Association vs. Prediction: The Impact of Cortical Surface Smoothing and Parcellation on Brain Age

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Association and prediction studies of the brain target the biological consequences of aging and their impact on brain function. Such studies are conducted using different smoothing levels and parcellations at the preprocessing stage, on which their results are dependent. However, the impact of these parameters on the relationship between association values and prediction accuracy is not established. In this study, we used cortical thickness and its relationship with age to investigate how different smoothing and parcellation levels affect the detection of age-related brain correlates as well as brain age prediction accuracy. Our main measures were resel numbers—resolution elements—and age-related variance explained. Using these common measures enabled us to directly compare parcellation and smoothing effects in both association and prediction studies.

In our sample of N = 608 participants with age range 18–88, we evaluated age-related cortical thickness changes as well as brain age prediction. We found a negative relationship between prediction performance and correlation values for both parameters. Our results also quantify the relationship between delta age estimates obtained based on different processing parameters. Furthermore, with the direct comparison of the two approaches, we highlight the importance of correct choice of smoothing and parcellation parameters in each task, and how they can affect the results of the analysis in opposite directions.

Keywords: brain aging, cortical thickness, prediction, delta age, smoothing, parcellation, association

INTRODUCTION

From a biological standpoint, aging is defined by the structural and functional alterations in living organisms (López-Otín et al., 2013). Traditionally, brain imaging studies have used neuroimaging data to find associations between age and tissue alterations across brain areas, using chronological age as the ground truth (Lemaître et al., 2005; Curiati et al., 2009; Takahashi et al., 2011; Ziegler et al., 2012; Booth et al., 2013; Hu et al., 2014). However, biological age might vary between individuals with identical chronological age as well as across different tissues within the same person (Horvath, 2013). To non-invasively measure the biological age of the brain, neuroimaging data is used to predict age. The difference between predicted age and chronological age is then defined as “delta” or brain age gap estimate i.e., “BrainAGE” to compare the subjects’ chronological age with the
predicted brain age in a given reference population (Franke et al., 2012; Cole and Franke, 2017; Franke and Gaser, 2019; Smith et al., 2019).

Both age-related brain alterations and delta age have been studied and used extensively in the neuroimaging literature. Age association studies translate and generalize easily across different datasets. These association studies are applied across brain regions and can distinguish the differential effect of age on different brain areas (Storsve et al., 2014). Furthermore, they directly relate to biological measures and mechanistic changes in the brain (Khundrakpam et al., 2015). More recently, it has been recognized that association studies are prone to overfitting and more studies focus on prediction as the main goal of the study (Yarkoni and Westfall, 2017; Bzdok et al., 2020). Brain age studies (i.e., age prediction studies based on neuroimaging data) rely on modeling and prediction accuracy. This goal is generally achieved by using a feature set that can capture the variability between and within subjects. On the other hand, prediction tasks face a trade-off between a more accurate whole brain model with no regional specificity vs. a model with lower accuracy and increased spatial resolution (Cole and Franke, 2017; Franke and Gaser, 2019). This limitation also results in a more indirect relationship between delta age and other phenotypes without a direct mechanistic and biological model. Nonetheless, the difference between brain age and chronological age is associated with cognitive decline (Gaser et al., 2013), predisposition to neuropsychiatric and neurodegenerative disorders (Kaufmann et al., 2019), and mortality (Cole et al., 2018). While evidence supports the application of delta age as a valuable measure to study aging in health and disease, it has been criticized due to its reliance on prediction accuracy (i.e., more accurate models result in lower delta values) (Cole and Franke, 2017).

The results of both association studies and delta estimation studies are impacted by processing steps such as data normalization, spatial resolution, and parcellation level (i.e., size of the parcels) of the analysis. Most association studies use smoothing to (i) normalize the distributions of cortical thickness across subjects, (ii) minimize registration and anatomical misalignment across subjects, (iii) reduce measurement noise, and (iv) increase statistical power (Worsley et al., 1999; Lerch and Evans, 2005; Lerch et al., 2006; Zhao et al., 2013). These advantages are gained at the cost of losing individual variability and spatial resolution. In fact, smoothing has been studied and optimized for best performance in association studies, using simulation as well as in real datasets. The smoothing level has been proposed as a dimension within the parameter space in the association analysis that needs to be searched for the given statistical contrast (Lerch and Evans, 2005; Zhao et al., 2013).

Brain age prediction studies have been conducted with various levels of data smoothing. Moreover, these studies rely on various dimension reduction techniques, brain parcellations, or a combination of the two approaches for feature extraction (Franke and Gaser, 2019; Smith et al., 2019). The optimal parcellation for a given task is an open research topic and it can vary between studies (Gorgolewski et al., 2016; Eickhoff et al., 2018; Salehi et al., 2020). While some studies have predicted brain age with multiple parcellation resolutions (Khundrakpam et al., 2015; Lewis J. D. et al., 2019), others have used a predetermined number of parcels. However, the effect of smoothing and parcellation in brain age prediction is not studied systematically. Furthermore, these changes in prediction accuracy also affect the delta estimate (i.e., the variable of interest), and it is not clear whether the delta estimates are robust or sensitive toward these initial choices.

In this study, we used cortical thickness as the brain measure of interest and examined the effect of smoothing and parcellation level on both brain associations with age and brain age prediction. Using different levels of parcellation and smoothing, we projected brain measures onto a lower dimension data representation space and investigated how this mapping affects the derived associations and predictions. We further examined the relationship between the two approaches. Finally, we examined how delta age estimates alter based on different smoothing and parcellation levels.

**METHODS**

**Data**

Data used in this study included subjects with T1-weighted MRI data available from the second stage of the Cambridge Centre for Ageing and Neuroscience (CamCAN, https://www.cam-can.org/index.php?content=dataset) dataset, described in more detail in Shafro et al. (2014) and Taylor et al. (2017). Subjects were screened for neurological and psychiatric conditions and those with such underlying disorders were excluded from the study.

**MRI Acquisition**

T1-weighted MRIs were acquired on a 3T Siemens TIM Trio, with a 32 channel head-coil using a 3D magnetization-prepared rapid gradient echo (MPRAGE) sequence (TR = 2,250 ms, TE = 2.99 ms, TI = 900 ms; FA = 9°; FOV = 256 × 240 × 192 mm; 1 mm isotropic; GRAPPA = 2; TA = 4 min 32 s). For detailed acquisition parameters see: https://camcan-archive.mrc-cbu.cam.ac.uk/dataaccess/pdfs/CAMCAN700_MR_params.pdf.

**MRI Processing**

We used CIVET 2.1.1 (http://www.bic.mni.mcgill.ca/ServicesSoftware/CIVET, release December 2019), a fully automated structural image analysis pipeline developed at the Montreal Neurological Institute, to perform surface extraction and cortical thickness estimation. Briefly, each subject's T1-weighted MRI is corrected for non-uniformity artifacts using the N3 algorithm (N3 distance = 125 mm) (Sled et al., 1998) and linearly registered to stereotaxic MNI152 space (voxel resolution = 0.5 mm) (Collins et al., 1994). The brain is extracted and undergoes tissue classification into three classes: white matter (WM) tissue, gray matter (GM) tissue, and cerebrospinal fluid (CSF) (Zijdenbos et al., 2002; Tohka et al., 2004). White and gray matter surfaces are extracted using the marching cube algorithm and constrained Laplacian-based automated segmentation with proximities (CLASP) algorithms, respectively (MacDonald et al., 2000; Kabani et al., 2001; Kim et al., 2005). Using the extracted surfaces, cortical thickness is measured as the distance between the white and gray cortical surfaces using the Laplace's equation (Jones et al., 2000). For blurring, a surface-based diffusion
smoothing kernel (not to be confused with volumetric kernels) is used, which generalizes Gaussian kernel smoothing and applies it to the curved cortical surfaces (Chung et al., 2002). We applied 6 different smoothing levels with FWHM = 0, 5, 10, 20, 30, and 40 mm. Cortical thickness was measured across the cortical surface for 81,924 vertices (40,962 vertices per hemisphere). The results underwent visual inspection, specifically subjects with major errors in extracted pial and gray–white surfaces were excluded.

Cortical Parcellations

We used the Schaefer functional MRI parcellations (Schaefer et al., 2018), a data-driven atlas based on the widely used seven large-scale functional network parcellations by Thomas Yeo et al. (2011). We used Schaefer parcellation with 100, 200, 400, and 1,000 regions (referred to as parcellation levels). All atlases were registered to the MNI cortical surface template and used in the MNI space (Lewis L. B. et al., 2019). Cortical thickness measurements with different smoothing levels were averaged across these parcellations. These parcellation based measures of cortical thickness were used alongside vertex-wise measurements to examine the interaction between the effect of brain parcellation averaging and smoothing on statistical associations as well as brain age prediction accuracies.

Cortical Resels and Effective Smoothing

In order to compare the findings between smoothing levels and different parcellations, first all obtained cortical thickness were projected to the brain surface. We used the number of resels (i.e., resolution elements) as the measure of interest, since it takes the statistical dependence of the brain map into consideration and is independent of the analysis resolution (at least from a theoretical standpoint) (Worsley et al., 1992, 1999; Worsley, 1996; Lerch et al., 2006). Using the statistical maps between aging and cortical thickness, we estimated the number of resels for each smoothing and parcellation level and used it to quantify the similarity between these conditions. Resels are the number of resolution elements approximated for a given search space [i.e., D(S2), S2 = brain surface] and a given smoothness level FWHM. While the effective FWHM measure varies across brain areas, we defined the overall effective smoothness of the brain map as the square root of the surface search space divided by the number of resels estimated across brain areas (Hayasaka et al., 2004). For the purpose of the current study, the main statistical maps considered are the linear associations between cortical thickness and the chronological age of the participants. All analysis were performed using SurfStat toolbox https://www.math.mcgill.ca/keith/surfstat/ (see Supplementary Methods for further details).

Statistical Methods

To examine the effect of the smoothing and parcellations, mean (µ) and standard deviation (σ) of cortical thickness for each vertex/parcel was calculated across the population. The coefficient of variation (CV), CV = σ/µ, was used as the main measure of variability. The CV was averaged across the 7 main cytoarchitectural brain regions (von Economo and Koskinas, 1927) in order to examine the effect of parcellation and smoothing across major cytoarchitectural regions and identify any differential impact on a given brain region. Finally, to measure the association between chronological age and cortical thickness across lifespan, correlation coefficient (r) for each vertex/region was calculated. Variance explained (r²) was used to visualize the results.

Brain Age Prediction

We used principal component analysis (PCA), a singular value decomposition based data factorization method, as the dimensionality reduction approach for our predictive variables (i.e., cortical thickness data) (Smith et al., 2019). This approach allowed us to use the same number of features across parcellation levels and smoothing kernels and therefore made it possible to compare model performance across these conditions. Our analysis for each condition included 1 to 100 first principal components as features to study different levels of dimensionality reduction. Hundred is used as the maximum possible number of independent components for the lowest number of parcels (i.e., Schaefer 100). To predict brain age, we used linear regression as the main prediction model, and to ensure generalizability and avoid overfitting, we used 10-fold cross validation. Finally, to increase robustness, results averaged over 100 repetitions are reported, however as discussed these repetitions are not necessary and had no impact on the conclusions. Root-mean-squared error (RMSE) was used as the natural cost function for linear regression models. Mean absolute error (MAE) and correlation between chronological age and predicted age (two other common error metrics in the age prediction literature; Franke and Gaser, 2019; Franke et al., 2020) are also reported in the Supplementary Materials. Finally, we have repeated the same procedure using a support vector machine (SVM) regression method with linear kernel as well as linear regression models with lasso and ridge regularization (results reported in the Supplementary Materials).

The Relationship Between Brain Age Prediction and Age Related Brain Association

To compare brain age association and age prediction, we used the variance explained between dependent and independent variables as the main measure of interest for each model. This common measure enabled us to quantify the two analyses in relation to each other. Furthermore, we examined how the number of resels affects whole brain associations with age as well as brain age prediction. To translate the age prediction error into variance explained, we used the predictive features in a linear model, calculating the variance explained for age using adjusted R². Finally, the overfitting bias between the variance explained (i.e., adjusted R²) using this linear model and the cross validated prediction (i.e., r² between predicted age and chronological age) is reported.

Delta Age

The main goal of brain age prediction studies is to calculate the deviation from chronological age based on the population
norm, also known as delta age. Here, we examined the effect of smoothing and parcellation on delta age estimation:

\[ Y = X\beta_1 - \delta_1 \]

where \( Y \) denotes chronological age, \( X \) denotes the neuroimaging features, and \( \delta_1 \) denotes the difference between predicted and chronological age. \( \delta_1 \) is a measure of brain state/health compared to the population with similar chronological age, and is used to study the predisposition to different brain disorders as well as individual cognitive abilities in neuroimaging literature.

\( \delta_1 \) being residual of the predictive model is by definition: (1) orthogonal to the predictive measures \( X \), and in the case of linear models (2) correlated with the output \( Y \) (i.e., chronological age) (Le et al., 2018; Liang et al., 2019; Smith et al., 2019). The first feature is unfavorable, since we are interested in brain related discrepancy between chronological and predicted age. The lack of association between \( \delta_1 \) and brain features predicting age undermines the interpretability of \( \delta_1 \) in relation to brain measures. The second property is also an adverse feature, since it makes it difficult to distinguish the effect of the chronological age from the additional biological delta age (due to their collinearity). Therefore, in the current study, we followed the recommendation of smith and colleagues (Smith et al., 2019) and used \( \delta_2 \), the orthogonalized residuals against chronological age:

\[ \delta_2 = \delta_1 - Y\beta_2 \]

\( \delta_2 \) is then used as the main measure of interest for association across conditions. The results for \( \delta_1 \) is provided in the Supplementary Materials. Note that \( \delta_2 \) is also consistently calculated using the same 10-fold cross validation with 100 repeats as \( \delta_1 \), however as discussed these repetitions are not necessary and had no impact on the conclusions. All statistical and prediction analyses were performed using MATLAB 2018a.

RESULTS

Cortical Thickness Aging, Resels, and Practical Smoothness

The parcellations have a considerable impact on the number of resels and function as region-based smoothing kernels applied across the brain (Figure 1A). This change in the number of resels affects the statistical power and the association as well as prediction results. Across parcellation levels from 100 to 1,000, the effect of the smaller smoothing kernels with FWHM 0–10 mm is negligible, while applying larger kernels reduces the number of resels dramatically. This equivalency plot also suggests that at the vertex level, the smoothing kernels act as a non-specific parcellation (from an anatomical perspective) across the brain.

Cortical Thickness Variability

While keeping the mean cortical thickness measure intact, smoothing resulted in underestimation of the cortical thickness in the gyri areas and overestimation in the sulci regions. The results are similar for parcellations in the case of uniformly sized parcels and balanced inclusion of gyri and sulci in each parcel (both criteria are met in Schaefer parcellations). Cortical thickness variability (i.e., CV) reduces significantly both as a result of using greater smoothing and larger parcels (Figure 2A).

The association cortices have the lowest CV across resolutions and parcellations. Both smoothing and parcellation result in the highest decrease in CV in limbic and insular cortices, while primary sensory and motor areas show the lowest change (Figure 2B). The results are shown for 0 mm smoothing across parcellations. The greatest change occurs with increasing the FWHM value from 10 to 20 mm, as well as decreasing the number of parcels from 400 to 200. The results for different smoothing kernels at vertex level were also similar (Supplementary Figure 1).

Statistical Association Between Cortical Thickness and Aging

Figure 3A shows the association between age and cortical thickness (using variance explained \( r^2 \)), calculated for each voxel/parcel for all conditions, after Bonferroni correction to account for the multiple comparisons at each level. The correlation increases with greater smoothing and larger parcels. Changing smoothing kernel size results in the highest variability in the correlation distribution across the brain at vertex level resolution (Figure 3B, top panel), whereas smoothing doesn’t change the results within Schaefer 100 parcellations (Figure 3B, bottom panel). The same pattern is evident between parcellation levels with 0 mm smoothing showing the highest variability, and 40 mm smoothing with lowest variability across parcellations. These findings are further explained with reference to the number of resels and effective smoothing in section The Relationship Between Prediction and Association. Finally, while present across all brain areas, the variability between correlation maps is the highest within association cortices, primary motor, and insular cortex.

Brain Age Prediction Based on Cortical Thickness

For age prediction, vertex-level data outperformed all parcellation-based data using the same (or a smaller) number of principal components as predictive features. The accuracy was also higher for lower smoothing kernel size. However, this effect was more pronounced for FWHMs >10 mm, and the results for FWHM values of 0, 5, and 10 mm showed a very similar performance in the vertex-level analysis. A similar pattern was present within each parcellation level. The best performing models (i.e., 0 and 5 mm smoothed vertex-wise), reach their minimum error using the first 20–30 principal components as features in the prediction model (i.e., a sample to feature ratio of 28–18). The pattern was similar for MAE and correlation between predicted age and chronological age (Supplementary Figures 2, 3).

The Relationship Between Prediction and Association

As expected, there was a negative relationship between the overall correlation between age and cortical thickness across
brain regions (measured by median $r^2$) and the number of resels within each condition (Figure 5A). Interestingly, we found a positive association between the number of resels and the overall ability of cortical thickness features to explain the variance of chronological age (as measured by adjusted $R^2$ of the linear model) shown in Figure 5B. These results suggest that the higher number of resels results in lower correlation values, but since resels are independent based on their relationship with age, they can explain different modes of chronological age within the population (hence the higher adjusted $R^2$), whereas, in conditions with lower resel numbers (i.e., higher smoothing and larger parcels) the correlation values are higher but homogenous across the brain and therefore explain a lower proportion of the age variance.

Finally, there was a strong linear relationship between (i) the overall variance explained (adjusted $R^2$) using a linear model with
age as dependent variable and PCs as independent variable and (ii) the predictive performance of the linear regression model, with a bias due to overfitting in the linear model (Figure 5C). Figure 5D shows the overfitting bias of the adjusted $R^2$ compared to the cross-validated prediction, as a function of the number of features in the model. Taken together, these results explain the opposing directions between correlation results and prediction accuracy across parcellation and smoothing conditions.

**The Effect of Smoothing and Parcellation on the Estimation of Brain Age Delta**

In this section, we present $\delta_2$ age prediction accuracy results with 10-fold cross validation. The prediction accuracy based on the modified $\delta_2$ is presented in Figure 6. One of the main assumptions in age prediction studies is that delta age measured in different studies using different processing parameters are similar and can be interpreted as the same measure. We have examined the relationship between the optimal $\delta_2$ across different parcellations and smoothing kernels (Figure 7). These results demonstrate the degree of sensitivity of $\delta_2$ as a function of our choice for parcellation and smoothing kernel. While there is high correlation for large smoothing kernels (20–40 mm) as well as lower number of parcels, these conditions have the lowest prediction accuracies. The correlations between these conditions and higher accuracy conditions (i.e., vertex-wise and 1,000 parcels with 0–10 mm smoothing) are lower ($r \sim 0.55$). See the results for $\delta_1$ in the Supplementary Figure 4.
DISCUSSION

In this article, we compared the effect of different smoothing and parcellation on associations between cortical thickness and chronological age as well as brain age prediction accuracy. We showed that the optimal choice for association analysis might indeed undermine age prediction accuracy, and vice versa. We further investigated this relationship and demonstrated the underlying differences that lead to this trade-off between the two analyses. Finally, we examined the effect of smoothing and parcellation on delta age estimation and showed that the initial smoothing and parcellation choices can change the delta estimation which in turn will affect any downstream analysis.

We used brain association with age and brain age prediction as our target analyses, since age is used as the main variable of interest or at least a confounding variable in most neuroimaging studies. We used cortical thickness as the main measure of interest due to the wide availability of T1-weighted MRI in research and clinical settings, cortical thickness is a suitable measure which has been widely used to study brain anatomy in general (Toga, 2015), and more specifically, brain aging and predicting brain age (Wang and Pham, 2011; Groves et al., 2012; Kandel et al., 2013; Liem et al., 2017). Finally, our results are presented based on a sample size of N~600 which is a common sample size for publicly available datasets in the field of neuroimaging.
Given the limited number of subjects in neuroimaging studies compared to potential features (number of vertices/voxels), most prediction studies apply dimension reduction as an initial step. We used PCA for dimension reduction of the cortical thickness data. Due to its simplicity and interpretability, PCA has been widely used in the brain age prediction literature. Furthermore, we employed linear regression with cross-validation as our prediction model (Smith et al., 2019). As expected, we observed an initial drop in the prediction error, followed by a plateau/increase in the error as the sample to feature ratio increases (Hastie et al., 2009). At each parcellation level, the accuracy drops with increased smoothing, and for each smoothing level, the accuracy decreases with larger parcels/regions.
FIGURE 6 | Root mean square error (RMSE) for Age prediction with $\delta_2$ as the error term. The x axis shows the number of principal components included as features. The results are grouped based on the parcellation resolution.

FIGURE 7 | $\delta_2$ age prediction error. The correlation between delta age (as measured by $\delta_2$) across parcellation resolutions (x and y axis labels) and smoothing kernels (represented by circle size).
The age prediction results presented in the main manuscript are based on linear regression analysis and PCA based features. We have also examined the performance of support vector machines with different kernel types as well as linear regression using lasso and ridge regularization methods. All these methods were applied on both raw cortical thickness values as well as PCA based features as predictors. In all cases, the PCA based features outperformed the same method using the raw cortical thickness values. These results can be due to the relatively small sample size and/or sample to feature ratio in the current study. Furthermore, with the exception of linear regression models with lasso regularization, the presented linear regression method outperformed all other methods. In the case of lasso, we optimized the method for the regularization weight (i.e., lambda parameter). With this optimization, we gained a 2% increase in accuracy. All the results based on the explained models are reported as Supplementary Tables 1–4 and Supplementary Figures 6–8. Although the accuracy might vary slightly between methods, the higher accuracy in smaller smoothing kernels (i.e., 0 and 5 mm smoothing) and smaller parcels (i.e., vertex based level) is consistent across all methods and the explained relationship between age-association and prediction holds true across these prediction models. Furthermore, the anatomical correlates of aging or the main anatomical features contributing to the brain age prediction were not the main target or in the scope of the current study. However, the mapping of the first 100 PCs (used in the prediction analysis) to the Schaeffer parcellation as well as whole brain vertices of CIVET is provided in the Supplementary Tables 5, 6. They can be used alongside the other Supplementary Tables to infer variables of interest and their anatomical distribution.

While 10-fold cross validation is enough in our case, for relatively small samples, the random assignment of data in the cross-validation partitions can lead to differences in the distribution of the training and test data in some of the folds, leading to highly variable performances across some folds. To exercise the best practice and ensure that the reported results are robust, the 10-fold cross validation procedure was repeated and the results were averaged so that such inhomogeneous assignments (however unlikely) do not impact the reported results. The randomized performances were very similar (mean correlation between repetitions was between 0.97 and 0.99) and the standard deviation of the repetitions is <1% of the reported value across repetitions, suggesting that our results are indeed robust and the repetitions were not necessary for the conclusions in the manuscript.

While not exceptionally high, the brain age prediction accuracy in this study is comparable to similar studies in the field (see Franke et al., 2020; Table 3). Furthermore, the accuracy of brain age prediction is dependent on two factors which can significantly impact prediction performance (1) Age range and variance: With the current population’s ages ranging between 18 and 88 years (mean age = 53.52, standard deviation = 18.07), CamCAN dataset is one of the more challenging datasets for prediction. (2) Distribution of age: Prediction models tend to favor values close to the mean of the population. Therefore, data with a Gaussian distribution (which is generally used in other similar brain age prediction studies) will result in a much better prediction performance compared to a rather uniform distribution of age which is the case for the CamCAN dataset.

In terms of variability within the cytoarchitectural regions, there is a distinction between the change in the insular cortex compared to the rest of the regions. The main shift occurs between 400 and 200 parcellation levels (where Insular cortex parcels are combined from 23 to 15 parcels). As a result, several regions (in both right and left hemispheres) with distinct cortical thickness values are combined and averaged together, resulting in a drop in variance and consequently coefficient of variation across regions. This might be due to the unique morphometric properties of insular regions as well as the limited number of parcels in the insular cortex compared to other cytoarchitectural regions. This misalignment might also be due to the functional nature of the Schaefer cortical parcellation, which doesn’t necessarily have a one-to-one correspondence with the structural variability in the same areas.

It is commonplace for neuroimaging studies to use smoothing and parcellation as the first step of their analysis to achieve higher statistical power with reducing the individual variability within the data. Furthermore, with increased availability of public neuroimaging datasets, it is commonplace to release a preprocessed version of the data with a fixed smoothing level and averaged based on a given parcellation. Many research groups in the field use preprocessed and parcellation-based data releases as the starting point for their analyses. In fact, in many cases, the raw data is not publicly distributed, and the preprocessed parcellated data is the only version of data available. For example, some of the most influential public datasets in the field of neuroimaging such as Adolescent Brain Cognitive Development (ABCD, for details see https://nda.nih.gov/abcd) Study and UKBiobank (for details see https://www.ukbiobank.ac.uk) provide cortical thickness data using Desikan-Killiany-Tourville parcellations (Klein and Tourville, 2012) with 62 regions (smoothing varies across studies) as one of their pre-calculated measures. Our findings can help provide a guide to interpret these available measures and shed light on the effect of these preselected parameters/parcellation when applied in aging studies.

Higher correlation values across brain regions (as a result of smoothing) can be explained by increased signal to noise ratio and reduced individual variability (Figure 2). The effect of smoothing on brain related associations has previously been studied (Lerch and Evans, 2005). Indeed, Zhao and colleagues propose smoothing as a scaling dimension which needs optimization for any given target analysis (Zhao et al., 2013). The effect of parcellation on brain association has been addressed in several studies. However, the optimal parcellation level is still an open question dependent on the specific case of interest (Eickhoff et al., 2018). Here, we showed that parcellation level has a similar impact, by reducing variability, using both CV (Figure 2) and number of resels (Figure 1).

Association/correlation analyses reflect the general patterns across the population (suitable for studies that investigate population specific trends), whereas prediction analyses aim to determine the likely value of a certain measure of interest at
the individual level (suitable for diagnosis/prognosis purposes). Association analyses in general benefit from averaging, since it lowers the levels of noise and improves the obtained correlations, allowing the analysis to draw out the overall trends of the population. In contrast, prediction is inherently a much more challenging task, since it aims to provide accurate estimates at the individual level. Averaging methods decrease individual variability and differences, preventing the prediction models from accurately capturing the individual variabilities.

Resel numbers are statistical constructs based on the association analysis, while at the same time informing our interpretation of prediction analysis. As such resels don’t have any inherent biological interpretation. Even within the same dataset and using the same metrics, the number of resels will differ between different statistical analyses (e.g., the number of resels will change if we use fluid intelligence or working memory measures instead of age) since it is a construct that evaluates the number of resolution elements by considering the dependency across regions/vertices with regards to a certain variable within an association or statistical contrast analysis over the region of interest (in our case entire brain surface). A higher number of resels reflects a higher number of independent features (with regards to age), which in turn captures the individual variability across the population and increases prediction accuracy (as confirmed by our findings presented in Figure 5). However, this higher level of individual variability represented in the features will result in lower correlation values across the brain (also shown in our results in Figure 5).

In neuroimaging, smoothing and parcellations are generally studied separately. In this study, we used a unified metric to directly compare the effect of smoothing and parcellation. Using resel numbers and variance explained in the model, we have calculated common measures for both association and prediction results. Our results show that with increased smoothing and larger parcels (i.e., lower number of resels), cortical thickness variability reduces. This will remove inter-individual differences across brain regions and result in higher associations between cortical thickness and aging (Figure 5A). However, while this improves the regional correlation with age, most of this general trend can be captured in a few PCs (mainly the first component) and the rest of the PCs do not explain the remaining variance of age. On the other hand, this relationship is reversed in the conditions with higher resel numbers (i.e., lower smoothing and higher spatial resolutions). While in these cases higher regional variability results in lower correlation with age, the age related associations capture different portions of age variance in different PCs and overall they have a higher adjusted \( R^2 \) (Figure 5B). There was a consistent bias in the adjusted \( R^2 \) across conditions (Figures 5C,D), however, the effects remained similar after removing the overfitting with cross-validation. Altogether, these analyses explain the seeming opposite direction of correlation values and prediction accuracies for different smoothing-parcellation levels in section Statistical Association Between Cortical Thickness and Aging and Brain Age Prediction Based on Cortical Thickness.

One should also consider that while the objective function in linear regression and its variants is based on RMSE (shown in Figure 4), considering the interdependence between the features, there is a close linear relationship between adjusted \( R^2 \) and the prediction accuracy based on the RMSE. Furthermore, our conclusions were independent of the use of RMSE and \( R^2 \) as shown in Supplementary Figure 3. With these considerations, without loss of generality, we have used \( r^2 \) from correlation analysis and adjusted \( R^2 \) from the linear regression model along with the resel numbers (as shown in Figure 5) to study the relationship between the association and prediction analysis.

While delta age in itself is not the target of the current study, it is important in so far as it is the main measure derived from age prediction studies. The discrepancy between predicted age and chronological age (i.e., delta age) is used to study other phenotypes (either demographic, biological, or clinical) (Cole and Franke, 2017). Based on this definition, subjects with higher delta age are assumed to have accelerated aging (i.e., their brain is similar to brains of older individuals). Several studies have found relationships between delta age and brain disorders including but not limited to traumatic brain injury, schizophrenia, epilepsy, mild cognitive impairment, and Alzheimer’s disease. (See Cole and Franke, 2017; Franke and Gaser, 2019; Franke et al., 2020 for a complete review of the topic). A recent study using 45,615 subjects simultaneously investigated the relationship between delta age and 10 different brain disorders and found that subjects with Schizophrenia, Multiple Sclerosis, Mild cognitive impairment, and dementia show higher delta age compared to the controls (Kaufmann et al., 2019). The increase in the studies of brain age emphasizes that not only a better understanding of the biological nature of delta age is needed, but also a systematic study of the effect of analytical and computational methods used to obtain delta age is necessary. However, the effect of the preprocessing condition on delta age estimation is not studied. Here we have examined the effect of parcellation and smoothing levels as an important factor that can change delta age estimation and consequently the aforementioned relationships with other measures. In the current manuscript, we found a range of associations (0.5–1) between \( \delta_{25} \) obtained in different conditions. These results suggest not only that each study needs to optimize their choice of the smoothing and parcellation level, but also when interpreting results from different studies in the field, these parameters should be considered.

One of the main limitations of the current study is the number of subjects (N~600), particularly given that their age spans across 70 years. This leads to overfitting as the number of features increase. In fact, for vertex-wise prediction (with 0 mm smoothing), the first 30 PCs only explain 20% of the variability in the data. This number is around 40% for 10 mm smoothing. In comparison, the first 30 PCs for 100 parcels explain 80 and 90% of the variance of the cortical thickness data for 0 mm and 40 mm smoothing levels, respectively (Supplementary Figure 5). Given the higher performance of the vertex-wise PCs at 0–10 mm smoothing, it is likely that with a larger sample size and increased sample to feature ratio, the accuracy can be further improved. It should be noted that in each case the variance explained corresponds to the total variability for the corresponding smoothing and parcellation condition. Another limitation in the current study is the use of functionally driven Schaefer...
parcellations. While this does not automatically suggest a disadvantage, multi-resolution anatomically driven parcellations have the theoretical advantage of a more relevant initial feature space for cortical thickness studies. Finally, CamCAN data used in our study is cross-sectional. This potentially decreases the detection power of our study, since we can only estimate the effect of time between subjects with individual variability as part of the measurement, whereas a longitudinal dataset can decrease variability by estimating the effect of aging within subjects.

Traditionally, neuroimaging studies have targeted brain related associations with a given phenotype/symptom or the statistical differences between different groups for a given brain region, followed up with the association of these differences with a given biological or behavioral variable of interest. More recently, there has been an ongoing conversation in the field toward prediction as an alternative approach. Along the same line, the field of brain aging, has pursued age related associations as well as age prediction. The relationship between the two approaches is often taken for granted (since in ideal settings, i.e., large sample size and low inter-individual variability or noise levels, the results would be equivalent) and ignored in practice. In this study, we have directly addressed both age association and prediction as a function of smoothing and parcellation levels. Within our sample size, we found an inverse relationship between regional age related associations and brain age prediction accuracy as a function of smoothing and parcellation level, highlighting the importance of the parameter selection based on the goal of the study.

DATA AVAILABILITY STATEMENT

Publicly available datasets were analyzed in this study. This data can be found here: https://camcan-archive.mrc-cbu.cam.ac.uk/dataaccess/datarequest.php.

ETHICS STATEMENT

This study is conducted in compliance with the Helsinki Declaration, and has been approved by the local ethics committee, Cambridgeshire 2 Research Ethics Committee (now East of England-Cambridge Central). The patients/participants provided their written informed consent to participate in this study.

AUTHOR CONTRIBUTIONS

YZ contributed to the study plan, analyzed the data, and wrote the manuscript. AE contributed to the study plan and revision of the manuscript. All authors contributed to the article and approved the submitted version.

FUNDING

This work was supported by funding from the Canada First Research Excellence Fund, awarded to McGill University for the Healthy Brains, Healthy Lives (HBHL) initiative, CBRAIN for Multidisciplinary Reproducible Science Grant (CANARIE R53-031), and Canadian Institutes of Health Research Operating Grant (CIHR PJT-173236).

ACKNOWLEDGMENTS

Data collection and sharing for this project was provided by the Cambridge Centre for Ageing and Neuroscience (CamCAN). CamCAN funding was provided by the UK Biotechnology and Biological Sciences Research Council (grant number BB/H008217/1), together with support from the UK Medical Research Council and University of Cambridge, UK. Authors thank Compute Canada (https://www.computecanada.ca/home) for the usage of the computing resources in the current work. Authors thank Dr. Lindsay B Lewis for providing the surface parcellations registered to the MNI surface space as well as Drs. Mahsa Dadar and Filip Morys for their inputs and comments regarding data analysis and the final manuscript.

SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fdata.2021.637724/full#supplementary-material

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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