Long-term prognosis and educational determinants of brain network decline in older adult individuals

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Older adults with lower education are at greater risk for dementia. It is unclear which brain changes lead to these outcomes. Longitudinal imaging-based measures of brain structure and function were examined in adult individuals (baseline age, 45–86 years; two to five visits per participant over 1–9 years). College degree completion differentiates individual-based and neighborhood-based measures of socioeconomic status and disadvantage. Older adults (~65 years and over) without a college degree exhibit a pattern of declining large-scale functional brain network organization (resting-state system segregation) that is less evident in their college-educated peers. Declining brain system segmentation predicts impending changes in dementia severity, measured up to 10 years past the last scan date. The prognostic value of brain network change is independent of Alzheimer’s disease (AD)-related genetic risk (APOE status), the presence of AD-associated pathology (cerebrospinal fluid phosphorylated tau, cortical amyloid) and cortical thinning. These results demonstrate that the trajectory of an individual’s brain network organization varies in relation to their educational attainment and, more broadly, is a unique indicator of individual brain health during older age.

Educational attainment differences are closely linked to health disparities across individuals (for example, ref. ¹). Adults with higher education live longer and healthier lives than their peers with less education. Conversely, lower education is associated with increased risk of mental health disorders and dementia during advanced age. Education-related differences in health outcomes during older age are likely mediated by a complex combination of socioeconomic factors that are realized via the opportunities that higher education affords over an individual’s adulthood. These factors include access to resources and environmental stimulation, health habits and exposure to different levels and types of stress (for example, ref. ¹ reviewed in ref. ¹). Critically, however, efforts to link an individual’s education and environment to their brain changes, including both brain structure (for example, refs. ¹¹; also see ref. ¹² for review) and measures of brain pathology (for example, refs. ¹³–¹⁵), have yielded mixed results. Establishing a link between educational attainment and specific brain changes during older age is not only an important step toward understanding environmental determinants of brain disease but could also catalyze discovery and incorporation of new brain health ‘biomarkers’ (for example, ref. ¹⁷).

Due to its devastating threat to older adults and to public health systems, there is an urgent need to elucidate the causes of AD. Except in rarer forms of the disease (for example, autosomal-dominant AD), there are no known direct determinants of AD, indicating an interaction between genetic risk and various environmental, psychosocial and lifestyle factors. AD brains are characterized by the presence of two types of pathology: abnormal levels of extracellular beta amyloid (Aβ) plaques and intracellular tau proteins in the form of neurofibrillary tangles. A recently proposed framework incorporates biomarkers sensitive to both of these neuropathologies (measured using positron emission tomography (PET) or from cerebrospinal fluid (CSF)) as well as measures of neurodegeneration (for example, brain atrophy, assessed by gray matter cortical thinning and volume loss, and hypometabolism, assessed by fluorodeoxyglucose uptake) to help classify and stage AD (that is, the amyloid, tau and neurodegeneration (A/T/N) framework). There has been substantial progress in understanding the trajectory of these AD-related biomarkers (for example, refs. ¹⁷–¹⁹) and their potential relationships with impending cognitive decline (for example, refs. ¹⁹–²⁰). However, it has also been clear that individuals with comparable biomarker profiles may still have different clinical profiles, suggesting that other moderators exist that have yet to be accounted for, which could also be more broadly informative in understanding trajectories of brain aging.

Progress in incorporating measures of brain function into models of aging, AD and dementia more generally has been slow. This is largely due to inherent constraints associated with characterizing brain signals in older and cognitively impaired populations with task-related functional imaging (that is, challenges with participant compliance and feasibility), but is also due to the complexities of accounting for brain variability associated with differing behavioral performance. However, brain function is also reflected in the correlation structure of brain region signals in the absence of overt task performance (that is, during the ‘resting state’ (ref. ²⁰)). Resting-state functional correlations (RSFCs) represent ‘Hebbian-like’ statistical histories of coactivation between areas of the brain. When sets of RSFC signals sampled across multiple brain areas are examined in aggregate, complex large-scale brain network organization is evident (review in ref. ²⁰). While RSFC networks remain relatively stable on a day-to-day basis and with variations of state...
(for example, ref. 39), they have been shown to differ across the lifespan (for example, refs. 32–34), reflecting protracted periods of changes in brain function that accompany childhood development and adult aging. Importantly, RSFC networks vary in relation to cognitive ability among healthy adults (for example, refs. 35,36) and also differ based on disease status (for example, refs. 37,38). The present work is motivated by the hypothesis that an individual’s large-scale brain network organization reflects their brain’s functional integrity and that functional brain network degradation may be prognostic of cognitive impairment beyond global measures of brain atrophy and pathological burden.

Large-scale RSFC networks are organized into a modular architecture (for example, ref. 39); modules correspond to functionally specialized brain systems, and the segregation of these systems supports brain function30,31. Multiple reports have now demonstrated that increasing adult age is associated with less-segregated brain networks30,31,34,42. This ‘dedifferentiation’ of functional systems has been associated with age-accompanied differences in patterns of brain activity, worse cognitive and motor ability, lower energy metabolism and altered neurotransmitter levels (for example, refs. 40–42). Cross-sectional comparisons have also revealed that middle-aged adults with lower socioeconomic status (SES) (35–64 years of age) exhibit lower brain system segregation than that of their peers of the same age with higher SES35, suggesting that there may exist environmental determinants of brain network aging. Finally, comparisons of patients with AD relative to age-matched healthier adults have provided some evidence that patterns of reduced segregation exist in the brain’s functional systems among individuals with dementia (for example, refs. 5,50). This collective work motivates the current study, which aims to determine (1) whether a link exists between an individual’s educational attainment and longitudinal changes in their brain system segregation during adulthood and (2) whether changes in brain network organization are prognostic of impending clinical decline in aging adult individuals.

To answer these questions, we have assembled and examined data from a diverse participant cohort of varying adult ages across multiple longitudinal magnetic resonance imaging (MRI) sessions and clinical visits (including clinical visits that were conducted up to 10 years after the participant’s last available MRI session). The richness of the dataset allows us to examine the relationship between educational attainment and changes in brain system segregation in individual adult participants over time while accounting for demographics and various measures of health and pathology. These brain network changes are then evaluated in relation to trajectories of clinical decline to examine the prognostic utility of changes in functional brain network organization over and above an individual’s AD-related genetic risk and presence of AD-related brain pathology.

Results

Longitudinal changes in brain system segregation were examined in predominantly cognitively normal middle-aged and older adult participants (n = 265; 45–86 years of age at baseline) enrolled in ongoing studies of normal aging and dementia at the Charles F. and Joanne Knight Alzheimer Disease Research Center (Knight ADRC) at Washington University in St. Louis. Each participant had two to five visits during which MRI was conducted, which included resting-state scans, collected over 323–3,372 d (0.88–9.24 years). Over 95% of longitudinal sessions were collected >1 year after previous sessions; all available sessions were included to maximize accuracy of individual change estimates (Fig. 1a). While a large portion of participants in the final sample came from the city of St. Louis and areas in the immediately surrounding St. Louis County, a sizable portion of participants came from other zip codes within the greater St. Louis metropolitan statistical area, which altogether encompassed reasonable geographical diversity (Fig. 1b).

College degree attainment differentiates individual-based and neighborhood-based measures of socioeconomic advantages. For each individual, educational attainment was considered categorically. The categorical distinction of having a college degree or not has been shown to confer significant health14 and socioeconomic advantages41 even after accounting for the economic costs to obtain the degree. Consistent with this, examination of the present study’s participant characteristics confirmed the relation between education and other socioeconomic outcomes. For participants for whom relevant information was available, a college education (‘college+’ versus ‘below college’) was associated with higher occupation-based socioeconomic index (SEI)34 (t = −10.791, P < 0.0001, CI95% = (−20.122, −13.888)) (Fig. 1c; CI95% of mean difference is reported), living in neighborhoods that had a higher median household income (t = −3.455, P < 0.001, CI95% = (−24,169.742, −6,591.351)) (Fig. 1d; after removing the ten participants that resided in two zip codes with a median household income above $150,000 per year (Ladue, MO and Chesterfield, MO), the estimated median household income remained marginally higher for ‘college+’ participants than that for ‘below college’ participants (t = −1.813, P = 0.072, CI95% = (−14,283.871, 618.663)); after outliers were removed, ‘below college’ mean = $79,620.19 and ‘college+’ mean = $86,452.80) and lower scores on an index that combines multiple environmental variables to characterize area deprivation (area deprivation index (ADI))34 (t = 3.197, P = 0.002, CI95% = (3.417, 14.443)) (a single statistical outlier was present in the ADI data, but removing it did not yield a qualitative difference in the statistical comparison; Fig. 1e).

Participants’ ages at their baseline scan session did not significantly differ between education groups; there was a higher proportion of females in the ‘below college’ group than the ‘college+’ group. Further, with the exception of mild depressive symptoms (measured by the Geriatric Depression Scale (GDS)), variables related to clinical status, AD-related pathology, cardiovascular health and history of traumatic brain injury (TBI) did not differ across education groups. We note that the participant sample was constructed from studies collected at the Knight ADRC, which typically have a higher proportion of participants with the APOE ε4 genotype than the general population; however, the proportion of participants positive for the APOE ε4 genotype did not significantly differ across education groups (see Table 1 for statistical descriptions and comparisons).

Older adults with less education exhibit greater declines in brain system segregation. Following rigorous quality control and data-cleaning procedures (Methods), each participant’s resting-state functional brain network (brain network graph) was constructed using a surface-based node set (brain area parcellation15), with each node labeled according to its functional system assignment30 (Fig. 2a). Generally, resting-state correlations between functional areas from the same systems are higher than those of different systems, reflecting a modular network organization (Fig. 2b). This modular organization promotes the functional specialization of distinct systems31, which can be effectively summarized using a measure of brain system segregation31.

Changes in brain system segregation were examined as a function of participants’ educational attainment and age, with follow-up analyses examining whether this relationship was independent of other health indicators that have established and hypothesized associations with education and brain function. A linear mixed-effects model revealed a significant main effect of education group (F = 5.073, P = 0.025, CI95% = (−1.973, −0.149); CI95% of the linear mixed-model term estimate is reported) and age at baseline (F = 7.029, P = 0.008, CI95% = (−0.032, −0.005)); participants with a college degree exhibited a higher level of brain system segregation, and older age was associated with lower brain system segregation. The statistical test also revealed significant interactions.
between education group and age at baseline ($F_{1,136}=4.949$, $P=0.027$, CI$_{95\%}$ = (0.002, 0.029)) and between time (that is, normalized from baseline) and education group ($F_{1,136}=5.756$, $P=0.017$, CI$_{95\%}$ = (0.139, 1.368)). Most importantly, the model revealed a significant three-way interaction between time, education group and age at baseline ($F_{1,136}=6.814$, $P=0.010$, CI$_{95\%}$ = (−0.021, −0.003)) after accounting for the effect of self-reported sex and in-scanner head motion on brain system segregation. The nature of this interaction can be appreciated in Fig. 3a; older adults (more than ~65 years of age) without a college degree exhibited reliable and declining brain system segregation over time, which was not uniformly evident in older adults with a college degree. These observations were reinforced in simple slope (Fig. 3b) and Johnson–Neyman (Fig. 3c) analyses. In the Johnson–Neyman analysis, it was evident that the predicted slopes in ‘below college’ adults became negative in older age, but the predicted slopes in ‘college+ adults’ remained relatively close to zero (that is, a flatter slope). In keeping with this, while the model indicated positive predicted slopes for some individuals in both education groups (for example, younger age for ‘below college’ and older age for ‘college+’), these positive slopes were relatively weaker and not statistically significant, as evidenced by the fact that the confidence intervals for these portions of participants included zero (shaded intervals in Fig. 3c).

Although the proportion of minority (non-white) participants in the present sample is limited ($n=26$, ~10%), we reanalyzed the model while accounting for participant self-reported race. The three-way interaction between time, education group and age at baseline remained significant for predicting brain system segregation ($F_{1,136}=6.399$, $P=0.012$, CI$_{95\%}$ = (−0.073, 0.021), CI$_{95\%}$ = (−0.002, 0.002)). Further, while the proportion of female participants across education groups differed (Table 2), including sex as an interaction term revealed no significant four-way interaction between time, education group, age at baseline and sex ($F_{1,136}=0.073$, $P=0.788$, CI$_{95\%}$ = (−0.014, 0.010)).
Education-related brain network decline is independent of clinical status, AD-related genetic risk and pathology, and general measures of health. The participants in the present dataset were recruited in research studies targeting populations with higher AD risk (for example, based on their age, family history or clinical status). Given that lower educational attainment is associated with greater risk of dementia in older age, it is important to determine whether the observed brain network changes are linked to baseline differences in clinical status and/or subclinical AD-related pathology but also other available measures of health more broadly. Multiple statistical models relating educational attainment to brain system-segregation changes were conducted while accounting for individual variability in measures related to baseline clinical status (clinical dementia rating (CDR); see section 1.1 in the Supplementary Information for an analysis excluding a small subset of individuals (n = 25) with a CDR of 0.5 at baseline, indicating very mild dementia), AD genetic risk (APOE status), baseline AD-related pathology (a categorical measure based on the presence of elevated CSF phosphorylated tau (CSF pTau) and/or elevated cortical amyloid levels measured using Pittsburgh compound B (Pib) PET), baseline cardiovascular health (an aggregate measure including body mass index (BMI), incidents of hypertension, hypercholesterolemia and other cardiovascular incidents), baseline depressive symptoms (as measured by the GDS) and history of traumatic brain injury. The results of statistical models incorporating these measures are summarized in Table 2. The observed relationships between educational attainment and changes in brain system segregation largely persisted after accounting for the various measures individually or in aggregate.

### Table 1 | Demographic, health and AD-related information

| Variables | Below college (n = 92) | College+ (n = 173) | Total (n = 265) | P value |
|-----------|------------------------|---------------------|-----------------|---------|
| **Demographic** | | | | |
| Age, mean, years (s.d.) | 68.07 (8.41) | 66.45 (9.66) | 67.01 (9.26) | 0.176 |
| Sex, n female (%) | 65 (70.7%) | 89 (51.4%) | 154 (58.1%) | 0.004 |
| Race, n white (%) | 82 (89.1%) | 157 (90.8%) | 239 (90.2%) | 0.837 |
| Education, mean, years (s.d.) | 12.77 (1.39) | 17.43 (1.43) | 15.82 (2.63) | <0.001 |
| **Cardiovascular health, neurological health and mental health** | | | | |
| BMI, mean, kg/m² (s.d.) | 27.47 (4.55) | 26.51 (4.48) | 26.85 (4.52) | 0.102 |
| Recent or remote hypertension, n (%) | 36 (39.1%) | 73 (42.2%) | 109 (41.1%) | 0.725 |
| Recent or remote hypercholesterolemia, n (%) | 44 (47.8%) | 65 (37.8%) | 109 (41.3%) | 0.148 |
| Recent or remote cardiovascular incidents, n (%) | 12 (13.0%) | 17 (9.9%) | 29 (11.0%) | 0.576 |
| Recent or remote traumatic brain injury, n (%) | 6 (6.7%) | 13 (7.8%) | 19 (7.5%) | 0.948 |
| GDS, mean score (s.d.) | 1.36 (1.56) | 0.91 (1.28) | 1.06 (1.40) | 0.012 |
| **Baseline clinical status, AD-related genetic risk and pathology** | | | | |
| CDR-5B, mean score (s.d.) | 0.19 (0.53) | 0.19 (0.64) | 0.19 (0.60) | 0.976 |
| CDR+ > 0, n (%) | 11 (12.0%) | 14 (8.1%) | 25 (9.4%) | 0.422 |
| APOE ε4, n (%) | 32 (35.2%) | 62 (36.7%) | 94 (36.2%) | 0.914 |
| CSF pTau, mean, pg/ml (s.d.) | 55.95 (34.23) | 60.12 (31.94) | 58.68 (32.74) | 0.352 |
| CSF pTau++, >67 pg/ml, n (%) | 15 (18.3%) | 46 (29.7%) | 62 (26.1%) | 0.080 |
| PiB amyloid, mean cortical SUVr (s.d.) | 1.30 (0.58) | 1.32 (0.64) | 1.31 (0.62) | 0.772 |
| PiB amyloid++, >1.42 SUVr, n (%) | 17 (23.3%) | 30 (22.4%) | 47 (22.7%) | 1.00 |
| **Other** | | | | |
| In-scanner motion, mean FD (s.d.) | 0.21 (0.07) | 0.21 (0.08) | 0.21 (0.07) | 0.478 |

The mean (s.d.) or counts (%) of numerical or categorical variables are shown for each education group and the entire sample. Statistical differences between the two education groups were calculated for continuous and categorical variables using t-tests and χ² tests, respectively. Missing data for ‘below college’ (BMI, n = 6; traumatic brain injury, n = 3; GDS, n = 1; CSF pTau, n = 10; PiB amyloid, n = 19; APOE, n = 1). Missing data for ‘college+’ (BMI, n = 1; hypercholesterolemia, n = 1; cardiovascular incidents, n = 2; traumatic brain injury, n = 7; APOE, n = 4; CSF pTau, n = 18; PiB amyloid, n = 39). Abbreviations: SUVr, standard uptake value ratio; FD, frame displacement. Variables that were measured at different times than the functional MRI (fMRI) scans. The closest point of data collection of each measure to that of the baseline fMRI scan was used (see text for details). *In-scanner motion from all available scanning sessions, compared across the two education groups.

Education-related changes in brain system segregation are not captured by measures of cortical thinning. Multiple studies have reported cross-sectional relationships between environmental variables and brain structure in adulthood (for example, refs. 16,48,55,56). Accordingly, it is important to determine whether the observed education-related brain network changes are captured by changes in brain structure. We first determined that there was a lack of relationship between brain system segregation and mean cortical thickness after controlling for age-related variance (partial correlation controlling for age, r = 0.066, P = 0.284, CI95% = (−0.055, 0.185); raw cross-sectional correlation, r = 0.176, P = 0.004, CI95% = (0.057, 0.291)). In keeping with this, the interaction between time, education group and age at baseline predicting brain system segregation remained significant after controlling for longitudinal measures of mean cortical thickness (F = 5.915, P = 0.016, CI95% = (−0.021, −0.002)).

Modeling longitudinal mean cortical thickness as a dependent measure resulted in a main effect of age at baseline (F = 82.683, P < 0.001, CI95% = (−0.062, −0.040)) and an interaction between time and age at baseline (F = 6.711, P = 0.010, CI95% = (−0.011, −0.002)). However, time, age and education group did not significantly interact (F = 0.003, P = 0.954, CI95% = (−0.005, 0.005)). This observation reveals that longitudinal decline in brain structure is more uniformly observed across participants, and, unlike brain system segregation, it does not vary in relation to educational attainment (see section 1.2 in the Supplementary Information and Supplementary Fig. 1 for simple slope and Johnson–Neyman analyses).

Aging-related system-specific changes vary as a function of educational attainment. Brain system segregation is a summary of the
The majority of participants had clinical data available beyond their last available MRI scan (Fig. 5a depicts when clinical sessions occurred with respect to MRI sessions). Including these clinical sessions allowed us to examine the long-term prognosis of changes in brain system segregation relative to their baseline networks, greater reductions in brain system segregation are observable in the participant on the right over the comparable time span (white dashed circle, reduced separation of brain systems).

While the exact topology of networks differs between individuals, examples of brain graphs are depicted here for two individuals of equivalent age at their first scan but with differing degrees of longitudinal change in brain system segregation. Relative to their baseline networks, greater reductions in brain system segregation are observable in the participant on the right as compared to global CDR (0–18 versus 0–3), thus providing improved capability to differentiate levels of impairment. Supplemental analyses using participant’s global CDR scores in place of CDR-SB yielded qualitatively comparable results (section 1.4.1 in the Supplementary Information and Supplementary Fig. 3).

A linear mixed-effects model predicting CDR-SB revealed a significant three-way interaction between time, age at baseline and education group. In older adults, greater declines in brain system segregation were associated with greater future cognitive impairment beyond known moderators of cognitive decline. Including measures of baseline AD-related pathology and APOE status in the model revealed that both AD-related pathology ($F_{1,258} = 3.277$, $P = 0.072$, $CI_{95\%} = (−0.041, 0.002)$) and APOE status ($F_{1,258} = 3.057$, $P = 0.082$, $CI_{95\%} = (−0.040, 0.002)$) marginally interacted with time and age in predicting changes in CDR-SB (Fig. 5c,d), while changes in brain system segregation significantly interacted with time and age in predicting changes in CDR-SB ($F_{1,260} = 7.957$, $P = 0.005$, $CI_{95\%} = (−0.052, −0.010)$; Fig. 5b). None of the higher-order interactions (four or five way) that included brain system segregation together with AD-related genetic risk and baseline pathology were statistically significant or marginally significant in predicting CDR-SB ($F < 2.608, P > 0.108$).

Together, the results suggest that changes in brain system segregation impact cognitive and functional status through a pathway independent of AD-related genetic risk or pathology. However, wider range of scores as compared to global CDR (0–18 versus 0–3), thus providing improved capability to differentiate levels of impairment. Supplemental analyses using participant’s global CDR scores in place of CDR-SB yielded qualitatively comparable results (section 1.4.1 in the Supplementary Information and Supplementary Fig. 3).

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Together, the results suggest that changes in brain system segregation impact cognitive and functional status through a pathway independent of AD-related genetic risk or pathology. However,
given the number of terms in the model that includes five independent variables (not including covariates), the present sample may be underpowered to detect higher-order interactions between these variables (for example, four-way or five-way interactions; Statistical analysis in the Methods). Based on the hypothesis that there may exist additive consequences across these variables that were not revealed in primary analyses, a supplemental analysis was conducted to depict possible relationships between these variables. These comparisons revealed some plausible additive effects that were not captured by the present statistical models (section 1.5 in the Supplementary Information and Supplementary Fig. 4).

In addition to amyloid and tau burden, another common biomarker for dementia is neurodegeneration\(^2\). Measures of neurodegeneration can be estimated using structural changes (for example, changes in gray matter cortical thickness or hippocampal volume) measured by in vivo structural imaging\(^5\). However, including either longitudinal changes in mean gray matter cortical thickness or hippocampal volume as a covariate did not qualitatively alter the...
interaction between time, age at baseline and change in brain system segregation in predicting CDR-SB ($F_1,264 = 7.969, P < 0.005$). Further, when change in brain system segregation was replaced by either change in cortical thickness or hippocampal volume as the independent variable interacting with time and age at baseline, these models did not significantly predict CDR-SB ($F_1,264 < 0.785, P > 0.377$).

Importantly, accounting for changes in CDR-SB observed during the period of time between an individual’s baseline MRI and last MRI scans did not qualitatively alter the relationship between changes in brain system segregation and impending cognitive decline among older adults (interaction between time, age at baseline and change in brain system segregation, $F_{1,255} = 7.273, P = 0.007, CI_{95\%} = (-0.037, -0.006)$). This demonstrates that changes in brain system segregation predicted future cognitive impairment beyond an individual’s clinical trajectory from the same time period that the brain measures were collected (for additional analyses, see section 1.4.2 in the Supplementary Information).

Notably, none of the observed relationships between changes in brain system segregation and changes in CDR-SB in any of the models were moderated by educational attainment (for example, primary model, interaction between education group, time, age at baseline and change in brain system segregation in predicting CDR-SB, $F_{1,264} = 0.317, P = 0.574, CI_{95\%} = (-0.014, 0.025)$). While declining brain system segregation was more prominent in older adults with less education, individuals who exhibited brain network decline were more likely to exhibit future cognitive and functional impairment, irrespective of their educational attainment.

**Discussion**

Functional brain network decline is greater in older adults without a college education than that in their college-educated peers. In addition, decreasing resting-state brain system segregation is predictive of impending cognitive and functional impairment (dementia severity) among older adult individuals, independent of known AD-related genetic risk factors and measures of brain pathology or cortical thinning. However, educational attainment does not moderate the link between brain network changes and changes in clinical outcomes. While an individual’s education relates to their brain network changes during older age, if and when brain network changes occur, the impact on cognition is equally devastating irrespective of educational attainment.

**Declining brain system segregation foreshadows impending cognitive decline in older adults.** The segregation of large-scale resting-state brain systems supports brain function throughout the lifespan\(^1\). Previous research has revealed that individual differences in measures of brain system segregation are related to differences in cognitive ability, such as episodic memory and processing speed, among normative adult samples\(^1,6,7\). It has been unclear whether changes in brain system segregation translate to any clinically meaningful changes in cognition or functional status as an individual grows older. The present study demonstrates that declining brain system segregation during older age is predictive of impending changes in dementia severity, independent of AD-related genetic risk and neuropathology or cortical thinning in older adults. Interestingly, this relationship was evident for impending decline beyond the available scan sessions, in some cases up to 10 years following an individual’s final MRI scan. Further analyses revealed that the predictive utility of changes in functional brain network organization on future clinical decline was also independent of the trajectory of clinical changes that coincided with scan sessions. It is important to acknowledge that clinical observations measured from an individual’s CDR-SB scores are not necessarily specific to AD but can also capture impairment related to other forms of dementia\(^8\). However, we focused here on AD given the targeted study population (recruited under studies from the Knight ADRC) and the richness of the dataset, which has available measures of AD-related genetic risk and biomarkers of AD-related neuropathology. APOE e4 is a genetic risk factor for AD that contributes to altered amyloid formation and clearance\(^9\) and was shown to be associated with cognitive decline in healthy middle-aged and older adults\(^10,11\) and during all stages of AD\(^12,13\). As expected, the presence of at least one e4 allele was predictive of impending decline across older adults in the present sample. Critically, however, declining brain system segregation explained cognitive decline independently of APOE status. This observation is consistent with the current understanding of AD prognosis, in which genetics alone do not completely predict disease development in typical AD (that is, late-onset AD). Instead, a combination of environmental, psychosocial and lifestyle factors\(^14,15\) likely interact to alter brain structure (and, as evidenced here, functional brain network organization), which can then manifest as cognitive and behavioral impairment when brain degradation is substantial.

**Multiple abnormal brain changes define AD and cognitive impairment, including Aβ deposition\(^16,17\), the presence of tau neurofibrillary tangles\(^18\) and neurodegeneration\(^19\). Validated proxies of these brain changes have been incorporated into an influential model of major biomarkers of AD, which summarizes the presence of pathology and neurodegeneration (A/T/N: abnormal levels of Aβ
The present results demonstrate that changes in functional brain network organization and its downstream consequence are, at least in part, independent of the cascade of known pathological and neurodegenerative burdens that form the basis of existing models of AD and should be incorporated into future models of AD. Future studies examining AD risk and dementia risk more broadly should aim at more targeted examination of the interactive effects and relationships between changes in functional brain network organization and changes in brain pathology and neurodegeneration, and their collective contributions to preclinical and clinical cognitive impairment (section 1.5 in the Supplementary Information).

Brain system segregation as a measure of ‘reserve’ in aging. Past studies have shown that, when compared to those with lower education, highly educated individuals can maintain cognitive and functional abilities despite harboring greater amounts of pathologic burden (for example, refs. 15,21,22). These observations helped motivate the ‘reserve’ theory of aging, which postulates the presence of an undetermined substrate that helps resist cognitive dysfunction despite the presence of neuropathology23. Reserve has typically been indexed by an individual’s educational attainment, which itself often serves as a proxy for SES and other environmental factors. But what aspects of brain structure or function actually allow an individual to seemingly resist the impacts of pathology? We previously hypothesized that an individual’s functional brain network organization, and brain system segregation more specifically, may be a brain measure of ‘reserve’ (refs. 41,48), and the present observations support this hypothesis. Lower education was associated with greater declines in brain system segregation, independently of baseline AD-related pathology or longitudinal measures of cortical thickness. However, educational attainment did not moderate the relationship between changes in brain system segregation and cognitive decline: some individuals with higher education also exhibited declining brain system segregation, and, when they did, they were not immune to the deleterious cognitive impact of this pattern of brain network change. The preceding observations are also consistent with the idea that education may be a crude index of other variables that modify an individual’s brain structure and function. For example, educational attainment has been linked to cognitive and intellectual engagement during adulthood, which were shown to relate to age-related cognitive differences (for example, refs. 24,79) and predict dementia risk78. However, given the lack of explicit measurement of cognitive engagement in the present participant sample, we were not able to directly determine whether and how aspects of engagement during adulthood related to changes in brain system segregation (see below for additional discussion on this and related issues).

In contrast to RSFC brain network organization, longitudinal structural changes (that is, mean gray matter cortical thinning) in older adults did not vary as a function of educational attainment (section 1.2 in the Supplementary Information). The absence of a relationship between education and longitudinal structural changes in adult individuals is consistent with a number of reports examining these relationships41,77. Accordingly, the present observations provide evidence that changes in functional network organization are more sensitive to an individual’s environment than changes in brain structure (at least globally defined) and motivate focus on resting-state network organization as a target for understanding environment-related and experience-related brain plasticity in studies of brain aging.

The mechanism by which changes in brain system segregation occur are uncertain41. One possibility is that focal degeneration of specific network nodes and their downstream outcomes (for example, refs. 74,79) may alter functional networks. There is some evidence that SES-related variables and cumulative experiences linked to an individual’s environments may relate to structural differences in specific brain regions80–82. Potentially in line with this idea, closer examination of the changes in functional brain network...
topology highlighted specific brain circuits that may be particularly vulnerable to environment-related changes during older age (Fig. 4). For example, differences in RSFC changes between education groups were most prominently observed across association systems that support integrative processing. In particular, changes existed in ‘below college’ older adults that were not evident in ‘college+’ older adults, including decreases in RSFC in the cingulo–opercular system and greater between-system RSFC among nodes of the default mode, memory-retrieval and frontal–parietal systems (the latter aligning with a previous cross-sectional study83; see section 1.3 in the Supplementary Information for subsets of system changes). It remains to be determined whether and how specific structural changes may relate to specific functional network changes.

Educational attainment is linked to social, economic and health disparities. Broadly, lower education is associated with greater incidence of dementia including AD but also greater incidence of mental health disorders. Education does not dictate these brain health outcomes directly or in isolation; education has been shown to differentiate many important aspects of health, resources and lifestyle (for example, refs. 1,4,5,11), a reality that was echoed in the present data (Fig. 1c–e and section 1.6 in the Supplementary Information). Because the complex relationship between education and economic opportunities broadly shapes an individual’s life course behaviors and environment, an important task is to understand what mediates the relationship between education and aging-accompanied brain changes. Multiple candidate processes exist, many of which converge on biological pathways reflecting chronic exposure to environmental stressors, in which chronic stress results in elevated cumulative allostatic load that causes deterioration of the body and brain. Admittedly, measures of physical and mental health included in the present study are neither clinically comprehensive nor complete. Further, absent in these measurements are deeper descriptions of lifestyle-level differences including nutrition and leisurely exercise (for example, refs. 6,7,8), health behaviors (for example, nicotine and alcohol use), cognitive and intellectual engagement (for example, refs. 9,10,11) and access to or utilization of healthcare resources, all of which likely play a critical role in stress and brain aging. Finally, an individual’s educational attainment can often be linked to environmental factors defining their childhood. Similar to the cumulative effects of childhood and adulthood adversity on health and mortality, early life experiences from infancy to adolescence relate to brain structure and cognition in older age. While the present observations do not differentiate early life from present experiences, an earlier study by members of our group reported that parental education (that is, childhood SES) did not attenuate the observed cross-sectional relationship between...
adult SES and brain system segregation\(^1\). Thus, it is likely that educational attainment and childhood experience have additive and unique effects on changes in brain network organization over the lifespan.

The present results fit with multiple lines of evidence demonstrating education-related disparities in brain health during advanced age (for example, refs. \(^1\)) but offer a measurable feature of brain function, changes in which also signal the risk of impending cognitive decline before its occurrence. An important goal of follow-up work will be to understand the complex relationships that intertwine social determinants of health, longevity, lifestyle factors and changes in brain network organization\(^1\). Ultimately, a deeper understanding about the interplay between one’s environment and the brain could fill in the missing links between broader psychosocial societal factors and dementia risk, prevalence and prevention (for example, ref. \(^1\)).

**Limitations.** Using graph theory to analyze brain networks necessitates selection of multiple preprocessing and analytic parameters\(^2\). Supplemental analyses confirmed that the reported results were not limited to our decisions regarding the trade-off between resting-state data quality and quantity (section 1.7.1 in the Supplementary Information) or graph construction (that is, possible age-related differences in node definition; section 1.7.2 in the Supplementary Information and Supplementary Fig. 5). However, continued examination of longitudinal resting-state data will be crucial not only to confirm but also to better understand the properties of brain network organization that underlie the reported results.

The present dataset drew from a broader Knight ADRC dataset pool that is relatively diverse and representative of the catchment area where the data were collected. However, as is the problem with many neuroimaging studies, the final participant sample that passed rigorous data quality control included few participants with very low education (less than high school) and also included fewer non-white participants than would be expected based on the broader Knight ADRC data sample. In addition to possible selection biases due to health or education, longitudinal measurement also exposes the data to attrition bias, in which individuals with poorer health and, by association, poorer cognition, are less likely to remain in the study. Altogether, the detrimental effect of lower education on the declining trajectory of brain system segregation and the latter’s relation to clinical decline are likely underestimated given the under-representation of adults with very low education and health disparities that are prevalent among individuals with lower education.

**Conclusion.** Older adults who never completed a college degree exhibit greater declines in resting-state brain system segregation, a measure of large-scale network organization and function. Declining brain system segregation predicts impending cognitive and functional impairment beyond known AD-related biomarkers of pathology and genetic risk and irrespective of an individual’s educational attainment. These observations demonstrate that changing functional network organization is an important preclinical warning signal of cognitive impairment that is not captured by measures of brain structure or pathology. Future studies should aim at both further elucidating the time course of brain network changes relative to clinical decline but also identifying environmental factors that mediate the relationship between an individual’s educational attainment and changes in their brain network organization. These developments would help establish causal pathways and identify modifiable targets for intervention. The urgency for continued work in this area cannot be overstated in the face of population aging, the prevalence of AD and other dementias that accompany increasing age, and growing health disparities among economically disadvantaged individuals.

**Methods**

**Participants.** Adult participant data were provided by the Knight ADRC at Washington University in St. Louis. Written informed consent was obtained from all participants in the study at the time of enrollment. Each of the datasets obtained from the Knight ADRC was collected under a study that had been approved by the Human Research Protections Office of Washington University in St. Louis. The data analysis included in the present work was approved by the Institutional Review Board of the University of Texas at Dallas. Participants’ data were only included in the present study if they had available (1) a minimum of two resting-state fMRI scans to allow longitudinal functional brain network analysis and (2) demographic (age and self-reported sex) and education information; a total of 417 participants from the Knight ADRC satisfied the criteria. Participants’ structural and resting-state scans underwent structural and fMRI preprocessing, motion processing and surface mapping. Following initial processing, 266 participants had two or more sessions of data that passed all neuropsychometric quality checks (QCs) listed in the Neuroimaging section. Participants who were excluded from subsequent analysis, 28 participants failed preprocessing QCs (for example, poor skull stripping or FreeSurfer surface estimates due to artifacts in their T1 images). 116 participants failed the motion-processing QC (that is, they did not have adequate data after motion ‘scrubbing’) and seven participants failed the surface-mapping QC. Lastly, a single participant with a baseline CDR score of 1 was excluded because of their clinical diagnosis of mild dementia at baseline.

The final sample of 265 participants were 45–86 years old (\(n = 154\)) at their baseline resting-state scan (mean age = 67.01 years, s.d. = 9.26 years). AD was associated with clinical diagnosis of AD-related pathology, cardiovascular health, neurological health and mental health were measured during separate data-collection sessions (for example, PET sessions). Data sessions closest to a participant’s baseline resting-state scan session were used for purposes of analysis (81% of data from these separate experimental sessions were collected within 1 year of the baseline resting-state scan session). The sample was largely cognitively normal, with 240 participants rated as cognitively normal at baseline on the CDR, corresponding to a CDR of 0; 25 participants had a CDR of 0.5, indicating the presence of very mild dementia. See Table 1 for a breakdown of other variables across education groups.

**Educational attainment and grouping.** Across participants, the range of self-reported education time was between 6 and 22 years (education time above 22 years was recoded as 22 years), with four participants reporting less than 12 years of education (that is, they did not complete high school). While education time was collected as a numerical variable, each education year is not a uniform measure. The difference between 13 and 14 years of education is likely minimal, as both represent an individual completing some college. However, the difference between 15 and 16 years of education typically reflects the difference between someone with or without a college degree.

Importantly, the categorical distinction of having a college degree or not was shown to confer significant socioeconomic advantages, even when accounting for formal college costs to obtain a degree\(^1\). This distinction is evident across large segments of adulthood, and this effect spans various fields of study, whereby even relatively less economically lucrative majors from a four-year degree translate into economic advantages\(^1\). Accordingly, for each participant, their self-reported time of formal education was converted into a variable coding for college degree attainment, for which 16 or more years of formal education was categorized as ‘college+’ as it approximates the time when most people complete a college degree. Those with fewer than 16 years of formal education were categorized as ‘below college’ (see section 1.8 in the Supplementary Information and Supplementary Figs. 6 and 7 for further analysis and discussion regarding categorization of education).

**Neuroimaging.** Each participant had two or more resting-state scan sessions available, enabling longitudinal comparisons of their functional brain networks. On average, a participant’s second scan was 3.22 years (s.d. = 0.43–1.62 years) after their baseline scan (see Fig. 1). This time difference is evident across large segments of adulthood, and this effect spans various fields of study, whereby even relatively less economically lucrative majors from a four-year degree translate into economic advantages\(^1\). Accordingly, for each participant, their self-reported time of formal education was converted into a variable coding for college degree attainment, for which 16 or more years of formal education was categorized as ‘college+’ as it approximates the time when most people complete a college degree. Those with fewer than 16 years of formal education were categorized as ‘below college’ (see section 1.8 in the Supplementary Information and Supplementary Figs. 6 and 7 for further analysis and discussion regarding categorization of education).

**Structural imaging acquisition and preprocessing.** T1-weighted images (magnetization-prepared rapid-acquisition gradient echo sequence) were processed with FreeSurfer 5.3 to obtain a cortical surface. FreeSurfer was used to perform cortical surface reconstruction and cortical thickness at each time point. Manual examination was performed on all FreeSurfer outputs, and, when necessary, manual editing (that is, control points, white and pial surface edits) was performed to ensure accurate construction of the cortical surface. Initial segmentation and manual editing were available with each of the structural images obtained from the Knight ADRC, every T1 segmentation (pial and white matter, and surface generation of pial and white surfaces) of every participant were rechecked, and additional manual editing was performed based on procedures developed by our laboratory\(^1\).
Resting-state fMRI acquisition and preprocessing. RSFCs were computed using functional brain images that are sensitive to blood-oxygen-level-dependent (BOLD) activity obtained using an echo-planar sequence (TR = 2.200 ms, TE = 22.2 ms, flip angle = 90°, FOV = 256 × 256 mm; 36 slices, interleaved acquisition; resolution = 4 × 4 × 4 mm). During each of the resting-state functional scans, participants were instructed to fixate on a visual cross-hair, remain still, keep their eyes open and not fall asleep. In general, each imaging session included two runs of resting-state scans, each consisting of 164 volumes. However, there were five sessions of data that included extra resting-state scans (four with three runs, and a single session had four runs); participant data for each of these sessions were inspected and processed along with the first two runs available for that session. BOLD images (resting state) corresponding to the same session as each of the structural images described above were processed using a standard fMRI preprocessing pipeline using SPM8 0.8.0, including the following steps: (1) slice-timing correction to remove odd–even volume phase errors; (2) head movement between volumes and (3) realignment to the T1-weighted image from the same session. All steps were performed using FSL 5.0.2.2, except for realignment between volumes and rigid body correction, for which SPM8 was used.

While a part of the global signal may contribute variance related to general levels of arousal and genuine neural activity (for example, refs. 10,11), there is considerable evidence that a large component of the global signal includes spatially nonspecific signal artifacts related to head motions12. Failure to explicitly remove the global signal prevents the control of these known sources of artifacts10,11,12. As no method presently exists for denoising known artifact signals while retaining ‘real’ signals13,14, the alternate option of retaining the global signal in each participant is likely to result in misestimation of correlations and the resultant network estimates. Accordingly, we employed data-censoring (‘scrubbing’) and bandpass filtering procedures, which, together with global signal regression, were shown to best reduce global and distance-dependent artifacts15,16,17,18.

Network estimates. Accordingly, we employed data-censoring (‘scrubbing’) and bandpass filtering.

Brain network construction. For each session of surface-mapped resting-state data, a functional correlation matrix was generated with 349 surface-based nodes that were defined from previous boundary-based analyses19,20, and were labeled based on their spatial overlap with the vertex-wise community map published by Power et al. (not the 264 nodes21). This is the same approach that we adopted in previous publications22 with an additional constraint (see step 3 below); as in refs. 19,20. Briefly, nodes were constructed using the following steps: (1) identifying putative core areas that covered at least 8 map points on a standard RSCC boundary19,20; (2) creating disks with a radius of 3 mm around the identified area centers to avoid area borders that may exhibit more variance between individuals, and (3) discarding nodes that were in areas of low signal intensity in the original data that were used to create the boundary map (<800 (ref. 6)). All vertices within a node disk were identified, calculated in their spatial overlap with an a priori vertex-wise community map in the same fs_LR space19,20, where each disk was labeled with a functional system based on a winner-take-all approach. The BOLD time series of all vertices across the cortical surface. The mean cortical thickness was calculated by averaging the cortical thickness measurement of the left and right hemispheres. Intracranial volume (ICV) was obtained from FreeSurfer to adjust the cortical thickness. Using non-ICV-adjusted cortical thickness yielded qualitatively similar results.

Hippocampal volume. At every time point when resting-state data were collected, using the edited FreeSurfer segmentations, gray matter cortical thickness was estimated as the distance (in mm) between pial and white matter surfaces across the vertices on the cortical surface. The mean cortical thickness was calculated using the following equation (as in ref. 33):
participants with elevated levels of amyloid deposits (amyloid PiB, \(n = 47\); PiB, \(n = 160\)). CSF pTau values were obtained by analyzing CSF samples using enzyme-linked immunosorbent assays (Innotest, Innogenetics\textsuperscript{110}). Participants with a value above 67 pg/ml were categorized as having elevated levels of pTau (pTau, \(n = 61\); pTau, \(n = 176\)). A large portion of the final sample had both of these variables available (n = 193 of 265 in the final sample); however, a subset was missing some form of AD-related pathology data (see Table 1 for missing data). Participants with one or more elevated AD-related pathology markers were categorized as positive for AD-related pathology.

APOE e4 status. APOE genotyping was performed following standard procedures for extracting DNA from peripheral blood samples (see detail for APOE genotyping in ref. \textsuperscript{111}). Participants with at least one copy of the e4 allele were categorized as APOE e4\textsuperscript{+} (n = 94, 36%).

Measures of cardiovascular health and mental health. Each participant's cardiovascular health was incorporated into analyses by including independent measures related to cardiovascular risk and cardiovascular incidents. Cardiovascular health was quantified as the proportion of cardiovascular-related variables that were available and met the following criteria: (1) BMI > 30, (2) recent or remote hypertension, (3) recent or remote hypercholesterolemia, (4) recent or remote incident of heart attack or cardiac arrest, atrial fibrillation, angioplasty or endarterectomy or stent, cardiac bypass procedure, pacemaker or congestive heart failure. All incident measures represented a binary distinction as to whether a participant had either an event remotely or the particular health issue or had never experienced it (that is, absent) at the time of their first scan. Measures of participant's neurologic and mental health included details of traumatic brain-injury incidents and depressive symptoms, respectively. Traumatic brain-injury incidents were categorized and included in analyses as a categorical variable. Participants with any recent or remote incident of the following were categorized as positive for incidence of traumatic brain injury: traumatic brain injury accompanied by (1) brief loss of consciousness (<5 min), (2) extended loss of consciousness (≥5 min) or (3) chronic deficit or dysfunction. All incident measures represented a binary distinction as to whether an event had occurred or never experienced it (that is, absent) at the time of their first scan.

One participant was missing GDS data. GDS score ranging from 0 to 8, with nine participants scoring between 5 and 8. Scale has a possible range from 0 to 15 (0–4, no depression; 5–8, mild depression; 9–11, moderate depression; 12–15, severe depression). The present sample had a mean GDS score of 4.8 (standard deviation = 2.8, range 0–10).

Measurement of alternate socioeconomic variables. Occupational socioeconomic index. Each participant's self-reported occupation was matched to a corresponding occupation code in the US census and then assigned a sex-specific SEI based on predicted occupation prestige, a composite score reflecting one’s occupational status. One participant was missing SEI data. As such, SEI values were available for 228 of 265 participants.

Neighborhood median household income (2011–2015 American Community Survey). Each participant's neighborhood income was estimated based on the median household income of the zip codes in which they resided. Beginning in 2010, the median household income of a zip code became available from the American Community Survey (ACS). Because some participants’ scans were collected before 2010, it was not possible to have individualized estimates based on the years in which the scan was collected for all participants. Instead, 5-year ACS data from 2011 to 2015 were used because they encompass the median scan date across all available scanning sessions (median scan date, 2011). Five-year ACS estimates were chosen instead of 1-year or 1-year ACS estimates because they covered a greater proportion of zip codes (that is, more areas are missing from 3-year or 1-year data) and used larger sample sizes to determine estimates. Not every zip code had 5-year ACS estimates available; therefore, some participants with zip code data did not have a matching neighborhood median household income estimate; thus they were excluded from the comparison in Fig. 1d. Neighborhood median household income data were estimated for 217 of 265 participants.

National area deprivation index (2011–2015 American Community Survey). The national ADI represents the percentile ranking of neighborhood SES disadvantages, calculated using multiple variables (for example, home value, gross rent, percent of families below poverty level, percent of households without a motor vehicle\textsuperscript{112}). ADI data used in the present study were obtained from https://www.neighborhoodatlas.medicine.wisc.edu\textsuperscript{113,114}, calculated using 5-year ACS data from 2011 to 2015. ADI scores were calculated on the block group level, for which data were first linked to nine-digit zip codes. Nine-digit zip codes are sub-areas of five-digit zip codes. To estimate the ADI for a participant from the Kidney ADI, ADI scores of all nine-digit zip codes within the participant's five-digit zip code were averaged together. ADI was available for 236 of 265 participants. Because ADI data are aggregated from nine-digit zip codes and multiple measures of ACS data, more participant ADI data were available than median household income data.

Statistical analysis. The present study used a longitudinal design, in which the measure of time (that is, days from baseline) was included as a within-participant variable. Both analyses examining brain changes and cognitive impairment changes used a linear mixed-effects approach. First, in the analysis predicting longitudinal brain changes, linear mixed-effects models were used to examine how the dependent variable (for example, primary analysis, brain system segregation; supplementary analyses, cortical thickness) was predicted by time (normalized time from baseline; within participant) and interaction with education group (between participant) and age at baseline (between participant). Age at each scan was included as a measure of time to allow us to investigate the interaction between age and time for the variable, the number of days that a scan was collected after that individual's baseline scan was normalized and adjusted to the point where 0 is the baseline.

In the primary model, in-scanner head motion (mean FD; within-participant variable collected at each longitudinal time point) and sex (between-participant variable) were included as covariates. The linear mixed-effects model was calculated as follows:

\[
Y_{ij} = \beta_0 + \beta_1 (\text{sex}) + \beta_2 (\text{age}) + \beta_3 (\text{edu grp}) + \beta_4 (\text{age} \times \text{edu grp}) + \beta_5 (\text{time}) + \beta_6 (\text{time} \times \text{edu grp}) + \beta_7 (\text{time} \times \text{age}) + \beta_8 (\text{time} \times \text{age} \times \text{edu grp}) + \beta_9 (\text{time} \times \text{age} \times \text{edu grp}) + \epsilon_{ij}
\]

where \(Y_{ij}\) denotes brain system segregation for each participant \(ij\) at time \(t\) \(\gamma\) denotes the estimated fixed-effect coefficients, \(\mu\) denotes the estimated random-effect coefficients, \(\epsilon\) denotes the residual for each participant \(ij\) at time point \(t\), and \(\text{edu grp}\) is the education group. Sex and its interaction with time was included to account for the fixed effect of sex on the random effect of time. Following a similar approach, subsequent models included between-participant covariates (for example, AD-related pathology, cardiovascular health factor) and their interaction with time by entering additional terms \(\gamma_{ij} (\text{covariate})\) and \(\gamma_{ij} (\text{time} \times \text{covariate})\), where \(k\) is the \(k\)-th covariate. When a covariate was collected longitudinally (for example, motion, cortical thickness), it was included simply as \(\text{edu grp}\).

Multiple-comparison correction was applied when examining the two unique longitudinal brain change measures (brain system segregation and cortical thickness). After correction, the comparison would require a \(P\) value of 0.025 to be considered significant. The interaction between age, time and education predicting brain system segregation surpassed this corrected \(P\) value, while the three-way interaction predicting cortical thickness remained insignificant. All reported \(P\) values in the text are raw (unadjusted).

In the analysis on longitudinal changes in cognitive impairment, similar linear mixed-effects models were constructed, with the dependent variable being CDR-SB or CDR (section 1.4.1 in the Supplementary Information). Changes in brain system segregation were defined as the observed difference between brain system segregation measured from the last time point and baseline, as opposed to estimated changes from mixed models. Although a fixed percentage of participants had more than two time points of resting-state data available, a notable portion only had two time points; therefore, estimates for these individuals were more prone to shrinkage (that is, coefficients shifted more toward population values than within-participant least square estimates). Accordingly, difference scores were used to ensure a consistent means to obtain within-person changes across all participants, which also meant that the conclusion from the model predicting CDR-SB was based on actual changes in brain system segregation instead of estimated changes from a separate mixed-effect model. Furthermore, when predicting changes in CDR-SB, in addition to sex and average head motion (basically or exactly same) between baseline and the last MRI scan was included to account for the varying length of time available to quantify changes in brain system segregation. Education group was also included as a covariate in all models except when its interaction was tested.

Normality of the dependent and independent variables was examined qualitatively using Q-Q plots, in which normality was relatively high for most variables. The normality of brain system segregation increased through quadratic transformation although not substantially. We repeated the analyses using squared brain system segregation as the dependent variable, and the results presented in Table 2 remained qualitatively the same. CDR-SB is not normally distributed due to the higher number of participants with 0 or relatively low CDR-SB scores. Following a previous study examining CDR-SB in relation to continuous time scale\textsuperscript{18}, when quadratic time was included in the model predicting CDR-SB, the interaction between time, age at baseline and brain system-segregation change remained qualitatively the same. Residuals of the linear mixed-effects models were examined to ensure that they were not correlated with the fitted values of the fixed effects portion of the model.
Block-level matrix comparisons across time. Longitudinal RSFC changes in older participants (65 years and over) within each education group were conducted by comparing their RSFC matrix from the first time point to that at the latest time point. Each node–by-node RSFC matrix at each time point was combined into ‘blocks’ based on predefined system labels19,20. Correlations among nodes from the same system were averaged together to form within-system blocks, and correlations among nodes across every pair of systems were averaged together to form between-system blocks. Observed block-wise comparisons across time were computed using paired t-tests (last scan versus first scan). Permutation was used to generate null distributions of block-level t statistics (permutation # n = 10,000). Each permutation iteration shuffled the matrices within an education group across participants and time points, and the value from a paired t-test using the permuted sample was recorded. For each block, the P value was calculated as the proportion of sampled statistics more extreme than the actual statistic. A two-tailed P value less than 0.05 was considered significant, and t values for each statistically significant block was visualized (blocks that survived false discovery-rate correction were further highlighted).

Software. Linear mixed models and data visualization were carried out in R 3.6.0 using the following packages: lme4 (1.1-26) for linear mixed-effects models; emmeans (1.3-5) for extracting model estimates; ggplot2 (3.3.3), tigris (0.9.4) and choreograph2 (0.5) for visualization. Block-level matrix permutation was conducted in MATLAB 2019a using in-house scripts. Longitudinal spring-embedded graphs were generated using SoNIA (1.2.2) and exported using Cytoscape (3.2.0) to generate high-resolution images. Visualization of nodes on cortical surfaces were generated using Connectome Workbench (1.3.2).

Reporting Summary. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability. Data include patient information and are private and unsuitable for public deposition. Imaging and behavioral data are available to investigators upon request and approval from the Knight ADRC Leadership Committee. The ADRC Leadership Committee meets on the second Monday of January, March, May, July, September, and November each year. Data requests are approved on a rolling basis. Requests involving neuroimaging data should be submitted to the ADRC for preliminary review before the leadership committee meeting (director of the imaging core, T. Benzinger, benzingt@wustl.edu). Detailed instructions for making a request can be found at https://knightradc.wustl.edu/Resources/RequestResource.htm.

Code availability. The calculation of brain system segregation uses custom code available at https://github.com/mychan24/system_matrix_tools. A modified version of the superheat package was used to generate matrix visualization (https://github.com/mychan24/superheat).

Received: 3 February 2021; Accepted: 7 September 2021; Published online: 11 November 2021

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**Acknowledgements**

This work was supported by the James S. McDonnell Foundation (G.S.W), NIH grant R01 AG063930 (G.S.W) and NIH grants P30 AG066444 (J.C. Morris), P01 AG03991 (J.C. Morris) and P01 AG026276 (J.C. Morris). G.S.W thanks S. Petersen and S. Fitzpatrick for valuable discussions related to this work.

**Author contributions**

M.Y.C., J.H. and G.S.W. designed the study; M.Y.C. analyzed data; M.Y.C., L.H., C.A.C. and Z.Z. processed data and checked data for quality; M.Y.C., C.A.C. and R.M.R. recorded occupation data; M.L. retrieved and entered archival data; G.S.W. supervised research and analysis; M.Y.C. and G.S.W. wrote the manuscript; and all co-authors provided comments on and edits to the manuscript.

**Competing interests**

The authors declare no competing interests.

**Additional information**

Supplementary information The online version contains supplementary material available at https://doi.org/10.1038/s43587-021-00125-4.

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Peer review information *Nature Aging* thanks Meadbh Brosnan, Cheryl Grady, Jorge Sepulcre and the other, anonymous, reviewer(s) for their contribution to the peer review of this work.

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For all statistical analyses, confirm that the following items are present in the figure legend, table legend, main text, or Methods section.

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☐ The exact sample size (n) for each experimental group/condition, given as a discrete number and unit of measurement

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Give P values as exact values whenever suitable.

☐ For Bayesian analysis, information on the choice of priors and Markov chain Monte Carlo settings

☐ For hierarchical and complex designs, identification of the appropriate level for tests and full reporting of outcomes

☐ Estimates of effect sizes (e.g. Cohen’s d, Pearson’s r), indicating how they were calculated

Our web collection on statistics for biologists contains articles on many of the points above.

Software and code

Policy information about availability of computer code

Data collection  No software was used for data collection.

Data analysis  All statistical analyses and plotting were conducted in R 3.6.0 using the following packages: lmer4 (v1.1-26) for linear mixed-effects models; emmeans (v1.3.5) for extracting model estimates; ggplot2 (v3.3.3), tigris (v0.9.4), and cowplot (v1.0.2), and ggpubr (v0.7.2) for plotting. Block-level matrix permutation was conducted in MATLAB 2019a using in-house scripts. Longitudinal spring-embedded graphs were generated using SoNIA (1.2.2), and exported through Cytoscape (3.2.0) to generate high resolution image. Visualization of nodes on the cortical surfaces were generated using Connectome Workbench (1.3.2).

Functional MRI (fMRI) BOLD images (resting-state) were processed through an in-house fMRI preprocessing pipeline using Nipype 0.8.0, FSL 5.0.2.2. Structural images were processed through FreeSurfer 5.3.

The calculation of brain network segregation uses custom code available on https://github.com/mychan24/system_matrix_tools. A modified version of superheat package was used to generate heatmaps in the supplemental material [https://github.com/mychan24/superheat].

For manuscripts utilizing custom algorithms or software that are central to the research but not yet described in published literature, software must be made available to editors and reviewers. We strongly encourage code deposition in a community repository (e.g. Github). See the Nature Research guidelines for submitting code & software for further information.
Data

Policy information about availability of data
All manuscripts must include a data availability statement. This statement should provide the following information, where applicable:
- Accession codes, unique identifiers, or web links for publicly available datasets
- A list of figures that have associated raw data
- A description of any restrictions on data availability

Data includes patient information and are private and unsuitable for public deposition. Imaging and behavioral data are available to investigators upon request and approval from the Knight ADRC Leadership Committee. The ADRC Leadership Committee meets the second Monday of January, March, May, July, September, and November each year. Data requests are approved on a rolling basis. Requests involving neuroimaging data should be submitted to the ADRC for preliminary review prior to the LeadershSHIP Committee meeting (Director of Imaging Core – Dr. Tammie Benzinger, benzinger@wustl.edu). Detailed instructions for making a request can be found on https://knightheadrc.wustl.edu/Research/ResourceRequest.htm

Field-specific reporting

Please select the one below that is the best fit for your research. If you are not sure, read the appropriate sections before making your selection.

☐ Life sciences  ☑ Behavioural & social sciences  ☐ Ecological, evolutionary & environmental sciences

For a reference copy of the document with all sections, see http://nature.com/documents/nr-reporting-summary-list.pdf

Behavioural & social sciences study design

All studies must disclose on these points even when the disclosure is negative.

| Study description | Quantitative longitudinal study. |
|-------------------|----------------------------------|
| Research sample   | The data was obtained from an existing dataset, collected at Washington University Knight Alzheimer's Disease Research Center (ADRC). The research questions necessitates a dataset with longitudinal resting-state data, neuropathology data, and clinical diagnosis of dementia. Furthermore, it was important that the participant sample exhibited heightened Alzheimer’s Disease-related risk (e.g., higher proportion of APOE e4 participants), resulting in adequate statistical power to examine the research question. The data included adult participants that were tested two times or more to form a longitudinal sample. The final sample, after quality control check included 265 unique participants (154F; aged 45-86y at baseline testing). The sample included middle-aged and older adults because this was the approximate age-range when the hypothesized brain changes would be observed. |
| Sampling strategy | This study uses an archival dataset. We conducted a power analysis prior to requesting the data based on previously published work reporting a SES by age interaction on brain system segregation with a partial eta squared of .03 (Chan et al. 2018). The statistical effect of interest (3-way interaction of a mixed-effects approach) required a minimum of 168 participants (achieving 95% power, alpha =0.05, effect size f=0.176 [converted from partial eta squared of 0.03]). The final sample size exceeded the minimum requirement. |
| Data collection   | Participants complete repeated (longitudinal) clinical assessments at the Knight ADRC. The clinical assessment includes a full neurological examination and administration of the Clinical Dementia Rating (CDR) by trained clinicians. Only the CDR and corresponding sum of boxes score were used in the present study. Neuropathology: As previously described in Fagan et al. (2006, Ann Neurol), cerebral spinal fluid (CSF) was collected at 8am after overnight fasting. A trained neurologist collects the sample in the polypropylene tube via gravity drip using 22-gauge Sprotte spinal needle. CSF sample was gently inverted and briefly centrifuged at low speed to pellet any cellular elements; and then aliquoted [500 μl] into polypropylene tubes before freezing at -84°C until time of assay. CSF samples were measured using commercial enzyme-linked immunosorbent assay (Innotest; Innogenetics, Ghent, Belgium). As previously described in Mintun et al. (2006, Neurology), amyloid posion tomography (PET) images were collected using [11C] Pittsburgh compound B (PiB) or Forbetapir (18FAV-45). Only PiB data were used in the present study. Standard uptake value ratios (SUVR) were calculated based on 30-60 minute post-injection window for PiB data. MRI: Participants were scanned using a Siemens 3T Trio or a Siemens Biograph mMR PET/MR scanner (Siemens, Malvern, PA) at the Center for Clinical Imaging Research located at the Washington University Medical Center. See magnetic resonance imaging section below for detailed sequence and processing information. Genetics: APOE genotyping was obtained from each participant's blood sample. APOE e4 positive status was cabled based on the presence of at least one e4 allele. Researchers are present when data are being collected. The study has no experimental vs. control condition. |
| Timing            | This study used a subset of data from the Knight ADRC data. Of the data that was obtained for this study, the earliest MRI scan date was 01/2007, and the latest scan date was 12/2016. The MRI data was collected on a rolling basis. Based on the earliest available MRI data, clinical data collected within a year of the baseline MRI data, from 05/2006 until 05/2019 were used. |
Data exclusions
A total of 417 participants from the Knight ADRC with two or more resting-state scans were preprocessed with a resting-state fMRI preprocessing pipeline. 28 participants failed preprocessing QC (e.g., poor skull-stripping, poor segmentation from FreeSurfer), 116 participants did not pass the QC for having adequate data after head-motion correction (’scrubbing’), 7 participants failed surface-mapping QC. Lastly, a single participant with a baseline Clinical Dementia Rating scale (CDR) of 1 was excluded. This resulted in a total of 265 unique participants with 2 or more resting-state scans.

Non-participation
Data-collection is on-going. Participant status or attrition data was not available as part of the data request.

Randomization
No experimental/control group assignment was used in the current study.

Reporting for specific materials, systems and methods
We require information from authors about some types of materials, experimental systems and methods used in many studies. Here, indicate whether each material, system or method listed is relevant to your study. If you are not sure if a list item applies to your research, read the appropriate section before selecting a response.

### Materials & experimental systems
| n/a | Involved in the study |
|-----|------------------------|
| X | Antibodies |
| X | Eukaryotic cell lines |
| X | Palaeontology and archaeology |
| X | Animals and other organisms |
| X | Human research participants |
| X | Clinical data |
| X | Dual use research of concern |

### Methods
| n/a | Involved in the study |
|-----|------------------------|
| X | ChiP-seq |
| X | Flow cytometry |
| X | MRI-based neuroimaging |

### Human research participants
Policy information about studies involving human research participants

#### Population characteristics
See above

#### Recruitment
Participants in the present study include a subset of participants recruited in studies under the Knight ADRC. Participants in Knight ADRC are recruited from the metropolitan St. Louis, Missouri region. Recruitment methods include word of mouth, physician referrals, and community recruitment activity.

The present dataset drew from a broader Knight ADRC dataset pool which is relatively diverse and representative of the catchment area where the data was collected. However, as is the problem with many neuroimaging studies, the final participant sample that passed rigorous data quality control included few participants with very low education (< high school) and also included fewer non-white participants than would be expected based on the broader Knight ADRC data sample. In addition to possible selection biases due to health or education, longitudinal measurement also exposes the data to attrition bias, where individuals with poorer health and by association, poorer cognition, are less likely to remain in the study. Altogether, the detrimental effect of lower education on the declining trajectory of brain system segregation and the latter’s relation to clinical decline are likely under-estimated given the health disparities that are prevalent among lower educated adults. These limitations add constraints to interpretation of the observed relationships.

#### Ethics oversight
The Knight ADRC study has been approved by the Human Research Protections Office of Washington University in St. Louis. Informed consent was obtained from all participants in the study. The analysis of Knight ADRC data included in the present work was approved by the Institutional Review Board of The University of Texas at Dallas.

Note that full information on the approval of the study protocol must also be provided in the manuscript.

Magnetic resonance imaging

### Experimental design

#### Design type
Resting-state

#### Design specifications
During each of these resting-state scans, participants were instructed to fixate on a visual cross-hair, remain still and keep their eyes open, and not fall asleep. In general, each imaging session included 2 runs of resting-state scans, each consisting of 164 volumes. However, there were 5 total sessions of data that included extra resting-state scans (four with 3 runs, and a single session had 4 runs); participant data for each of these sessions was inspected and processed along with the first two runs available for that session.

#### Behavioral performance measures
Resting-state fMRI is a task-free protocol and does not have behavioral performance measures.
Acquisition

Imaging type(s)  resting-state fMRI, T1 MPRAGE

Field strength  3

Sequence & imaging parameters  T1-weighted anatomical images were obtained using a magnetization-prepared rapid-acquisition gradient echo sequence (MPRAGE); repetition time [TR] = 2400ms, echo time [TE] = 3.16ms; flip angle = 8°; field of view [FOV] = 256 × 256mm; slices = 176; resolution = 1 × 1 × 1mm). Resting-state functional correlations were computed using functional brain images that are sensitive to blood oxygen level-dependent (BOLD) activity obtained using an echo-planar sequence (TR = 2200ms, TE = 27ms; flip angle = 90°; FOV = 256 × 236mm; slices = 36, interleaved acquisition; resolution = 4 × 4 × 4mm).

Area of acquisition  whole brain

Diffusion MRI  Not used

Preprocessing

Preprocessing software  T1 images were processed using FreeSurfer 5.3 to obtain the participant’s cortical surface, cortical thickness measure, and hippocampal volume at each time point. Manual examination was performed on all FreeSurfer outputs, and when necessary, manual editing (i.e., control points, white/pial surface edits) was performed to ensure accurate construction of the cortical surface.

BOLD images (resting-state) were processed through an in-house fMRI preprocessing pipeline using Nipype 0.8.0: (i) slice-time correction to remove odd-even slice intensity differences due to interleaved acquisition, (ii) rigid body correction for estimating and correcting head movement between volumes, and (iii) realignment to the the T1 weighted image from the same session. All steps were done using FSL 5.0.2.2 except for realignment between volumes and rigid body correction, where SPMM was used as it provided more accurate estimates in our sample. Functional data in volume-space was not smoothed since the data were surface-mapped to the fs_LR surface template, and smoothed using a Gaussian smoothing kernel (σ=2.55) on the surface.

Normalization  Participants’ anatomical image was used to construct their cortical surface. Anatomical image was deformed to the fsaverage surface using FreeSurfer, version 5.3. The left and right fsaverage surfaces were then registered to a hybrid left-right fsaverage atlas (fs_LR).

Functional data were registered to the fs_LR [32k] surface-based atlas for analysis. Using the transformation matrix and deformation maps generated during the preprocessing of the corresponding anatomical data, the volumetric functional data were resampled to the fs_LR surfaces through a one-step transformation.

Normalization template  The fs_LR atlas is a hybrid left–right fsaverage surface. The fsaverage-registered left and right hemisphere surfaces were brought into register with each other using deformation maps from a landmark-based registration of the left and right fsaverage surfaces (Van Essen et al. 2012).

Noise and artifact removal  Multiple regression to remove variance related to whole-brain signal, ventricular signal, white matter signal, their derivatives, and the “Friston24” motion regressors.

Volume censoring  In-house scripts written in MATLAB was used to censor volumes that exceed frame-wise displacement of 0.3mm, or were between two contaminated volumes that were less than 5 volumes apart. Participants with less than 100 volumes of data after censoring were excluded from further analysis.

Statistical modeling & inference

Model type and settings  N/A. Graph theory based technique was used to extract network summary index for each participant.

Effect(s) tested  Resting-state fMRI does not have a task or stimulus condition. Linear mixed effects models were used to analyze changes in network summary measure.

Specify type of analysis:  Both

Anatomical location(s)  Functionally defined resting-state ROIs across the entire cortex (349 3mm radius disks) were used to construct a network graph (Chan et al. 2014; 2017).

Statistic type for inference  Voxel wise and cluster-wise analyses were not performed. Linear mixed effects models were used to analyze changes in network summary measure. Permutation (10,000) was used to examine block level system changes.

Correction  FDR correction was applied to block level system change analysis.
### Models & Analysis

|               |   |
|---------------|---|
| n/a | Involved in the study |
|     | □ | Functional and/or effective connectivity |
|     | □ | Graph analysis |
| | | □ | Multivariate modeling or predictive analysis |

**Functional and/or effective connectivity**

- Fisher’s z-transformed Pearson’s correlation

**Graph analysis**

- A global summary measure, brain system segregation [Chan et al. 2014], was calculated using weighted graph on the subject-level.