#### RESEARCH ARTICLE

## WILEY

## Sex- and age-specific associations between cardiometabolic risk and white matter brain age in the UK Biobank cohort

Sivaniya Subramaniapillai<sup>1,2,3</sup> Sana Suri<sup>4,5</sup> Claudia Barth<sup>6,7</sup> | Ivan I. Maximov<sup>3,6,8</sup> | Irene Voldsbekk<sup>3,6</sup> | Dennis van der Meer<sup>6,9</sup> | Tiril P. Gurholt<sup>6</sup> | Dani Beck<sup>3,6,7</sup> | Bogdan Draganski<sup>1,10</sup> | Ole A. Andreassen<sup>6,11</sup> | Klaus P. Ebmeier<sup>4</sup> | Lars T. Westlye<sup>3,6,11</sup> | | Ann-Marie G. de Lange<sup>1,3,4</sup>

<sup>1</sup>LREN, Centre for Research in Neurosciences, Department of Clinical Neurosciences, Lausanne University Hospital (CHUV) and University of Lausanne, Lausanne, Switzerland

<sup>2</sup>Department of Psychology, Faculty of Science, McGill University, Montreal, Quebec, Canada

<sup>3</sup>Department of Psychology, University of Oslo, Oslo, Norway

<sup>4</sup>Department of Psychiatry, University of Oxford, Oxford, UK

<sup>5</sup>Wellcome Centre for Integrative Neuroimaging, University of Oxford, Oxford, UK

<sup>6</sup>Norwegian Centre for Mental Disorders Research (NORMENT), Division of Mental Health and Addiction, Oslo University Hospital and University of Oslo, Oslo, Norway

<sup>7</sup>Department of Psychiatric Research, Diakonhjemmet Hospital, Oslo, Norway

<sup>8</sup>Department of Health and Functioning, Western Norway University of Applied Sciences, Bergen, Norway

<sup>9</sup>School of Mental Health and Neuroscience, Faculty of Health Medicine and Life Sciences, Maastricht University, Maastricht, The Netherlands

<sup>10</sup>Department of Neurology, Max Planck Institute for Human Cognitive and Brain Sciences, Leipzig, Germany

<sup>11</sup>KG Jebsen Centre for Neurodevelopmental Disorders, University of Oslo, Oslo, Norway

#### Correspondence

Sivaniya Subramaniapillai and Ann-Marie G. de Lange, LREN, Centre for Research in Neurosciences, Department of Clinical Neurosciences, Lausanne University Hospital (CHUV) and University of Lausanne, Lausanne, Switzerland.

Email: sivaniya.subramaniapillai@mail.mcgill.ca and ann-marie.de-lange@chuv.ch

#### **Funding information**

Alzheimer's Society, Grant/Award Number: 441; Fondation Leenaards; H2020 European Research Council, Grant/Award Numbers: 732592, 802998, 847776; Healthy Brains Healthy Lives; Helse Sør-Øst RHF, Grant/ Award Numbers: 2017-112, 2018076, 2019-108, 2019101, 2022080; Norges Forskningsråd, Grant/Award Numbers: 223273, 249795, 273345, 2837989, 283799, 298646, 300768; NSERC Michael Smith Foreign Study Supplement; Schweizerischer Nationalfonds zur Förderung der

#### Abstract

Cardiometabolic risk (CMR) factors are associated with accelerated brain aging and increased risk for sex-dimorphic illnesses such as Alzheimer's disease (AD). Yet, it is unknown how CMRs interact with sex and apolipoprotein E- $\epsilon$ 4 (*APOE4*), a known genetic risk factor for AD, to influence brain age across different life stages. Using age prediction based on multi-shell diffusion-weighted imaging data in 21,308 UK Biobank participants, we investigated whether associations between white matter Brain Age Gap (BAG) and body mass index (BMI), waist-to-hip ratio (WHR), body fat percentage (BF%), and *APOE4* status varied (i) between males and females, (ii) according to age at menopause in females, and (iii) across different age groups in males and females. We report sex differences in associations between BAG and all three CMRs, with stronger positive associations among males compared to females. Independent of *APOE4* status, higher BAG (older brain age relative to chronological age) was associated with greater BMI, WHR, and BF% in males, whereas in females, higher BAG was associated with greater WHR, but not BMI and BF%. These

This is an open access article under the terms of the Creative Commons Attribution License, which permits use, distribution and reproduction in any medium, provided the original work is properly cited.

© 2022 The Authors. Human Brain Mapping published by Wiley Periodicals LLC.

Wissenschaftlichen Forschung, Grant/Award Numbers: 32003B\_135679, 32003B\_159780, 324730\_192755, CRSK- 3\_190185, PZ00P3\_193658; The Academy of Medical Sciences/the Wellcome Trust/the Government Department of Business, Energy and Industrial Strategy/the British Heart Foundation/ Diabetes UK Springboard Award, Grant/Award Number: SBF006\1078; the HDH Wills 1965 Charitable Trust, Grant/Award Number: 1117747; Wellcome Trust, Grant/Award Number: 203139/Z/16/Z; Women in Cognitive Science-Canada

divergent associations were most prominent within the oldest group of females (66-81 years), where greater BF% was linked to lower BAG. Earlier menopause transition was associated with higher BAG, but no interactions were found with CMRs. In conclusion, the findings point to sex- and age-specific associations between CMRs and brain age. Incorporating sex as a factor of interest in studies addressing CMR may promote sex-specific precision medicine, consequently improving health care for both males and females.

#### KEYWORDS

APOE genetic risk, brain age, cardiometabolic health, menopause, sex differences

#### 1 | INTRODUCTION

Cardiometabolic risk (CMR) factors such as obesity are associated with adverse health outcomes including accelerated brain aging (Beck, de Lange, Pedersen, et al., 2021) and increased risk for Alzheimer's disease (AD; Livingston et al., 2020; Qiu & Fratiglioni, 2015). The impact of both CMRs and genetic risk for AD, such as apolipoprotein  $E-\epsilon 4$  (APOE4), are known to differ between males and females (Algarni et al., 2021; Bretsky et al., 1999; Geerlings et al., 2010; Gerdts & Regitz-Zagrosek, 2019; Neu et al., 2017; Schorr et al., 2018). Yet, it is unknown whether these risk factors interact in sex-specific ways to influence brain health across different age periods or endocrine life stages. While males experience greater incidence and prevalence of cardiometabolic disease in early midlife, the greatest risk of cardiometabolic disease in females is observed up to 10 years later, coinciding with the menopausal transition (Dubnov et al., 2003; Maas & Appelman, 2010). Postmenopause, female APOE4 carriers are also at greater AD risk than their male counterparts (Bretsky et al., 1999; Neu et al., 2017), but it is unknown whether a reduction in neuroprotective ovarian hormones, combined with an elevated cardiometabolic and genetic risk profile, may accelerate brain aging and subsequent AD risk in females compared to males. Investigating how CMRs interact with APOE genotype to influence the brain at different life stages, whether age- or menopause-related, may thus clarify sexspecific risk profiles for accelerated brain aging and illuminate critical periods for preventive interventions.

Poor cardiometabolic health is associated with changes in brain microvasculature, which may be reflected in magnetic resonance imaging (MRI)-derived indices of white matter (WM) microstructure (Alfaro et al., 2018). CMRs such as adiposity, hypertension, smoking, and diabetes have been associated with lower fractional anisotropy (Birdsill et al., 2017; Mueller et al., 2011; Stanek et al., 2011), lower myelin and iron content (Trofimova et al., 2021), and greater WM hyperintensity (WMH) burden (Alqarni et al., 2021; Griffanti et al., 2018; Habes et al., 2018; Lampe et al., 2019; Raffield et al., 2016; Sachdev et al., 2009) in healthy older adults. Although females usually have higher volumes of WMH than males (de Leeuw et al., 2001; DeCarli et al., 2005; Fatemi et al., 2018; Sachdev et al., 2009; van den Heuvel et al., 2004), extensive literature indicates

that males with CMRs (adiposity, hypertension, diabetes, and atherosclerosis) are more likely to develop WMH compared to females with similar levels of risk (Algarni et al., 2021; Assareh et al., 2014; Filomena et al., 2015; Geerlings et al., 2010; Jongen et al., 2007). Hence, cardiometabolic health may influence WM microstructure differently in males and females, and risk profiles may also vary with age. Despite the consensus that high blood pressure and cholesterol levels are associated with accelerated brain aging and elevated dementia risk (Qiu et al., 2005; Solomon et al., 2009; van Vliet, 2012), the role of body fat is inconclusive (Fitzpatrick et al., 2009; Stewart et al., 2005). While mixed findings may be a result of selection or survivor bias (Heffernan et al., 2016; Jacobsen et al., 2021; Munafò et al., 2018; Salthouse, 2014), or variation in body fat indices used across studies (e.g., body mass index [BMI] versus waist-to-hip ratio [WHR] or body fat percentage [BF%]; Huxley et al., 2010; Lavie et al., 2012; Tchernof & Després, 2013: Tomiyama et al., 2016), they could also reflect a variable role of body composition throughout the lifespan. For example, one study found that BMI had a positive association with dementia risk when measured >20 years before dementia diagnosis, and a negative association when measured <10 years before dementia diagnosis (Kivimäki et al., 2018). Low BMI at later life stages may indicate frailty, sarcopenia (muscle loss), or preclinical dementia (Buchman et al., 2005; Hassan et al., 2019; Johnson et al., 2006; Subramaniapillai et al., 2021). Hence, while high BMI in mid-adulthood may largely reflect obesity, higher BMI in senescence may reflect overall physical fitness or lack of degenerative diseases. Furthermore, body fat may act as a source of estrogen in postmenopausal females (Simpson, 2003), potentially protecting against WM decline (Klosinski et al., 2015). However, only a few studies have tried to disentangle the role of body fat composition in endocrine versus chronological aging (Sowers et al., 2007; Trikudanathan et al., 2013). Since risk profiles for adverse brain health may vary by sex across certain life stages, it is relevant to investigate specific age windows at which CMRs may have sex- and genotype-specific effects on the brain.

One strategy for detecting atypical brain aging, particularly if it does not involve visible pathognomonic hallmarks of degenerative brain disease, is to use machine learning to predict an individual's age based on neuroimaging-derived measures (Cole & Franke, 2017; Cole, Marioni, et al., 2019; Kaufmann et al., 2019). Brain Age Gap (BAG)

<sup>2</sup>\_\_\_\_WILEY-

provides a measure of deviation from expected age trajectories, and has been used to identify differences in patients with neurological and psychiatric disorders relative to healthy controls (Cole, Raffel, et al., 2019; Franke & Gaser, 2019; Han et al., 2020; Kaufmann et al., 2019; Rokicki et al., 2020; Tønnesen et al., 2020), as well as predicting future dementia risk (Wang et al., 2019) and prognosis (Biondo et al., 2020; Franke & Gaser, 2012; Gaser et al., 2013; Löwe et al., 2016). While individual variation in BAG reflects a combination of genetic and environmental factors (Elliott et al., 2019; Kaufmann et al., 2019; Vidal-Piñeiro et al., 2021), clinical studies indicate that an older "brain age" relative to what is expected for an individual's chronological age (i.e., positive BAG) may in part reflect accelerated neural aging processes (Cole, Raffel, et al., 2019; Han et al., 2020; Kaufmann et al., 2019; Kolenic et al., 2018; Rokicki et al., 2020; Tønnesen et al., 2020; van Gestel et al., 2019). Positive BAG values have also been associated with negative outcomes in population-based studies, including cardiovascular risk, cognitive impairments, and dementia risk (Biondo et al., 2021; de Lange, Anatürk, et al., 2020; Egorova et al., 2019; Franke & Gaser, 2012; Gaser et al., 2013; Kolbeinsson et al., 2020; Löwe et al., 2016; Wang et al., 2019). Previous studies have shown accurate age prediction based on diffusion-weighted imaging measures (Beck, de Lange, Maximov, et al., 2021; Cole, 2020; Richard et al., 2018; Voldsbekk et al., 2021), as well as associations between WM BAG and CMRs (Beck, de Lange, Alnaes, et al., 2021; Beck, de Lange, Pedersen, et al., 2021). However, these previous studies did not assess sex-specific effects, or whether CMRs interact with APOE genotype to influence WM BAG during certain life phases.

In this study, we examined the associations between WM BAG and key CMRs, including BMI, WHR, and BF% (Bowman et al., 2017; Bradbury et al., 2017: UI-Hag et al., 2014), and APOE4 status in males (N = 10,605) and females (N = 10,703). We further assessed whether these risk factors had salient effects in middle age (44-55 years) and different stages of older adulthood (56-65 years and 66-82 years) (Grady et al., 2006; Subramaniapillai, Rajagopal, et al., 2019; Subramaniapillai, Rajah, et al., 2019). Participants' brain ages were computed using a prediction model based on WM measures derived from three diffusion imaging modes: diffusion tensor imaging (DTI) (Basser et al., 1994), diffusional kurtosis imaging (DKI) (Jensen et al., 2005), and WM tract integrity (WMTI) (Fieremans et al., 2011). While DTI is commonly used to estimate WM indices that are highly sensitive to age (Beck, de Lange, Maximov, et al., 2021; Krogsrud et al., 2016; Storsve et al., 2016; Westlye et al., 2010), biophysical diffusion models such as WMTI (Fieremans et al., 2011), which is derived from DKI (Jensen et al., 2005), may more accurately capture WM tissue structure complexity (Jelescu & Budde, 2017; Jensen et al., 2005), thus providing greater biological specificity (Novikov et al., 2018). Given previous work indicating structural and functional sex differences in the human brain (Armstrong et al., 2019; Kaczkurkin et al., 2019; Ritchie et al., 2018; Scheinost et al., 2015), the diffusion metrics were used as input features to three separate prediction models: (1) mixed sex, (2) female only, and (3) male only, in order to improve the accuracy of the sex-specific analyses (Biskup et al., 2019; Cirillo et al., 2020; Kaufmann et al., 2019). We used linear regression

models to assess whether associations between BAG and BMI, WHR, BF%, and *APOE* genotype varied (i) between males and females, (ii) according to age at menopause in females, and (iii) across different age groups in males and females.

#### 2 | METHODS AND MATERIALS

#### 2.1 | Sample characteristics

The initial sample was drawn from the UK Biobank cohort (www. ukbiobank.ac.uk), and included 39,232 participants with diffusionweighted imaging and demographic data. We excluded 3379 participants with diagnosed brain disorders based on ICD10 (chapter V and VI, field F; mental and behavioral disorders, including F00-F03 for AD and dementia, and F06.7 "Mild cognitive disorder," and field G; diseases of the nervous system, including inflammatory and neurodegenerative diseases (except G55-59: "Nerve, nerve root and plexus disorders"). Diagnostic details are provided in the UK Biobank online resources (http://biobank.ndph.ox.ac.uk/showcase/field.cgi?id=41270), and in the ICD10 diagnostic manual (https://www.who.int/classifications/icd/ icdonlineversions). In addition, 113 participants were excluded based on MRI outliers (see Section 2.2), leaving a total of 35,740 participants with diffusion-weighted imaging data that were included in the brain age models. Only participants with complete data on demographic factors, APOE genotype, BMI, WHR, and BF% from the MRI assessment time point were included in the subsequent analyses, yielding a final sample of 21,308 (male = 10,605, female = 10,703). Sample demographics are provided in Table 1. Sex of participants refers to binary data on biological sex acquired from the NHS registry at recruitment. but in some cases updated by the participant (see https://biobank. ndph.ox.ac.uk/showcase/field.cgi?id=31).

#### 2.2 | MRI data acquisition and processing

A detailed overview of the UK Biobank data acquisition and protocols is available in (Alfaro-Almagro et al., 2018; Miller et al., 2016). Briefly, we processed diffusion-weighted imaging data using an optimized diffusion pipeline as described in detail in Maximov et al., 2019. We included metrics derived from DTI (Basser et al., 1994), DKI (Jensen et al., 2005), and WMTI (Fieremans et al., 2011) as input features in the age prediction models, as described in Voldsbekk et al., 2021. The metrics for each model are listed in Supporting Information (SI) Section 1. The metrics were extracted based on subject-specific skeletonized images (Smith et al., 2006), and Johns Hopkins University (JHU) atlases for WM tracts (with 0 thresholding; Mori et al., 2005) were used to provide global mean values and regional measures for 12 tracts used in previous aging and development studies (Krogsrud et al., 2016; Storsve et al., 2016; Voldsbekk et al., 2021; Westlye et al., 2010); anterior thalamic radiation, corticospinal tract, cingulate gyrus, cingulum hippocampus, forceps major, forceps minor, inferior fronto-occipital fasciculus, inferior longitudinal fasciculus, superior

## ₄ WILEY-

#### TABLE 1 Sample demographics

Variable		Male	Female	p-value	Test
Ν		10,605	10,703		
Age	Mean ± SD	64.58 ± 7.65	63.12 ± 7.33	<.001	KW
	Range (years)	44.57-81.30	45.48-81.89		
Ethnic background	% White	97.33	97.39	<.001	X <sup>2</sup>
	% Black	0.47	0.62		
	% Mixed	0.35	0.46		
	% Asian	1.31	0.63		
	% Chinese	0.20	0.30		
	% Other	0.34	0.61		
Education	% University/college degree	50.00	47.81	<.001	X <sup>2</sup>
	% A levels or equivalent	12.08	15.00		
	% O levels/GCSE or equivalent	17.06	20.41		
	% NVQ or equivalent	11.32	6.67		
	% Professional qualification	3.92	5.56		
	% None of the above	5.62	4.56		
Assessment location (N)	Newcastle	1916	1999	.04	X <sup>2</sup>
	Cheadle	7280	7178		
	Reading	1409	1526		
APOE4 status	% carrier	25.48	26.65	.05	X <sup>2</sup>
	% noncarrier	74.52	73.35		
BMI	Mean ± SD	26.72 ± 3.60	25.65 ± 4.18	<.001	KW
	Range	16.38-39.95	14.55-39.99		
WHR	Mean ± SD	0.93 ± 0.06	0.81 ± 0.07	<.001	KW
	Range	0.53-1.26	0.60-1.16		
BF%	Mean ± SD	25.12 ± 5.47	35.57 ± 6.44	<.001	КW
	Range	5.20-42.50	7.40-57.5		

Note: Mean ± standard deviation (SD) and ranges for age, body mass index (BMI), waist-to-hip ratio (WHR), and body fat percentage (BF%), and % in each group for ethnic background, education, assessment location, and APOE4 status.

Abbreviations: GCSE, general certificate of secondary education; KW, Kruskal-Wallis; NVQ, national vocational qualification; X<sup>2</sup>, chi-squared test.

longitudinal fasciculus, uncinate fasciculus, superior longitudinal fasciculus temporal, and corpus callosum. The included diffusion MRI data passed tract-based spatial statistics (TBSS) post-processing quality control using the YTTRIUM algorithm (Maximov et al., 2021), and were residualized with respect to scanning site using linear models. To remove further outliers, participants with SD ± 4 on the global mean FA measure were excluded, yielding a final sample of 35,740 participants with MRI data (male = 16,909, female = 18,831). To optimize prediction accuracy, the full MRI sample was included in the brain age models (Section 2.3), while the subsequent analyses (Sections 2.5 and 2.6) included only participants with complete data on CMRs and APOE genotype (N = 21,308; Table 1).

#### 2.3 | Brain age prediction

We ran three age prediction models: (1) mixed sex, (2) female only, and (3) male only, to obtain sex-specific BAG values (Biskup

et al., 2019; Kaufmann et al., 2019) as well as general BAG estimates based on the mixed sample. The prediction models were run using the XGBoost regression algorithm (eXtreme Gradient Boosting; https:// github.com/dmlc/xgboost). XGboost includes advanced regularization to reduce over-fitting, and has shown superior performance in machine learning competitions (Chen & Guestrin, 2016). Parameters were tuned in a nested cross-validation using five inner folds for grid search, and 10 outer folds for model validation. Feature importance rankings for each model were extracted using gain scores, which are calculated based on each feature's contribution to each tree in the model and thus indicate the relative contribution of each feature to the prediction. BAG values were calculated by subtracting chronological age from predicted brain age. The age and BAG distributions for each of the age prediction models are shown in SI Figure 1. To ensure that associations with the variables of interest were not driven by age-dependence in the BAG estimations (Li et al., 2018; Liang et al., 2019), chronological age was regressed out of the BAG values before they were used in subsequent analyses (de Lange &

Cole, 2020; Le et al., 2018). As a cross-check, we performed a supplementary analysis for the mixed-sex model where we used 10% of the data as a held-out validation sample (N = 3574) for model optimization (grid search), and derived the best-fit parameters to run a separate age prediction model in the rest of the sample (N = 32,166) with 10-fold cross-validation.

#### 2.4 | APOE genotyping

Participants' APOE genotype was extracted using the UK Biobank version 3 imputed data, which has been rigorously assessed for quality control by the UK Biobank genetics team (Bycroft et al., 2018). The two APOE single-nucleotide polymorphisms—rs7412 and rs429358 (Lyall et al., 2016) were used to estimate APOE genotype. APOE  $\epsilon$ 4 status was labeled *carrier* for  $\epsilon$ 3/ $\epsilon$ 4 and  $\epsilon$ 4/ $\epsilon$ 4 combinations, and *noncarrier* for  $\epsilon$ 2/ $\epsilon$ 2,  $\epsilon$ 2/ $\epsilon$ 3, and  $\epsilon$ 3/ $\epsilon$ 3 combinations (Lyall et al., 2019). We removed participants with the homozygous  $\epsilon$ 2/ $\epsilon$ 4 allele combination due to its ambiguity with  $\epsilon$ 1/ $\epsilon$ 3 (Lyall et al., 2016; Wisdom et al., 2011). For more information on the genotyping process, refer to (Bycroft et al., 2018).

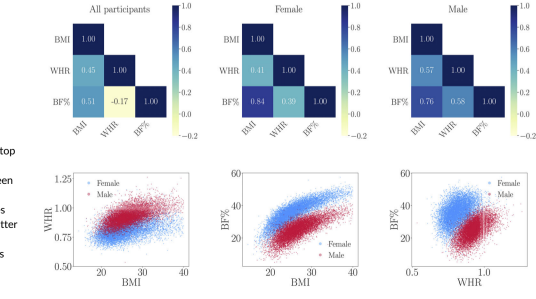
#### 2.5 | CMR factors: BMI, WHR, and BF%

CMRs included BMI (kg/m<sup>2</sup>), WHR (waist circumference/hip circumference), and BF% based on body composition by impedance measurement. All assessment procedures are described in detail in the UK Biobank protocol (Elliott & Peakman, 2008). As compared to BMI, which is a general measure of body adiposity, BF% distinguishes fat from muscles, while WHR is a more specific measure of abdominal obesity. Participants with BMI >40 were excluded (N = 196), since these values indicate morbid obesity and risk for serious health complications and comorbidities (Jarolimova et al., 2013; Schelbert, 2009). The correlations between BMI, WHR, and BF% are shown in Figure 1, indicating shared variance corresponding to previous studies (Chen et al., 2010; Myint et al., 2014; Ranasinghe et al., 2013; UI-Haq et al., 2014; Wiltink et al., 2013).

## 2.6 | Categorizing groups based on (i) age at menopause and (ii) chronological age

To investigate whether associations of BAG with CMRs and APOE genotype varied according to age at menopause, we analyzed data from a subset of menopausal females who had complete information on age at menopause (N = 9693). Based on previous studies binning age at menopause in approximately 5-year group bins (Gordon et al., 1978; Wasti et al., 1993), we applied similar age at menopause (i.e., Menopause Age Group) categories to our sample using the following bins: 40-45 years, 46-50 years, 51-55 years, and 56-62 years (Table 2). Since the average age of menopause is typically around 51.5 years and perimenopause lasts on average 4 years (Brinton et al., 2015; Harlow et al., 2012), we excluded females with age at menopause <40 years (N = 205) and >62 years (N = 4) from our female-specific analyses to ensure that results were not driven by extreme values. We also excluded participants who had undergone a hysterectomy and/or oophorectomy (N = 714) before natural menopause, as these females often experience premature menopause associated with their surgery and may be at elevated risk of dementia (Rocca et al., 2007; total N of participants included = 8770). As a cross-check, we repeated the interaction analyses including all participants. Distributions for BMI, WHR, and BF% within each Menopause Age Group are shown in SI Figure 2.

To investigate whether effects of CMRs and APOE risk varied across different age groups, we categorized participants' ages (i.e., Age Group) using the following bins: 45–55 years, 56–65 years, and 66–82 years (Table 2). The bins were selected to take the full cohort age range into account, and to enable comparisons of effects in middle-age and different stages of older adulthood in line with



**FIGURE 1** The matrices (top row) show the correlations (Pearson's coefficients) between BMI, WHR, and BF% for all participants, as well as females and males separately. The scatter plots (bottom row) show the correlations for all participants with males plotted in red and females in blue

**TABLE 2**Number of participants in each Menopause Age Groupand each Age Group, separated by sex and with % of APOE4 carriersin brackets

	Female	
Menopause Age Group	N (APOE4)	
40-45 years	986 (27.48%)	
46-50 years	2842 (26.39%)	
51–55 years	4059 (27.32%)	
56-62 years	883 (25.93%)	
	Female	Male
Age Group	N (APOE4)	N (APOE4)
45-55 years	1049 (29.26%)	1651 (26.95%)
56-65 years	4127 (27.16%)	3716 (25.61%)
66-82 years	3594 (25.90%)	5238 (24.91%)

previous studies examining participants throughout the adult lifespan (Grady et al., 2006; Subramaniapillai, Rajagopal, et al., 2019; Subramaniapillai, Rajah, et al., 2019). Distributions for BMI, WHR, and BF % within each Age Group are shown in SI Figure 2.

To test if results were consistent between Menopause Age Group and Age Group bins and the continuous measures of these variables, we conducted supplementary analyses using the continuous variables of Age at Menopause and Age.

#### 2.7 | Statistical analyses

The statistical analyses were conducted using Python 3.7.6 and R version 3.5. All variables were standardized (subtracting the mean and dividing by the SD) before being entered into the analyses. *p*-values were corrected for multiple comparisons using false discovery rate (FDR) correction (Benjamini & Hochberg, 1995). We report the *F*-test significance of each main effect and each interaction, as *F* is useful for interpreting models containing categorical variables with more than two levels. The *F*-statistics were generated in R by adding the *Anova* wrapper function to the linear model (*Im*) of interest.

We first determined whether there were sex differences in the effects of CMRs and APOE genotype on WM BAG, with the dependent variable representing BAG values based on the mixed-sex age prediction model (BAG<sub>ms</sub>). In order to adjust for age-dependence in CMRs and APOE4 status, age was included as a covariate. The following *lm* was used for these analyses, with *x* representing each CMR (BMI, WHR, and BF%) or APOE4 status, respectively:

$$BAG_{ms} \sim x \times Sex + Age.$$
 (1)

To test if interactions between sex and CMRs varied according to APOE4 status, the following *Im* was used with *CMR* representing each risk factor (BMI, WHR, and BF%):

$$BAG_{ms} \sim CMR \times Sex \times APOE + Age.$$
 (2)

icipants in each M

We then used the BAG values from the female-specific model to determine whether CMR and APOE effects on BAG varied according to age at menopause in females (Menopause Age Group). The following *lm* was used for these analyses, with  $BAG_{ss}$  representing BAG estimates based on the sex-specific model, and *x* representing each CMR or APOE4 status, respectively:

$$BAG_{ss} \sim x \times Menopause Age Group + Age.$$
 (3)

To test if interactions between Menopause Age Group and CMR varied according to APOE genotype, the following *Im* was used with *CMR* representing each risk factor:

$$BAG_{ss} \sim CMR \times Menopause Age Group \times APOE + Age.$$
 (4)

To determine whether effects in females and males varied across age bins, we ran analyses within each sex assessing the interactions of CMRs, APOE genotype, and Age Group on BAG. Sex-specific BAG values were used as dependent variables in the following *Im*, with *x* representing each CMR or *APOE4* status, respectively:

$$BAG_{ss} \sim x \times Age Group.$$
 (5)

To test if interactions between Age Group and CMRs varied according to APOE genotype, the following *lm* was used with *CMR* representing each risk factor:

$$BAG_{ss} \sim CMR \times Age Group \times APOE.$$
 (6)

As a follow-up, we ran the following *lm* within each of the Age Groups (for each sex) adjusting for effects of age within each Age Group bin:

$$BAG_{ss} \sim CMR \times APOE + Age.$$
 (7)

#### 3 | RESULTS

#### 3.1 | Brain age prediction

The accuracy of the age prediction models are shown in Table 3. SI Tables 1 and 2 depict the top 10 WM features for the mixed-sex and sex-specific age predictions, with the majority of the features overlapping between the three models. The model accuracy and prediction values were highly consistent when using a held-out validation sample for model optimization, as shown in SI Table 3.

## 3.2 | Sex differences in effects of CMRs and APOE genotype on WM BAG

To assess sex differences in the associations between BAG and BMI, WHR, BF%, and APOE4 status, we tested for sex-interactions as

# **TABLE 3** Age prediction accuracy for the mixed-sex and sex-specific models, including average $R^2$ , root mean square error (RMSE), mean absolute error (MAE), and correlations (*r*) between predicted and chronological age

Model	R <sup>2</sup>	RMSE	MAE	r [95% Cl]	р
Mixed (N = 35,740)	0.51 ± 0.009	5.27 ± 0.007	4.23 ± 0.005	0.72[0.71, 0.72]	<.0001
Male (N = 16,909)	0.50 ± 0.017	5.39 ± 0.115	4.34 ± 0.080	0.71[0.70, 0.72]	<.0001
Female (N = 18,831)	0.49 ± 0.015	5.28 ± 0.081	4.27 ± 0.070	0.69[0.69, 0.70]	<.0001

Abbreviation: CI, confidence interval.

**TABLE 4**Sex differences in the associations between Brain AgeGap and body mass index (BMI), waist-to-hip-ratio (WHR), body fatpercentage (BF%), and APOE4 status based on Formula (1) inSection 2.7

Interaction	F	p	pcorr
$Sex\timesBMI$	29.15	$\textbf{6.76}\times\textbf{10}^{-8}$	$\textbf{1.35}\times\textbf{10}^{-7}$
$Sex\timesWHR$	11.45	$7.2  imes 10^{-4}$	$9.56  imes 10^{-4}$
$Sex\timesBF\%$	43.37	$4.65\times10^{-11}$	$\textbf{1.86}\times\textbf{10}^{-\textbf{10}}$
$Sex \times APOE$	0.93	0.34	0.34

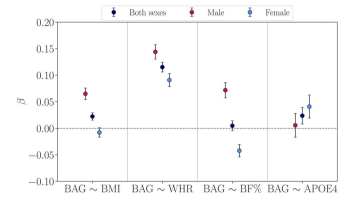
Note: Degrees of freedom = (1, 21, 303).

**TABLE 5**The interaction between Menopause Age Group and<br/>body mass index (BMI), waist-to-hip-ratio (WHR), body fat percentage<br/>(BF%), and APOE4 status on Brain Age Gap in females (Formula 3,<br/>Section 2.7)

Interaction	F	р	pcorr
$\text{BMI} \times \text{Menopause Age Group}$	2.00	.11	.22
WHR $\times$ Menopause Age Group	1.50	.21	.28
$BF\% \times Menopause$ Age Group	2.08	.10	.22
$APOE \times Menopause \ Age \ Group$	0.73	.53	.53

Note: Degrees of freedom = (3, 8761).

described in Section 2.7. The analyses revealed significant interaction effects of Sex  $\times$  BMI, Sex  $\times$  WHR, and Sex  $\times$  BF% on BAG, as shown in Table 4 and Figure 2. Greater WHR was associated with higher BAG (older brain age relative to chronological age) across sexes, but the effect was more prominent in males compared to females. BF% showed diverging associations, with higher values related to higher BAG in males and higher values related to lower BAG (younger brain age relative to chronological age) in females. Higher BMI values were associated with higher BAG in males, while no significant effect of BMI was seen in females. We observed no significant main effects of APOE genotype, and no significant interactions between APOE genotype and Sex or CMRs (see SI Table 4 for full results). Figure 2 shows the beta values for the BAG associations with each CMR and APOE4 status for both sexes, and for males and females separately. For comparison, we produced the same plot using estimates based on the sex-specific models (BAG values estimated relative to sex-specific age trajectories), as shown in SI Figure 3. The results showed similar patterns, with associations between higher BAG and greater BMI, WHR, and BF% in males, and between higher BAG and greater WHR, but not BMI and BF%, in females.



**FIGURE 2** Associations between WM Brain Age Gap (BAG) and each cardiometabolic risk factor as well as APOE4 status for both sexes, and for males and females separately.  $\beta$  (y-axis) represents the beta value (slope) for each association, for example, a positive  $\beta$  value indicates an association between greater cardiometabolic measures or APOE4 and higher BAG (older brain age relative to chronological age). The error bars represent standard errors on the  $\beta$ . BF%, body fat percentage; BMI, body mass index; WHR, waist-to-hip ratio

## 3.3 | Sex- and age-specific effects of CMRs and APOE genotype on WM BAG

To assess whether effects of CMRs and *APOE4* status varied according to age at menopause or across specific age periods, we next performed a series of regressions testing for interactions with Menopause Age Group in females and Age Group in males and females separately, as described in Sections 2.6 and 2.7.

## 3.3.1 | Effects of Menopause Age Group $\times$ CMRs and APOE genotype on WM BAG

In females, no interactions of CMR measures and APOE4 status with Menopause Age Group were found, as shown in Table 5 (see SI Table 5 for full results). The results were consistent when using the continuous Age at Menopause variable, as shown in SI Table 6, and when including participants with age at menopause <40 and >62 years, hysterectomy and/or oophorectomy, as shown in SI Tables 7–9. Across models, there was a main effect of Menopause Age Group such that a lower age at menopause was associated with higher WM BAG (see SI Table 5 and SI Figure 2). This effect was consistent when using the continuous Age at Menopause variable in a regression model including hormone replacement therapy use, education, income, Townsend Deprivation Index, alcohol intake, physical

7

WILFY⊥

8

**TABLE 6**The interaction between Age Group and BMI, WHR, BF%, and APOE genotype on Brain Age Gap (Formula 5, Section 2.7)

	Female			Male		
Interaction	F	р	pcorr	F	р	pcorr
$BMI\timesAge\:Group$	1.96	.14	.25	2.33	.10	.39
$WHR\timesAge\:Group$	1.68	.19	.25	0.72	.49	.49
$\text{BF\%} \times \text{Age Group}$	1.76	.17	.25	1.27	.28	.49
$APOE \times Age \ Group$	0.31	.73	.73	0.87	.42	.49

Note: Degrees of freedom = (2, 8764) and (2, 10,599) for females and males, respectively.

activity, and number of childbirths in addition to age, BMI, and APOE4 status as covariates ( $\beta = -0.036 \pm 0.010$ ,  $p = 1.40 \times 10^{-4}$ ; see SI Section 6 for details about the covariates).

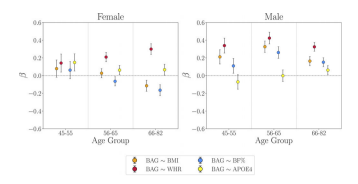
## 3.3.2 | Effects of age group $\times$ CMRs and APOE genotype on WM BAG

To assess whether effects of CMRs and APOE4 status varied across specific age periods, we applied regressions including interaction terms with Age Group as described in Formula (5), Section 2.7. For both females and males, no significant effects were found for Age Group  $\times$  CMR measures/APOE4 status on BAG (Table 6 and SI Table 10), indicating that the BAG associations with each CMR did not vary significantly between age groups. The results were consistent when using the continuous Age variable (SI Table 11). Figure 3 shows the associations between BAG and each CMR within each Age Group, and SI Figure 5 shows the associations grouped by APOE genotype.

While the BAG associations with each CMR were not significantly different between age groups (Table 6), the follow-up regression analyses within each age group indicated that the divergence between the associations with WHR versus BMI/BF% increased with older age in females (Figure 3; see SI Tables 12 and 13 for full results). As a post hoc test, we estimated the differences between the WHR versus BMI/BF% associations for females within each age group using a *Z* test for correlated samples, as described in SI Section 7. The results confirmed that the divergence between the BAG associations with WHR versus BMI/BF% increased with age, and was most prominent in the oldest age group (SI Table 14).

#### 4 | DISCUSSION

This study investigated whether associations between BAG and CMRs and APOE genotype varied (i) between males and females, (ii) according to age at menopause in females, and (iii) across different age groups in males and females. In summary, the results showed sex differences in associations between BAG and all three CMRs, with stronger positive associations among males compared to females. Higher BAG (older brain age relative to chronological age) was



**FIGURE 3** Associations between WM BAG and cardiometabolic risk factors as well as APOE4 status within each Age Group bin (Formula 7, Section 2.7).  $\beta$  (y-axis) represents the beta value (slope) for each association. The error bars represent standard errors on the  $\beta$ . BF%, body fat percentage; BMI, body mass index; WHR, waist-to-hip ratio

associated with greater BMI, WHR, and BF% in males, whereas in females, higher BAG was associated with greater WHR, but not BMI and BF%. Earlier age at menopause was linked to higher BAG in females, but no interactions were found between age at menopause and CMRs. While none of the associations between BAG and each CMR were significantly different between age groups, follow-up analyses indicated that the divergence between the WHR and BMI/BF% associations observed in females was most prominent within the oldest age group (66–81 years). APOE4 status showed no significant main effects on BAG, no age- or age at menopause-specific effects, and no significant interactions with CMRs. The findings demonstrate sex-specific associations between body fat composition and brain age, emphasizing the importance of analyzing males and females separately in studies addressing CMR in aging.

The analyses including sex as an interaction term showed that all three CMRs were associated with higher BAG in males relative to females. These findings support a recent study, which revealed that males may be more vulnerable than females to WM brain aging in the presence of greater BMI (Algarni et al., 2021) and, more broadly, greater cardiometabolic burden (Assareh et al., 2014; Filomena et al., 2015; Geerlings et al., 2010; Jongen et al., 2007). Our findings may shed some light on sex differences in the timing of cardiometabolic disease, emerging on average earlier in males than females (Maas & Appelman, 2010). Unlike females, it is possible that the combined effect of all three CMRs on males' higher BAG as early as midlife may, in turn, be associated with their earlier cardiometabolic disease risk. However, these measures of body adiposity may also be more sensitive to males (Foret et al., 2021), while other unexplored CMRs, such as hypertension, may be more robust in capturing females' health risk earlier in adulthood (Gilsanz et al., 2017; Wei et al., 2017). Furthermore, sex differences in adipose fat distribution may differentially contribute to brain age, with males more likely having fat distributed in the visceral adipose tissue surrounding the abdominal organs, while females tend to have more subcutaneous adipose tissue (Bredella, 2017; Chang et al., 2018). Compared to the latter, greater visceral adipose fat distribution is associated with

elevated risk of cardiometabolic disease. In females, differential effects of BF% and BMI compared to WHR were observed, with divergent associations particularly prominent in the oldest age group (66-82 years) where lower BF% was linked to higher BAG. This could reflect that lower BF% in older females may be an indicator of frailty and preclinical dementia, and/or indicate a protective role of certain sources of adipose fat in females at later ages (Buchman et al., 2005; Johnson et al., 2006; Klosinski et al., 2015; Subramaniapillai et al., 2021). While our results showed that earlier menopause transition was associated with greater BAG, we found no significant interactions between age at menopause and CMRs. Further longitudinal work is required to clarify the role of adipose tissue in female brain health, and how it may relate to sex-specific factors including the menopausal transition.

The evidence for the role of endogenous estrogen exposure in neurodegeneration and AD is controversial, with some studies reporting an association between shorter reproductive span and greater dementia risk (e.g., Gilsanz et al., 2019), while others have found that a longer reproductive span (i.e., later age of menopause) did not confer protective effects (e.g., Geerlings et al., 2001; Mosconi et al., 2018; Najar et al., 2020). Some studies have reported genotypespecific effects, with a longer reproductive span conferring greater risk in APOE4 carriers (Geerlings et al., 2001), and older brain age linked to greater estradiol levels in APOE4 carriers and lower levels in noncarriers (de Lange, Barth, et al., 2020). These studies point to a possible modulatory role of estrogen exposure (Barth & de Lange, 2020), typically having beneficial effects, but potentially becoming neurotoxic in the context of greater AD pathology (Jack Jr et al., 2010) or diseased cell populations (i.e., "healthy cell" hypothesis: Brinton, 2005). While future studies including detailed data about the menopausal transition (i.e., pre-, peri, postmenopause) as well as specific measures of AD-related brain pathology (Rahman et al., 2020) are required to test these hypotheses, our findings showed a small but significant association between earlier menopause transition and higher WM BAG in the current cohort, in line with studies linking a shorter reproductive span to risk for neurodegeneration (Fox et al., 2013; Geerlings et al., 2001; Gilsanz et al., 2019; Mishra & Brinton, 2018; Schelbaum et al., 2021; Scheyer et al., 2018).

Unlike females, in which menses cessation is a marker of menopause status, males have a more gradual endocrine aging process, with no significant marker for age at andropause. Although low levels of body fat in older age may be an indicator of frailty in both sexes (Buchman et al., 2005; Johnson et al., 2006), our results show that in this sample, greater WHR, BMI, and BF% were consistently linked to higher BAG in males. This corresponds with previous studies reporting negative effects of higher body fat levels on brain health in males across midlife and older ages (Dekkers et al., 2019; Taki et al., 2008). The male endocrine aging process typically involves gradual declines in testosterone levels. Greater adipose tissue may increase levels of aromatase, an enzyme that converts testosterone to estrogen, which in males may be associated with their accelerated endocrine and consequently, brain aging process (Blouin et al., 2006; Meyer et al., 2011; Vosberg et al., 2021). Therefore, while greater adipose tissue can potentially act as a source of estrogen in postmenopausal females, it can be detrimental to males, in whom this may result in reduced testosterone levels (see Vosberg et al., 2021 demonstrating that the genetic architecture of testosterone contributes to sex differences in cardiometabolic traits in the UK Biobank). Importantly, increasing evidence points to the role of sex hormones in mediating cerebrovascular function, in which dysregulation is linked to cerebrovascular diseases, cognitive impairment, and dementia (see reviews by Gannon et al., 2019; Robison et al., 2019). Further research on the links between sex hormones and cardiometabolic factors in endocrine aging may help inform sex-specific health interventions for both males and females.

No significant effects were found for APOE genotype. This is in line with our previous study showing no effects of APOE4 status or polygenic risk for AD on gray-matter based brain age in the UK Biobank sample (de Lange, Barth, et al., 2020), as well as a recent UK Biobank study showing that APOE4 genotype was associated with WM hyperintensities, but not with FA or MD in WM tracts (Lvall et al., 2020). While our age prediction was based on several diffusion models known to be sensitive to WM aging (Beck, de Lange, Maximov, et al., 2021; Jelescu & Budde, 2017; Jensen et al., 2005), it is possible that specific estimates of WM hyperintensities could vield APOE-sensitive CMR associations. Furthermore, a recent UK Biobank study revealed region- and metric-specific effects of age and sex on WM microstructure (Lawrence et al., 2021). Although age prediction models combine a rich variety of WM characteristics into single estimates, global BAG estimates do not provide specific information about regional WM connections. Hence, future studies may aim to investigate regional and diffusion metric-specific estimates of brain aging in relation to APOE genotype and CMRs. Modality-specific BAG estimates are also relevant for identifying differences in brain tissue affected by a specific condition or disease (Beck, de Lange, Pedersen, et al., 2021; Cole, 2020; de Lange, Anatürk, et al., 2020; Rokicki et al., 2020). For example, one of our previous studies found that BMI interacted with AD risk to influence gray-matter based BAG, such that females with greater AD risk benefited more from a higher BMI (Subramaniapillai et al., 2021). While the current study focused on WM measures given their susceptibility to CMRs, future studies may aim to include several brain modalities to directly compare sex- and age-specific effects. More detailed measures of fat distribution obtained with body MRI (Beck, de Lange, Alnaes, et al., 2021; Gurholt et al., 2021; Leinhard et al., 2008; Linge et al., 2018) may also clarify the divergent associations observed in females, and provide a more complete understanding of adipose tissue distribution in relation to cardiometabolic disease (Linge et al., 2018; Linge et al., 2019), AD risk (Diehl-Wiesenecker et al., 2015), and endocrine aging processes (El Khoudary et al., 2015). Due to sample size restrictions in relation to our study goals of determining sex- and age-specific effects, we could not currently probe body MRI in our subgroups, but ongoing data collection of this measure from UK Biobank participants will render future analyses of these measures feasible.

Although the large UK Biobank cohort enabled us to investigate whether effects were sensitive to specific age- or age at menopause ⊥WILEY-

periods, the sample sizes were limited by probing variables with different subgroups (i.e., dividing our sample across sex, APOE genotype, and Menopause/Age Group levels). However, our sub-group samples (n > 250) are still large compared to the majority of previous studies investigating sex- or age-specific effects on brain structure (Szucs & Ioannidis, 2020). The results also highlight potential causes of mixed findings in the literature: variations in associations reported across studies could be due to not separating analyses by sex (see Figure 2), investigating samples with different age ranges (see Figure 3 and SI Table 8), and/or the use of different CMRs. As participant re-testing of the UK Biobank baseline cohort is actively underway, our future work will aim to integrate longitudinal investigations of brain age, as the current cross-sectional analyses prevent any conclusions about causality. Longitudinal designs may also enable differentiation between age-specific effects and effects that emerge as a result of a selective attrition or survival bias (Heffernan et al., 2016; Jacobsen et al., 2021; Munafò et al., 2018; Salthouse, 2014). While UK Biobank provides an excellent resource of open-access population health data. the cohort is homogeneous with regard to ethnic background and education, and characterized by a "healthy volunteer effect" (Fry et al., 2017), indicating that it is not representative of the general population (Keyes & Westreich, 2019). Although the current results may not generalize to populations beyond those represented in this cohort, our findings may prompt further study into sex- and age-specific effects of CMR as well as endocrine aging. Lastly, since neural aging processes are multi-factorial, single risk factors can only explain parts of the individual variation. Hence, future studies may aim to go beyond investigating risk factors in isolation and adopt approaches that can model complex relationships between a variety of geneenvironment interactions and brain health in aging (Mulugeta et al., 2021; Wang et al., 2020).

In conclusion, this study demonstrates notable sex differences in associations between body fat indices and WM brain age, underlining the importance of stratifying samples by sex in population-based and clinical studies (Clayton, 2018; Ewelina et al., 2020; Ferretti et al., 2018; Miller et al., 2017; Shansky & Murphy, 2021; Shansky & Woolley, 2016). Independent of cardiometabolic profile, earlier menopause transition was associated with higher BAG in females. Hence, considering effects of both chronological and endocrine aging may increase our understanding of sex-specific brain aging trajectories and disease prevalence (Jacobs & Goldstein, 2018; Taylor et al., 2019). Given the historical lack of research into sex-specific influences on brain health and disease (Cirillo et al., 2020; de Lange, Jacobs, & Galea, 2020; Ferretti & Santuccione Chadha, 2021; Taylor et al., 2019), future studies incorporating sex as a variable of interest may provide valuable contributions to precision medicine research, consequently improving health care for both males and females.

#### ACKNOWLEDGMENTS

This research has been conducted using the UK Biobank under Application 27412. UKB has received ethics approval from the National Health Service National Research Ethics Service (ref 11/NW/0382). The work was performed on the Service for Sensitive Data (TSD)

platform, owned by the University of Oslo, operated and developed by the TSD service group at the University of Oslo IT-Department (USIT). Computations were also performed using resources provided by UNINETT Sigma2-the National Infrastructure for High Performance Computing and Data Storage in Norway. While working on this study, the authors received funding from the Women in Cognitive Science-Canada (Sivaniya Subramaniapillai), NSERC Michael Smith Foreign Study Supplement (Sivaniya Subramaniapillai), Healthy Brains Healthy Lives (Sivaniya Subramaniapillai), the Alzheimer's Society (Sana Suri; 441), the Academy of Medical Sciences/the Wellcome Trust/the Government Department of Business, Energy and Industrial Strategy/the British Heart Foundation/Diabetes UK Springboard Award (Sana Suri: SBF006\1078), the Research Council of Norway (Lars T. Westlye; 273345, 249795, 298646, 300768, 223273; Ole A. Andreassen; 223273, 283799, 2837989), the South-Eastern Norway Regional Health Authority (Lars T. Westlye; 2018076, 2019101, Ole A. Andreassen; 2019-108, 2017-112, Tiril P. Gurholt; 2022080), the European Research Council under the European Union's Horizon 2020 research and innovation programme (Lars T. Westlve: 802998. Ole A. Andreassen: 847776. Klaus P. Ebmeier: 732592), the HDH Wills 1965 Charitable Trust (Klaus P. Ebmeier; 1117747), the Leenaards Foundation (Bogdan Draganski), and the Swiss National Science Foundation (Ann-Marie G. de Lange; PZ00P3\_193658; Bogdan Draganski; NCCR Synapsy, project grants Nr 32003B 135679, 32003B 159780, 324730 192755, and CRSK-3 190185). The Wellcome Centre for Integrative Neuroimaging is supported by core funding from the Wellcome Trust (203139/ Z/16/Z).

#### DATA AVAILABILITY STATEMENT

The data that support the findings of this study are available through the UK Biobank data access procedures (https://www.ukbiobank.ac. uk/researchers). Code for running the age prediction models is available at https://github.com/amdelange/brainage\_women.

#### ORCID

Sivaniya Subramaniapillai https://orcid.org/0000-0003-3207-0700 Sana Suri https://orcid.org/0000-0002-5136-8302 Claudia Barth https://orcid.org/0000-0001-6544-0945 Ivan I. Maximov https://orcid.org/0000-0001-6319-6774 Irene Voldsbekk https://orcid.org/0000-0001-9393-5857 Dennis van der Meer https://orcid.org/0000-0002-0466-386X Tiril P. Gurholt https://orcid.org/0000-0002-1272-7616 Dani Beck https://orcid.org/0000-0002-1272-7616 Dani Beck https://orcid.org/0000-0002-0974-9304 Bogdan Draganski https://orcid.org/0000-0002-5159-5919 Ole A. Andreassen https://orcid.org/0000-0002-5159-5919 Ole A. Andreassen https://orcid.org/0000-0002-5190-7038 Lars T. Westlye https://orcid.org/0000-0001-8644-956X Ann-Marie G. de Lange https://orcid.org/0000-0002-5150-6656

#### REFERENCES

Alfaro, F. J., Gavrieli, A., Saade-Lemus, P., Lioutas, V.-A., Upadhyay, J., & Novak, V. (2018). White matter microstructure and cognitive decline in metabolic syndrome: A review of diffusion tensor imaging. *Metabolism*, 78, 52–68.

- Alfaro-Almagro, F., Jenkinson, M., Bangerter, N. K., Andersson, J. L., Griffanti, L., Douaud, G., Sotiropoulos, S. N., Jbabdi, S., Hernandez-Fernandez, M., Vallee, E., Vidaurre, D., Webster, M., McCarthy, P., Rorden, C., Daducci, A., Alexander, D. C., Zhang, H., Dragonu, I., Matthews, P. M., ... Smith, S. M. (2018). Image processing and quality control for the first 10,000 brain imaging datasets from UK Biobank. *NeuroImage*, 166, 400–424.
- Alqarni, A., Jiang, J., Crawford, J. D., Koch, F., Brodaty, H., Sachdev, P., & Wen, W. (2021). Sex differences in risk factors for white matter hyperintensities in non-demented older individuals. *Neurobiology of Aging*, 98, 197–204.
- Armstrong, N. M., An, Y., Beason-Held, L., Doshi, J., Erus, G., Ferrucci, L., Davatzikos, C., & Resnick, S. M. (2019). Sex differences in brain aging and predictors of neurodegeneration in cognitively healthy older adults. *Neurobiology of Aging*, 81, 146–156.
- Assareh, A. A., Mather, K. A., Crawford, J. D., Wen, W., Anstey, K. J., Easteal, S., Tan, X., Mack, H. A., Kwok, J. B., Schofield, P. R., & Sachdev, P. S. (2014). Renin-angiotensin system genetic polymorphisms and brain white matter lesions in older Australians. *American Journal of Hypertension*, 27, 1191–1198.
- Barth, C., & de Lange, A.-M. G. (2020). Towards an understanding of women's brain aging: The immunology of pregnancy and menopause. *Frontiers in Neuroendocrinology*, 100850, 100850.
- Basser, P. J., Mattiello, J., & LeBihan, D. (1994). MR diffusion tensor spectroscopy and imaging. *Biophysical Journal*, 66, 259–267.
- Beck, D., de Lange, A.-M., Maximov, I. I., Richard, G., Andreassen, O. A., Nordvik, J. E., & Westlye, L. T. (2021). White matter microstructure across the adult lifespan: A mixed longitudinal and cross-sectional study using advanced diffusion models and brain-age prediction. *NeuroImage*, 224, 117441.
- Beck, D., de Lange, A.-M. G., Alnaes, D., Maximov, I. I., Pedersen, M. L., Leinhard, O. D., Linge, J., Simon, R., Richard, G., Dorum, E. S., Kolskår, K. K., Sanders, A.-M., Winterton, A., Gurholt, T. P., Kaufmann, T., Steen, N. E., Nordvik, J. E., Andreassen, O. A., & Westlye, L. T. (2021). Adipose tissue distribution from body MRI is associated with cross-sectional and longitudinal brain age in adults. *NeuroImage: Clinical*, 33, 102949.
- Beck, D., de Lange, A.-M. G., Pedersen, M. L., Alnæs, D., Maximov, I. I., Voldsbekk, I., Richard, G., Sanders, A.-M., Ulrichsen, K. M., Dørum, E. S., Kolskår, K. K., Høgestøl, E. A., Steen, N. E., Djurovic, S., Andreassen, O. A., Nordvik, J. E., Kaufmann, T., & Westlye, L. T. (2021). Cardiometabolic risk factors associated with brain age and accelerate brain ageing. *Human Brain Mapping*, 43, 700–720.
- Benjamini, Y., & Hochberg, Y. (1995). Controlling the false discovery rate: A practical and powerful approach to multiple testing. *Journal of the Royal Statistical Society: Series B (Methodological)*, 57, 289–300.
- Biondo, F., Jewell, A., Pritchard, M., Aarsland, D., Steves, C. J., Mueller, C., & Cole, J. H. (2021). Brain-age predicts subsequent dementia in memory clinic patients. *medRxiv*.
- Biondo, F., Jewell, A., Pritchard, M., Mueller, C., Steves, C. J., & Cole, J. (2020). Brain-age predicts subsequent dementia in memory clinic patients: Neuroimaging/optimal neuroimaging measures for early detection. *Alzheimer's & Dementia*, 16, e037378.
- Birdsill, A. C., Oleson, S., Kaur, S., Pasha, E., Ireton, A., Tanaka, H., & Haley, A. (2017). Abdominal obesity and white matter microstructure in midlife. *Human Brain Mapping*, 38, 3337–3344.
- Biskup, E., Quevenco, F.-C., Ferretti, M. T., & Santuccione-Chadha, A. (2019). Sex differences in brain metabolic activity: Beyond the concept of brain age. *Proceedings of the National Academy of Sciences*, 116, 10630–10631.
- Blouin, K., Richard, C., Brochu, G., Hould, F.-S., Lebel, S., Marceau, S., Biron, S., Luu-The, V., & Tchernof, A. (2006). Androgen inactivation

and steroid-converting enzyme expression in abdominal adipose tissue in men. *Journal of Endocrinology*, 191, 637–649.

- Bowman, K., Atkins, J. L., Delgado, J., Kos, K., Kuchel, G. A., Ble, A., Ferrucci, L., & Melzer, D. (2017). Central adiposity and the overweight risk paradox in aging: Follow-up of 130,473 UK Biobank participants. *The American Journal of Clinical Nutrition*, 106, 130–135.
- Bradbury, K. E., Guo, W., Cairns, B. J., Armstrong, M. E., & Key, T. J. (2017). Association between physical activity and body fat percentage, with adjustment for BMI: A large cross-sectional analysis of UK Biobank. *BMJ Open*, 7, e011843.
- Bredella, M. A. (2017). Sex differences in body composition. Advances in Experimental Medicine and Biology, 1043, 9–27.
- Bretsky, P., Buckwalter, J., Seeman, T., Miller, C., Poirier, J., Schellenberg, G., Finch, C., & Henderson, V. (1999). Evidence for an interaction between apolipoprotein e genotype, gender, and Alzheimer disease. *Alzheimer Disease and Associated Disorders*, 13, 216–221.
- Brinton, R. D. (2005). Investigative models for determining hormone therapy-induced outcomes in brain: Evidence in support of a healthy cell bias of estrogen action. Annals of the New York Academy of Sciences, 1052, 57–74.
- Brinton, R. D., Yao, J., Yin, F., Mack, W. J., & Cadenas, E. (2015). Perimenopause as a neurological transition state. *Nature Reviews Endocrinology*, 11, 393–405.
- Buchman, A. S., Wilson, R. S., Bienias, J. L., Shah, R. C., Evans, D. A., & Bennett, D. A. (2005). Change in body mass index and risk of incident Alzheimer disease. *Neurology*, 65, 892–897.
- Bycroft, C., Freeman, C., Petkova, D., Band, G., Elliott, L. T., Sharp, K., Motyer, A., Vukcevic, D., Delaneau, O., O'Connell, J., Cortes, A., Welsh, S., Young, A., Effingham, M., McVean, G., Leslie, S., Allen, N., Donnelly, P., & Marchini, J. (2018). The UK Biobank resource with deep phenotyping and genomic data. *Nature*, *562*, 203–209.
- Chang, E., Varghese, M., & Singer, K. (2018). Gender and sex differences in adipose tissue. *Current Diabetes Reports*, 18, 1–10.
- Chen, T. & Guestrin, C. (2016). XGBoost: A scalable tree boosting system. Proceedings of the 22nd ACM SIGKDD International Conference on Knowledge Discovery and Data Mining; 2016; pp. 785–794.
- Chen, W., Xu-Hong, H., Zhang, M.-L., Yu-Qian, B., Yu-Hua, Z., Zhong, W.-H., Xiang, K.-S., & Wei-Ping, J. (2010). Comparison of body mass index with body fat percentage in the evaluation of obesity in Chinese. *Biomedical and Environmental Sciences*, 23, 173–179.
- Cirillo, D., Catuara-Solarz, S., Morey, C., Guney, E., Subirats, L., Mellino, S., Gigante, A., Valencia, A., Rementeria, M. J., Chadha, A. S., & Mavridis, N. (2020). Sex and gender differences and biases in artificial intelligence for biomedicine and healthcare. NPJ Digital Medicine, 3, 1–11.
- Clayton, J. A. (2018). Applying the new SABV (sex as a biological variable) policy to research and clinical care. Physiology & Behavior, 187, 2–5.
- Cole, J., Raffel, J., Friede, T., Eshaghi, A., Brownlee, W., Chard, D., De Stefano, N., Enzinger, C., Pirpamer, L., Filippi, M., C Gasperini, MA Rocca, A Rovira, S Ruggieri, J Sastre-Garriga, ML Stromillo, BMJ Uitdehaag, H Vrenken, F Barkhof, R Nicholas, O Ciccarelli, the MAGNIMS study group. (2019). Accelerated brain ageing and disability in multiple sclerosis. *bioRxiv.* 584888.
- Cole, J. H. (2020). Multi-modality neuroimaging brain-age in UK Biobank: Relationship to biomedical, lifestyle and cognitive factors. *Neurobiology* of Aging, 92, 34–42.
- Cole, J. H., & Franke, K. (2017). Predicting age using neuroimaging: Innovative brain ageing biomarkers. *Trends in Neurosciences*, 40, 681–690.
- Cole, J. H., Marioni, R. E., Harris, S. E., & Deary, I. J. (2019). Brain age and other bodily 'ages': Implications for neuropsychiatry. *Molecular Psychiatry*, 24, 266–281.
- de Lange, A., Jacobs, E. G., & Galea, L. A. (2020). The scientific body of knowledge: Whose body does it serve? A spotlight on women's brain health. *Frontiers in Neuroendocrinology*, 60, 100898.

## <sup>12</sup> WILEY-

- de Lange, A.-M. G., Anatürk, M., Suri, S., Kaufmann, T., Cole, J. H., Griffanti, L., Zsoldos, E., Jensen, D. E., Filippini, N., Singh-Manoux, A., Kivimäki, M., Westlye, L. T., & Ebmeier, K. P. (2020). Multimodal brainage prediction and cardiovascular risk: The Whitehall II MRI sub-study. *NeuroImage*, 222, 117292.
- de Lange, A.-M. G., Barth, C., Kaufmann, T., Maximov, I. I., van der Meer, D., Agartz, I., & Westlye, L. T. (2020). Women's brain aging: Effects of sex-hormone exposure, pregnancies, and genetic risk for Alzheimer's disease. *Human Brain Mapping*, 41, 5141–5150.
- de Lange, A.-M. G., & Cole, J. H. (2020). Commentary: Correction procedures in brain-age prediction. *NeuroImage: Clinical*, 26, 102229.
- de Leeuw, F., de Groot, J. C., Achten, E., Oudkerk, M., Ramos, L., Heijboer, R., Hofman, A., Jolles, J., van Gijn, J., & Breteler, M. (2001). Prevalence of cerebral white matter lesions in elderly people: A population based magnetic resonance imaging study. The Rotterdam scan study. *Journal of Neurology, Neurosurgery & Psychiatry*, 70, 9–14.
- DeCarli, C., Massaro, J., Harvey, D., Hald, J., Tullberg, M., Au, R., Beiser, A., D'Agostino, R., & Wolf, P. A. (2005). Measures of brain morphology and infarction in the Framingham heart study: Establishing what is normal. *Neurobiology of Aging*, 26, 491–510.
- Dekkers, I. A., Jansen, P. R., & Lamb, H. J. (2019). Obesity, brain volume, and white matter microstructure at MRI: A cross-sectional UK Biobank study. *Radiology*, 291, 763–771.
- Diehl-Wiesenecker, E., von Arnim, C. A., Dupuis, L., Mueller, H.-P., Ludolph, A. C., & Kassubek, J. (2015). Adipose tissue distribution in patients with Alzheimer's disease: A whole body MRI case-control study. *Journal of Alzheimer's Disease*, 48, 825–832.
- Dubnov, G., Brzezinski, A., & Berry, E. M. (2003). Weight control and the management of obesity after menopause: The role of physical activity. *Maturitas*, 44, 89–101.
- Egorova, N., Liem, F., Hachinski, V., & Brodtmann, A. (2019). Predicted brain age after stroke. *Frontiers in Aging Neuroscience*, 11, 348.
- El Khoudary, S. R., Shields, K. J., Janssen, I., Hanley, C., Budoff, M. J., Barinas-Mitchell, E., Everson-Rose, S. A., Powell, L. H., & Matthews, K. A. (2015). Cardiovascular fat, menopause, and sex hormones in women: The swan cardiovascular fat ancillary study. *The Journal of Clinical Endocrinology & Metabolism*, 100, 3304–3312.
- Elliott, M. L., Belsky, D. W., Knodt, A. R., Ireland, D., Melzer, T. R., Poulton, R., Ramrakha, S., Caspi, A., Moffitt, T. E., & Hariri, A. R. (2019). Brain-age in midlife is associated with accelerated biological aging and cognitive decline in a longitudinal birth cohort. *Molecular Psychiatry*, *26*, 3829–3838.
- Elliott, P., & Peakman, T. C. (2008). The UK Biobank sample handling and storage protocol for the collection, processing and archiving of human blood and urine. *International Journal of Epidemiology*, 37, 234–244.
- Ewelina, B., Julie, M., & Teresa, F. M. (2020). Gender medicine: Towards a gender-specific treatment of neuropsychiatric disorders. In *Handbook* of clinical neurology (Vol. 175, pp. 437–448). Elsevier.
- Fatemi, F., Kantarci, K., Graff-Radford, J., Preboske, G. M., Weigand, S. D., Przybelski, S. A., Knopman, D. S., Machulda, M. M., Roberts, R. O., Mielke, M. M., Petersen, R. C., Jack, C. R., Jr., & Vemuri, P. (2018). Sex differences in cerebrovascular pathologies on flair in cognitively unimpaired elderly. *Neurology*, 90, e466–e473.
- Ferretti, M. T., Iulita, M. F., Cavedo, E., Chiesa, P. A., Schumacher Dimech, A., Santuccione Chadha, A., Baracchi, F., Girouard, H., Misoch, S., Giacobini, E., Depypere, H., & Hampel, H. (2018). Sex differences in Alzheimer disease—The gateway to precision medicine, nature reviews. *Neurology*, 14, 457–469.
- Ferretti, M. T., & Santuccione Chadha, A. (2021). The missing x factor in Alzheimer disease. Nature Reviews. Neurology, 17, 1–2.
- Fieremans, E., Jensen, J. H., & Helpern, J. A. (2011). White matter characterization with diffusional kurtosis imaging. *NeuroImage*, 58, 177–188.
- Filomena, J., Riba-Llena, I., Vinyoles, E., Tovar, J. L., Mundet, X., Castañé, X., Vilar, A., López-Rueda, A., Jiménez-Baladó, J., Cartanyà, A., Montaner, J., Delgado, P., & ISSYS Investigators. (2015).

Short-term blood pressure variability relates to the presence of subclinical brain small vessel disease in primary hypertension. *Hypertension*, *66*, 634–640.

- Fitzpatrick, A. L., Kuller, L. H., Lopez, O. L., Diehr, P., O'Meara, E. S., Longstreth, W., & Luchsinger, J. A. (2009). Midlife and late-life obesity and the risk of dementia: Cardiovascular health study. *Archives of Neurology*, 66, 336–342.
- Foret, J. T., Dekhtyar, M., Cole, J. H., Gourley, D. D., Caillaud, M., Tanaka, H., & Haley, A. P. (2021). Network modeling sex differences in brain integrity and metabolic health. *Frontiers in Aging Neuroscience*, 13, 329.
- Fox, M., Berzuini, C., & Knapp, L. A. (2013). Cumulative estrogen exposure, number of menstrual cycles, and Alzheimer's risk in a cohort of british women. *Psychoneuroendocrinology*, 38, 2973–2982.
- Franke, K., & Gaser, C. (2012). Longitudinal changes in individual brainage in healthy aging, mild cognitive impairment, and Alzheimer's disease. *GeroPsych*, 25, 235–245.
- Franke, K., & Gaser, C. (2019). Ten years of brainage as a neuroimaging biomarker of brain aging: What insights have we gained? *Frontiers in Neurology*, 10, 789.
- Fry, A., Littlejohns, T. J., Sudlow, C., Doherty, N., Adamska, L., Sprosen, T., Collins, R., & Allen, N. E. (2017). Comparison of sociodemographic and health-related characteristics of UK Biobank participants with those of the general population. *American Journal of Epidemiology*, 186, 1026– 1034.
- Gannon, O., Robison, L., Custozzo, A., & Zuloaga, K. (2019). Sex differences in risk factors for vascular contributions to cognitive impairment & dementia. *Neurochemistry International*, 127, 38–55.
- Gaser, C., Franke, K., Klöppel, S., Koutsouleris, N., Sauer, H., & Alzheimer's Disease Neuroimaging Initiative. (2013). Brainage in mild cognitive impaired patients: Predicting the conversion to Alzheimer's disease. *PLoS One*, 8, e67346.
- Geerlings, M. I., Appelman, A. P., Vincken, K. L., Algra, A., Witkamp, T. D., Mali, W. P., van der Graaf, Y., & SMART Study Group. (2010). Brain volumes and cerebrovascular lesions on MRI in patients with atherosclerotic disease. The smart-MR study. *Atherosclerosis*, 210, 130–136.
- Geerlings, M. I., Ruitenberg, A., Witteman, J. C., van Swieten, J. C., Hofman, A., van Duijn, C. M., Breteler, M. M., & Launer, L. J. (2001). Reproductive period and risk of dementia in postmenopausal women. JAMA, 285, 1475–1481.
- Gerdts, E., & Regitz-Zagrosek, V. (2019). Sex differences in cardiometabolic disorders. *Nature Medicine*, *25*, 1657–1666.
- Gilsanz, P., Lee, C., Corrada, M. M., Kawas, C. H., Quesenberry, C. P., & Whitmer, R. A. (2019). Reproductive period and risk of dementia in a diverse cohort of health care members. *Neurology*, 92, e2005–e2014.
- Gilsanz, P., Mayeda, E. R., Glymour, M. M., Quesenberry, C. P., Mungas, D. M., DeCarli, C., Dean, A., & Whitmer, R. A. (2017). Female sex, early-onset hypertension, and risk of dementia. *Neurology*, *89*, 1886–1893.
- Gordon, T., Kannel, W. B., Hjortland, M. C., & McNamara, P. M. (1978). Menopause and coronary heart disease: The Framingham study. *Annals of Internal Medicine*, 89, 157–161.
- Grady, C. L., Springer, M. V., Hongwanishkul, D., McIntosh, A. R., & Winocur, G. (2006). Age-related changes in brain activity across the adult lifespan. *Journal of Cognitive Neuroscience*, 18, 227–241.
- Griffanti, L., Jenkinson, M., Suri, S., Zsoldos, E., Mahmood, A., Filippini, N., Sexton, C. E., Topiwala, A., Allan, C., Kivimäki, M., Singh-Manoux, A., Ebmeier, K. P., Mackay, C. E., & Zamboni, G. (2018). Classification and characterization of periventricular and deep white matter hyperintensities on MRI: A study in older adults. *NeuroImage*, 170, 174–181.
- Gurholt, T. P., Kaufmann, T., Frei, O., Alnæs, D., Haukvik, U. K., van der Meer, D., Moberget, T., O'Connell, K. S., Leinhard, O. D., Linge, J., Simon, R., Smeland, O. B., Sønderby, I. E., Winterton, A., Steen, N. E., Westlye, L. T., & Andreassen, O. A. (2021). Population-based body-

brain mapping links brain morphology with anthropometrics and body composition. *Translational Psychiatry*, 11, 1–12.

- Habes, M., Sotiras, A., Erus, G., Toledo, J. B., Janowitz, D., Wolk, D. A., Shou, H., Bryan, N. R., Doshi, J., Völzke, H., Schminke, U., Hoffmann, W., Resnick, S. M., Grabe, H. J., & Davatzikos, C. (2018). White matter lesions: Spatial heterogeneity, links to risk factors, cognition, genetics, and atrophy. *Neurology*, *91*, e964–e975.
- Han, L. K., Dinga, R., Hahn, T., Ching, C. R., Eyler, L. T., Aftanas, L., Aghajani, M., Aleman, A., Baune, B. T., Berger, K., Brak, I., Filho, G. B., Carballedo, A., Connolly, C. G., Couvy-Duchesne, B., Cullen, K. R., Dannlowski, U., Davey, C. G., Dima, D., ... Schmaal, L. (2020). Brain aging in major depressive disorder: Results from the ENIGMA major depressive disorder working group. *Molecular Psychiatry*, *26*, 5124– 5139.
- Harlow, S. D., Gass, M., Hall, J. E., Lobo, R., Maki, P., Rebar, R. W., Sherman, S., Sluss, P. M., de Villiers, T. J., & STRAW + 10 Collaborative Group. (2012). Executive summary of the stages of reproductive aging workshop+ 10: Addressing the unfinished agenda of staging reproductive aging. *The Journal of Clinical Endocrinology & Metabolism*, 97, 1159–1168.
- Hassan, E. B., Szoeke, C., Vogrin, S., Phu, S., Venkatraman, V., Desmond, P., Steward, C., & Duque, G. (2019). Association between structural changes in brain with muscle function in sarcopenic older women: The women's healthy ageing project (WHAP). Journal of Musculoskeletal & Neuronal Interactions, 19, 136–141.
- Heffernan, A. L., Chidgey, C., Peng, P., Masters, C. L., & Roberts, B. R. (2016). The neurobiology and age-related prevalence of the *e*4 allele of apolipoprotein E in Alzheimer's disease cohorts. *Journal of Molecular Neuroscience*, 60, 316–324.
- Huxley, R., Mendis, S., Zheleznyakov, E., Reddy, S., & Chan, J. (2010). Body mass index, waist circumference and waist: Hip ratio as predictors of cardiovascular risk—A review of the literature. *European Journal of Clinical Nutrition*, 64, 16–22.
- Jack, C. R., Jr., Knopman, D. S., Jagust, W. J., Shaw, L. M., Aisen, P. S., Weiner, M. W., Petersen, R. C., & Trojanowski, J. Q. (2010). Hypothetical model of dynamic biomarkers of the Alzheimer's pathological cascade. *The Lancet Neurology*, 9, 119–128.
- Jacobs, E. G., & Goldstein, J. M. (2018). The middle-aged brain: Biological sex and sex hormones shape memory circuitry. *Current Opinion in Behavioral Sciences*, 23, 84–91.
- Jacobsen, E., Ran, X., Liu, A., Chang, C.-C. H., & Ganguli, M. (2021). Predictors of attrition in a longitudinal population-based study of aging. *International Psychogeriatrics*, 33, 767–778.
- Jarolimova, J., Tagoni, J., & Stern, T. A. (2013). Obesity: Its epidemiology, comorbidities, and management. *The Primary Care Companion for CNS Disorders*, 15, PCC.12f01475.
- Jelescu, I. O., & Budde, M. D. (2017). Design and validation of diffusion MRI models of white matter. *Frontiers in Physics*, 5, 61.
- Jensen, J. H., Helpern, J. A., Ramani, A., Lu, H., & Kaczynski, K. (2005). Diffusional kurtosis imaging: The quantification of non-Gaussian water diffusion by means of magnetic resonance imaging. *Magnetic Resonance in Medicine*, 53, 1432–1440.
- Johnson, D. K., Wilkins, C. H., & Morris, J. C. (2006). Accelerated weight loss may precede diagnosis in Alzheimer disease. Archives of Neurology, 63, 1312–1317.
- Jongen, C., van der Grond, J., Kappelle, L., Biessels, G., Viergever, M., & Pluim, J. (2007). Automated measurement of brain and white matter lesion volume in type 2 diabetes mellitus. *Diabetologia*, 50, 1509–1516.
- Kaczkurkin, A. N., Raznahan, A., & Satterthwaite, T. D. (2019). Sex differences in the developing brain: Insights from multimodal neuroimaging. *Neuropsychopharmacology*, 44, 71–85.
- Kaufmann, T., van der Meer, D., Doan, N. T., Schwarz, E., Lund, M. J., Agartz, I., Alnæs, D., Barch, D. M., Baur-Streubel, R., Bertolino, A., Bettella, F., Beyer, M. K., Bøen, E., Borgwardt, S., Brandt, C. L., Buitelaar, J., Celius, E. G., Cervenka, S., Conzelmann, A., ...

Westlye, L. T. (2019). Common brain disorders are associated with heritable patterns of apparent aging of the brain. *Nature Neuroscience*, *22*, 1617–1623.

- Keyes, K. M., & Westreich, D. (2019). UK Biobank, big data, and the consequences of non-representativeness. *The Lancet*, 393, 1297.
- Kivimäki, M., Luukkonen, R., Batty, G. D., Ferrie, J. E., Pentti, J., Nyberg, S. T., Shipley, M. J., Alfredsson, L., Fransson, E. I., Goldberg, M., Knutsson, A., Koskenvuo, M., Kuosma, E., Nordin, M., Suominen, S. B., Theorell, T., Vuoksimaa, E., Westerholm, P., Westerlund, H., ... Jokela, M. (2018). Body mass index and risk of dementia: Analysis of individual-level data from 1.3 million individuals. *Alzheimer's & Dementia*, 14, 601–609.
- Klosinski, L. P., Yao, J., Yin, F., Fonteh, A. N., Harrington, M. G., Christensen, T. A., Trushina, E., & Brinton, R. D. (2015). White matter lipids as a ketogenic fuel supply in aging female brain: Implications for Alzheimer's disease. *eBioMedicine*, 2, 1888–1904.
- Kolbeinsson, A., Filippi, S., Panagakis, Y., Matthews, P. M., Elliott, P., Dehghan, A., & Tzoulaki, I. (2020). Accelerated MRI-predicted brain ageing and its associations with cardiometabolic and brain disorders. *Scientific Reports*, 10, 1–9.
- Kolenic, M., Franke, K., Hlinka, J., Matejka, M., Capkova, J., Pausova, Z., Uher, R., Alda, M., Spaniel, F., & Hajek, T. (2018). Obesity, dyslipidemia and brain age in first-episode psychosis. *Journal of Psychiatric Research*, 99, 151–158.
- Krogsrud, S. K., Fjell, A. M., Tamnes, C. K., Grydeland, H., Mork, L., Due-Tønnessen, P., Bjørnerud, A., Sampaio-Baptista, C., Andersson, J., Johansen-Berg, H., & Walhovd, K. B. (2016). Changes in white matter microstructure in the developing brain—A longitudinal diffusion tensor imaging study of children from 4 to 11 years of age. *NeuroImage*, 124, 473–486.
- Lampe, L., Zhang, R., Beyer, F., Huhn, S., Kharabian Masouleh, S., Preusser, S., Bazin, P.-L., Schroeter, M. L., Villringer, A., & Witte, A. V. (2019). Visceral obesity relates to deep white matter hyperintensities via inflammation. *Annals of Neurology*, 85, 194–203.
- Lavie, C. J., De Schutter, A., Patel, D. A., Romero-Corral, A., Artham, S. M., & Milani, R. V. (2012). Body composition and survival in stable coronary heart disease: Impact of lean mass index and body fat in the "obesity paradox". *Journal of the American College of Cardiology*, 60, 1374–1380.
- Lawrence, K. E., Nabulsi, L., Santhalingam, V., Abaryan, Z., Villalon-Reina, J. E., Nir, T. M., Gari, I. B., Zhu, A. H., Haddad, E., Muir, A. M., Jahanshad N, & Thompson P. (2021). Age and sex effects on advanced white matter microstructure measures in 15,628 UK Biobank participants. *bioRxiv* 2020–09.
- Le, T. T., Kuplicki, R. T., McKinney, B. A., Yeh, H.-W., Thompson, W. K., Paulus, M. P., & Tulsa 1000 Investigators. (2018). A nonlinear simulation framework supports adjusting for age when analyzing brainAGE. *Frontiers in Aging Neuroscience*, 10, 317.
- Leinhard, O. D., Johansson, A., Rydell, J., Smedby, O., Nystrom, F., Lundberg, P., Borga, M. (2008). Quantitative abdominal fat estimation using MRI, in: 2008 19th international conference on pattern recognition. *IEEE*; pp. 1–4.
- Li, H., Satterthwaite, T. D., & Fan, Y. (2018). Brain age prediction based on resting-state functional connectivity patterns using convolutional neural networks. 2018 IEEE 15th International Symposium on Biomedical Imaging (ISBI 2018), IEEE, pp. 101–104.
- Liang, H., Zhang, F., & Niu, X. (2019). Investigating systematic bias in brain age estimation with application to post-traumatic stress disorders. *Human Brain Mapping*, 40, 3143–3152.
- Linge, J., Borga, M., West, J., Tuthill, T., Miller, M. R., Dumitriu, A., Thomas, E. L., Romu, T., Tunón, P., Bell, J. D., & Dahlqvist Leinhard, O. (2018). Body composition profiling in the UK Biobank imaging study. *Obesity*, 26, 1785–1795.
- Linge, J., Whitcher, B., Borga, M., & Leinhard, O. D. (2019). Subphenotyping metabolic disorders using body composition: An

### <sup>14</sup> WILEY\_

individualized, nonparametric approach utilizing large data sets. *Obesity*, 27, 1190–1199.

- Livingston, G., Huntley, J., Sommerlad, A., Ames, D., Ballard, C., Banerjee, S., Brayne, C., Burns, A., Cohen-Mansfield, J., Cooper, C., Costafreda, S. G., Dias, A., Fox, N., Gitlin, L. N., Howard, R., Kales, H. C., Kivimäki, M., Larson, E. B., Ogunniyi, A., ... Mukadam, N. (2020). Dementia prevention, intervention, and care: 2020 report of the lancet commission. *The Lancet*, *396*, 413–446.
- Löwe, L. C., Gaser, C., Franke, K., & Alzheimer's Disease Neuroimaging Initiative. (2016). The effect of the APOE genotype on individual brainAGE in normal aging, mild cognitive impairment, and Alzheimer's disease. *PLoS One*, 11, e0157514.
- Lyall, D. M., Cox, S. R., Lyall, L. M., Celis-Morales, C., Cullen, B., Mackay, D. F., Ward, J., Strawbridge, R. J., McIntosh, A. M., Sattar, N., Smith, D. J., Cavanagh, J., Deary, I. J., & Pell, J. P. (2019). Association between APOE e4 and white matter hyperintensity volume, but not total brain volume or white matter integrity. *Brain Imaging and Behavior*, 14, 1–9.
- Lyall, D. M., Cox, S. R., Lyall, L. M., Celis-Morales, C., Cullen, B., Mackay, D. F., Ward, J., Strawbridge, R. J., McIntosh, A. M., Sattar, N., Smith, D. J., Cavanagh, J., Deary, I. J., & Pell, J. P. (2020). Association between APOE e4 and white matter hyperintensity volume, but not total brain volume or white matter integrity. *Brain Imaging and Behavior*, 14, 1468–1476.
- Lyall, D. M., Ward, J., Ritchie, S. J., Davies, G., Cullen, B., Celis, C., Bailey, M. E., Anderson, J., Evans, J., Mckay, D. F., Mcintosh, A. M., Sattar, N., Smith, D. J., Deary, I. J., & Pell, J. P. (2016). Alzheimer disease genetic risk factor APOE e4 and cognitive abilities in 111,739 UK Biobank participants. *Age and Ageing*, 45, 511–517.
- Maas, A. H., & Appelman, Y. E. (2010). Gender differences in coronary heart disease. *Netherlands Heart Journal*, 18, 598–603.
- Maximov, I. I., Alnaes, D., & Westlye, L. T. (2019). Towards an optimised processing pipeline for diffusion magnetic resonance imaging data: Effects of artefact corrections on diffusion metrics and their age associations in UK Biobank. *Human Brain Mapping*, 40, 4146–4162.
- Maximov, I. I., van der Meer, D., de Lange, A.-M. G., Kaufmann, T., Shadrin, A., Frei, O., Wolfers, T., & Westlye, L. T. (2021). Fast quality control method for derived diffusion metrics (YTTRIUM) in big data analysis: UK Biobank 18,608 example. *Human Brain Mapping*, 42, 3141–3155.
- Meyer, M. R., Clegg, D. J., Prossnitz, E. R., & Barton, M. (2011). Obesity, insulin resistance and diabetes: Sex differences and role of oestrogen receptors. Acta Physiologica, 203, 259–269.
- Miller, K. L., Alfaro-Almagro, F., Bangerter, N. K., Thomas, D. L., Yacoub, E., Xu, J., Bartsch, A. J., Jbabdi, S., Sotiropoulos, S. N., Andersson, J. L., Griffanti, L., Douaud, G., Okell, T. W., Weale, P., Dragonu, I., Garratt, S., Hudson, S., Collins, R., Jenkinson, M., ... Smith, S. M. (2016). Multimodal population brain imaging in the UK Biobank prospective epidemiological study. *Nature Neuroscience*, *19*, 1523–1536.
- Miller, L. R., Marks, C., Becker, J. B., Hurn, P. D., Chen, W.-J., Woodruff, T., McCarthy, M. M., Sohrabji, F., Schiebinger, L., Wetherington, C. L., Makris, S., Arnold, A. P., Einstein, G., Miller, V. M., Sandberg, K., Maier, S., Cornelison, T. L., & Clayton, J. A. (2017). Considering sex as a biological variable in preclinical research. *The FASEB Journal*, *31*, 29–34.
- Mishra, A., & Brinton, R. D. (2018). Inflammation: Bridging age, menopause and APOE e4 genotype to Alzheimer's disease. Frontiers in Aging Neuroscience, 10, 312.
- Mori, S., Wakana, S., Van Zijl, P. C., & Nagae-Poetscher, L. (2005). MRI atlas of human white matter. Elsevier.
- Mosconi, L., Rahman, A., Diaz, I., Wu, X., Scheyer, O., Hristov, H. W., Vallabhajosula, S., Isaacson, R. S., de Leon, M. J., & Brinton, R. D. (2018). Increased Alzheimer's risk during the menopause transition: A 3-year longitudinal brain imaging study. *PLoS One*, 13, e0207885.

- Mueller, K., Anwander, A., Möller, H. E., Horstmann, A., Lepsien, J., Busse, F., Mohammadi, S., Schroeter, M. L., Stumvoll, M., Villringer, A., & Pleger, B. (2011). Sex-dependent influences of obesity on cerebral white matter investigated by diffusion-tensor imaging. *PLoS One*, *6*, e18544.
- Mulugeta, A., Lumsden, A., & Hyppönen, E. (2021). Unlocking the causal link of metabolically different adiposity subtypes with brain volumes and the risks of dementia and stroke: A mendelian randomization study. *Neurobiology of Aging*, 102, 161–169.
- Munafò, M. R., Tilling, K., Taylor, A. E., Evans, D. M., & Smith, G. D. (2018). Collider scope: When selection bias can substantially influence observed associations. *International Journal of Epidemiology*, 47, 226–235.
- Myint, P. K., Kwok, C. S., Luben, R. N., Wareham, N. J., & Khaw, K.-T. (2014). Body fat percentage, body mass index and waist-to-hip ratio as predictors of mortality and cardiovascular disease. *Heart*, 100, 1613–1619.
- Najar, J., Östling, S., Waern, M., Zettergren, A., Kern, S., Wetterberg, H., Hällström, T., & Skoog, I. (2020). Reproductive period and dementia: A 44-year longitudinal population study of swedish women. *Alzheimer's & Dementia*, 16, 1153–1163.
- Neu, S. C., Pa, J., Kukull, W., Beekly, D., Kuzma, A., Gangadharan, P., Wang, L.-S., Romero, K., Arneric, S. P., Redolfi, A., Orlandi, D., Frisoni, G. B., Au, R., Devine, S., Auerbach, S., Espinosa, A., Boada, M., Ruiz, A., Johnson, S. C., ... Toga, A. W. (2017). Apolipoprotein e genotype and sex risk factors for Alzheimer disease: A meta-analysis. JAMA Neurology, 74, 1178–1189.
- Novikov, D. S., Kiselev, V. G., & Jespersen, S. N. (2018). On modeling. Magnetic Resonance in Medicine, 79, 3172–3193.
- Qiu, C., & Fratiglioni, L. (2015). A major role for cardiovascular burden in age-related cognitive decline. *Nature Reviews Cardiology*, 12, 267–277.
- Qiu, C., Winblad, B., & Fratiglioni, L. (2005). The age-dependent relation of blood pressure to cognitive function and dementia. *The Lancet Neurol*ogy, 4, 487–499.
- Raffield, L. M., Cox, A. J., Freedman, B. I., Hugenschmidt, C. E., Hsu, F.-C., Wagner, B. C., Xu, J., Maldjian, J. A., & Bowden, D. W. (2016). Analysis of the relationships between type 2 diabetes status, glycemic control, and neuroimaging measures in the diabetes heart study mind. Acta Diabetologica, 53, 439–447.
- Rahman, A., Schelbaum, E., Hoffman, K., Diaz, I., Hristov, H., Andrews, R., Jett, S., Jackson, H., Lee, A., Sarva, H., Pahlajani, S., Matthews, D., Dyke, J., de Leon, M. J., Isaacson, R. S., Brinton, R. D., & Mosconi, L. (2020). Sex-driven modifiers of Alzheimer risk: A multimodality brain imaging study. *Neurology*, 95, e166–e178.
- Ranasinghe, C., Gamage, P., Katulanda, P., Andraweera, N., Thilakarathne, S., & Tharanga, P. (2013). Relationship between body mass index (BMI) and body fat percentage, estimated by bioelectrical impedance, in a group of Sri Lankan adults: A cross sectional study. BMC Public Health, 13, 1–8.
- Richard, G., Kolskår, K., Sanders, A.-M., Kaufmann, T., Petersen, A., Doan, N. T., Monereo Sánchez, J., Alnæs, D., Ulrichsen, K. M., Dørum, E. S., Andreassen, O. A., Nordvik, J. E., & Westlye, L. T. (2018). Assessing distinct patterns of cognitive aging using tissue-specific brain age prediction based on diffusion tensor imaging and brain morphometry. *PeerJ*, *6*, e5908.
- Ritchie, S. J., Cox, S. R., Shen, X., Lombardo, M. V., Reus, L. M., Alloza, C., Harris, M. A., Alderson, H. L., Hunter, S., Neilson, E., Liewald, D. C. M., Auyeung, B., Whalley, H. C., Lawrie, S. M., Gale, C. R., Bastin, M. E., McIntosh, A. M., & Deary, I. J. (2018). Sex differences in the adult human brain: Evidence from 5216 UK Biobank participants. *Cerebral Cortex*, 28, 2959–2975.
- Robison, L. S., Gannon, O. J., Salinero, A. E., & Zuloaga, K. L. (2019). Contributions of sex to cerebrovascular function and pathology. *Brain Research*, 1710, 43–60.

- Rocca, W., Bower, J., Maraganore, D., Ahlskog, J., Grossardt, B., De Andrade, M., & Melton, L. R. (2007). Increased risk of cognitive impairment or dementia in women who underwent oophorectomy before menopause. *Neurology*, *69*, 1074–1083.
- Rokicki, J., Wolfers, T., Nordhøy, W., Tesli, N., Quintana, D. S., Alnæs, D., Richard, G., de Lange, A.-M. G., Lund, M. J., Norbom, L., Agartz, I., Melle, I., Nærland, T., Selbæk, G., Persson, K., Nordvik, J. E., Schwarz, E., Andreassen, O. A., Kaufmann, T., & Westlye, L. T. (2021). Multimodal imaging improves brain age prediction and reveals distinct abnormalities in patients with psychiatric and neurological disorders. *Human Brain Mapping*, *42*, 1714–1726.
- Sachdev, P. S., Parslow, R., Wen, W., Anstey, K., & Easteal, S. (2009). Sex differences in the causes and consequences of white matter hyperintensities. *Neurobiology of Aging*, 30, 946–956.
- Salthouse, T. A. (2014). Selectivity of attrition in longitudinal studies of cognitive functioning. Journals of Gerontology Series B: Psychological Sciences and Social Sciences, 69, 567–574.
- Scheinost, D., Finn, E. S., Tokoglu, F., Shen, X., Papademetris, X., Hampson, M., & Constable, R. T. (2015). Sex differences in normal age trajectories of functional brain networks. *Human Brain Mapping*, *36*, 1524–1535.
- Schelbaum, E., Loughlin, L., Jett, S., Zhang, C., Jang, G., Malviya, N., Hristov, H., Pahlajani, S., Isaacson, R., Dyke, J. P., Kamel, H., Brinton, R. D., & Mosconi, L. (2021). Association of reproductive history with brain MRI biomarkers of dementia risk in midlife. *Neurology*, 97, e2328–e2339.
- Schelbert, K. B. (2009). Comorbidities of obesity. *Primary Care: Clinics in Office Practice*, 36, 271–285.
- Scheyer, O., Rahman, A., Hristov, H., Berkowitz, C., Isaacson, R., Brinton, R. D., & Mosconi, L. (2018). Female sex and Alzheimer's risk: The menopause connection. *The Journal of Prevention of Alzheimer's Disease*, 5, 225–230.
- Schorr, M., Dichtel, L. E., Gerweck, A. V., Valera, R. D., Torriani, M., Miller, K. K., & Bredella, M. A. (2018). Sex differences in body composition and association with cardiometabolic risk. *Biology of Sex Differences*, 9, 1–10.
- Shansky, R. M., & Murphy, A. Z. (2021). Considering sex as a biological variable will require a global shift in science culture. *Nature Neuroscience*, 24, 457–464.
- Shansky, R. M., & Woolley, C. S. (2016). Considering sex as a biological variable will be valuable for neuroscience research. *Journal of Neuroscience*, 36, 11817–11822.
- Simpson, E. R. (2003). Sources of estrogen and their importance. *The Journal of Steroid Biochemistry and Molecular Biology*, *86*, 225–230.
- Smith, S. M., Jenkinson, M., Johansen-Berg, H., Rueckert, D., Nichols, T. E., Mackay, C. E., Watkins, K. E., Ciccarelli, O., Cader, M. Z., Matthews, P. M., & Behrens, T. E. (2006). Tract-based spatial statistics: Voxelwise analysis of multi-subject diffusion data. *NeuroImage*, *31*, 1487–1505.
- Solomon, A., Kivipelto, M., Wolozin, B., Zhou, J., & Whitmer, R. A. (2009). Midlife serum cholesterol and increased risk of Alzheimer's and vascular dementia three decades later. *Dementia and Geriatric Cognitive Dis*orders, 28, 75–80.
- Sowers, M., Zheng, H., Tomey, K., Karvonen-Gutierrez, C., Jannausch, M., Li, X., Yosef, M., & Symons, J. (2007). Changes in body composition in women over six years at midlife: Ovarian and chronological aging. *The Journal of Clinical Endocrinology & Metabolism*, 92, 895–901.
- Stanek, K. M., Grieve, S. M., Brickman, A. M., Korgaonkar, M. S., Paul, R. H., Cohen, R. A., & Gunstad, J. J. (2011). Obesity is associated with reduced white matter integrity in otherwise healthy adults. *Obesity*, 19, 500–504.
- Stewart, R., Masaki, K., Xue, Q.-L., Peila, R., Petrovitch, H., White, L. R., & Launer, L. J. (2005). A 32-year prospective study of change in body weight and incident dementia: The Honolulu-Asia aging study. Archives of Neurology, 62, 55–60.

- Storsve, A. B., Fjell, A. M., Yendiki, A., & Walhovd, K. B. (2016). Longitudinal changes in white matter tract integrity across the adult lifespan and its relation to cortical thinning. *PLoS One*, 11, e0156770.
- Subramaniapillai, S., Rajagopal, S., Elshiekh, A., Pasvanis, S., Ankudowich, E., & Rajah, M. N. (2019). Sex differences in the neural correlates of spatial context memory decline in healthy aging. *Journal* of Cognitive Neuroscience, 31, 1895–1916.
- Subramaniapillai, S., Rajagopal, S., Snytte, J., Otto, A. R., Einstein, G., Rajah, M. N., the PREVENT-AD Research Group, Einstein, G., & Rajah, M. N. (2021). Sex differences in brain aging among adults with family history of Alzheimer's disease and APOE4 genetic risk. *NeuroImage: Clinical*, 30, 102620.
- Subramaniapillai, S., Rajah, M. N., Pasvanis, S., & Titone, D. (2019). Bilingual experience and executive control over the adult lifespan: The role of biological sex. *Bilingualism: Language and Cognition*, 22, 733–751.
- Szucs, D., & Ioannidis, J. P. (2020). Sample size evolution in neuroimaging research: An evaluation of highly-cited studies (1990–2012) and of latest practices (2017–2018) in high-impact journals. *NeuroImage*, 221, 117164.
- Taki, Y., Kinomura, S., Sato, K., Inoue, K., Goto, R., Okada, K., Uchida, S., Kawashima, R., & Fukuda, H. (2008). Relationship between body mass index and gray matter volume in 1,428 healthy individuals. *Obesity*, 16, 119–124.
- Taylor, C., Pritschet, L., Yu, S., & Jacobs, E. G. (2019). Applying a women's health lens to the study of the aging brain. Frontiers in Human Neuroscience, 13, 224.
- Tchernof, A., & Després, J.-P. (2013). Pathophysiology of human visceral obesity: An update. *Physiological Reviews*, 93, 359–404.
- Tomiyama, A. J., Hunger, J. M., Nguyen-Cuu, J., & Wells, C. (2016). Misclassification of cardiometabolic health when using body mass index categories in NHANES 2005–2012. *International Journal of Obesity*, 40, 883–886.
- Tønnesen, S., Kaufmann, T., de Lange, A.-M., Richard, G., Doan, N. T., Alnaes, D., van der Meer, D., Rokicki, J., Moberget, T., Maximov, I. I., Agartz, I., Aminoff, S. R., Beck, D., Barch, D. M., Beresniewicz, J., Cervenka, S., Fatouros-Bergman, H., Craven, A. R., Flyckt, L., ... Westlye, L. T. (2020). Brain age prediction reveals aberrant brain white matter in schizophrenia and bipolar disorder: A multi-sample diffusion tensor imaging study. *bioRxiv* 607754.
- Trikudanathan, S., Pedley, A., Massaro, J. M., Hoffmann, U., Seely, E. W., Murabito, J. M., & Fox, C. S. (2013). Association of female reproductive factors with body composition: The Framingham heart study. *The Journal of Clinical Endocrinology & Metabolism*, 98, 236–244.
- Trofimova, O., Loued-Khenissi, L., DiDomenicantonio, G., Lutti, A., Kliegel, M., Stringhini, S., Marques-Vidal, P., Vollenweider, P., Waeber, G., Preisig, M., Kherif, F., & Draganski, B. (2021). Brain tissue properties link cardio-vascular risk factors, mood and cognitive performance in the CoLaus|PsyCoLaus epidemiological cohort. *Neurobiology* of Aging, 102, 50–63.
- UI-Haq, Z., Smith, D. J., Nicholl, B. I., Cullen, B., Martin, D., Gill, J. M., Evans, J., Roberts, B., Deary, I. J., Craddock, N., Mackay, D. F., & Pell, J. P. (2014). Gender differences in the association between adiposity and probable major depression: A cross-sectional study of 140,564 UK Biobank participants. *BMC Psychiatry*, 14, 1–10.
- van den Heuvel, D., Admiraal-Behloul, F., ten Dam, V., Olofsen, H., Bollen, E., Murray, H., Blauw, G., Westendorp, R., de Craen, A., van Buchem, M., & for the PROSPER Study Group. (2004). Different progression rates for deep white matter hyperintensities in elderly men and women. *Neurology*, *63*, 1699–1701.
- van Gestel, H., Franke, K., Petite, J., Slaney, C., Garnham, J., Helmick, C., Johnson, K., Uher, R., Alda, M., & Hajek, T. (2019). Brain age in bipolar disorders: Effects of lithium treatment. *Australian & New Zealand Journal of Psychiatry*, 53, 1179–1188.
- van Vliet, P. (2012). Cholesterol and late-life cognitive decline. Journal of Alzheimer's Disease, 30, S147–S162.

## <sup>16</sup> ↓ WILEY-

- Vidal-Pineiro, D., Wang, Y., Krogsrud, S. K., Amlien, I. K., Baaré, W. F., Bartres-Faz, D., Bertram, L., Brandmaier, A. M., Drevon, C. A., Düzel, S., Ebmeier, K. P., Henson, R. N., Junqué, C., Kievit, R. A., Kühn, S., Leonardsen, E., Lindenberger, U., Madsen, K. S., Magnussen, F., ... Fjell, A. (2021). Individual variations in "brain age" relate to early life factors more than to longitudinal brain change. *eLife*, 10, e69995.
- Voldsbekk, I., Barth, C., Maximov, I. I., Kaufmann, T., Beck, D., Richard, G., Moberget, T., Westlye, L. T., & de Lange, A.-M. G. (2021). A history of previous childbirths is linked to women's white matter brain age in midlife and older age. *Human Brain Mapping*, 42, 4372–4386.
- Vosberg, D. E., Parker, N., Shin, J., Pausova, Z., & Paus, T. (2021). The genetics of testosterone contributes to "femaleness/maleness" of cardiometabolic traits and type 2 diabetes. *International Journal of Obesity*, 46, 1–3.
- Wang, H.-T., Smallwood, J., Mourao-Miranda, J., Xia, C. H., Satterthwaite, T. D., Bassett, D. S., & Bzdok, D. (2020). Finding the needle in a high-dimensional haystack: Canonical correlation analysis for neuroscientists. *NeuroImage*, 216, 116745.
- Wang, J., Knol, M. J., Tiulpin, A., Dubost, F., de Bruijne, M., Vernooij, M. W., Adams, H. H., Ikram, M. A., Niessen, W. J., & Roshchupkin, G. V. (2019). Gray matter age prediction as a biomarker for risk of dementia. *Proceedings of the National Academy of Sciences*, 116, 21213–21218.
- Wasti, S., Robinson, S., Akhtar, Y., Khan, S., & Badaruddin, N. (1993). Characteristics of menopause in three socioeconomic urban groups in Karachi, Pakistan. *Maturitas*, 16, 61–69.
- Wei, Y.-C., George, N. I., Chang, C.-W., & Hicks, K. A. (2017). Assessing sex differences in the risk of cardiovascular disease and mortality per increment in systolic blood pressure: A systematic review and metaanalysis of follow-up studies in the United States. *PLoS One*, 12, e0170218.

- Westlye, L. T., Walhovd, K. B., Dale, A. M., Bjørnerud, A., Due-Tønnessen, P., Engvig, A., Grydeland, H., Tamnes, C. K., Østby, Y., & Fjell, A. M. (2010). Life-span changes of the human brain white matter: Diffusion tensor imaging (DTI) and volumetry. *Cerebral Cortex*, 20, 2055–2068.
- Wiltink, J., Michal, M., Wild, P. S., Zwiener, I., Blettner, M., Münzel, T., Schulz, A., Kirschner, Y., & Beutel, M. E. (2013). Associations between depression and different measures of obesity (BMI, WC, WHtR, WHR). *BMC Psychiatry*, 13, 1–7.
- Wisdom, N. M., Callahan, J. L., & Hawkins, K. A. (2011). The effects of apolipoprotein E on non-impaired cognitive functioning: A meta-analysis. *Neurobiology of Aging*, 32, 63–74.

#### SUPPORTING INFORMATION

Additional supporting information may be found in the online version of the article at the publisher's website.

How to cite this article: Subramaniapillai, S., Suri, S., Barth, C., Maximov, I. I., Voldsbekk, I., D. van der Meer, Gurholt, T. P., Beck, D., Draganski, B., Andreassen, O. A., Ebmeier, K. P., Westlye, L. T., A.-M. G. de Lange (2022). Sex- and age-specific associations between cardiometabolic risk and white matter brain age in the UK Biobank cohort. *Human Brain Mapping*, 1–16. https://doi.org/10.1002/hbm.25882