 AD signature regions were averaged together to form a composite cortical thickness score for subsequent analyses. Cortical thickness as a function of ability discrepancy is displayed in Fig. 2 (top left). (One subject did have a mean cortical thickness value about 4 standard deviations below the group mean thickness level $\left[ \text{subject mean thickness} = 2.00 \text{ mm}, \text{ group mean thickness} = 2.70 \text{ mm} \right]$.) Visual inspection of the FreeSurfer segmentation did not reveal clear errors in the segmentation but rather verified substantial atrophy throughout the cortex. Nevertheless, the previously mentioned analyses were re-run excluding this subject with nearly identical results, suggesting that this person did not inflate the relationships between ability discrepancy and cortical thickness. This subject also did not undergo PET scanning and so was not included in the subsequent PET analyses. Notably, thickness in the control region (primary visual cortex) was not associated with ability discrepancy $\left( p = 0.38 \right)$.

### 3.4. Effects of ability discrepancy on cortical Aβ deposition

We conducted an ability discrepancy × ROI ANCOVA on Aβ deposition. This ANCOVA yielded a main effect of ability discrepancy $\left( F[1, 39] = 6.40, p = 0.016, \eta^2_p = 0.14 \right)$, which was significant because, as predicted, subjects with a higher ability discrepancy score had more Aβ deposition. The main effect of ROI was not significant $\left( p > 0.38 \right)$. However, an ability discrepancy × ROI interaction was significant $\left( F[7, 273] = 4.38, p < 0.001, \eta^2_p = 0.10 \right)$, suggesting that the relationship between ability discrepancy and Aβ deposition differed across the ROIs. The parameter estimates from the ANCOVA indicated that the ability discrepancy effect was observed across only 4 of the 8 ROIs: precuneus, temporal cortex, posterior cingulate, and anterior cingulate (Table 2). Marginal significance was observed for dorsolateral prefrontal cortex $\left( p = 0.053 \right)$, parietal $\left( p = 0.077 \right)$, and orbitofrontal cortex $\left( p = 0.10 \right)$, and nonsignificance was observed for occipital cortex $\left( p > 0.16 \right)$. (Controlling for APOE status did not change the significance of the results nor was APOE status a significant factor $\left[ p = 0.64 \right]$.) Controlling for education changed the significance between ability discrepancy and Aβ in the anterior cingulate from $p = 0.044$ to $p = 0.058$, although education itself was not significant $\left[ p > 0.05 \right]$. Ability discrepancy did not interact with age $\left[ \text{all } p's > 0.50 \right]$.

Because of these observed regional differences, Aβ values were averaged across only the 4 significant regions to form a composite cortical Aβ score for subsequent analysis. Fig. 2 (top right) displays the aggregated values for Aβ as a function of ability discrepancy. Using a threshold approach, the Aβ+ group had larger ability discrepancy scores than the Aβ- group $\left[ M_{Aβ+} = 0.91 \text{ and } M_{Aβ-} = 0.54; t[41] = 2.16, \text{ standard error of the mean} = 0.18, p = 0.037 \right]$. When including subjects in the full sample of 112 individuals,
ability discrepancy was numerically larger in the Aβ+ than Aβ− group (M_{Aβ+} = 0.30 and M_{Aβ−} = −0.089) but did not reach significance (t(73) = 1.78, standard error of the mean = 0.22, p = 0.078).

3.5. Independent contributions of cortical thickness and Aβ on ability discrepancy

Using backward regression, we entered chronological age, sex, mean cortical thickness, mean cortical Aβ, and mean cortical thickness × mean cortical Aβ interaction to predict ability discrepancy. As shown in Table 3, the final regression model was significant (F[4, 38] = 5.74, mean squared error = 0.24, p = 0.001) and explained 37.7% of the variance in ability discrepancy. Mean cortical thickness (p = 0.05) and mean cortical Aβ (p = 0.03) were each significantly related to ability discrepancy and accounted for 6.5% and 8.1% unique variance in the model, respectively. The interaction term (Thickness × Aβ) was not significant (p = 0.57) and was removed from the model. Age (p = 0.02) also accounted for unique variance (9.7%) but sex did not (p = 0.83). (Adding APOE status in the model did not change the significance of the results nor was APOE status a significant factor [p > 0.05].)

4. Discussion

The present study sought to assess whether a positive difference between crystallized and fluid ability in cognitively normal adults was associated with 2 brain markers of preclinical AD—cortical thinning and Aβ deposition. Results indicated that (1) decreased thickness and greater Aβ deposition in AD-related regions were associated with a larger discrepancy score; (2) both cortical thickness and Aβ deposition were significant and independent factors associated with a greater positive ability discrepancy score; (3) these relationships were significant in both positive and negative discrepancy groups for Aβ deposition, but only in the positive discrepancy group for cortical thickness; and (4) age was independently associated with ability discrepancy after accounting for cortical thickness and Aβ deposition.

The cortical thickness regions selected as ROIs have previously been found to be specific to mild AD in 4 independent samples by Dickerson and colleagues (2009). In Dickerson et al. (2009), cortical thickness was reduced in AD patients compared with cognitively normal controls and was associated with the severity of cognitive symptoms in the AD patients. Moreover, cortical thickness in these regions also was negatively associated with an in vivo measure of Aβ deposition in cognitively normal older adults (e.g., Bakkour et al., 2009; Dickerson et al., 2009). Our finding that cortical thinning in these same regions was associated with higher ability discrepancy scores was only found in older adults with more exaggerated discrepancies in ability (i.e., the positive ability discrepancy group), suggesting that this marker is sensitive in later stages of preclinical AD, when neurodegeneration has already begun. We also report that, consistent with Dickerson et al. (2009), the relationship between ability discrepancy and thickness did not occur in primary visual cortex, providing specificity of the thickness effects to AD-sensitive regions. Although recent models of AD focus on hippocampal volume as the primary structural measure that declines in the cascade of events towards AD (e.g., Jack et al., 2012, 2013), the present findings support the notion that cortical thinning in these AD-related regions might be included as a marker of preclinical AD as well (e.g., Bakkour et al., 2009; Dickerson et al., 2009).

Next, we turn to the findings regarding Aβ deposition. The presence of Aβ at autopsy is 1 requirement for a diagnosis of AD (McKhann et al., 2011). There is growing evidence that Aβ deposition in cognitively normal adults is associated with subtle fluid processing decline (e.g., Hedden et al., 2013; Rodriguez et al., 2012; Snitz et al., 2013; Wirth et al., 2013a,b) and is later related to the progression to mild cognitive impairment (e.g., Blasko et al., 2008; Okello et al., 2009), resulting in the categorization of such individuals as having preclinical AD. In the present study, we report that a greater magnitude of ability discrepancy was consistently associated with more Aβ deposition. We found this relationship both in the positive and negative ability discrepancy groups, suggesting this marker is sensitive to both early and later stages of preclinical AD.

This Aβ effect on ability discrepancy was specific to regions associated with AD such as the precuneus (e.g., Braak and Braak, 1991) but was absent in the occipital cortex, a region that shows little accumulation of Aβ in the earliest stages of AD (Rodrigue, 2012; Vlassenko et al., 2012). This specificity nicely dovetails with the cortical thickness findings that also showed the same
dissociation with the occipital cortex. The convergence of the cortical thickness and Aβ deposition findings strengthen evidence for the 2 markers together serving as a behavioral index of preclinical AD.

4.1. Ability discrepancy scores: normal aging or neuropathology?

The results suggest that an ability discrepancy score might provide unique information, in the absence of expensive imaging data, regarding the increased likelihood that an individual is harboring Aβ and is already exhibiting signs of neurodegeneration. In the context of this conclusion, it is important to consider whether ability discrepancy is a specific indicator of neuropathology or merely a general characteristic of normal aging. There is some evidence that ability discrepancy increases with age (e.g., Eisdorfer et al., 1959; Rabbit, 1993). It is well-recognized that cross-sectional studies of aging typically yield a decrease in fluid abilities from young adulthood into old age, whereas crystallized ability remains stable or even increases with age (e.g., Baltes, 1987; Park et al., 1996). Recently, an impressive study sampling over 48,000 adults found that performance on fluid measures such as processing speed peaked as early as late teens/early 20’s, whereas crystallized abilities peaked later in the 50’s (Hartsbome and Germine, 2015). These different peaks in performance, on the surface, suggest that ability discrepancies are characteristic of normal aging and may not confer important information regarding early stages of pathology.

However, there are 2 important reasons that the present findings support the use of a greater discrepancy score as a potential indicator of preclinical AD. First, many subjects in presumably “normal” samples will have latent neuropathology, including Aβ deposition, and are on an initial trajectory of pathological decline (cf. Sliwinski et al., 1996). In such cases, those individuals might actually drive the discrepancies between fluid and crystallized abilities found in other studies, rather than the cognitively normal adults without latent neuropathology. Second, even if such discrepancies do exist in normal aging, the present results show that the magnitude of ability discrepancy might separate normal from pathological aging. Just as a moderate memory decline occurs almost universally with advanced age (e.g., Hertzog et al., 2003; Park et al., 1996) and a precipitous memory decline is diagnostic of AD, the same appears to be true for an ability discrepancy: a moderate ability discrepancy occurs with advanced age, but a precipitous increase in discrepancy is diagnostic of latent pathology and early neurodegeneration. The fact that we found the effects of Aβ deposition and cortical thickness on ability discrepancy after chronological age was statistically controlled further bolsters this idea.

We are unaware of any studies that have related AD risk factors to ability discrepancy, but the extant literature does show that patients with AD evidence greater ability discrepancies than cognitively normal adults, and these discrepancies increase as disease severity worsens (e.g., Dierckx et al., 2008; Lezak, 1995; McCarthy et al., 2005). In addition, longitudinal evidence has shown that the combination of increases in Aβ and greater cortical thinning is associated with steeper declines in cognition (e.g., Dickerson and Wolk, 2012; Landau et al., 2012; Resnick et al., 2010) and greater likelihood to convert to AD (e.g., Dickerson et al., 2011; Julkunen et al., 2010; Landau et al., 2012; Okello et al., 2009; Querbes et al., 2009). Together, these sources of information converge on the conclusion that a large ability discrepancy is, at the least, an additional signal to researchers and clinicians that an individual harbors AD brain markers and therefore might be in the preclinical stages of AD. In the presence of cognitive complaints or other symptoms, further investigation into an individual’s condition is warranted.

Using an ability discrepancy score has an important advantage over other measures of cognition such as memory. Because crystallized ability often is used as a premorbid measure of ability, the difference between fluid and crystallized ability could serve as a proxy for longitudinal decline in a cross-sectional sample. Thus, unlike cross-sectional measures of memory, intraindividual variability is not confounded with longitudinal decline. As more longitudinal data become available, we would predict that older adults with an exaggerated ability discrepancy would exhibit faster rates of cognitive decline and eventual clinical symptomology of AD. However, until these predictions can be confirmed, it is premature to diagnose a cognitively normal individual with a high ability discrepancy score as having preclinical AD, but it is more reasonable to consider a high discrepancy score as evidence for increased risk that an individual harbors Aβ deposition and cortical thinning in AD signature regions.

5. Conclusion

One of the most pressing questions in aging research is determining whether some older adults are on a pathological aging trajectory toward dementia or on a normal aging trajectory. The present study uses theoretical models of cognitive aging to direct attention toward discrepancies between fluid and crystallized abilities as a significant risk factor that has not been considered in the recent past. These discrepancies might be important in predicting who is more likely to be in the preclinical stages of AD in a cross-sectional sample.

Disclosure statement

The authors declare no competing financial interests or any conflicts to disclose.

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