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Robust estimation of sulcal morphology

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2 Abstract

- While it is well established that cortical morphology differs in relation to a variety of
- inter-individual factors, it is often characterized using estimates of volume, thickness,
- surface area, or gyrification. Here we developed a computational approach for
- estimating sulcal width and depth that relies on cortical surface reconstructions output
- by FreeSurfer. While other approaches for estimating sulcal morphology exist, studies
- often require the use of multiple brain morphology programs that have been shown to
- 19 differ in their approaches to localize sulcal landmarks, yielding morphological
- estimates based on inconsistent boundaries. To demonstrate the approach, sulcal
- 21 morphology was estimated in three large sample of adults across the lifespan, in
- ²² relation to aging. A fourth sample is additionally used to estimate test-retest reliability
- of the approach. This toolbox is now made freely available as supplemental to this
- paper: https://cmadan.github.io/calcSulc/.
- ²⁶ **Keywords**: sulcal width; sulcal depth; age; cortical structure; atrophy; gyrification;
- 27 cerebral sulci

Robust estimation of sulcal morphology

1 Introduction

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Cortical structure differs between individuals. It is well known that cortical thickness
   generally decreases with age (Fjell et al., 2009; Hogstrom et al., 2013; Hutton et al., 2009;
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   Lemaitre et al., 2012; Madan & Kensinger, 2016, 2018; McKay et al., 2014; Salat et al.,
   2004; Sowell et al., 2003, 2007); however, a more visually prominent difference is the
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   widening of sulci, sometimes described as "sulcal prominence" (Coffey et al., 1992;
   Drayer, 1988; Jacoby et al., 1980; Laffey et al., 1984; Tomlinson et al., 1968; Yue et al.,
   1997). In the literature, this measure has been referred to using a variety of names,
   including sulcal width, span, dilation, and enlargement, as well as fold opening. With
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   respect to aging and brain morphology, sulcal width has been assessed qualitatively by
   clinicians as an index of cortical atrophy (Coffey et al., 1992; Drayer, 1988; Laffey et al.,
   1984; Pasquier et al., 1996; Scheltens et al., 1997; Tomlinson et al., 1968). An illustration
   of age-related differences in sulcal morphology is shown in Figure 1.
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         Using quantitative approaches, sulcal width has been shown to increase with age
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   (Kochunov et al., 2005, 2008; Liu et al., 2010, 2013) likely relating to subsequent
   findings of age-related decreases in cortical gyrification (Cao et al., 2017; Hogstrom et
   al., 2013; Madan & Kensinger, 2016, 2018; Madan, 2018a). Sulcal widening has also
   been shown to be associated with decreases in cognitive abilities (Liu et al., 2011) and
   physical activity (Lamont et al., 2014). With respect to clinical conditions, increased
   sulcal width has been found in dementia patients relative to healthy controls
   (Andersen et al., 2015; Hamelin et al., 2015; Huckman et al., 1975; Liu et al., 2012; Ming
   et al., 2015; Plocharski & Østergaard, 2016; Reiner et al., 2012), as well as with
   schizophrenia patients (Largen et al., 1984; Palaniyappan et al., 2015; Rieder et al., 1979)
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   and mood disorders (Elkis et al., 1995).
        One of the most common programs for conducting cortical surface analyses is
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   FreeSurfer (Fischl, 2012). Unfortunately, though FreeSurfer reconstructs cortical
   surfaces, it does not estimate sulcal width or depth, leading researchers to use
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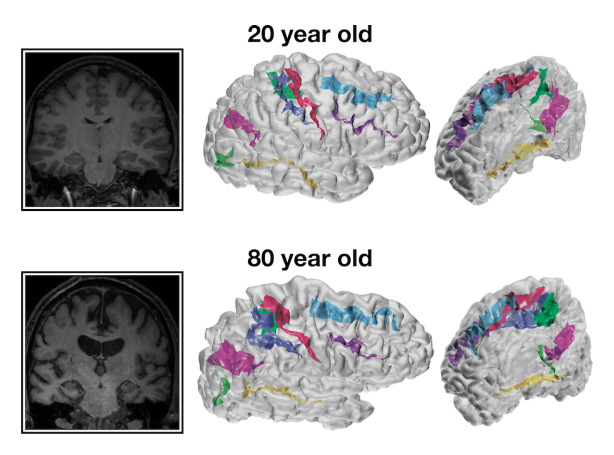


Figure 1. Representative coronal slices and cortical surfaces with sulcal identification for 20- and 80-year-old individuals.

FreeSurfer along with another surface analysis program, BrainVISA (Kochunov et al., 2012; Mangin, Rivière, et al., 2004; Mangin, Riviere, et al., 2004; Rivière et al., 2002), to characterize cortical thickness along with sulcal morphology (e.g., Cai et al., 2017; Lamont et al., 2014; Liu et al., 2011, 2013; Pizzagalli et al., 2017). While this combination allows for the estimation of sulcal morphology in addition to standard measures such 60 as cortical thickness, FreeSurfer and BrainVISA rely on different anatomical landmarks (Mikhael et al., 2018) which can yield differences in their resulting cortical surface reconstructions (Lee et al., 2006). Admittedly, determining the boundaries for an 63 individual sulcus and incorporating individual cortical variability is difficult (John et 64 al., 2006; Mikhael et al., 2018; Ono et al., 1990; Welker, 1990). While an ennumerate 65 amount of other methods have already been proposed to identify and characterize sulcal morphology (e.g., Andreasen et al., 1994; Auzias et al., 2015; Beeston & Taylor, 67 2000; Behnke et al., 2003; Eskildsen et al., 2005; Im et al., 2010; Jones et al., 2000; Le

Goualher et al., 1996, 1998; Li et al., 2008; Lohmann & von Cramon, 2000; Lohmann et al., 2008; Nowinski et al., 1996; Oguz et al., 2008; Perrot et al., 2011; Royackkers et al., 1999; Thompson et al., 1996; Vaillant & Davatzikos, 1997; Yun et al., 2013), ultimately 71 these all are again using different landmarks than FreeSurfer uses for cortical parcellations (i.e., volume, thickness, surface area, gyrification). Note that, though 73 FreeSurfer itself does compute sulcal maps, these are computed as normalized depths, not in real-world units (e.g. Kippenhan et al., 2005), furthermore, these are also independent of sulcal width information. Here we describe a procedure for estimating sulcal morphology and report 77 age-related differences in sulcal width and depth using three large samples of adults across the lifespan: two of these datasets are from Western samples, Dallas Lifespan Brain Study (DLBS) and Open Access Series of Imaging Studies (OASIS), as well as one 80 East Asian sample, Southwest University Adult Lifespan (SALD), as potential differences between populations have been relatively understudied (Leong et al., 2017; 82 Madan, 2017). To further validate the method, test-retest reliability was also assessed using a sample of young adults who were scanned ten times within the span of a 84 month (Chen et al., 2015; Madan & Kensinger, 2017b). All four of these datasets are open-access and have sufficient sample sizes to be suitable for brain morphology research (Madan, 2017). This procedure has been implemented as a MATLAB toolbox, calcSulc, that calculates sulcal morphology-both width and depth-using files generated as part of the standard FreeSurfer cortical reconstruction and parcellation pipeline. This toolbox is now made freely available as supplemental to this paper:

2 Materials and Methods

https://cmadan.github.io/calcSulc/.

3 2.1 Datasets

- OASIS. This dataset consisted of 314 healthy adults (196 females), aged 18–94, from
- 95 the Open Access Series of Imaging Studies (OASIS) cross-sectional dataset

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(http://www.oasis-brains.org) (Marcus et al., 2007). Participants were
   recruited from a database of individuals who had (a) previously participated in MRI
   studies at Washington University, (b) were part of the Washington University
   Community, or (c) were from the longitudinal pool of the Washington University
   Alzheimer Disease Research Center. Participants were screened for neurological and
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   psychiatric issues; the Mini-Mental State Examination (MMSE) and Clinical Dementia
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   Rating (CDR) were administered to participants aged 60 and older. To only include
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   healthy adults, participants with a CDR above zero were excluded; all remaining
   participants scored 25 or above on the MMSE. Multiple T1 volumes were acquired
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   using a Siemens Vision 1.5 T with a MPRAGE sequence; only the first volume was used
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   here. Scan parameters were: TR=9.7 ms; TE=4.0 ms; flip angle=10°;
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   voxel size=1.25×1×1 mm. Age-related comparisons for volumetric and fractal
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   dimensionality measures from the OASIS dataset were previously reported in Madan
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   and Kensinger (2017a), Madan and Kensinger (2018), and Madan (2018b) 1.
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            This dataset consisted of 315 healthy adults (198 females), aged 20–89, from
   wave 1 of the Dallas Lifespan Brain Study (DLBS), made available through the
   International Neuroimaging Data-sharing Initiative (INDI) (Mennes et al., 2013) and
   hosted on the Neuroimaging Informatics Tools and Resources Clearinghouse (NITRC)
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   (Kennedy et al., 2016)
   (http://fcon_1000.projects.nitrc.org/indi/retro/dlbs.html).
   Participants were screened for neurological and psychiatric issues. No participants in
   this dataset were excluded a priori. All participants scored 26 or above on the MMSE.
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   T1 volumes were acquired using a Philips Achieva 3 T with a MPRAGE sequence. Scan
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Age-related comparisons for volumetric and fractal dimensionality measures from the

parameters were: TR=8.1 ms; TE=3.7 ms; flip angle=12°; voxel size=1×1×1 mm. See

Kennedy et al. (2015) and Chan et al. (2014) for further details about the dataset.

DLBS dataset were previously reported in Madan and Kensinger (2017a), Madan and

1 Note that analyses reported in these previous papers were based on preprocessing in FreeSurfer 5.3.0, rather than FreeSurfer 6.0.

Kensinger (2018), and Madan (2018b) 1 .

- SALD. This dataset consisted of 483 healthy adults (303 females), aged 19–80, from the Southwest University Adult Lifespan Dataset (SALD) (Wei et al., 2018), also made available through INDI and hosted on NITRC (http://fcon_1000.projects.nitrc.org/indi/retro/sald.html). No participants in this dataset were excluded *a priori*. T1 volumes were acquired using a Siemens Trio 3 T with a MPRAGE sequence. Scan parameters were: TR=1.9 s; TE=2.52 ms; flip angle=9°; voxel size=1×1×1 mm.
- This dataset consisted of 30 healthy adults (15 females), aged 20–30, from the CCBD. 131 Center for Cognition and Brain Disorders (CCBD) at Hangzhou Normal University 132 (Chen et al., 2015). Each participant was scanned for 10 sessions, occurring 2-3 days 133 apart over a one-month period. No participants in this dataset were excluded a priori. 134 T1 volumes were acquired using a SCANNER with a FSPGR sequence. Scan 135 parameters were: TR=8.06 ms; TE=3.1 ms; flip angle=8°; voxel size: $1\times1\times1$ mm. This 136 dataset is included as part of the Consortium for Reliability and Reproducibility 137 (CoRR) (Zuo et al., 2014) as HNU1. Test-retest comparisons for volumetric and fractal 138 dimensionality measures from the CCBD dataset were previously reported in Madan 139 and Kensinger (2017b)¹.

41 2.2 Procedure

Data were analyzed using FreeSurfer 6.0

(https://surfer.nmr.mgh.harvard.edu) on a machine running Red Hat
Enterprise Linux (RHEL) 7.4. FreeSurfer was used to automatically volumetrically
segment and parcellate cortical and subcortical structures from the T1-weighted
images (Fischl, 2012; Fischl & Dale, 2000) FreeSurfer's standard pipeline was used (i.e.,
recon-all). No manual edits were made to the surface meshes, but surfaces were
visually inspected. Cortical thickness is calculated as the distance between the white

- matter surface (white-gray interface) and pial surface (gray-CSF interface) .
- Gyrification was also calculated using FreeSurfer, as described in Schaer et al. (2012).
- Cortical regions were parcellated based on the Destrieux et al. (2010) atlas, also part of
- the standard FreeSurfer analysis pipeline.

3 Calculation

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Here we outline a novel, simple yet robust, automated approach for estimating sulcal
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   width and depth, based on intermediate files generated as part of the standard
   FreeSurfer analysis pipeline. This procedure and functionality has been implemented
   in an accompanying MATLAB toolbox, calcSulc. The toolbox is supplemental
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   material to this paper and is made freely available:
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   https://cmadan.github.io/calcSulc/.
         For each individual sulcus (for each hemisphere and participant), the following
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   approach was used to characterize the sulcal morphology. The procedure has been
   validated and is supported for the following sulci: central, post-central, superior
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   frontal, inferior frontal, parieto-occipital, occipito-temporal, middle occipital and
   lunate, and marginal part of the cingulate (S_central, S_postcentral,
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165 S_front_sup, S_front_inf, S_parieto_occipital,

166 S_oc-temp_med&Lingual, S_oc_middle&Lunatus, S_cingul-Marginalis).

All of the sulci are labeled in Figure 2.

First the pial surface and Destrieux et al. (2010) parcellation labels were read into MATLAB by using the FreeSurfer-MATLAB toolbox provided alongside FreeSurfer (calcSulc_load), this consists of the ?h.pial (FreeSurfer cortical surface mesh) ?h.aparc.a2009s.annot (FreeSurfer parcellation annotation) files. Using this, the faces associated with the individual sulcus were isolated as a 3D mesh (calcSulc_isolate).

The width of each sulcus (calcSulc_width) was calculated by determining
which vertices lay on the boundary of the sulcus and the adjacent gyrus. An iterative
procedure was then used to determine the 'chain' of edges that would form a

contiguous edge-loop that encircle the sulcal region (calcSulc_getEdgeLoop). This provided an exhaustive list of all vertices that were mid-way between the peak of the respective adjacent gyri and depth of the sulcus itself. For each vertex in this edge-loop, 179 the nearest point in 3D space that was *not* neighbouring in the loop was determined, with the goal of finding the nearest vertex in the edge that was on the opposite side of 181 the sulcus-i.e., a line between these two vertices would 'bridge' across the sulcus. Since these nearest vertices in the edge loop are not necessarily the nearest vertex along the 183 opposite sulcus wall, an exhaustive search (walk) was performed, moving up to a 4 184 edges from the initially determined nearest vertex (configurable as 185 options.setWidthWalk). The sulcal width was then taken as the median of these 186 distances that bridged across the sulcus. 187

The depth of each sulcus (calcSulc_depth) additionally used FreeSurfer's sulcal maps (?h.sulc) to determine the relative inflections in the surface mesh, which would be in alignment with the gyral crown. The deepest points of the sulcus, i.e., the sulcal fundus, were taken as the 100 vertices within the sulcus with the lowest values in the sulcal map. For these 100 vertices, the shortest distance to the smoothed enclosing surface was calculated (generated by FreeSurfer's built-in gyrification analysis [?h.pial-outer-smoothed], Schaer et al., 2012), and the median of these was then taken as the sulcal depth.

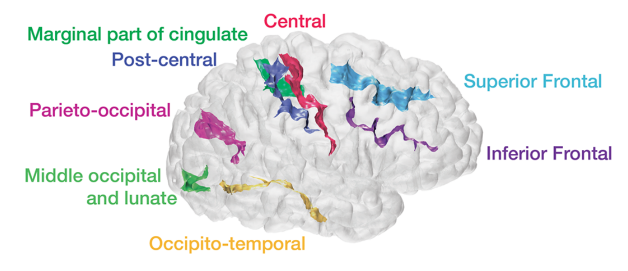


Figure 2. Example cortical surface with estimated sulci identified and labelled.

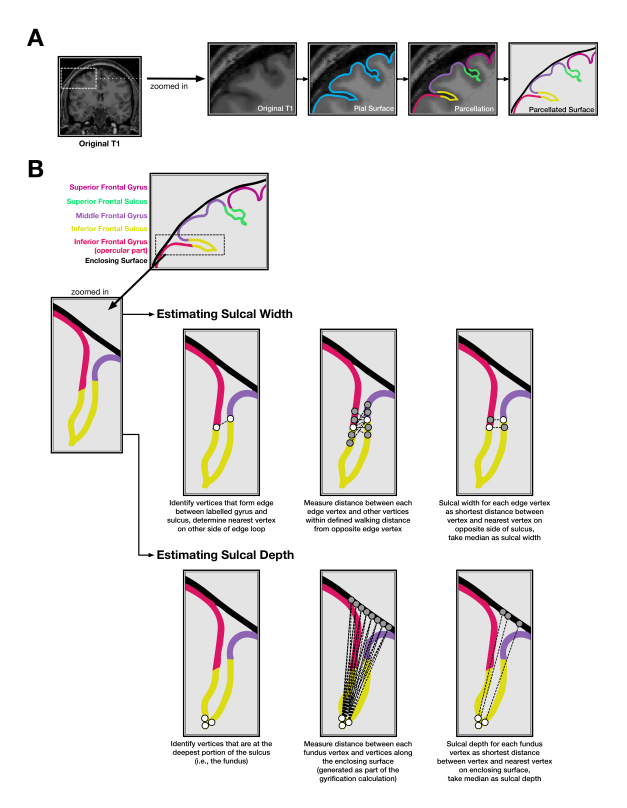


Figure 3. Illustration of the sulcal morphology method. (A) Cortical surface estimation and sulcal identification, as output from FreeSurfer. (B) Sulcal width and depth estimation procedure. Note that the surface mesh and estimation algorithm use many more vertices than shown here.

Sulcal morphology, with and depth, was estimated for eight major sulci in each 196 hemisphere: central, post-central, superior frontal, inferior frontal, parieto-occipital, 197 occipito-temporal, middle occipital and lunate, and marginal part of the cingulate 198 (S_central, S_postcentral, S_front_sup, S_front_inf, S_parieto_occipital, S_oc-temp_med&Lingual, S_oc_middle&Lunatus, 200 S_cingul-Marginalis). Preliminary analyses additionally included superior and 201 inferior temporal sulci and intraparietal sulcus but these were removed from further 202 analysis when the sulci width estimation was found to fail to determine a closed 203 boundary edge-loop at an unacceptable rate (> 10%) for at least one hemisphere. This 204 edge boundary determination failed when parcellated regions were labeled by 205 FreeSurfer to comprise at least two discontinuous regions, such that they could not be 206 identified using a single edge loop. Nonetheless, sulcal measures failed to be estimated for some participants, resulting in final samples of 310 adults from the OASIS dataset, 208 312 adults from the DLBS dataset, 481 adults from the SALD dataset, and 30 adults 209 from the CCBD dataset.

3.1 Test-retest reliability

Test-retest reliability was assessed as intraclass correlation coefficient (*ICC*), which can be used to quantify the relationship between multiple measurements (Asendorpf & Wallbott, 1979; Bartko, 1966; Chen et al., 2018; Hallgren, 2012; Koo & Li, 2016; Madan & Kensinger, 2017b; Rajaratnam, 1960; Shrout & Fleiss, 1979). McGraw and Wong (1996) provide a comprehensive review of the various *ICC* formulas and their applicability to different research questions. *ICC* was calculated as the one-way random effects model for the consistency of single measurements, i.e., *ICC*(1, 1). As a general guideline, *ICC* values between .75 and 1.00 are considered 'excellent,' .60–.74 is 'good,' .40–.59 is 'fair,' and below .40 is 'poor' (Cicchetti, 1994).

4 Results & Discussion

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4.1 Age-related differences in sulcal morphology

Scatter plots showing the relationships between each individual sulcal width and depth and age, for the OASIS dataset, are shown in Figure 4; the corresponding correlations for all datasets are shown in Tables 1 and 2. The width and depth of the central and post-central sulci appear to be particularly correlated with age, with wider and shallower sulci in older adults. Age-related differences in sulcal width and depth and generally present in other sulci as well, but are generally weaker.

Age-related relationships for each sulcus were relatively consistent between the two Western lifespan datasets (OASIS and DLBS), but age-related differences in sulcal width (but not depth) were markedly weaker in the East Asian lifespan dataset (SALD). This finding will need to be studied further, but may be related to gross differences in anatomical structure (Kochunov et al., 2003; Tang et al., 2010). Importantly, test-retest reliability, ICC(1,1), was particularly good for the sulcal depth across individual sulci.

			OASIS	DLBS	SALD	CCBD	
Sulci Name	FreeSurfer Label†	Hemi.	r(Age)	r(Age)	r(Age)	ICC(1,1)	95% CI of ICC
Central	S_central	L	.586	.486	.322	.858	[0.785, 0.918]
		R	.632	.523	.294	.842	[0.764, 0.908]
Post-central	S_postcentral	L	.413	.391	.198	.764	[0.660, 0.858]
		R	.460	.436	.213	.864	[0.794, 0.922]
Superior Frontal	S_front_sup	L	.281	.421	.055	.797	[0.703, 0.880]
		R	.205	.291	.035	.843	[0.764, 0.909]
Inferior Frontal	S_front_inf	L	.217	.323	037	.775	[0.675, 0.865]
		R	.043	.222	036	.831	[0.748, 0.901]
Parieto-occipital	S parieto occipital	L	.348	.279	.145	.616	[0.486, 0.753]
•		R	.257	.357	.213	.682	[0.561, 0.802]
Occipito-temporal	S oc-temp med&Lingual	L	.227	.270	055	.660	[0.535, 0.786]
•		R	.168	.189	.017	.692	[0.572, 0.808]
Middle occipital and lunate	S oc middle&Lunatus	L	.306	.271	.145	.605	[0.474, 0.744]
_		R	.212	.177	.023	.625	[0.496, 0.760]
Marginal part of cingulate	S cingul-Marginalis	L	.340	.275	.075	.783	[0.685, 0.871]
		R	.430	.382	.161	.757	[0.651, 0.853]
Mean			.636	.592	.227	.907	[0.856, 0.947]

Table 1 Correlations between sulcal width and age for each sulci and hemisphere, for each of the three lifespan datasets examined. Test-retest reliability, ICC(1,1), is also included from the CCBD dataset. $^{\dagger}FreeSurfer$ labels in version 6.0; labels are named slightly different in version 5.3.

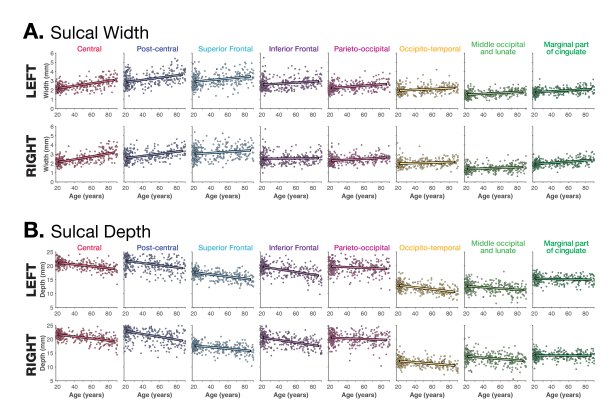


Figure 4. Relationship between (A) sulcal depth and (B) width for each of the sulci examined, based on the OASIS dataset.

			OASIS	DLBS	SALD	CCBD	
Sulci Name	FreeSurfer Label†	Hemi.	r(Age)	r(Age)	r(Age)	ICC(1,1)	95% CI of ICC
Central	S_central	L	517	205	346	.848	[0.772, 0.912]
		R	505	256	348	.860	[0.789, 0.919]
Post-central	S_postcentral	L	371	264	268	.965	[0.944, 0.981]
		R	436	246	330	.890	[0.831, 0.937]
Superior Frontal	S_front_sup	L	523	454	397	.899	[0.844, 0.943]
		R	413	465	444	.886	[0.825, 0.935]
Inferior Frontal	S front inf	L	517	490	491	.932	[0.893, 0.962]
		R	496	480	490	.915	[0.868, 0.952]
Parieto-occipital	S parieto occipital	L	145	093	241	.979	[0.966, 0.989]
•		R	124	.059	229	.970	[0.952, 0.984]
Occipito-temporal	S oc-temp med&Lingual	L	509	323	263	.953	[0.926, 0.974]
•		R	404	316	281	.913	[0.864, 0.951]
Middle occipital and lunate	S oc middle&Lunatus	L	290	167	150	.949	[0.919, 0.972]
_		R	288	120	132	.922	[0.879, 0.956]
Marginal part of cingulate	S cingul-Marginalis	L	092	035	268	.952	[0.925, 0.974]
		R	032	017	156	.918	[0.872, 0.954]
Mean			465	645	600	.972	[0.955, 0.985]

Table 2 Correlations between sulcal depth and age for each sulci and hemisphere, for each of the three lifespan datasets examined. Test-retest reliability, ICC(1,1), is also included from the CCBD dataset. $^{\dagger}FreeSurfer$ labels in version 6.0; labels are named slightly different in version 5.3.

To obtain a coarse summary measure across sulci, we averaged the sulcal width 235 across the 16 individual sulci for each individual, and with each dataset, and examined 236 the relationship between mean sulcal width with age. These correlations, shown in 237 Table 1, indicate that the mean sulcal width was generally a better indicator of 238 age-related differences in sulcal morphology than individual sulci, and had increased 239 test-retest reliability. Mean sulcal depth was similarly more sensitive to age-related differences than for an individual sulcus (e.g., it is unclear why the relationship 241 between age and width of the central sulcus differed between samples) and the magnitude of this relationship was more consistent across datasets. Reliability was 243 even higher for mean sulcal depth than mean sulcal width.

4.2 Comparison with other age-related structural differences

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Within each dataset, mean sulcal depth and width correlated with age, as shown in
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   Tables 1 and 2. Of course, other measures of brain morphology also differ with age,
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   such as mean (global) cortical thickness [OASIS: r(308) = -.793, p < .001; DLBS:
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   r(310) = -.759, p < .001; SALD: r(479) = -.642, p < .001]. Additionally, volume of the
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   third ventricle (ICV-corrected) has been previously shown to significantly related to
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   age (Madan & Kensinger, 2017a; Walhovd et al., 2011), and was found to be true in
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   each of the examined lifespan datasets here as well [OASIS: r(308) = .665, p < .001;
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   DLBS: r(310) = .677, p < .001; SALD: r(479) = .328, p < .001]. Previous studies have
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   demonstrated that both of these measures are robust estimates of age-related
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   differences in brain structure.
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         To test if these mean sulcal measures served as distinct measures of age-related
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   differences in brain morphology, beyond those provided by other measures, such as
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   mean cortical thickness and volume of the third ventricle, we conducted partial
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   correlations that controlled for these two other measures of age-related atrophy. Mean
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   sulcal width [OASIS: r_p(306) = .188, p < .001; DLBS: r_p(308) = .177, p = .002; SALD:
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   r(477) = .003, p = .96] and depth [OASIS: r_p(306) = -.443, p < .001; DLBS:
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   r_p(308) = -.397, p < .001; SALD: r_p(477) = -.534, p < .001] both explained unique
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variance in relation to age. Thus, even though more established measures of 263 age-related differences in brain morphology were replicated here, the additional sulcal 264 measures captured aspects of aging that are not accounted for by these extant 265 measures, indicating that these sulcal measures are worth pursuing further and are not redundant with other measures of brain structure. Providing additional support for 267 this, mean sulcal width and depth were only weakly related to each other [OASIS: 268 r(308) = -.192, p < .001; DLBS: r(310) = .092, p = .104; SALD: r(479) = .119, p = .009].269 As with the individual sulci measures, we did observe a difference between samples where some age-related measures were less sensitive in the East Asian lifespan 27 sample (SALD), here in the ventricle volume correlation and the unsurprisingly weaker 272 age relationship in the partial correlation using sulcal width. These sample differences 273 are puzzling, though there is a general correspondence between the two Western 274 samples. Given that much of the literature is also based on Western samples, we think 275 further research with East Asian samples, and particularly comparing samples with the same analysis pipeline, is necessary to shed further light on this initial finding. 277

5 Conclusion

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Differences in sulcal width and depth are quite visually prominent, but are not often 279 quantified when examining individual differences in cortical structure. Here we 280 examined age-related differences in both sulcal measures as a proof-of-principle to demonstrate the utility of the calcSulc toolbox that accompanies this paper and is 282 designed to closely compliments the standard FreeSurfer pipeline. This allows for the 283 additional measurement of sulcal morphology, to add to the extant measures of brain 284 morphology such as cortical thickness, area, and gyrification. Critically, this approach 285 uses the same landmarks and boundaries as in the Destrieux et al. (2010) parcellation 286 atlas, in contrast to all previous approaches to characterize sulcal features. This toolbox 287 is now made freely available as supplemental to this paper: 288

https://cmadan.github.io/calcSulc/.

Using this approach, here we demonstrate age-related differences in sulcal width

and depth, as well as high test-retest reliability. Since individual differences in sulcal morphology are sufficiently distinct from those characterized by other brain morphology measures, this approach should complement extant work of investigating factors that influence brain morphology, e.g., see Figure 3 of Madan and Kensinger (2018). Given the flexibility in the methodological approach, these measures can be readily applied to other samples after being initially processed with FreeSurfer.

Acknowledgments

MRI data used in the preparation of this article were obtained from several sources, 298 data were provided in part by: (1) the Open Access Series of Imaging Studies (OASIS) 299 (Marcus et al., 2007); (2) wave 1 of the Dallas Lifespan Brain Study (DLBS) led by Dr. 300 Denise Park and distributed through INDI (Mennes et al., 2013) and NITRC (Kennedy 301 et al., 2016); (3) the Southwest University Adult Lifespan Dataset (SALD) (Wei et al., 302 2018), also made available through INDI and hosted on NITRC; and (4) the Center for 303 Cognition and Brain Disorders (CCBD) (Chen et al., 2015) as dataset HNU1 in the 304 Consortium for Reliability and Reproducibility (CoRR) (Zuo et al., 2014). 305

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