

TECHNICAL REPORT

BrainAGE: Revisited and reframed machine learning workflow

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Abstract

Since the introduction of the BrainAGE method, novel machine learning methods for brain age prediction have continued to emerge. The idea of estimating the chronological age from magnetic resonance images proved to be an interesting field of research due to the relative simplicity of its interpretation and its potential use as a biomarker of brain health. We revised our previous BrainAGE approach, originally utilising relevance vector regression (RVR), and substituted it with Gaussian process regression (GPR), which enables more stable processing of larger datasets, such as the UK Biobank (UKB). In addition, we extended the global BrainAGE approach to regional BrainAGE, providing spatially specific scores for five brain lobes per hemisphere. We tested the performance of the new algorithms under several different conditions and investigated their validity on the ADNI and schizophrenia samples, as well as on a synthetic dataset of neocortical thinning. The results show an improved performance of the reframed global model on the UKB sample with a mean absolute error (MAE) of less than 2 years and a significant difference in BrainAGE between healthy participants and patients with Alzheimer's disease and schizophrenia. Moreover, the workings of the algorithm show meaningful effects for a simulated neocortical atrophy dataset. The regional BrainAGE model performed well on two clinical samples, showing disease-specific patterns for different levels of impairment. The results demonstrate that the new improved algorithms provide reliable and valid brain age estimations.

KEYWORDS

Alzheimer's disease, brain age, Gaussian process regression, machine learning, mean absolute error, pre-processing, schizophrenia, structural MRI, UK Biobank

1 | INTRODUCTION

Our original BrainAGE framework (Franke et al., 2010) is a supervised machine learning method that employs relevance vector regression (RVR) in order to predict the chronological age of an individual based on a single T1-weighted magnetic resonance (MR) scan of their brain. The difference between the apparent age of the individual's brain as

predicted by the algorithm and their chronological age is the so-called brain age gap estimation (BrainAGE). A positive gap implies more prominent structural brain changes that commonly occur with ageing progression, for example, decrease of grey matter volume (Bethlehem et al., 2022), whereas a negative difference signifies a slower ageing in comparison to other subjects within a sample. The BrainAGE method was introduced more than a decade ago and was successfully applied

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to various subject groups, such as adolescents (Franke et al., 2012), elderly people with mild cognitive impairment (MCI) and Alzheimer's disease (AD; Franke et al., 2010; Franke & Gaser, 2012), musicians (Rogenmoser et al., 2018), meditators (Luders et al., 2016), and even other animal species, such as rodents and non-human primates (Franke et al., 2017). The method proved to be a reliable biomarker of AD (Franke & Gaser, 2012), Parkinson's disease (Eickhoff et al., 2021), and is associated with brain alterations in schizophrenia (Nenadić et al., 2017) and type 2 diabetes (Franke et al., 2013). A complete review can be found in Franke and Gaser (2019) or Cole and Franke (2017).

Since the introduction of the concept of brain age as a biomarker of ageing brains, numerous machine learning and deep learning models of estimating the age of the brain based on different MR imaging modalities have emerged (e.g., Beheshti et al., 2020; Cole et al., 2018; Hahn et al., 2021; Smith et al., 2019; Varzandian et al., 2021). The advancement of deep learning methods in the field of brain age prediction (e.g., Abrol et al., 2021; Gong et al., 2021; Hahn et al., 2021; Lombardi et al., 2021; Peng et al., 2021) and the availability of large multimodal imaging data collection, such as the UK Biobank (UKB; <https://www.ukbiobank.ac.uk/>), pushed the performance of the brain age prediction to unprecedented levels, managing to decrease mean absolute errors (MAE; i.e., a metric commonly used to compare the accuracy of prediction) to as low as 2.14 years (Peng et al., 2021).

The aim of the study was to revise the original BrainAGE machine learning framework to be able to keep pace with the advancement of deep learning algorithms in dealing with large datasets, while retaining the parsimonious and straightforward nature of voxel-based morphometry (VBM). Studies testing the performance of multiple algorithms in brain age estimation showed a good performance of Gaussian process regression (GPR) (Baecker et al., 2021; Beheshti et al., 2022; Cole et al., 2015, 2018; More et al., 2023), which can be seen as an alternative approach to artificial neural networks (Chen & Ren, 2009). We therefore implemented GPR in the new BrainAGE framework and utilised model ensembling to limit overfitting and improve prediction accuracy (van Veen et al., 2015). We replaced the initial BrainAGE framework (Franke et al., 2010; Franke & Gaser, 2012) with a combination of models using grey- or white matter tissue segments (of varying spatial resolutions and smoothing sizes) associated by (weighted) averaging and stacking. In addition to the global BrainAGE, we further implemented a regional BrainAGE approach to obtain greater spatial specificity for future applications to various clinical groups.

In order to evaluate the performance of our novel BrainAGE algorithms and to allow for straightforward comparison to results of other groups, we first focused on the UK Biobank cohort. The UKB is a very large biomedical database and research resource that contains genetic, lifestyle and health information from half a million UK Biobank participants. We investigated the effects of various conditions (e.g., regularisation, varying training sample size) on prediction accuracy and examined the reliability of the predictions. In the second part of the article, we tested the performance of the global and regional

models on external datasets. We applied the new global model to a novel synthetic dataset of global neocortical thinning (Rusak et al., 2022), and to two samples from clinical populations, namely MCI and AD patients from the ADNI cohort as well as schizophrenia patients (SZ).

2 | METHODS

2.1 | Datasets

This research has been conducted using data from UK Biobank under the application number 41655. The UK Biobank has ethical approval from the North West Multi-Centre Research Ethics Committee (MREC) and is in possession of informed consents from the study cohort participants. Participants with pre-existing neurological or psychiatric diagnoses were excluded from the analyses (UKB data-fields #41202-0.0 to 41202-0.78 Diagnoses—main ICD10: presence of F0-F99 Mental and behavioural disorders). Overall, we analysed 36,840 T1-weighted MR scans (52% women; age range: 45–82; $M_{\text{Age}} = 64.09$, $SD_{\text{Age}} = 7.53$ years). Some of the analyses were run on a pseudo random subsample of 7241 scans (52% women; age range: 45–82; $M_{\text{Age}} = 64.08$, $SD_{\text{Age}} = 7.54$ years), including only subject IDs starting with 1* to avoid the introduction of selection bias. For the cross-site prediction, we randomly sampled a fifth of the full sample ($n = 7368$) to obtain two size-matched subsamples from the scanning site 1 (Cheadle) and the scanning site 3 (Newcastle). The tests performed on the UKB sample served as a benchmark to compare our different BrainAGE models to the ones from other groups (e.g., Baecker et al., 2021; Peng et al., 2021).

Although the UKB is one of the largest neuroimaging datasets, the strict data policy and restricted age range present a major drawback for using it as a normative sample for training the BrainAGE models. We therefore combined images from IXI, OASIS-3, Cam-CAN, SALD, and Enhanced NKI-RS databases to create an independent normative dataset of healthy adults. An overview of the datasets used in this study can be seen in Table 1.

Additionally, 695 T1-weighted images from Alzheimer's Disease Neuroimaging Initiative (ADNI, <https://adni.loni.usc.edu/>) were used to validate the model on a sample including healthy control subjects (HC, $n = 231$, $M_{\text{Age}} = 76.01 \pm 5.01$), subjects with stable MCI (sMCI, $n = 100$, $M_{\text{Age}} = 75.40 \pm 7.27$), subjects who progressed from MCI to Alzheimer's dementia (pMCI, $n = 164$, $M_{\text{Age}} = 74.57 \pm 7.03$), and AD patients ($n = 200$, $M_{\text{Age}} = 75.64 \pm 7.72$).

We also used a synthetic global neocortical atrophy dataset by Rusak et al. (2022), which was derived from ADNI baseline scans of 20 individuals without AD ($n = 20$; $M_{\text{Age}} = 70.65 \pm 5.39$) and used to simulate neocortical thinning, progressing from 0 to 0.1 mm or 1 mm thickness loss, with steps of 0.01 or 0.1 mm between the time points, respectively, resulting in total of 400 synthetic images.

In order to validate the (regional) model on clinical data, a sample of schizophrenia patients and healthy controls previously described in Nenadić et al. (2015) was used.

TABLE 1 Characteristics of the datasets used in this study.

Dataset		N (sessions)	Age range	M $\pm$ SD	Males/females
UKB		36840	45–82	64.09 $\pm$ 7.53	17557/19283
IXI		547	19–86	48.12 $\pm$ 16.61	242/305
OASIS-3		549	42–97	69.53 $\pm$ 8.86	208/341
Cam-CAN		652	18–88	54.30 $\pm$ 18.59	322/330
SALD		494	19–80	45.18 $\pm$ 17.44	187/307
NKle		629	18–85	38.74 $\pm$ 21.27	233/396
ADNI	HC	231	60–90	76.01 $\pm$ 5.01	119/112
	sMCI	100	58–88	75.40 $\pm$ 7.27	66/34
	pMCI	164	55–88	74.57 $\pm$ 7.03	97/67
	AD	200	55–91	75.64 $\pm$ 7.72	103/97
SZ	HC	108	20–59	32.16 $\pm$ 9.99	68/40
	Positive	33	19–65	35.54 $\pm$ 11.18	19/14
	Negative	31	18–54	35.54 $\pm$ 11.17	15/16
	Disorg.	23	19–59	35.39 $\pm$ 11.13	14/9
Synthetic dataset		20 (20)	59–78	70.65 $\pm$ 5.39	10/10

2.2 | Data acquisition

The T1-weighted MR images of the UKB dataset were acquired on Siemens Skyra 3 T scanners with a 32-channel head coil. A sagittal MPRAGE sequence was used with 1 mm isotropic resolution, inversion/repetition/echo time (TI/TR/TE) = 880/2000/na ms, flip angle (FA) = na, field-of-view (FOV) = 208 $\times$ 256 $\times$ 256, acceleration factor (R) = 2, acquisition time: 5 min (https://biobank.ctsu.ox.ac.uk/crystal/crystal/docs/brain_mri.pdf).

In the ADNI sample used, T1-weighted images were acquired on 1.5 T scanners manufactured by Siemens, Philips, and GE with single-channel coils. The sagittal MPRAGE sequences typically had about 0.95 mm in-plane resolution and 1.2 mm slice-thickness, TI/TR/TE = 853–1000/2300–3000/na ms, FA = 8–9°, FOV = 240–260 $\times$ 240 mm, R = 1, acquisition time: 7:11–9:38 min (Jack et al., 2008; see also <https://adni.loni.usc.edu/methods/mri-tool/mri-acquisition/>).

The IXI sample comprises data from 3 different scanners (GE 1.5 T, Philips 1.5 and 3 T) with about 0.95 mm in-plane resolution and 1.2 mm slice-thickness, TI/TR/TE time = na/9.600–9.813/4.603 ms, FA = 8°, FOV = 256 mm, R = na (<https://brain-development.org/ixi-dataset>).

The MRI data from the OASIS-3 dataset used in this study were acquired on two Siemens TIM Trio and one BioGraph mMR PET-MR 3 T scanners with 20-channel head coil. The MPRAGE scans were acquired with (i) 1 mm isotropic TI/TR/TE = 1000/2400/3.16 ms, FA = 8°, FOV phase = 100%, R = 2 and (ii) 1.20 $\times$ 1.05 $\times$ 1.05 mm resolution, TI/TR/TE = 900/2300/2.95 ms, FA = 9°, FOV phase = 93.75 mm, R = 2 (LaMontagne et al., 2019).

In the SALD dataset, a sagittal MPRAGE sequence on a 3 T Siemens Trio MRI scanner was used with 1 mm isotropic resolution, TI/TR/TE = 900/1900/2.52 ms, FA = 9°, FOV = 256 mm, R = 2 (Wei et al., 2018).

The Cam-CAN dataset was obtained on a 3 T Siemens TIM Trio scanner with a 32-channel head coil, using an MPRAGE sequence with 1 mm isotropic resolution, TI/TR/TE = 900/2250/2.98 ms, FA = 9°, FOV = 256 $\times$ 240 $\times$ 192 mm, R = 2, acquisition time = 4:32 min (Taylor et al., 2017).

T1 MPRAGE images from the NKI Enhanced dataset were acquired on a 3 T Siemens TIM Trio scanner with 32-channel head coil with 1 mm isotropic resolution, TI/TR/TE = 1200/2500/3.5 ms, FA = 8° (https://fcon_1000.projects.nitrc.org/indi/pro/eNKI_RS_TRT/FrontPage.html).

We used a sample of schizophrenia patients and healthy controls previously described in Nenadić et al. (2015). The sagittal MPRAGE images were acquired on a 1.5 T Philips Gyroscan ASCII scanner with 1 mm isotropic resolution, TI/TR/TE = na/13/5 ms, FOV = 256 mm.

2.3 | Data preprocessing

The UKB T1-weighted images were segmented into grey matter (GM) and white matter (WM) and (affinely/non-linearly) normalised by using default preprocessing of the CAT12.7 Toolbox (Gaser et al., 2022) as standalone version compiled under Matlab 2019b and SPM12 (Wellcome Centre for Human Neuroimaging, <https://www.fil.ion.ucl.ac.uk/spm/>) massively parallelized on the JURECA High Performance Computing system (Jülich Supercomputing Center, 2018). The preprocessing utilises the unified segmentation (Ashburner & Friston, 2005) to remove B0-inhomogeneities and create an initial segmentation that is used for (local) intensity scaling and adaptive non-local means denoising (Manjon et al., 2010). An adaptive maximum a posteriori (AMAP; Rajapakse et al., 1997) segmentation with a hidden Markov random field (Cuadra et al., 2005) and partial volume effect model (Tohka et al., 2004) is utilised to create the final segmentation. For the optional non-linear registration, the Shooting method (Ashburner & Friston, 2011) with modulation was used.

The synthetic dataset, IXI, Cam-CAN, NKle, OASIS-3, ADNI, and the SZ sample were processed with the CAT12.8 version with the same parameters.

2.4 | Global BrainAGE models estimation

In order to keep the process as parsimonious as possible, a linear affine registration of tissue segments to the MNI152Nlin2009cAsym template was used as default, except for a single non-linearly registered GM test case. Eight combinations of single tissue class models (GM/WM) with varying parameterisations of spatial resolution (4 mm/8 mm) and Gaussian smoothing (FWHM: 4 mm/8 mm) were trained and processed in the same way. With the exception of the first testing condition (see the test conditions below), principal component analysis (PCA) using singular value decomposition was applied to all the models to regularise the data. We employed a GPR model instead of the previously used RVR (see Figure 1a for the comparison of performance of both algorithms). This GPR uses a linear covariance function, a constant mean function, and a Gaussian likelihood function with the hyperparameters set to 100 for the constant mean function and -1 for the likelihood function (Rasmussen & Williams, 2006). A conjugate gradient method was used for numerical optimization in the GPR.

A 10-fold cross-validation (CV) was performed with stratified chronological age to preserve the age distribution in each fold. The estimated brain age was age-bias adjusted by a linear term as described in Smith et al. (2019). Single tissue class models were assembled by averaging (unless stated otherwise).

The estimations were run in Matlab 2021a (Mathworks Inc., Natick, MA, USA).

2.5 | Regional BrainAGE models estimation

Regional BrainAGE models were trained on a large sample of healthy adult participants from 5 databases (IXI, OASIS-3, NKle, SALD, Cam-CAN) and tested on the ADNI sample, spanning from healthy control subjects to AD patients, as well as on the SZ dataset. The training samples (AD: $n = 1947$, SZ: $n = 2639$) were age-matched to the ADNI and SZ test samples.

To obtain regional BrainAGE estimations, the GM segments were divided into 10 lobes (5 per hemisphere) derived from the Brain lobe atlas (Toro et al., 2009). Only GM segments were used as both previously mentioned diseases were shown to predominantly affect GM (Haijma et al., 2013; Wang et al., 2015). For each lobe, independent GPR models were estimated as described above. Only lobular regions were defined to support a useful amount of GM voxels for the

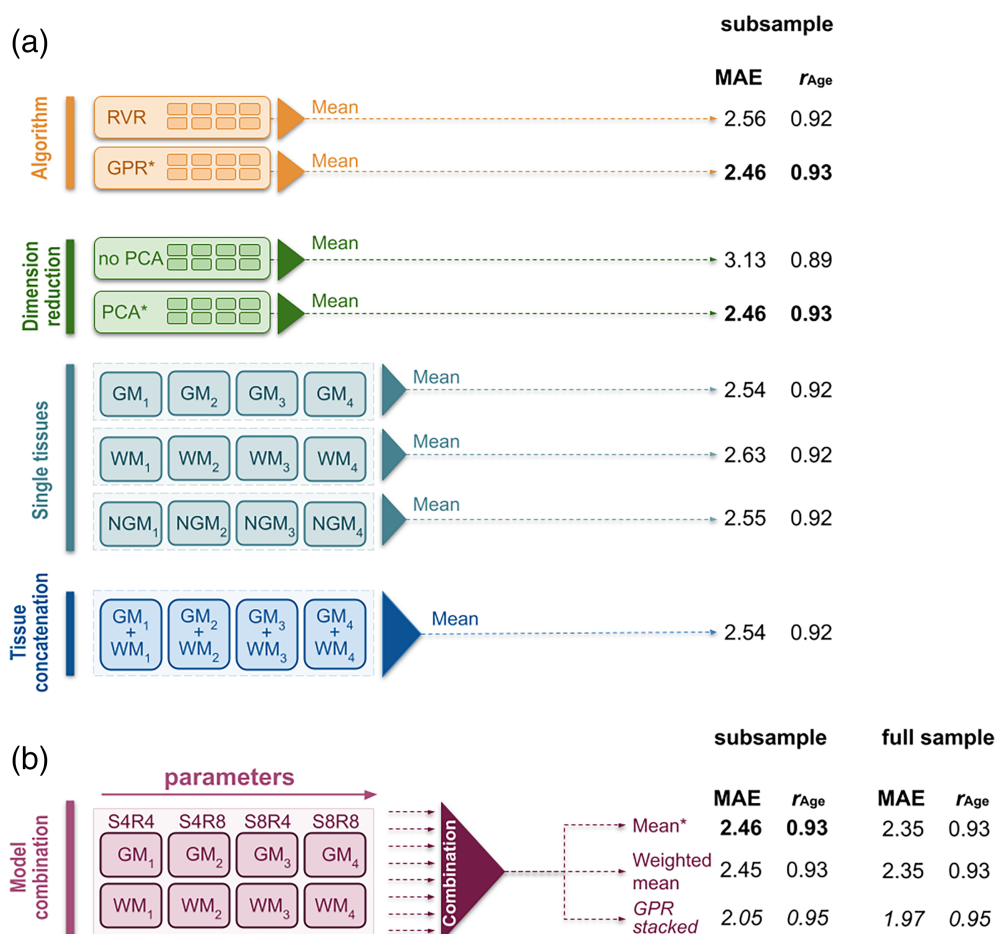


FIGURE 1 (a) Various single models differing in spatial resolution (R: 4 mm/8 mm), smoothing kernel (S: 4 mm/8 mm), and tissue segments (GM, WM, NGM: non-linearly registered modulated GM) were estimated and combined. The results of model estimations combined by averaging are presented as age-bias corrected MAEs and the Pearson's correlation coefficients between predicted- and chronological age for analyses run on a UKB 1 subsample. The comparison of the ML algorithms (RVR/GPR), as well as the effects of dimension reduction and the choice of brain tissues or their concatenation are shown. The results in bold represent the same combined model. (b) Eight single models were ensembled by averaging, weighted averaging, or GPR stacking. The results are presented for the subsample and the full UKB sample.

models. A linear age-bias correction was used globally as described in Smith et al. (2019).

3 | RESULTS

With respect to the change in the algorithm, we tested the novel BrainAGE framework performance under different conditions outlined in the following section, jointly reporting methods and results. We show the MAEs for the average-ensembled age-bias corrected models and Pearson's correlation coefficients between the predicted age and the chronological age. See the Supporting Information (Tables S1 and S2) for the performance of all tested models without age-bias correction.

3.1 | The effect of principal component analysis

In this step, we tested the effect of implementing PCA using singular value decomposition. The method was applied to the training data only, and the resulting transformation was used on the test sample to avoid data leakage. The performance of the model greatly improved with the drop from MAE of 3.13 years ($r_{\text{Age}} = 0.89$) to MAE of 2.46 years ($r_{\text{Age}} = 0.93$). All subsequent estimations on the full UKB dataset were therefore run on regularised data.

3.2 | Tissue class combination by model ensembling

Single tissue class models using only one linearly registered brain tissue class (either GM, WM, or non-linearly registered modulated GM), with varying parameterisations of spatial resolution (4 mm/8 mm) and Gaussian smoothing (FWHM: 4 mm/8 mm) were ensembled in order to minimise the MAE of the brain age prediction (Figure 1b). Three different ensembling approaches were tested, namely (i) their average performance, (ii) the weighted mean, and (iii) GPR stacking of the estimated models. The weighted mean was calculated with weighting w.r.t. squared MAE. The models with higher MAE were assigned a lower weight. A nested 10×10 CV was used in the GPR stacking,

which utilised the predictions from the first level models as features for the second GPR learner.

We additionally tested the performance of mean-ensembled single-tissue-class models (either GM-, non-linearly registered modulated GM-, WM-, and concatenated GM- and WM vectors), which performed satisfactorily.

The results of ensembling different brain tissue classes are presented in Figure 1a,b for the subsample and the full sample, respectively. Model ensembling provides a more accurate BrainAGE estimation, with the most striking improvement obtained by stacking. Using the full UKB sample, GPR stacking resulted in 2.18 years of MAE for uncorrected BrainAGE prediction, MAE = 1.97 for linear age-bias correction as in Smith et al. (2019), and 2.34 years age-corrected as in Cole et al. (2018). Nevertheless, a simple mean-ensembled GM model was finally chosen for the application to the clinical datasets, as previous studies have shown that the most accurate models are not the best in differentiating between (clinical) phenotypes (Bashyam et al., 2020).

3.3 | Sample dependence and the effect of training sample size

We compared the performance of the algorithm between different subsamples of the UKB dataset. The subsamples were of comparable sizes and age/sex distributions and showed similar results (see Table 2). We further tested the effect of varying training sample size by using 20%–100% of the full dataset, resulting in only minor improvements in the performance of the algorithm (Figure S1).

3.4 | Cross-scanner prediction reliability

We investigated the cross-site prediction by training the algorithm on a sample of 6631 T1-weighted MRI scans (using 90% of the sample to keep the size of training sample constant) from scanning site 1 (Cheadle, $M_{\text{Age}} = 63.45 \pm 7.44$, age range: 45–81) and tested its performance on a dataset from scanning site 3 (Newcastle, $M_{\text{Age}} = 64.86 \pm 7.45$, age range: 47–81), and vice versa. When predicting from

TABLE 2 Sample characteristics of different UKB subsamples with MAEs of the mean-ensembled model performance and Pearson's correlation coefficients between the predicted and chronological age.

Sample	N training	N testing	Males/females	Age range	M $\pm$ SD	MAE (years)	r_{Age}
UKB 1	6517	724	3503/3738	45–82	64.08 $\pm$ 7.54	2.46	0.93
UKB 2	6745	749	3618/3876	46–82	64.10 $\pm$ 7.50	2.47	0.92
UKB 3	6564	729	3406/3887	45–81	64.15 $\pm$ 7.53	2.51	0.92
UKB 4	6665	740	3523/3882	44–81	64.05 $\pm$ 7.58	2.48	0.92
UKB 5	6667	740	3507/3900	45–81	64.05 $\pm$ 7.52	2.50	0.92
UKB full	33158	3682	17557/19283	44–82	64.09 $\pm$ 7.53	2.35	0.93

scanning site 1–3, we obtained a MAE of 2.4 years ($r_{\text{Age}} = 0.93$), while the reverse resulted in a MAE of 2.48 years ($r_{\text{Age}} = 0.92$).

4 | VALIDATION OF THE GLOBAL MODEL ON EXTERNAL DATASETS

Due to the limited age range of the UKB sample, we trained the GM models on a previously described normative dataset that matched the age range of the training and validation samples. As the most prominent differences are expected in the GM tissue, only the results of the mean-ensembled GM model are presented in the following section.

4.1 | Synthetic dataset of global neocortical thinning

We applied the global GM model to a synthetic dataset of global neocortical thinning from Rusak et al. (2022), who simulated neocortical thickness loss from 0 to 0.1 mm in steps of 0.01 mm per time point. The increment reflects a yearly rate of cortical loss after the age of 70 (Fjell et al., 2009; Fox & Schott, 2004). The authors also simulated the hypothetical loss from 0.1 to 1 mm in 0.1 mm increments, which would represent about 100 years of global neocortical thinning.

Our aim was to investigate how the BrainAGE algorithm performs on simulated data with well-defined neocortical atrophy. We wanted to test whether the BrainAGE algorithm utilises the global pattern of neocortical thinning for age prediction, and see to what extent the BrainAGE scores reflect the putative annual changes in neocortical thickness (Figure 2).

As can be seen in the figure, the BrainAGE scores reflect the loss of global cortical thickness, although the values do not represent the expected annual rate of cortical thinning as only neocortical atrophy was simulated in the dataset.

4.2 | Application to the ADNI sample

Furthermore, the global GM model was used to predict the BrainAGE in the ADNI sample of healthy controls, subjects with stable MCI and the ones who later progressed from MCI to AD, as well as AD patients. The ADNI test samples were matched with regard to age distribution of the normative database. The results are presented in Figure 3a.

One-way ANOVA showed statistically significant differences between the mean BrainAGE scores of the four groups ($F(3, 691) = 55.69, p < .001$). Post hoc comparisons using Tukey HSD test showed significant differences in mean BrainAGE scores of healthy controls and subjects with AD ($p < .001$), healthy controls and pMCI ($p < .001$), as well as sMCI and pMCI ($p < .001$) and sMCI and AD patients ($p < .001$) at baseline.

4.3 | Application to the schizophrenia dataset

We applied the global GM model trained on a normative sample to the SZ dataset (Figure 3b). Statistically significant differences were found between the groups of patients with different symptoms ($F(3, 191) = 9.64, p < .001$). Tukey's HSD *post hoc* test revealed statistically significant differences in mean BrainAGE scores of healthy controls and patients with negative ($p < .001$) and positive symptoms ($p = .001$), but not with the disorganised group ($p = .715$).

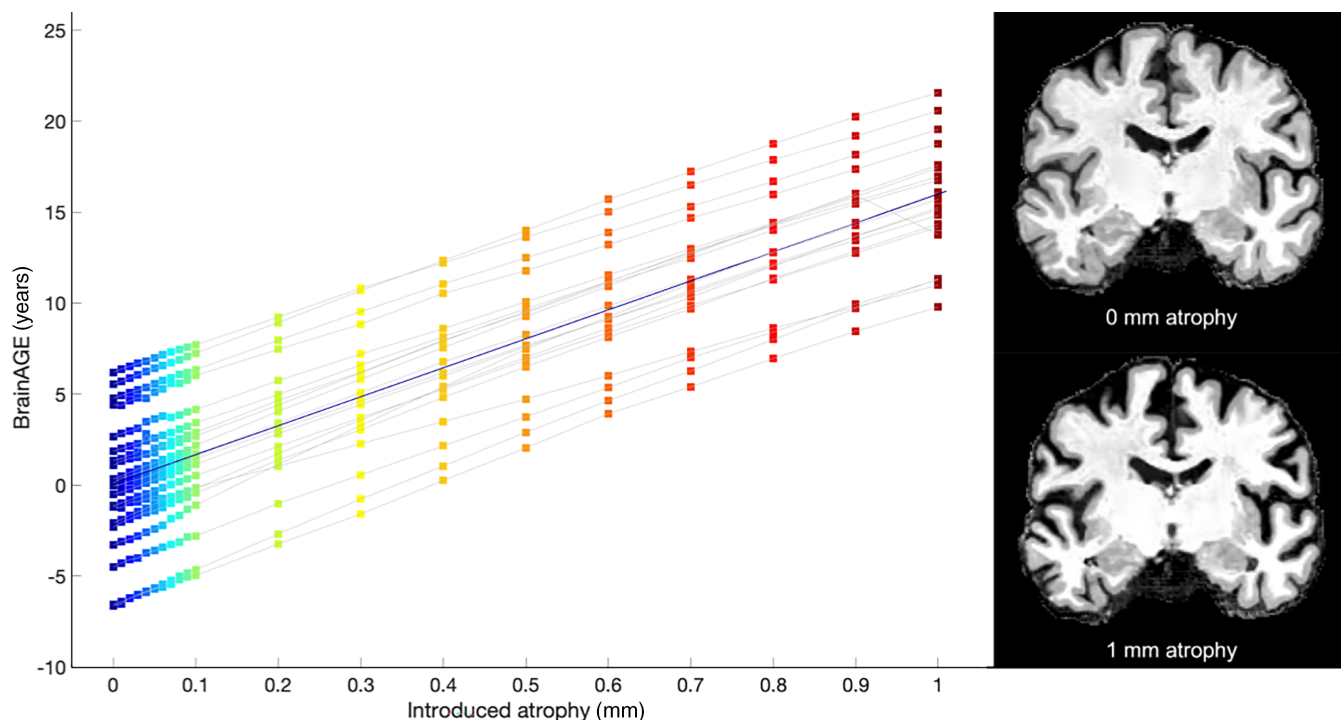


FIGURE 2 BrainAGE scores of the simulated neocortical thinning dataset with 0 and 1 mm atrophy examples on the right side.

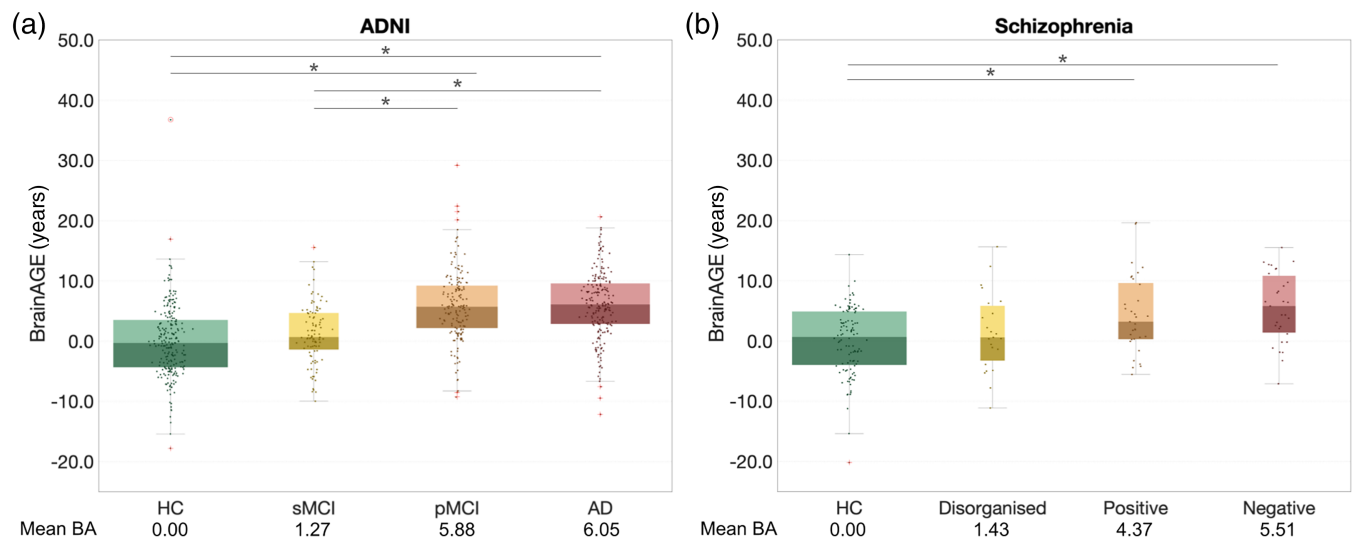


FIGURE 3 (a) Mean BrainAGE scores for healthy individuals and patients with differing levels of neurocognitive impairment from the ADNI sample. (b) Mean BrainAGE scores for healthy control participants and groups of patients with varying symptoms from the SZ sample.

TABLE 3 MAEs of the mean-ensembled model performance and Pearson's correlation coefficients between the predicted and chronological age for regional BrainAGE models on the large sample of UKB with the linear age-bias correction as in Smith et al. (2019).

	Frontal		Parietal		Occipital		Temporal		Subcortical/Cerebellar	
	L	R	L	R	L	R	L	R	L	R
MAE (years)	2.70	2.70	2.65	2.63	2.55	2.55	2.67	2.63	2.67	2.64
r_{Age}	0.92	0.92	0.91	0.91	0.90	0.90	0.92	0.92	0.94	0.94

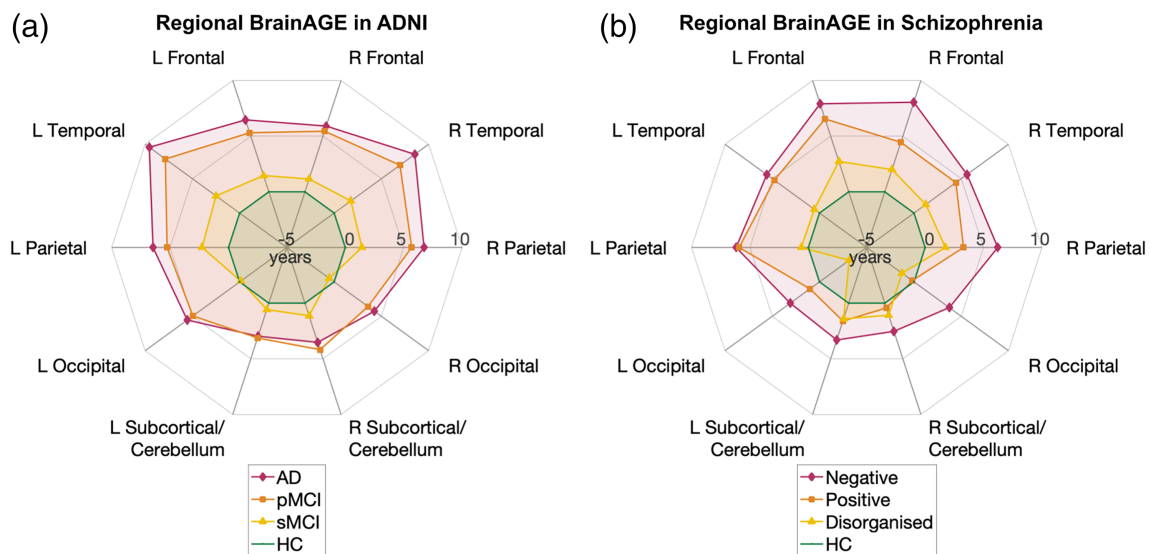


FIGURE 4 (a) Regional BrainAGE scores in healthy control subjects (HC), patients with stable MCI (sMCI), progressive MCI (pMCI), and AD patients from the ADNI dataset. (b) Regional BrainAGE scores for control subjects and three subgroups of SZ patients.

5 | REGIONAL BrainAGE IN HEALTH AND DISEASE

To support the spatial specificity of the BrainAGE model, we developed a regional approach, parcellating the brain into 10 regions

(5 per hemisphere). Separate GPR models were applied to a large sample of UKB and two clinical samples of AD and schizophrenia. The regional results from the large UKB sample are shown in Table 3. Regional BrainAGE scores show an anterior-posterior pattern with higher frontal and lower occipital values, and a slight left-

right difference with higher BrainAGE scores on the left side compared to the right side. See Tables S4 and S5 for the results without age-bias correction and with correction as in Cole (2020). The results of the models applied to AD and SZ patients are shown in Figure 4.

Higher BrainAGE scores can be observed especially in the temporal regions for AD and progressive MCI patients. In contrast, the sMCI group showed only minor differences overall and none in the occipital regions. The results for SZ patients are evident in the frontal, temporal, and parietal regions, most notably for the subjects with negative symptoms. The nearly unaffected disorganised group only had slightly increased brain age values, mostly in frontal regions.

6 | DISCUSSION

In our present work, we have revised the BrainAGE model that was introduced over a decade ago (Franke et al., 2010). We replaced RVR with a GPR algorithm, which has proven to be more stable when dealing with large datasets. This was demonstrated through extensive evaluation of the performance of the new framework under different conditions, using the large neuroimaging sample of the UKB. We explored the contribution of grey and white matter tissue segments and their ensembles to BrainAGE prediction accuracy. By employing the model on the ADNI and SZ samples, we tested its external validity in dealing with pathological brain tissue changes. In addition, the application of the model on a synthetic dataset showed how the algorithm reflects uniform decreases in global neocortical thickness. Furthermore, we have developed a regional BrainAGE model to increase the spatial specificity of the BrainAGE prediction. The advantages and limitations of our new BrainAGE frameworks are discussed in the following sections.

6.1 | Dimension reduction and training sample size

The results revealed that reducing highly individual features, which can be seen as anatomical noise, is one of the most effective ways to improve brain age estimation. In our case, this was conducted by smoothing, downsampling, and finally a PCA. As already noted in Franke et al. (2010), not all voxels of structural images are of equal importance to the brain age estimation, therefore, dimensionality reduction is one of the strategies to obtain more accurate results in case of highly dimensional feature space (Baecker et al., 2021; More et al., 2023). It does not only reduce the computational costs, but also prevent method-specific over-fitting of machine learning models (Franke & Gaser, 2019).

While RVR shows its superiority over GPR on smaller sample sizes (Baecker et al., 2021), training of a GPR model on a large enough sample yielded slightly better results than the RVR approach. However, increasing training sample size over 13,000 subjects resulted in plateauing of BrainAGE accuracies.

6.2 | Tissue class combination and model ensembling

Based on our analyses, a combination of models trained on affine registered GM and WM segments with varying preprocessing parameterization jointly contribute to the best prediction in our performance test on the UKB. Interestingly, the use of non-linearly registered GM data—typically used in VBM studies—did not result in a better performance in comparison to the models that utilised affine registered GM. We speculate that linear registration provides more spatial information of the individual brain anatomy and its atrophy pattern, being more analogous to raw brain anatomy used for deep learning methods, whereas the segmentation (with its denoising and bias-correction steps) helps to remove protocol-based image properties (Ashburner & Friston, 2000).

An important factor that strongly improved the prediction accuracy of BrainAGE estimation was combining various single BrainAGE models. Model ensembling is an effective way of improving the brain age estimation as it serves as a regularisation step to prevent overfitting and thus improves prediction accuracy (van Veen et al., 2015). Although the models we ensemble were quite similar, we do not consider this a strong drawback. Similar models can still provide additional variation (van Veen et al., 2015), and our results show improvement without a significant increase in computational costs. As can be seen from our results, employing GPR stacking substantially decreased the MAE.

Additional improvements of age prediction may be possible by ensembling more independent brain age models, for example, by using different image modalities (e.g., DWI, T2-weighted MR, rs-fMRI), morphometric metrics (e.g., thickness, sulcal depth), or other brain-age frameworks (e.g., Cole et al., 2018; Smith et al., 2019). Although (slight) improvements by adding modalities were shown before (e.g., Cole, 2020; Niu et al., 2020), we focused only on broadly available T1-weighted imaging data and a standard VBM paradigm in order to keep our framework parsimonious. Moreover, the incorporation of multiple modalities generally enhances prediction accuracy, but does not necessarily improve discrimination of relevant phenotypes (Jirsaraie et al., 2023). Nevertheless, the inclusion of other modalities may prove beneficial in cases where structural changes are less pronounced.

6.3 | Generalisability and validity

Our results show a comparable performance of the algorithm based on training and testing on different UKB subsamples. A repeated 10×10 CV performance demonstrated high consistency between the predicted BrainAGE scores across different folds and repetitions (Table S3). Moreover, the cross-scanner prediction within two UKB sample sites shows consistent performance. Further testing of generalisability of the BrainAGE approach was done using the ADNI and SZ samples. The BrainAGE model showed meaningful differences between the diagnostic groups.

The application of the algorithm to the synthetic global neocortical atrophy dataset proved that the BrainAGE algorithm interprets cortical thinning as ageing, however, the obtained values do not reflect the expected annual changes. This discrepancy may be attributed to the properties of the synthetic dataset, which only reflects the neocortical tissue loss, but does not simulate changes in white matter, ventricles, cerebellum, deep grey matter, hippocampus and amygdala (Rusak et al., 2022). These structures are also known to drive brain age prediction (Levakov et al., 2020). Despite the artificial character of simulated atrophy pattern, such datasets allow for testing specific structural changes (Aubert-Broche et al., 2006, Lerch & Evans, 2005, Rusak et al., 2022) and their interaction with the brain age algorithms. Simulated datasets can potentially serve as a complementary approach to explainable AI by providing information on the influence of anatomical characteristics on brain age prediction.

Overall, the revised BrainAGE algorithm provides promising and generalisable results. Future studies might benefit from harmonising data over multiple imaging studies (Pomponio et al., 2020), thus allowing the application of brain age prediction to more diverse data.

6.4 | Regional BrainAGE

Global BrainAGE models fail to grasp neurodegenerative- and psychiatric disease-specific structural brain changes (Gianchandani et al., 2023; Kaufmann et al., 2019). To enhance the spatial specificity of the BrainAGE model, we developed a regional BrainAGE approach that estimates brain age in five lobular brain regions per hemisphere. On average, the regional predictions were less accurate than the global measure, which is expected due to the lower amount of information provided by a single lobe. Nevertheless, the regional BrainAGE scores provided meaningful information about brain ageing in clinical samples. When applied to the ADNI sample, an increase in BrainAGE scores was observed in the temporal lobe of AD patients and MCI subjects converting to AD, which corresponds to the involvement of (medial) temporal lobe in AD (Migliaccio & Cacciamani, 2022). The results of the SZ sample were consistent with the findings of the previous study on the same sample, where Nenadić et al. (2015) found the most prominent neocortical thinning in the negative symptoms group and the least in the disorganised group. Similar to the studies of Kaufmann et al. (2019) and Zhu et al. (2023), our sample of SZ patients had the highest BrainAGE scores in the frontal and temporal regions. Additional studies are required to validate the regional BrainAGE model in various clinical groups.

7 | CONCLUSION

Our BrainAGE algorithm underwent a thorough revision that enabled its application on the very large UK Biobank dataset. Moreover, we extended the framework to support regional brain age estimation and increase the spatial specificity of the method. Application of the BrainAGE models on subjects with varying levels of neuropsychiatric- and

neurocognitive impairment showcases the model's clinical validity and the BrainAGE scores from a synthetic dataset reflect a meaningful pattern according to known neocortical thickness loss with age. Future research can profit from the use of synthetic data to better understand the inner workings of ML algorithms.

AUTHOR CONTRIBUTIONS

Polona Kalc: Writing—original draft; visualisation. **Robert Dahnke:** Writing—review and editing; visualisation; data curation; software. **Felix Hoffstaedter:** Data curation; writing—review and editing. **Christian Gaser:** Conceptualization; methodology; formal analysis; software; writing—review and editing; visualisation; supervision; funding acquisition.

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CONFLICT OF INTEREST STATEMENT

The authors declare no conflict of interest.

DATA AVAILABILITY STATEMENT

The data used for this work were obtained from the UK Biobank Resource (Project Number 41655). Due to the nature of the data sharing agreement, we are not allowed to publish the data.

The ADNI dataset is publicly available at <https://ida.loni.usc.edu/login.jsp?project=ADNI> and IXI at <https://brain-development.org/ixi-dataset/>, Cam-CAN is available under <https://camcan-archive.mrc-cbu.cam.ac.uk/dataaccess/>, OASIS-3 can be found on <https://www.oasis-brains.org/#data>, Enhanced NKI-RS dataset is available at https://fcon_1000.projects.nitrc.org/indi/pro/eNKI_RS_TRT/FrontPage.html, and the SALD dataset can be downloaded from https://fcon_1000.projects.nitrc.org/indi/retro/sald.html. The synthetic dataset from Rusak et al. (2022) is available at the CSIRO data access portal.

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SUPPORTING INFORMATION

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