

Locus coeruleus MRI contrast is associated with cortical thickness in older adults

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1 **Abstract**

2 There is growing evidence that neuronal integrity of the noradrenergic locus coeruleus (LC) is important
3 for later-life cognition. Less understood is how LC integrity relates to brain correlates of cognition, such
4 as brain structure. Here, we examined the relationship between cortical thickness and a measure reflecting
5 LC integrity in older (n = 229) and younger adults (n = 67). Using a magnetic resonance imaging
6 sequence which yields high signal intensity in the LC, we assessed the contrast between signal intensity
7 of the LC and that of neighboring pontine reference tissue. The Freesurfer software suite was used to
8 quantify cortical thickness. LC contrast was positively related to cortical thickness in older adults, and
9 this association was prominent in parietal, frontal, and occipital regions. Brain regions where LC contrast
10 was related to cortical thickness include portions of the frontoparietal network which have been
11 implicated in noradrenergically modulated cognitive functions. These findings provide novel evidence for
12 a link between LC structure and cortical brain structure in later adulthood.

13 *Keywords:*

14 Aging; brain structure; cortical thickness; frontoparietal network; locus coeruleus; norepinephrine

1 **1. Introduction**

2 A central goal of aging research is to identify factors which protect against age-related cognitive
3 decline. Recently, the locus coeruleus-norepinephrine (LC-NE) system has been postulated as one such
4 factor shaping later-life cognition (Mather & Harley, 2016; Robertson, 2013; Weinshenker, 2018). NE, an
5 arousal-related neurotransmitter, facilitates neural and cognitive processes involved in vigilance,
6 attention, and memory (Sara, 2009). The brain's primary source of NE is the LC, a brainstem nucleus
7 located adjacent to the lateral floor of the fourth ventricle (Sara & Bouret, 2012). The LC is densely
8 packed with neurons which release NE throughout the brain (Schwarz & Luo, 2015); through these
9 distributed projections, the LC stimulates neural responses which promote processing of relevant or
10 salient information (Berridge & Waterhouse, 2003; Mather et al., 2016). Initial evidence that LC neuronal
11 integrity relates to cognitive performance came from a longitudinal study which indicated that LC
12 neuronal density – quantified through post-mortem brain autopsies – was higher in individuals who
13 exhibited slower cognitive decline over a 6-year period before death (Wilson et al., 2013).

14 Studying the LC in vivo in humans has historically been challenging due to the small size of this
15 nucleus. In recent years, however, the human LC has been quantified using magnetic resonance imaging
16 (MRI) protocols, including turbo spin echo (TSE) and magnetization transfer sequences (Betts et al.,
17 2019b; Liu et al., 2017; Sasaki et al., 2006), in which the LC is visible as a hyperintense region which is
18 distinguishable from surrounding tissue (Keren et al., 2009; Shibata et al., 2006). Signal intensity contrast
19 between the LC and surrounding pontine tissue in these sequences – henceforth referred to as LC contrast
20 – has been validated through histological examination, with locations of high LC contrast corresponding
21 to those of cells within the LC (Cassidy et al., 2019; Keren et al., 2015).

22 Studies using MRI to assess LC contrast have pointed to the importance of the LC in cognitive
23 aging. In one study, LC contrast was positively related to cognitive reserve (Clewett et al., 2016), one
24 indicator of the brain's resilience against age-related pathology (Stern, 2009). LC contrast has also been
25 linked to memory ability, with older adults with higher LC contrast having better episodic memory
26 performance than those with lower contrast (Dahl et al., 2019; Hämmerer et al., 2018; Liu et al., 2020).

1 Additionally, individuals with Alzheimer’s disease (Betts et al., 2019a) and Parkinson’s disease (Liu et
2 al., 2017) exhibited reduced LC contrast compared to healthy older adults. Together, these studies suggest
3 a potentially neuroprotective role of LC integrity, as indexed by LC contrast, in older adulthood.

4 Of relevance to later-life cognition, NE released from the LC may protect against age-related
5 neuropathology. For example, NE protects against neuroinflammation (Feinstein et al., 2016), regulates
6 microglial functions which facilitate the clearance of amyloid beta (Heneka et al., 2010), contributes to
7 increased expression of neurotrophic factors (Braun et al., 2014), reduces damage from neurotoxicity
8 (Madrigal et al., 2007), and alleviates oxidative stress (Troadec et al., 2001). In addition, NE modulates a
9 host of cognitive processes which typically decline with age, including selective attention (Mather et al.,
10 2016; Sara, 2009), working and episodic memory (Berridge & Waterhouse, 2003; O’Dell et al., 2015),
11 and cognitive flexibility (Lapiz & Morilak, 2006). Thus, for older adults, having relatively preserved
12 neuronal integrity of the LC could lead to enhanced protection against brain pathology and enhanced
13 noradrenergic modulation of cognition (Mather & Harley, 2016; Robertson, 2013).

14 If the LC has a neuroprotective role in older adulthood, LC integrity should be associated with
15 cortical integrity. Older adults with greater cortical thickness have better fluid cognitive ability compared
16 to those with lower cortical thickness (Fjell et al., 2006), and reductions in cortical thickness and volume
17 explain a large amount of the variance in age-related cognitive decline (Fjell & Walhovd, 2010). To test
18 the possibility that LC integrity is associated with brain structure, we examined the relationship between
19 LC contrast quantified from TSE MRI scans and cortical thickness in older and younger adults. We
20 predicted that LC contrast would be positively associated with cortical thickness and that this association
21 would be strongest in brain regions which are recruited by NE modulated cognitive functions, including
22 attention, cognitive flexibility, and memory (Corbetta et al., 2008; Robertson, 2014; Sara & Bouret,
23 2012).

24

1 **2. Methods**

2 *2.1. Participants*

3 Data were collected as part of the Berlin Aging Study II (BASE-II; Bertram et al., 2014;
4 <https://www.base2.mpg.de>). A MR-eligible subsample of the BASE-II cohort for whom TSE scans were
5 collected during the study's second MR measurement-timepoint was selected for this study (n = 323).
6 This subset included older and younger adults with no history of neurological disorder, psychiatric
7 disorder, or head injury. All participants were right-handed with normal or corrected-to-normal vision.
8 Data for eligible participants were acquired at the Max Planck Institute for Human Development in
9 Berlin, Germany. The Ethics Committees of the German Psychological Society and of the Max Planck
10 Institute for Human Development approved the MRI procedure. All participants provided written,
11 informed consent prior to participation.

12 Participants were excluded if they did not have complete neural (n = 19) or demographic (n = 1)
13 data. Following visual inspection of TSE scans, additional participants were excluded due to incorrect
14 scan positioning (n = 3), motion artefact (n = 2), or incorrect placement of the LC search space (n = 1; see
15 Section 2.3). One additional participant was excluded for having an inadequate cortical reconstruction.
16 The final sample, described in Table 1, consisted of 229 older adults (82 females) and 67 younger adults
17 (22 females). Older and younger adults did not differ significantly in terms of gender distribution, $\chi^2(1,$
18 $296) = 0.092, p = .762$, or mean years of education, $t(104.4) = -0.347, p = .729$. The older adult cohort
19 had a significantly higher mean body mass index than did the younger adults, $t(82.3) = 6.83, p < .001$.
20 Mini Mental State Examination (MMSE) scores for the older adult cohort ranged from 22 to 30 (mean =
21 28.6, SD = 1.30). No exclusions were performed based on MMSE scores. Excluding the two older adult
22 participants who scored less than 25 on the MMSE did not change the pattern of results. The final sample
23 exhibited a 99% overlap with the sample of BASE-II analyzed by Dahl et al. (2019).

24 *2.2. MRI Data Acquisition*

25 MRI data were acquired using a 3T Siemens Magnetom TIM Trio scanner with a 12-channel
26 head coil. Only sequences used for the present analyses are described in this section. A three-dimensional,

1 T₁-weighted magnetization prepared rapid acquisition gradient-echo (MPRAGE) MRI sequence was
2 applied in the sagittal plane (TR = 2500 ms, TE = 4.77 ms, TI = 1100 ms, scan flip angle = 7°; bandwidth
3 = 140 Hz/pixel, field of view = 256 mm, number of slices = 192, isometric voxel size = 1 mm³, echo
4 spacing = 10.9 ms; duration = 9:20 min).

5 Based on the MPRAGE sequence, a two-dimensional TSE sequence was applied by aligning the
6 field of view orthogonally with respect to the anatomic axis of each participant's brainstem (TR = 600
7 ms; TE = 11 ms; refocusing flip angle = 120°; no explicit MT saturation; bandwidth = 287 Hz/pixel; field
8 of view = 256 mm; voxel size = 0.5 × 0.5 × 2.5 mm; echo spacing = 10.9 ms; duration = 2 × 5:54 min).
9 This sequence included ten axial slices and a 20% gap between slices in the z-dimension, encompassing
10 the entire pons. For some participants, fewer slices were acquired due to specific absorption rate limits
11 being exceeded (for details, see Dahl et al., 2019; for a discussion, see Betts et al., 2019b). Four online
12 averages were performed, yielding two TSE scans per participant. TSE scans from randomly selected
13 younger and older adult participants are displayed in Figure S1.

14 *2.3. LC Signal Intensity Assessment*

15 We obtained LC contrast estimates using a semi-automated method, as described by Dahl et al.
16 (2019). This approach yielded a LC location probability map which corresponded to published LC masks
17 (Betts et al., 2017; Keren et al., 2009; Liu et al., 2019) as well as intensity estimates which corresponded
18 to those determined through manual LC delineation. In this approach, Advanced Normalization Tools
19 (Version 2.1; Avants et al., 2009) was used to generate a template brainstem volume by aligning and
20 pooling TSE scans across participants (Figure 1A-C). The template was thresholded based on the signal
21 intensity of a reference region in the neighboring dorsal pontine tegmentum (DPT), yielding a binarized
22 LC search space (Figure 1D, 1E). After verifying that this search space encompassed the LC on a group
23 level, this search space was applied as a mask on individual, template-coregistered TSE scans. The
24 intensity and location of the maximum-intensity voxel within the masked region was extracted for each
25 slice in the z (rostrocaudal, in whole-body coordinates) dimension of each masked volume. Likewise, a
26 binarized mask of the DPT reference region was applied to template-coregistered TSE scans, and the

1 intensity and location of the maximum-intensity DPT voxel was extracted for each slice. LC contrast
2 values were calculated for each slice as a ratio (Betts et al., 2019b; Sasaki et al., 2006):

$$3 \quad \frac{\textit{intensity of peak LC voxel} - \textit{intensity of peak DPT voxel}}{\textit{intensity of peak DPT voxel}}$$

4 LC contrast values for each hemisphere were assessed in each participant's two TSE scans
5 separately, and contrast values were subsequently averaged within participants. Estimates of left and right
6 LC contrast, reflecting the peak LC contrast value across all MRI dimensions in each hemisphere, were
7 extracted. A structural equation model was used to estimate an overall value of LC contrast for each
8 participant using left and right LC contrast as indicators (Dahl et al., 2019); the resulting estimates will
9 henceforth be referred to as overall LC contrast values. In addition, for analyses of contrast along the
10 LC's rostrocaudal axis, we averaged LC contrast values across hemispheres within each slice to obtain
11 slice-wise values of LC contrast for each participant.

12 *2.4. Cortical Thickness Assessment*

13 Cortical reconstruction for all T₁-weighted anatomical images was performed with the Freesurfer
14 software suite (Version 5.3.0; <https://surfer.nmr.mgh.harvard.edu>). Freesurfer's automated processing
15 pipeline has been described previously and entails motion correction, brain extraction, registration to
16 Talairach coordinates, bias field estimation and correction, and intensity normalization (Dale et al., 1999;
17 Fischl et al., 1999). These operations were applied, and pial and white matter surfaces were generated
18 based on intensity gradients across tissue classes. One researcher (SB) blinded to participants' age
19 reviewed all resulting surfaces overlaid onto their respective anatomical images to identify segmentation
20 inaccuracies which could bias subsequent thickness estimates (McCarthy et al., 2015). Only segmentation
21 errors which did not resolve within 5 slices in any direction were manually corrected. A total of 122 (95
22 older adult, 27 younger adult) reconstructions contained inaccuracies on the pial surface and required that
23 voxels be manually erased or included. The proportion of edited reconstructions did not differ by age
24 group, $\chi^2(1, 296) = 0.001, p = .974$. For group analysis, all surfaces were tessellated and registered to a
25 spherical atlas. Neuroanatomical labels corresponding to sulcal and gyral regions were automatically

1 assigned to each vertex (Desikan et al., 2006; Fischl et al., 2004). At each vertex, cortical thickness was
2 calculated as the shortest distance between the pial and white matter surfaces (Fischl & Dale, 2000).
3 Thickness maps were smoothed with a circularly symmetric Gaussian kernel of 10 mm at full width half
4 maximum, which was chosen to balance the increase in signal-to-noise afforded by smoothing with the
5 risk of false alarms and loss of spatial precision which comes along with larger smoothing kernels. To
6 probe relations between LC contrast and thickness of the entire cortical surface, a single, global thickness
7 value was computed for each participant by averaging thickness over the left and right hemispheres and
8 using the surface area of each hemisphere as a weighting factor.

9 *2.5. Statistical Analysis*

10 *2.5.1. LC contrast and global cortical thickness in the sample.*

11 Sample characteristics of younger and older adults were examined using independent samples *t*-
12 tests with equal variances not assumed, chi-square independence tests, and Pearson correlation analyses.
13 Freesurfer's group analysis stream was used to examine cortical regions in which thickness differed
14 between older and younger adults. Previous studies have reported age-related differences in LC contrast
15 along the LC's rostrocaudal extent (Betts et al., 2017; Dahl et al., 2019; Liu et al., 2019; Manaye et al.,
16 1995). To confirm that the slight sample difference in this study did not change the topographic results
17 reported in Dahl et al. (2019), we reimplemented the nonparametric, cluster-wise permutation test to
18 identify clusters of slices where LC contrast differed significantly between younger and older adults
19 (Maris & Oostenveld, 2007). This approach is described in detail in the Supplementary Methods (Section
20 1.2). To test for reliable age differences in LC contrast according to topography, we then performed a
21 mixed analysis of variance with age group (younger, older) and topography (one level per identified
22 cluster) as factors.

23 *2.5.2. Analysis of associations between LC contrast and global cortical thickness*

24 We used multiple linear regression to determine whether LC contrast was associated with global
25 cortical thickness and whether this association depended on age. Specifically, we constructed a regression
26 model with overall LC contrast and chronological age as predictors of global thickness. In addition, to

1 determine whether the effect of LC contrast on global thickness differed between older and younger
2 adults, we included the interaction effect of overall LC contrast and age group (coded as 1=older adults,
3 -1=younger adults) as a predictor. Multiple linear regression analyses were subsequently conducted to
4 examine the effects of overall LC contrast and age on global thickness in each age group separately. Each
5 predictor was standardized by mean centering and dividing by its standard deviation; thus, results of these
6 analyses are reported as standardized regression coefficients and standard error values. For analyses
7 involving both age groups, predictors were standardized based on values from the entire sample; for
8 analyses of each age group separately, predictors were standardized for each group separately. To
9 determine whether associations between LC contrast and thickness differed based on LC topography, we
10 repeated these analyses for each set of cluster-specific LC contrast values. We also performed
11 supplementary analyses for left and right LC contrast to examine potential effects of LC laterality. In
12 addition, exploratory analyses indicated significant sex differences in overall LC contrast and global
13 cortical thickness, so we performed separate regression analyses accounting for sex and its interaction
14 effects with other predictors on thickness (Supplementary Methods, Section 1.3). All descriptive statistics
15 and regression analyses were performed in R (Version 3.6.2; R Core Team, 2019).

16 *2.5.3. Vertex-wise analysis of associations between LC contrast and cortical thickness*

17 To examine locations on the cortical surface where LC contrast was associated with thickness, we
18 fit a series of vertex-wise general linear models using Freesurfer's group analysis stream. First, we tested
19 whether there were cortical regions where the association between overall LC contrast and thickness
20 differed in older and younger adults, while regressing out the effect of chronological age. Then, for each
21 age group separately, we modeled cortical thickness at each vertex as a function of overall LC contrast
22 while regressing out the effect of chronological age. To determine whether associations with thickness
23 depended on LC laterality, we performed supplementary vertex-wise analyses for left and right LC
24 contrast separately. In addition, to determine whether regionally specific associations between cortical
25 thickness and LC contrast differed based on LC topography, we performed these vertex-wise analyses
26 again using each set of cluster-specific LC contrast values. Finally, in line with analyses of global

1 thickness, we conducted supplementary vertex-wise analyses accounting for sex (Supplementary
2 Methods, Section 1.4). For all vertex-wise analyses, significance maps were thresholded at vertex-wise p
3 < 0.05 , and cluster-wise correction for multiple comparisons was performed using a Monte Carlo Null-Z
4 simulation with 10000 iterations (Hagler et al., 2006). This entailed generating a distribution of the
5 maximum cluster size under the null hypothesis, which was used to compute cluster-wise p -values. A
6 cluster-wise threshold of $p < 0.05$ was applied to include only those clusters unlikely to appear by chance.

7 **3. Results**

8 *3.1. LC contrast & global cortical thickness in the sample*

9 Overall LC contrast was comparable in younger and older adults, $t(88.6) = -1.24$, $p = .218$, and
10 was not significantly correlated with chronological age in either age group (Figure S2). Across the
11 sample, contrast was higher for the left compared to the right LC, $t(295) = 6.72$, $p < .001$ (Figure S3).
12 Consistent with the findings of Dahl et al. (2019), we detected age group differences in LC contrast along
13 the LC's rostrocaudal extent (Figure 1F). Specifically, we detected a cluster of 6 caudal slices where older
14 adults had greater LC contrast than younger adults, $p = .002$, as well as a cluster of 3 rostral slices where
15 there was a trend towards older adults having lower LC contrast than younger adults, $p = .083$. A
16 subsequent analysis of variance indicated a significant interaction effect of age group and topography
17 (rostral/caudal) on LC contrast, $F(1,294) = 25.1$, $p < .001$. As anticipated, global cortical thickness was
18 significantly lower in older adults compared to younger adults, $t(108.9) = -9.61$, $p < .001$, and was
19 negatively correlated with age in both age groups (Figure S4A). Thickness was lower in older relative to
20 younger adults in many cortical regions (Figure S4B). Exploratory analyses indicated significant sex
21 differences in both overall LC contrast and global cortical thickness (Figure S5).

22 *3.2. Analysis of the association between LC contrast and global cortical thickness*

23 Figure 2 depicts associations between LC contrast and cortical thickness in older and younger
24 adults. Multiple linear regression analysis indicated a significant interaction effect between overall LC
25 contrast and age group on global thickness, $\beta = 0.138$, $p = .009$, indicating that the association between
26 overall LC contrast and global thickness differed significantly in older and younger adults (Table 2A). In

1 older adults, higher overall LC contrast, $\beta = 0.216$, $p < .001$, and lower age, $\beta = -0.153$, $p = .018$, were
2 significantly associated with greater global thickness (Table 2B). In younger adults, neither overall LC
3 contrast nor age was significantly associated with global thickness (Table 2C). Exploratory analyses
4 indicated that in older adults, global thickness was positively related to both left and right LC contrast
5 (Supplementary Results, Section 2.3.1).

6 To examine effects of LC topography on the association between LC contrast and thickness, we
7 performed analogous regression analyses using rostral and caudal values of LC contrast, respectively, as
8 predictors instead of overall LC contrast. An analysis of the full sample indicated a trend towards an
9 interaction effect between rostral LC contrast and age group, $\beta = 0.103$, $p = .082$ (Table 3A). Although
10 this result did not indicate a significant difference in the rostral LC contrast-global thickness association
11 by age group, for consistency with analyses of overall LC contrast, we performed subsequent regression
12 analyses in each age group separately. These analyses demonstrated that the rostral LC-thickness
13 association was driven by older adults, with rostral LC contrast demonstrating a significantly positive
14 association with global thickness in older adults, $\beta = 0.214$, $p < .001$ (Table 3B), but not in younger
15 adults, $\beta = 0.002$, $p = .990$ (Table 3C).

16 A regression analysis examining the association between caudal LC contrast and global cortical
17 thickness indicated a trend towards caudal LC contrast being negatively associated with global thickness,
18 $\beta = -0.128$, $p = .071$, as well as a marginally significant interaction effect between caudal LC contrast and
19 age group on global thickness, $\beta = 0.135$, $p = .057$ (Table 4A). In older adults, caudal LC contrast was not
20 significantly associated with global cortical thickness, $\beta = 0.006$, $p = .922$ (Table 4B). However, in
21 younger adults, higher caudal LC contrast was associated with lower global cortical thickness, $\beta = -0.275$,
22 $p = .023$ (Table 4C). Results of analyses examining associations between LC contrast and global
23 thickness accounting for sex are included in the Supplementary Results (Section 2.4.1).

24 *3.3. Analysis of the association between LC contrast and regional cortical thickness*

25 Using vertex-wise analyses, we detected ten cortical clusters where the association between
26 overall LC contrast and thickness was more positive in older than younger adults; cluster details are

1 presented in Table 5A. In an analysis of older adults only, we found a significant association between
2 overall LC contrast and cortical thickness in fourteen clusters (Table 5B); in each of these clusters, overall
3 LC contrast was positively associated with thickness. As shown in Figure 3, clusters identified in both
4 analyses included regions in frontal, parietal, occipital, and temporal cortices. In an analysis of younger
5 adults only, there were no clusters surviving multiple comparison correction in which overall LC contrast
6 was associated with thickness. Uncorrected (vertex-wise $p < .05$) significance maps from analyses of
7 overall LC contrast are displayed in Figure S6. Exploratory analyses indicated more widespread
8 associations with thickness for left versus right LC contrast (Supplementary Results, Section 2.3.2).

9 Analogous vertex-wise analyses indicated that, after correction for multiple comparisons, the
10 association between rostral LC contrast differed in older and younger adults in one cluster in left superior
11 parietal cortex (Table 6A); in this cluster, thickness was more positively associated with rostral LC
12 contrast in older adults. In an analysis of older adults only, rostral LC contrast was positively associated
13 with thickness in fifteen cortical clusters (Table 6B). Clusters where rostral LC contrast was associated
14 with thickness in older adults, which are displayed in Figure 4A, included portions of parietal, frontal, and
15 occipital cortices, and many overlapped with clusters where overall LC contrast was associated with
16 thickness in older adults. In younger adults, there were no cortical clusters in which rostral LC contrast
17 was significantly associated with thickness. Next, we found that the association between caudal LC
18 contrast and thickness was more positive in older than younger adults in four cortical clusters (Table 6C).
19 Analysis of older adults separately indicated that caudal LC contrast was only associated with thickness in
20 one cluster in rostral middle frontal cortex (Table 6D), and in this cluster, caudal LC contrast was
21 negatively associated with thickness (Figure 4B). Furthermore, in younger adults, caudal LC contrast was
22 negatively associated with thickness in three clusters, which included regions in parietal and occipital
23 cortices (Table 6E). Figures S7 and S8 depict uncorrected (vertex-wise $p < .05$) significance maps from
24 analyses of rostral and caudal LC contrast, respectively, before cluster-wise correction for multiple
25 comparisons. Results of supplementary vertex-wise analyses examining associations between LC contrast
26 and thickness accounting for sex are included in the Supplementary Results (Section 2.4.2).

1 4. Discussion

2 A growing body of literature suggests that the LC plays a role in cognition in later life, with
3 recent studies linking LC MRI contrast (as an in vivo estimate of LC neuronal integrity) to cognitive
4 outcomes in older adults (Clewett et al., 2016; Dahl et al., 2019; Hämmerer et al., 2018; Liu et al., 2020).
5 In this study, we examined whether LC contrast was associated with cortical thickness, one aspect of
6 brain structure and a documented brain indicator of cognitive ability in older adulthood (Fjell et al., 2006;
7 Lee et al., 2016). We found that higher LC contrast was associated with greater average cortical thickness
8 in older adults, even after regressing out effects of age on thickness. When we examined where this
9 association was most evident in older adults, we found widespread regions in parietal, frontal, and
10 occipital cortices in which higher LC contrast was associated with greater thickness. Together, these
11 findings constitute novel evidence for a link between LC contrast and brain structure.

12 What biological processes might the relationship between LC contrast and cortical thickness
13 reflect? One possibility is that having greater MR-indexed LC integrity could lead to higher levels of NE
14 release, allowing the beneficial effects of β -adrenergic receptor activation to be realized throughout the
15 brain and in turn to the preservation of brain structure. Many of the brain regions where LC contrast was
16 associated with thickness belong to sensory, motor, association, and prefrontal cortices, all of which are
17 innervated by the LC (Bouret & Sara, 2002; Chandler et al., 2014; Hirschberg et al., 2017). Critically, a
18 number of regions where thickness was associated with LC contrast are contained within the brain's
19 frontoparietal network (FPN), which maintains flexible representations of environmental priority and
20 internally- and externally-guided goals (Ptak, 2012; Zanto & Gazzaley, 2013). Frontoparietal regions
21 show strong resting-state functional connectivity with the LC (Jacobs et al., 2018), and during arousal and
22 attention, phasic LC activity can promote neural gain, modulating signal-to-noise ratio of neuronal
23 activity within the FPN to direct processing toward salient and/or task-relevant stimuli (Aston-Jones &
24 Cohen, 2005; Bouret & Sara, 2005; Corbetta et al., 2008; Lee et al., 2018). Thus, reduced or diminished
25 LC-NE activity later in life, resulting from reduced LC neuronal integrity, could contribute to
26 impairments in processes such as selective attention and working and episodic memory (Corbetta et al.,

1 2008). Reduced noradrenergic signaling within brain regions implicated in NE-mediated functions, such
2 as the FPN but also including occipital regions, could reduce NE's protective effects in these brain
3 regions, eventually permitting gray matter atrophy. On the other hand, preserved LC integrity could
4 maintain the NE available to act on the FPN and other brain regions and, concurrently, more of NE's
5 protective effects being realized throughout the cortex.

6 We also found that the association between LC contrast and thickness depended on LC
7 topography, as contrast of the rostral LC – but not the caudal LC – demonstrated associations with
8 thickness in older adulthood. Previous work has demonstrated that cortical and thalamic projections from
9 the LC – particularly those to frontal, sensory, and occipital cortices – tend to arise in its more rostral
10 portion (Schwarz & Luo, 2015; Waterhouse et al., 1983). Likewise, projections from the LC to the basal
11 forebrain tend to originate more from rostral than from caudal LC (España & Berridge, 2006). Supporting
12 this idea, Dahl et al., (2019) found that rostral, but not caudal, LC contrast was associated with memory
13 performance in older adults in a largely overlapping sample from BASE-II. Furthermore, consistent with
14 findings of Liu et al. (2019), we found that rostral but not caudal LC exhibited age-related decline in
15 contrast. On the other hand, contrast of the caudal LC demonstrated markedly different associations with
16 thickness compared to contrast of the rostral LC. Whereas in older adults caudal LC contrast was largely
17 unrelated to thickness, it was negatively associated with thickness in younger adults. As previous studies
18 have not reported associations between cognition and caudal LC contrast, we take caution in interpreting
19 these associations. Further, the imaging method may be less reliable in the LC's caudal aspect, as this
20 aspect is more diffusely organized relative to the rostral aspect (Fernandes et al., 2012).

21 The present study has several additional limitations. For one, we cannot conclude that the patterns
22 we observed for the LC are unique compared to those for other brain regions. However, the pattern of
23 results did not change when accounting for total ventricular volume, which demonstrates non-specific
24 age-related increases, suggesting that LC associations with thickness may extend beyond those of regions
25 which change regularly with age. In addition, we have not included cognitive measures as the BASE-II
26 cognitive battery was designed with a focus on episodic memory rather than tasks associated with the

1 frontoparietal network. However, using a 99% overlapping sample from BASE-II, we recently reported
2 an association between LC contrast and initial recall on the Rey Auditory Verbal Learning Task (Dahl et
3 al., 2019), which may also depend on attention (Chun & Turke-Browne, 2007). Another limitation is that
4 due to the smaller size of the younger relative to the older cohort, results of analyses performed on the
5 entire sample may be driven by older adults. In addition, we cannot rule out the potential influence of
6 external factors such as time of day which may explain some variability in measurements of brain
7 structure (Karch et al., 2019). Finally, both the younger and older adults who participated in BASE-II
8 constituted a highly educated convenience sample (Bertram et al., 2014), meaning that results may not
9 generalize to the general population.

10 We have offered hypotheses about processes underlying a relationship between LC neuronal
11 integrity and cortical thickness, but these possibilities remain to be tested. In addition, it remains to be
12 understood precisely what is reflected by MRI-based measures of LC contrast. Measures of LC contrast
13 are interpreted as indices of LC neuronal integrity based on findings of Keren et al. (2015) demonstrating
14 that locations of high LC contrast correspond to locations of cells in the LC which contain neuromelanin,
15 a pigment byproduct of catecholamine breakdown (Fedorow et al., 2005; Mann & Yates, 1974). One
16 other possibility is that the LC contrast measure reflects other features of noradrenergic neurons (e.g.,
17 water content), besides neuromelanin (Watanabe et al., 2019a; Watanabe et al., 2019b). Furthermore,
18 there has been limited work relating MRI-based LC contrast measures to LC function. One study
19 demonstrated that LC contrast was associated with LC activity during goal-relevant processing (Clewett
20 et al., 2018). Another recent study found that functional connectivity between LC and FPN brain regions
21 during an auditory oddball task was associated with LC contrast (Mather et al., 2020). However, more
22 work is needed to understand the LC contrast-LC function relationship.

23 To conclude, we found that LC MRI contrast, particularly that of the rostral LC, was associated
24 with thickness of widespread cortical regions in older adults. These findings constitute novel evidence for
25 a link between LC contrast and brain structure in older adulthood and are consistent with previous reports
26 of LC contrast being associated with cognition in older adulthood. Furthermore, our results provide strong

1 motivation for studying the relationship between the LC and FPN in older adulthood, as these brain
2 regions are implicated in attention and memory functions that undergo regular age-related decline.

3

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25

26

1 **Disclosure Statement**

2 The authors have no conflicts of interest to disclose.

3 **Data Statement**

4 Data will be shared upon approved request to the BASE-II Steering Committee

5 (<https://www.base2.mpg.de/en/project-information/team>).

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Tables

Table 1

Sample characteristics

	Younger adults (<i>n</i> = 67; 22 females)				Older adults (<i>n</i> = 229; 82 females)			
	Mean	SD	Min	Max	Mean	SD	Min	Max
Age	32.6	3.62	25	40	72.3	4.10	63	83
BMI	23.0	3.75	11.6	31.9	26.7	3.49	19.1	37.9
Education	14.3	2.54	10	18	14.1	2.94	7	18
MMSE	-	-	-	-	28.6	1.30	22	30

Note. Age and education are expressed in years. BMI = body mass index; MMSE = Mini Mental State Examination; SD = standard deviation.

Table 2

Results of multiple linear regression analyses examining the relationship between overall locus coeruleus contrast and global cortical thickness

Predictor	β	SE	95% CI	<i>t</i>	<i>p</i>
A. Full sample, $F(3, 292)=39.38, p<.001, R^2=0.288, \text{adjusted } R^2 = 0.281$					
Overall LC contrast	0.069	0.052	-0.033, 0.172	1.33	.185
Age	-0.506	0.050	-0.604, -0.408	-10.2	<.001
Overall LC contrast * Age Group	0.138	0.052	0.035, 0.241	2.64	.009
B. Older adults, $F(2, 226)=9.14, p<.001, R^2=0.075, \text{adjusted } R^2=0.067$					
Overall LC contrast	0.216	0.064	0.090, 0.342	3.37	<.001
Age	-0.153	0.064	-0.280, -0.027	-2.39	.018
C. Younger adults, $F(2, 64)=1.72, p=.187, R^2=0.051, \text{adjusted } R^2=0.021$					
Overall LC contrast	-0.087	0.123	-0.332, 0.158	-0.71	.482
Age	-0.198	0.123	-0.443, 0.047	-1.62	.111

Note. Regression coefficients and standard errors are standardized values. Age group was coded as: older adults = 1, younger adults = -1. CI = confidence interval; LC = locus coeruleus; SE = standard error.

Table 3

Results of multiple linear regression analyses examining the relationship between rostral locus coeruleus contrast and global cortical thickness

Predictor	β	SE	95% CI	<i>t</i>	<i>p</i>
A. Full sample, $F(3, 292)=38.98, p<.001, R^2=0.286, \text{adjusted } R^2=0.279$					
Rostral LC contrast	0.088	0.060	-0.029, 0.204	1.48	.139
Age	-0.503	0.051	-0.602, -0.403	-9.93	<.001
Rostral LC contrast * Age Group	0.103	0.059	-0.013, 0.220	1.75	.082
B. Older adults, $F(2, 226)=9.08, p<.001, R^2=0.074, \text{adjusted } R^2=0.066$					
Rostral LC contrast	0.214	0.064	0.088, 0.341	3.35	<.001
Age	-0.163	0.064	-0.289, -0.036	-2.54	.012
C. Younger adults, $F(2, 64)=1.46, p=.240, R^2=0.044, \text{adjusted } R^2=0.014$					
Rostral LC contrast	0.002	0.124	-0.246, 0.250	0.01	.990
Age	-0.209	0.124	-0.457, 0.039	-1.69	.097

Note. Regression coefficients and standard errors are standardized values. Age group was coded as: older adults = 1, younger adults = -1. CI = confidence interval; LC = locus coeruleus; SE = standard error.

Table 4

Results of multiple linear regression analyses examining the relationship between caudal LC contrast and global cortical thickness

Predictor	β	SE	95% CI	<i>t</i>	<i>p</i>
A. Full sample, $F(3, 292)=35.77, p<.001, R^2=0.269, \text{adjusted } R^2=0.261$					
Caudal LC contrast	-0.128	0.071	-0.266, 0.011	-1.81	.071
Age	-0.461	0.055	-0.570, -0.352	-8.32	<.001
Caudal LC contrast * Age Group	0.135	0.071	-0.004, 0.275	1.91	.057
B. Older adults, $F(2, 226)=3.31, p=.038, R^2=0.028, \text{adjusted } R^2=0.020$					
Caudal LC contrast	0.006	0.066	-0.123, 0.136	0.10	.922
Age	-0.168	0.066	-0.298, -0.039	-2.56	.011
C. Younger adults, $F(2, 64)=4.29, p=.018, R^2=0.118, \text{adjusted } R^2=0.091$					
Caudal LC contrast	-0.275	0.118	-0.511, -0.039	-2.33	.023
Age	-0.241	0.118	-0.477, -0.005	-2.04	.046

Note. Regression coefficients and standard errors are standardized values. Age group was coded as: older adults = 1, younger adults = -1. CI = confidence interval; LC = locus coeruleus; SE = standard error.

Table 5.

Cortical clusters where the association between overall LC contrast and thickness was more positive in older than in younger adults (A) and positive in older adults (B)

Region	Hemi- sphere	Size (mm ²)	Peak vertex*			CWP**
			X	Y	Z	
A. Overall LC - thickness association more positive in OA vs. YA						
Superior parietal	Left	1377.59	-16.0	-89.4	21.9	0.0005
Rostral middle frontal	Left	1028.72	-35.5	31.4	14.8	0.0106
Paracentral	Left	953.99	-8.1	-22.4	65.1	0.0184
Middle temporal	Right	1447.35	51.1	-12.5	-20.7	0.0008
Postcentral	Right	1301.42	14.4	-33.8	64.5	0.0021
Fusiform	Right	1282.95	36.0	-43.5	-13.9	0.0023
Lateral occipital	Right	1230.74	48.2	-75.0	5.9	0.0032
Paracentral	Right	905.96	5.9	-12.1	52.7	0.0251
Banks superior temporal sulcus	Right	903.84	46.0	-42.9	13.9	0.0259
Lateral occipital	Right	811.95	23.9	-90.1	18.9	0.0475
B. Overall LC positively associated with thickness in OA						
Inferior parietal	Left	5334.79	-44.4	-70.2	9.4	0.0001
Pars triangularis	Left	2829.49	-47.9	34.8	-8.5	0.0001
Superior frontal	Left	2590.56	-8.9	2.1	67.1	0.0001
Inferior parietal	Left	1350.49	-43.3	-63.9	38.9	0.0013
Precentral	Left	1039.94	-34.7	-21.0	49.8	0.0102
Superior temporal	Left	910.53	-59.4	-12.0	-3.3	0.0244
Precentral	Right	5682.29	40.8	-10.3	58.2	0.0001
Precuneus	Right	2556.13	6.7	-57.4	32.3	0.0001
Inferior parietal	Right	2260.05	37.0	-53.9	38.1	0.0001
Lateral occipital	Right	1783.10	40.8	-85.3	-13.0	0.0001
Lingual	Right	1602.38	20.4	-74.2	-8.8	0.0002
Pars orbitalis	Right	1389.28	45.8	30.2	-12.3	0.0006
Bank superior temporal sulcus	Right	1154.07	48.8	-40.6	12.5	0.0056
Rostral middle frontal	Right	1007.60	31.0	41.6	20.2	0.0138

Note. LC = locus coeruleus; OA = older adults; YA = younger adults. There were no clusters where overall LC contrast was associated with thickness in younger adults.

*Talairach coordinates of vertex with peak CWP value.

**Cluster-wise probability (CWP) resulting from cluster-wise Monte Carlo correction for multiple comparisons, reflecting the probability of the cluster appearing by chance. Only clusters with CWP < .05 are listed.

Table 6.

Cortical clusters where the association between rostral LC contrast and thickness was more positive in older than in younger adults (A) and positive in older adults (B) and where the association between caudal LC contrast and thickness was more positive in older than in younger adults (C) and negative in older adults (D) and younger adults (E)

Region	Hemi- sphere	Association with thickness	Size (mm ²)	Peak vertex*			CWP
				X	Y	Z	
A. Rostral LC contrast - thickness association more positive in OA vs. YA							
Superior parietal	Left	OA > YA	842.14	-22.9	-74.4	29.9	0.0387
B. Rostral LC contrast associated with thickness in OA							
Pars triangularis	Left	Positive	2633.77	-48.1	35.4	-6.3	0.0001
Caudal middle frontal	Left	Positive	2288.88	-39.6	11.4	51.4	0.0001
Superior frontal	Left	Positive	1958.83	-10.1	39.7	47.9	0.0001
Superior parietal	Left	Positive	1811.48	-10.2	-87.1	29.1	0.0001
Postcentral	Left	Positive	1472.10	-47.0	-25.4	33.9	0.0002
Precuneus	Left	Positive	1391.17	-8.2	-45.3	44.3	0.0005
Inferior parietal	Left	Positive	1091.19	-43.3	-64.6	40.3	0.0070
Precentral	Left	Positive	974.53	-45.2	-10.9	39.7	0.0166
Precentral	Right	Positive	4717.72	41.0	-9.8	58.5	0.0001
Supramarginal	Right	Positive	3809.56	57.9	-39.9	35.6	0.0001
Lateral occipital	Right	Positive	2799.21	47.8	-77.2	5.1	0.0001
Isthmus cingulate	Right	Positive	2070.17	8.9	-53.3	9.6	0.0001
Rostral middle frontal	Right	Positive	1424.36	42.7	33.6	28.2	0.0002
Superior frontal	Right	Positive	1338.58	6.8	40.8	46.9	0.0008
Pars triangularis	Right	Positive	1096.91	54.2	26.6	8.1	0.0047
C. Caudal LC contrast - thickness association more positive in OA vs. YA							
Superior parietal	Left	OA > YA	3135.14	-19.1	-86.0	18.8	0.0001
Supramarginal	Right	OA > YA	1278.17	45.1	-28.6	37.7	0.0014
Superior parietal	Right	OA > YA	928.04	25.8	-79.4	21.4	0.0227
Paracentral	Right	OA > YA	841.08	15.6	-38.4	53.3	0.0409
D. Caudal LC contrast associated with thickness in OA							
Rostral middle frontal	Left	Negative	857.23	-19.1	57.7	-11.9	0.0351
E. Caudal LC contrast associated with thickness in YA							
Lateral occipital	Left	Negative	2959.26	-34.0	-87.6	9.6	0.0001
Paracentral	Left	Negative	1075.54	-5.4	-18.8	52.4	0.0065
Superior parietal	Right	Negative	2419.58	20.6	-41.3	58.2	0.0001

Note. LC = locus coeruleus; OA = older adults; YA = younger adults. There were no clusters in younger adults where rostral LC contrast was associated with thickness.

*Talairach coordinates of vertex with peak CWP value.

**Cluster-wise probability (CWP) resulting from cluster-wise Monte Carlo correction for multiple comparisons, reflecting the probability of the cluster appearing by chance. Only clusters with CWP < .05 are listed.

Figures

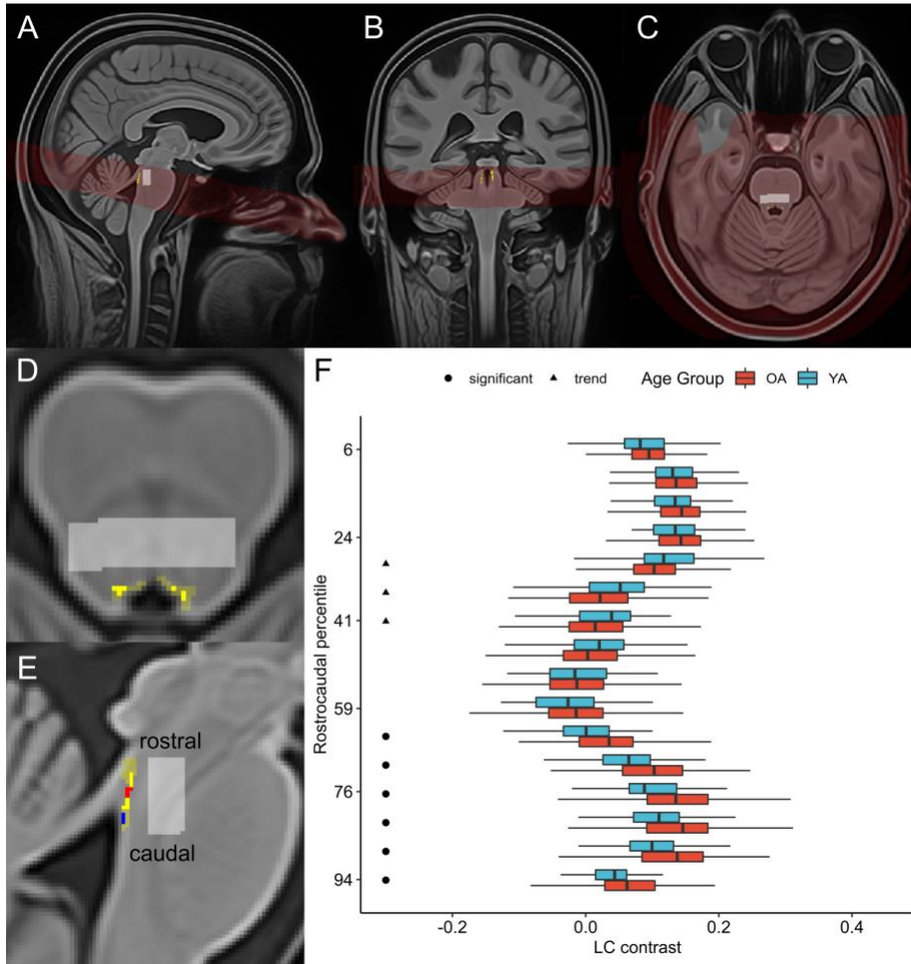


Figure 1. (A) Sagittal, (B) coronal, and (C) axial views of brainstem template volume (in red; generated from all participants' turbo spin echo scans) overlaid onto whole-brain template (generated from all participants' MPRAGE scans), with locus coeruleus (LC) probability map in common template space (Dahl et al., 2019) displayed in yellow. Pontine reference region in common template space is displayed in light gray. (D) Axial and (E) sagittal views of template brainstem volume. LC probability map (opaque yellow) is overlaid onto LC search space (translucent yellow), both of which are shown in common template space. Pontine reference region in common template space is displayed in light gray. In (E), rostral and caudal clusters within LC probability map are depicted in red and blue, respectively. (F) Boxplots depicting LC contrast values along the LC's rostrocaudal extent are displayed for older adults (OA; red) and younger adults (YA; blue). The y-axis is arranged from most caudal (100) to most rostral (0) LC segment. For visual clarity, outliers – defined as data points falling outside the interval defined by 1.5 times the interquartile range (i.e. below the 25% quantile or above the 75% quantile) – are omitted. A cluster-wise permutation test was performed to identify clusters of slices where contrast differed significantly by age group. Circles indicate a caudal cluster of slices where older adults had significantly higher contrast than younger adults, and triangles indicate a rostral cluster of slices where there was a trend towards older adults having lower contrast than younger adults.

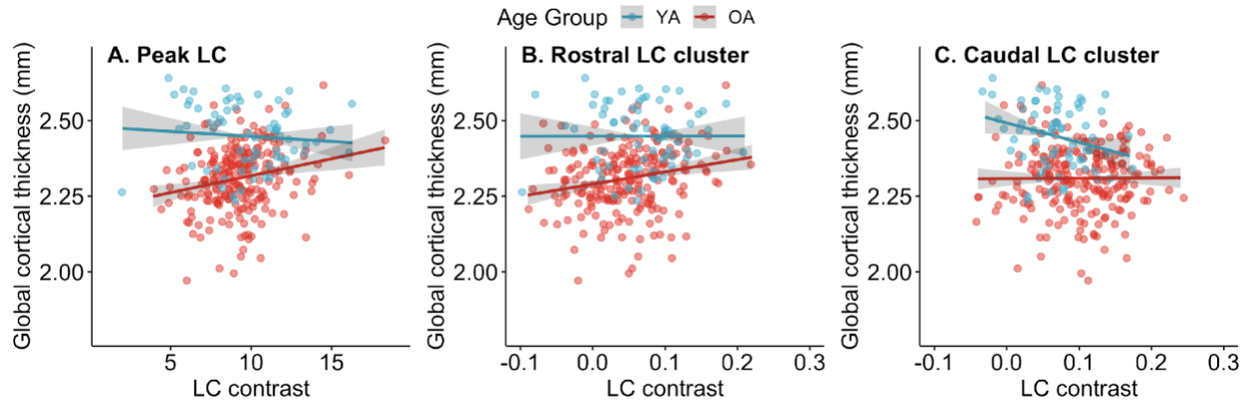


Figure 2. Associations between locus coeruleus (LC) contrast and global cortical thickness. Associations with global cortical thickness are displayed separately for older adults (OA) and younger adults (YA) for (A) overall LC contrast (reflecting contrast of peak LC voxels across x-, y-, and z-dimensions), (B) contrast of the rostral LC, and (C) contrast of the caudal LC. Points indicate raw data. Model fit lines reflect the association between LC contrast and global cortical thickness after regressing out the effect of chronological age on thickness. 95% confidence intervals are shown in gray. Fit lines and confidence intervals were calculated using the mean value of chronological age within each age group. Differences in scales on the x-axes are due to overall LC being calculated as a latent variable in a structural equation modeling framework, whereas rostral and caudal LC contrast were calculated by averaging signal intensity contrast ratios.

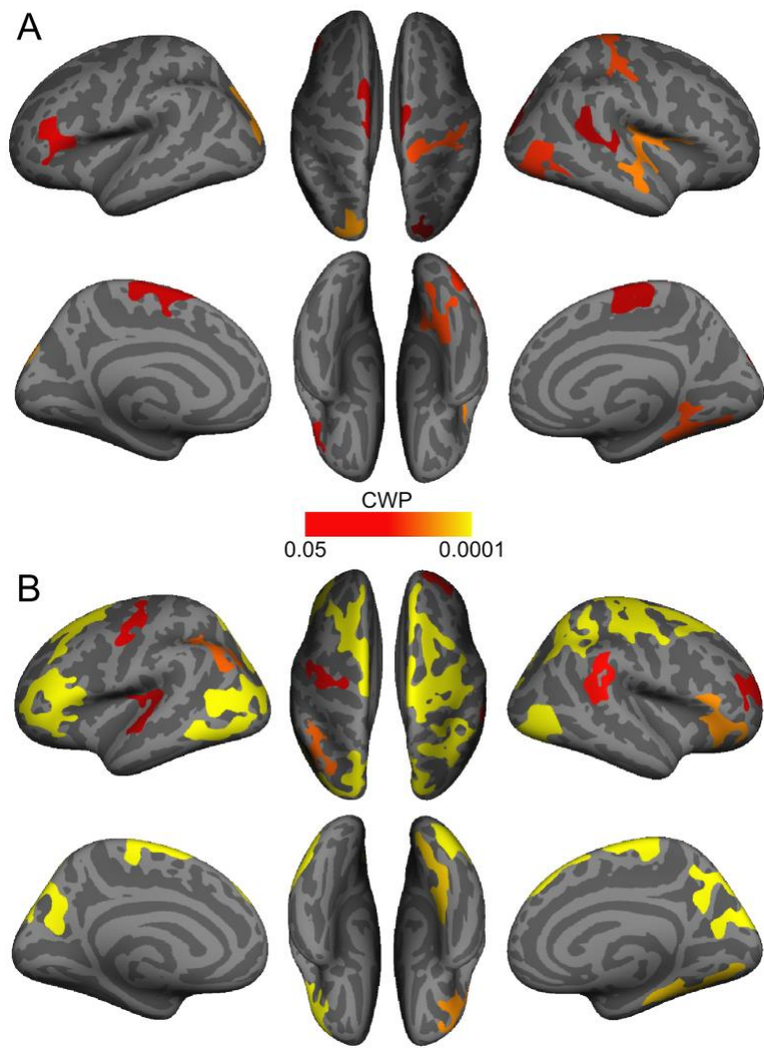


Figure 3. Cortical regions where the association between overall locus coeruleus contrast and thickness was more positive in older than in younger adults (A) and positive in older adults (B). Only clusters that survived cluster-wise correction for multiple comparisons (cluster-wise probability (CWP) < .05) are shown.

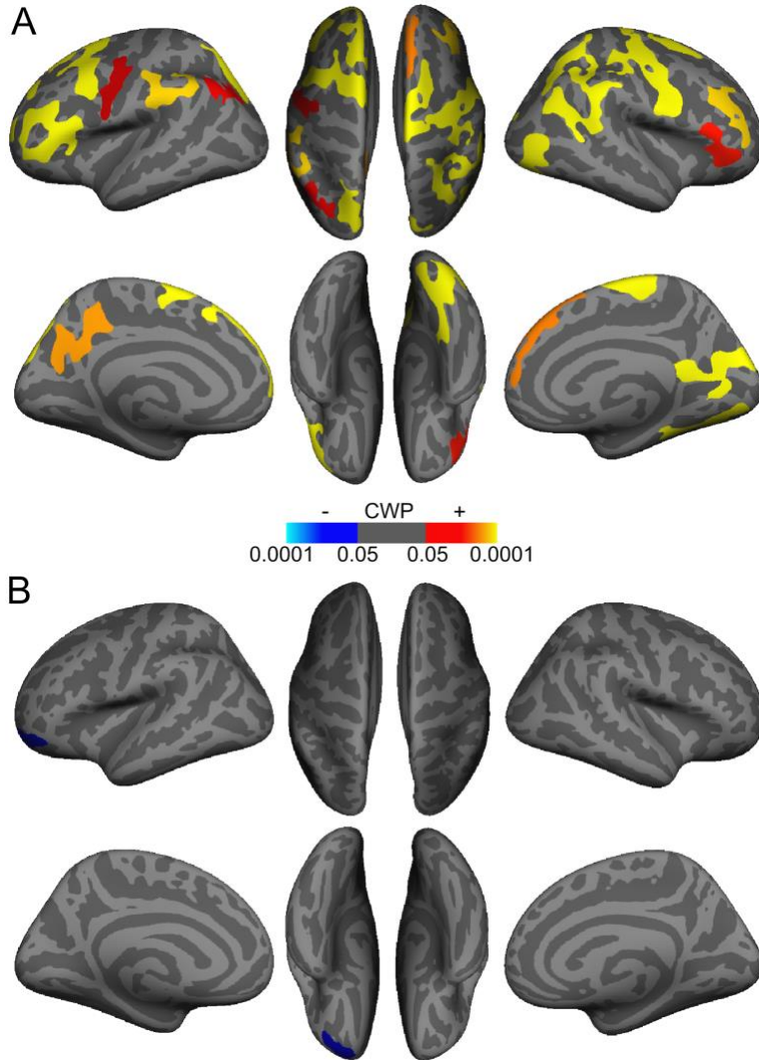


Figure 4. Cortical regions demonstrating an association between rostral (A) and caudal (B) locus coeruleus (LC) contrast and thickness in older adults. Warmer colors indicate clusters with a positive association between LC contrast and cortical thickness, and cooler colors indicate clusters with a negative association. Only clusters that survived cluster-wise correction for multiple comparisons (cluster-wise probability (CWP) < .05) are shown.

SUPPLEMENTARY MATERIAL

Locus coeruleus MRI contrast is associated with cortical thickness in older adults

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Table S5. Results of regression analyses examining the association between overall LC contrast and global cortical thickness, after controlling for age and sex

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2.4.2. Vertex-wise analysis of LC contrast and regional cortical thickness

Table S8. Cortical clusters where the association between overall LC contrast and thickness was greater in older than younger adults (A) and positive in older adults (B)

Table S9. Cortical clusters where the association between rostral LC contrast and thickness was greater in older than younger adults (A) and positive in older adults (B) and where caudal LC contrast was associated with thickness in older adults (C).

3. Supplementary References

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1. Supplementary Methods

1.1. Visualization of turbo spin echo scans

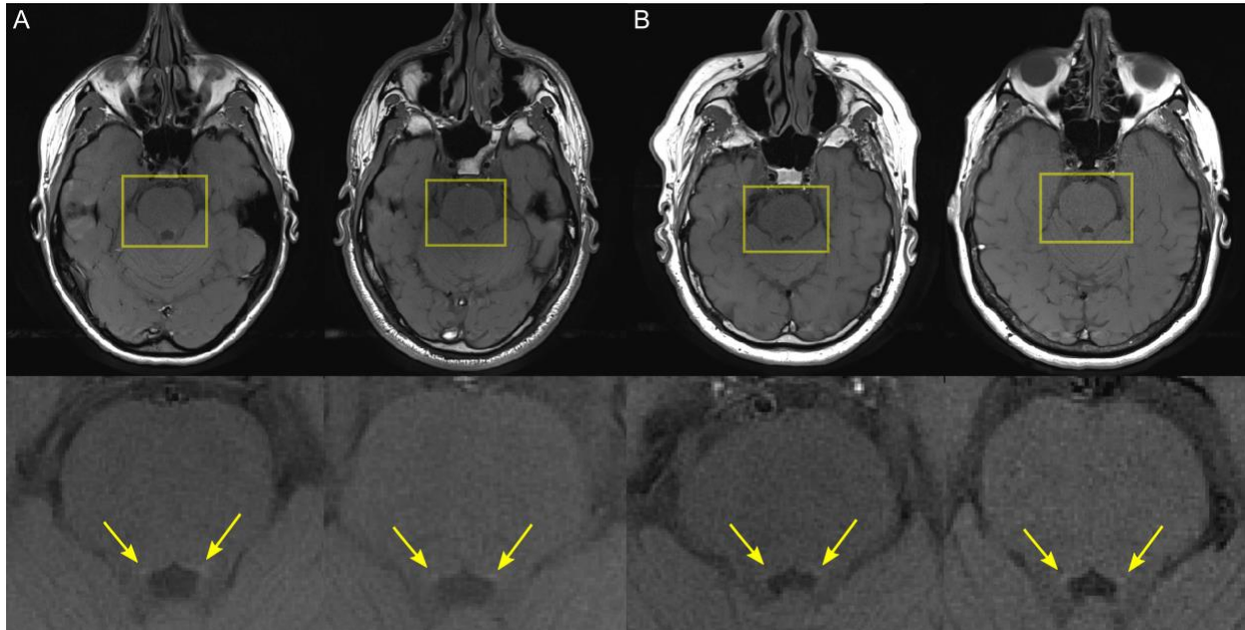


Figure S1. Turbo spin echo scans from two randomly selected younger (A) and two randomly selected older (B) adult participants. Axial view of scans is shown in upper panel. Closer view of pontine region in the same scans, outlined in yellow, is shown in lower panel. In lower panel, yellow arrows indicate hyperintensities bordering the fourth ventricle.

1.2. Cluster-wise permutation test of age differences in contrast according to LC topography

To identify clusters of slices where LC contrast differed significantly between younger and older adults, we reimplemented the nonparametric, cluster-wise permutation test also performed in Dahl et al. (2019), using the FieldTrip toolbox (Oostenveld et al., 2011; <http://www.fieldtriptoolbox.org>). This entailed first performing an independent samples t -test comparing LC contrast of older and younger adults for each slice where all participants exhibited reliable LC signal intensity ($n=16$ slices). Next, clusters were formed by grouping adjacent slices where $p < 0.05$, and a cluster mass statistic for each cluster was calculated by summing the t -statistic values of all slices comprising each cluster. A Monte Carlo method was used to generate a permutation distribution of mass statistics for the summed t -statistics in each cluster under the assumption of no effect of age group on LC contrast; specifically, in each of 100,000 permutations, age group labels were shuffled, slice-wise independent samples t -tests were performed, and

cluster mass statistic(s) were computed. To implement a two-tailed test, we considered a cluster significant, if its cluster mass statistic fell in the lowest or highest 2.5% of the respective permutation distribution. For each resulting cluster(s) where LC contrast differed significantly between older and younger adults, we averaged LC contrast values over the slices comprising each cluster to obtain cluster-specific LC contrast estimates for each participant.

1.3. Analysis of associations between LC contrast and global cortical thickness, controlling for sex

Exploratory analyses of our data revealed significant sex differences in both LC contrast (Figure S3A) and global cortical thickness (Figure S3B), with females demonstrating significantly greater values of both measures than males. Consequently, in an attempt to test whether the LC contrast-thickness associations depended on sex, we performed a series of supplemental analyses including sex as a predictor of thickness. However, we advise caution in the interpretation of these results, because LC contrast and sex were collinear in both age groups. Having correlated (e.g. non-independent) predictors in a regression analysis can lead to unreliable estimates of the associations between the individual predictors and the response variable (James et al., 2013).

Similar to the approach taken in the main text, we first performed a multiple linear regression analysis using data from all participants to examine the extent to which overall LC contrast and chronological age predicted global thickness and to what extent the overall LC contrast-thickness association differed by age group. We added sex as a predictor, as well as interaction effects between (a) sex and overall LC contrast, (b) sex and age group, and (c) a three-way interaction effect between sex, age group, and overall LC contrast. For these analyses, sex was coded as female = 1, male = -1 so that regression coefficients for overall LC contrast, age, and the overall LC contrast-by-age interaction would reflect values averaged across females and males. As with all other analyses involving age group, age group was coded as older adults = 1, younger adults = -1. We then repeated this analysis separately for rostral and caudal LC contrast values.

1.4. Analysis of associations between LC contrast and global cortical thickness, controlling for sex

To examine cortical regions where the association between LC contrast and thickness differed in older and younger adults after accounting for effects of chronological age and sex on thickness, we performed a vertex-wise analysis using Freesurfer's group analysis stream. For this analysis, four groups were defined (older adult females, older adult males, younger adult females, and younger adult males), and the LC contrast slope of the former two groups was contrasted with that of the latter two groups, controlling for effects of chronological age. Consistent with analyses of global thickness, we then examined cortical regions where LC contrast was associated with thickness after accounting for age and sex in each group separately. The procedure was similar to that used in main analyses, but in addition to age being included as a covariate, sex was included as a fixed factor. Cluster-wise correction for multiple comparisons was performed in the same method as described in the main text. These analyses were repeated separately to identify cortical regions where thickness was associated with each rostral and caudal LC contrast.

2. Supplementary Results

2.1. LC contrast and cortical thickness in the sample

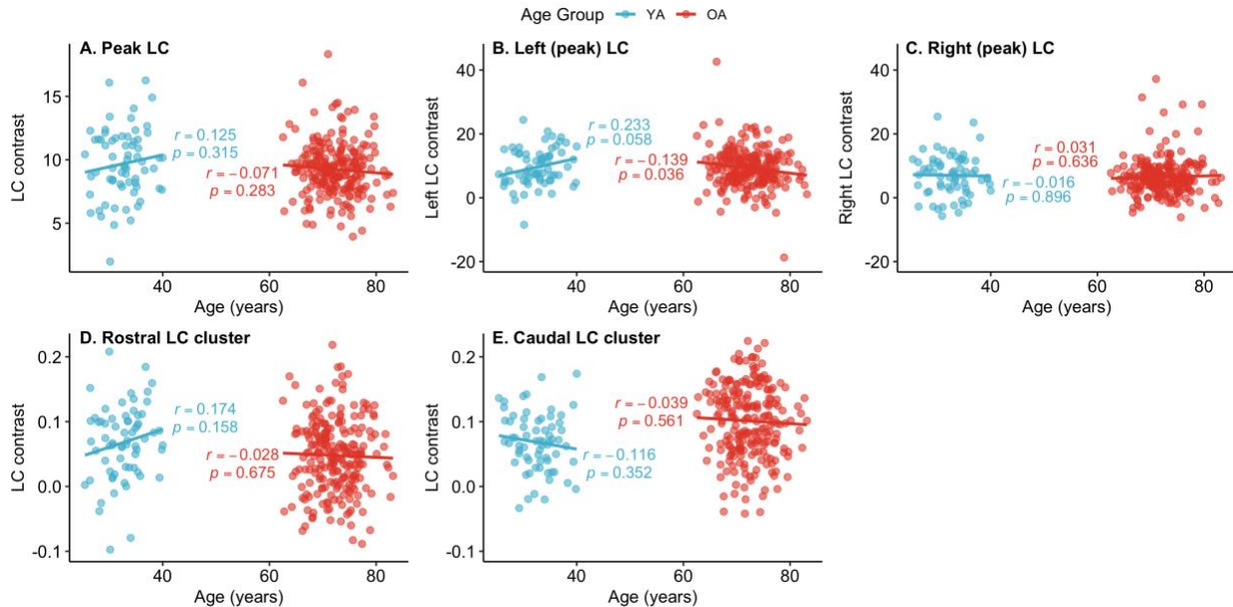


Figure S2. Locus coeruleus (LC) contrast versus chronological age in younger adults (YA) and older adults (OA). Pearson's correlation analyses indicated that chronological age was not significantly correlated with overall LC contrast in either younger adults, $r(65) = 0.125$, $p = .315$, or older adults, $r(227) = -0.071$, $p = .283$ (A). Left LC contrast was negatively correlated with age in older adults, $r(227) = -0.139$, $p = .036$, but was not significantly correlated with age in younger adults, $r(65) = 0.233$, $p = .058$ (B). Right LC contrast was not significantly correlated with age in older adults, $r(227) = 0.031$, $p = .636$, or younger adults, $r(65) = -0.016$, $p = .896$ (C). Rostral LC contrast was not significantly correlated with age in younger adults, $r(65) = 0.174$, $p = .158$, or older adults, $r(227) = -0.028$, $p = .675$ (D). Caudal LC contrast was also not significantly correlated with age in younger adults, $r(65) = -0.116$, $p = .352$, or older adults, $r(227) = -0.039$, $p = .561$ (E). Fisher's r -to- z transformations indicated that the correlation between age and overall LC contrast differed significantly for older and younger adults, $Z = 2.38$, $p = .020$, as did the correlation between rostral LC contrast and age, $Z = 2.47$, $p = .010$, and the correlation between left LC contrast and age, $Z = 4.56$, $p < .001$. The correlation between age and caudal LC contrast did not differ significantly between older and younger adults, $Z = 0.940$, $p = .350$, nor did the correlation between age and right LC contrast, $Z = 0.580$, $p = .560$. While the correlations between LC contrast and age did not reach statistical significance, the trends observed for overall and rostral LC contrast are in line with the results reported by Liu et al., 2019 (Figures 5a,b), wherein LC contrast ratios increased with age in younger adults and decreased with age in older adults.

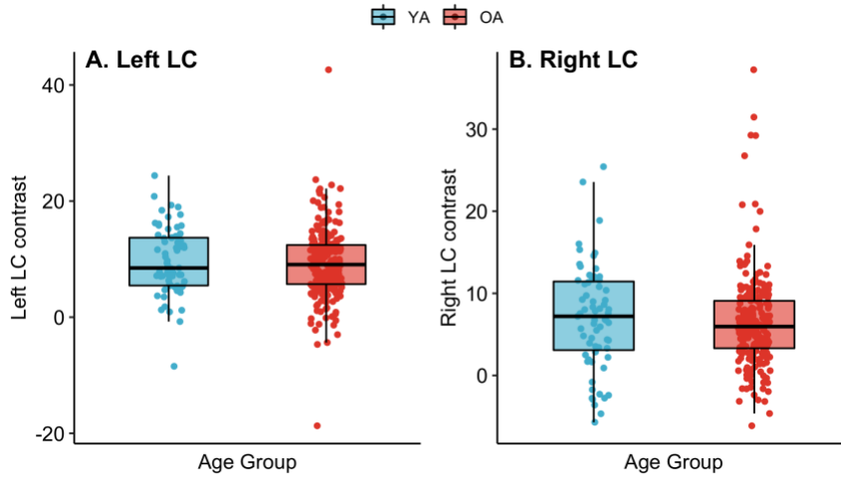


Figure S3. Left and right locus coeruleus (LC) contrast in younger adults (YA) and older adults (OA). A 2 (laterality: left, right) x 2 (age group: OA, YA) mixed-design ANOVA indicated a significant main effect of laterality on LC contrast, $F(1, 294) = 45.1, p < .001$, with left LC contrast being greater than right LC contrast. There was no significant main effect of age group on LC contrast, $F(1,294) = 0.45, p = .503$, and no significant interaction effect between laterality and age group, $F(1,294) = 0.025, p = .874$.

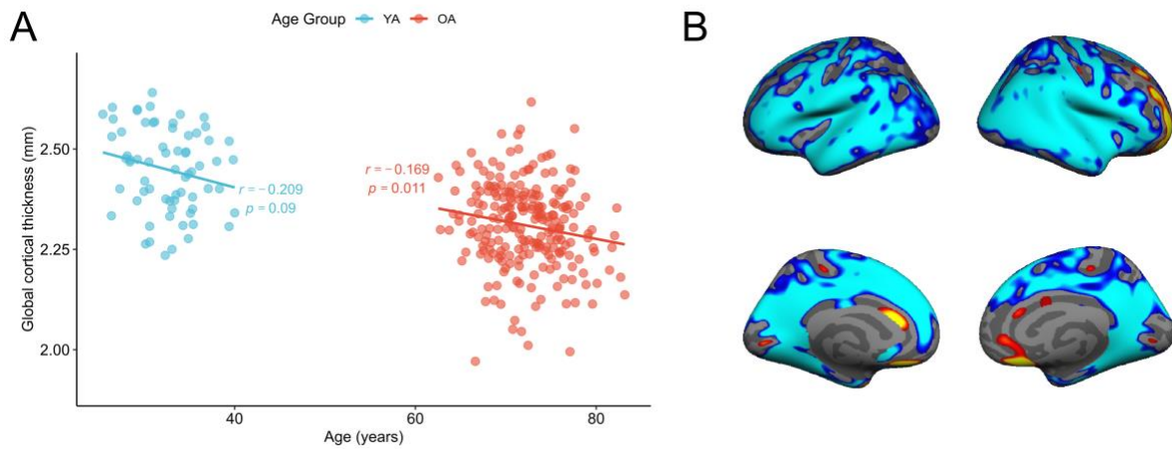


Figure S4. (A) Global cortical thickness versus chronological age in younger adults (YA) and older adults (OA). Pearson's correlation analyses demonstrated that chronological age was negatively correlated with global cortical thickness in older adults, $r(227) = -0.169$, $p = .011$. Further, we observed a trend towards a negative correlation between age and thickness in younger adults, $r(65) = -0.209$, $p = .090$. A Fisher's r -to- z transformation revealed that the correlation between age and thickness did not differ significantly between older and younger adults, $Z = 0.510$, $p = .610$. (B) Uncorrected significance maps (vertex-wise $p < .05$) depicting cortical vertices where thickness differed in older versus younger adults; cooler colors (blue-turquoise) depict vertices where thickness was lower in older than in younger adults, and warmer colors (red-yellow) indicate vertices where thickness was higher in older than in younger adults.

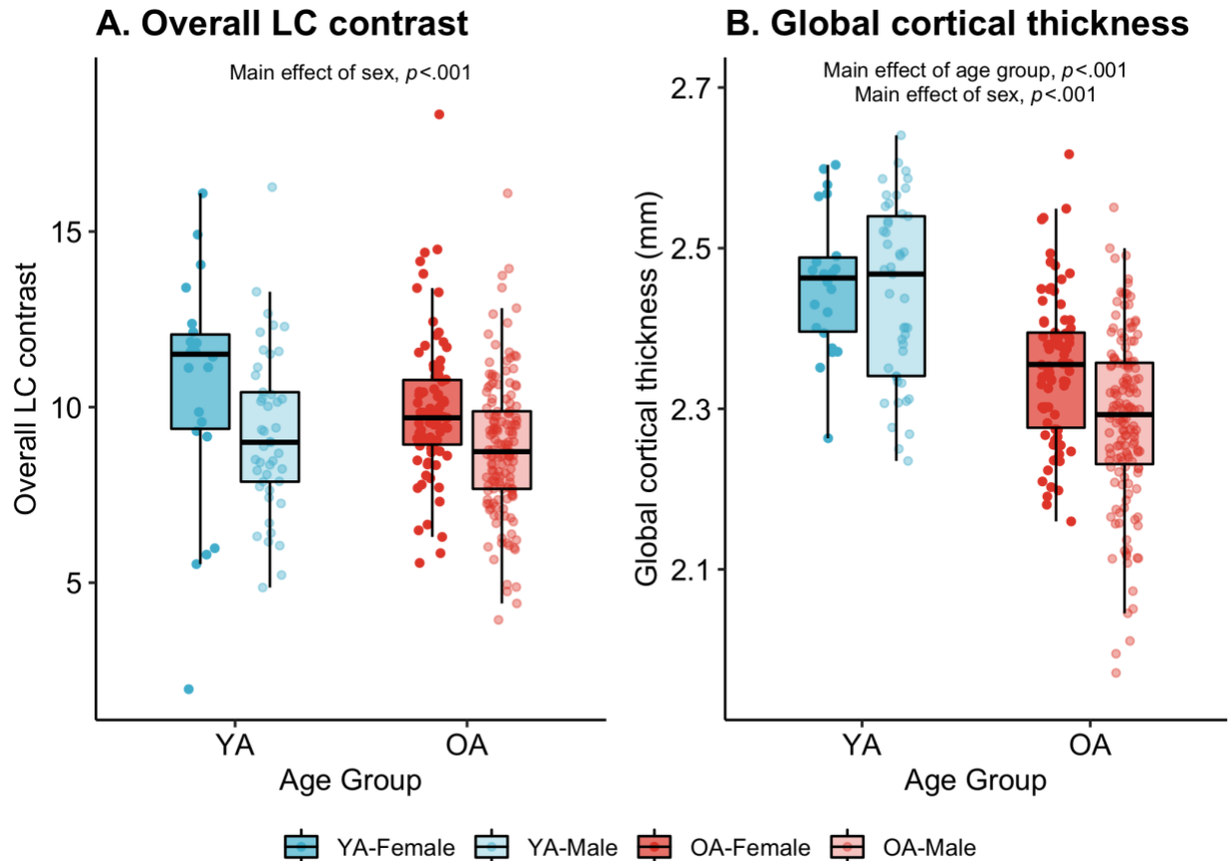


Figure S5. Overall locus coeruleus (LC) contrast and global cortical thickness by age group and sex. As an exploratory analysis of whether sex had an effect on overall LC contrast in either age group, we performed a series of 2 (age group: OA, YA) \times 2 (sex: female, male) analyses of variance (ANOVAs). (A) The first ANOVA revealed a main effect of sex on overall LC contrast, $F(1, 292) = 18.5, p < .001$, with females having greater contrast than males. This analysis indicated no significant main effect of age group, $F(1, 292) = 2.23, p = .136$, or interaction effect between age group and sex on overall LC contrast, $F(1, 292) = 0.126, p = .723$. (B) The second ANOVA indicated a main effect of sex, $F(1, 292) = 17.8, p < .001$, a main effect of age group, $F(1, 292) = 96.5, p < .001$, and a trend towards an interaction effect of age group and sex on global thickness, $F(1, 292) = 2.87, p = .092$. On boxplots, upper and lower hinges correspond to first and third quartiles, respectively, and medians are indicated with horizontal bars. OA = older adults; YA = younger adults.

2.2. Uncorrected significance maps from vertex-wise analyses of associations between LC contrast and cortical thickness

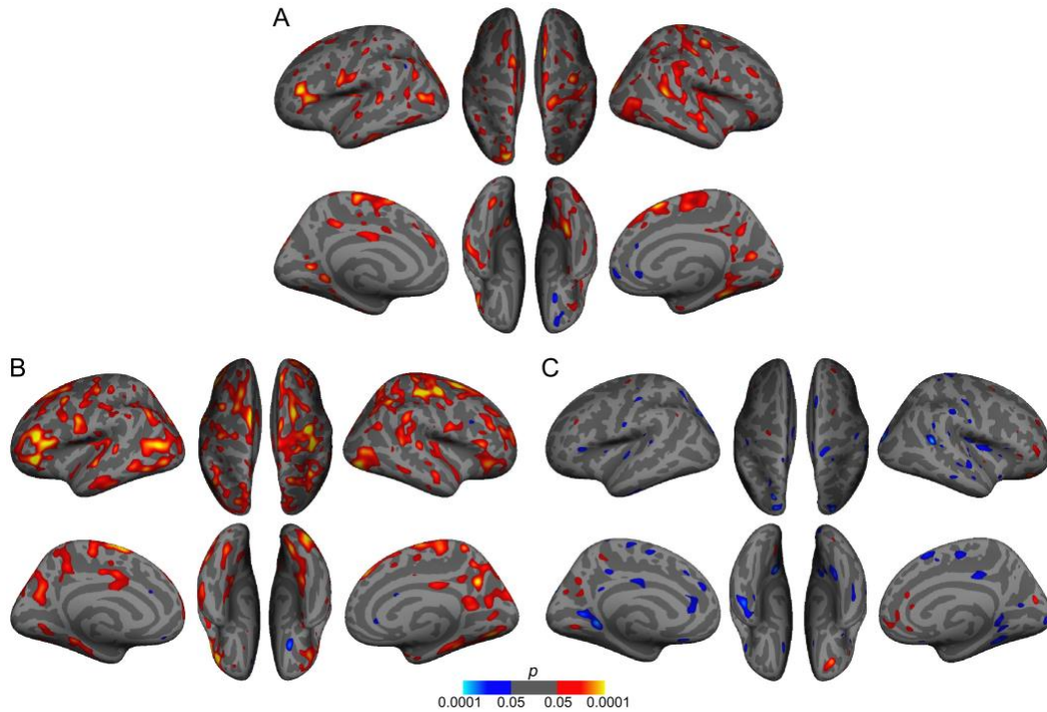


Figure S6. Uncorrected significance maps from analyses of overall locus coeruleus (LC) contrast and thickness. (A) shows vertices where the association between overall LC contrast and thickness differed significantly in older and younger adults; in (A), warmer colors (red-yellow) indicate a more positive association in older than younger adults, and cooler colors (blue-turquoise) indicate a more positive association in younger than in older adults. Vertices exhibiting a significant association between overall LC contrast and thickness in older and younger adults are shown in (B) and (C), respectively. In (B) and (C), warmer colors (red-yellow) indicate a positive association between overall LC contrast and cortical thickness, and cooler colors (blue-turquoise) indicate a negative association between overall LC contrast and cortical thickness. All maps were thresholded at vertex-wise $p < .05$.

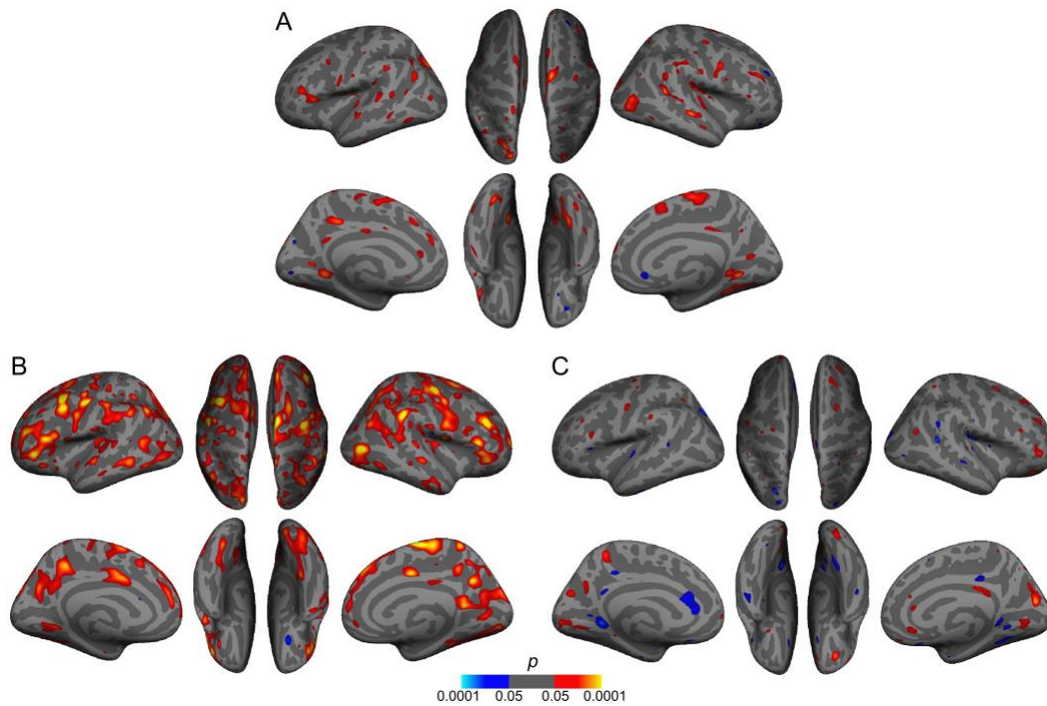


Figure S7. Uncorrected significance maps from analyses of rostral locus coeruleus (LC) contrast and thickness. (A) shows vertices where the association between rostral LC contrast and thickness differed significantly in older and younger adults; in (A), warmer colors (red-yellow) indicate a more positive association in older than younger adults, and cooler colors (blue-turquoise) indicate a more positive association in younger than in older adults. Vertices exhibiting a significant association between rostral LC contrast and thickness in older and younger adults are shown in (B) and (C), respectively. In (B) and (C), warmer colors (red-yellow) indicate a positive association between rostral LC contrast and cortical thickness, and cooler colors (blue-turquoise) indicate a negative association between rostral LC contrast and cortical thickness. All maps were thresholded at vertex-wise $p < .05$.

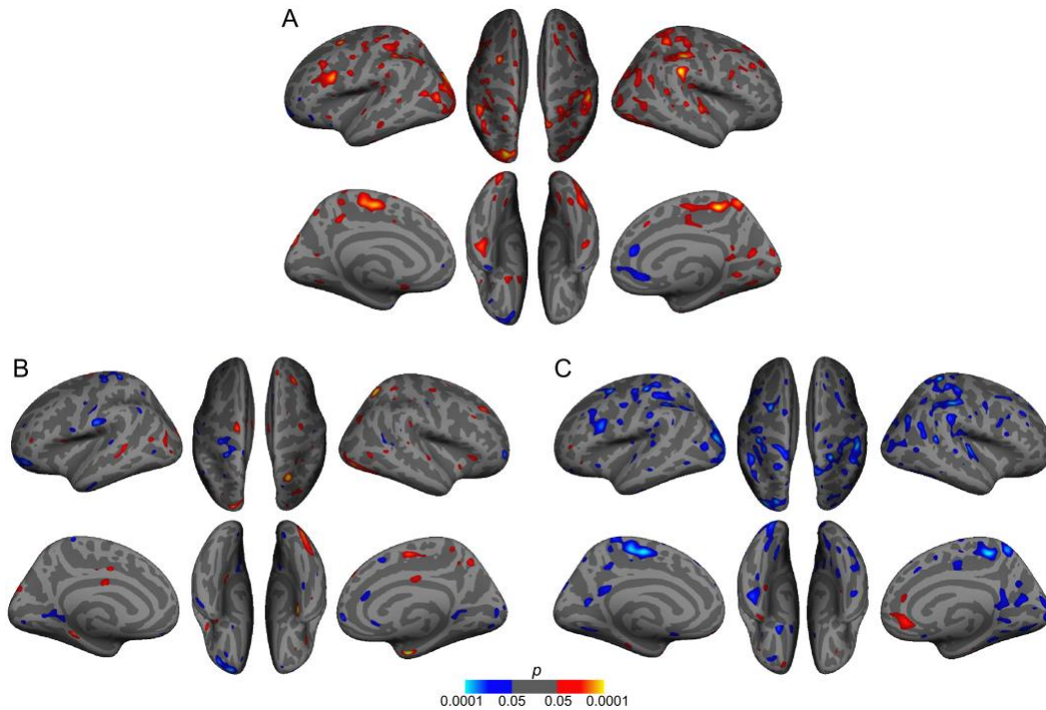


Figure S8. Uncorrected significance maps from analyses of caudal locus coeruleus (LC) contrast and thickness. (A) shows vertices where the association between caudal LC contrast and thickness differed significantly in older and younger adults; in (A), warmer colors (red-yellow) indicate a more positive association in older than younger adults, and cooler colors (blue-turquoise) indicate a more positive association in younger than in older adults. Vertices exhibiting a significant association between caudal LC contrast and thickness in older and younger adults are shown in (B) and (C), respectively. In (B) and (C), warmer colors (red-yellow) indicate a positive association between caudal LC contrast and cortical thickness, and cooler colors (blue-turquoise) indicate a negative association between caudal LC contrast and cortical thickness. All maps were thresholded at vertex-wise $p < .05$.

2.3. Analysis of associations between left and right LC contrast and thickness

2.3.1. Analysis of left and right LC contrast and global cortical thickness

A series of exploratory multiple regression analyses were performed to examine how each left and right LC contrast were associated with global cortical thickness, after accounting for age. Results of analyses of left and right LC contrast are presented in Tables S1 and S2, respectively. We found a significant interaction effect between left LC contrast and age group on global thickness, indicating the left LC-thickness association differed in older and younger adults (Table S1A). Subsequent analysis in each age group separately indicated that this interaction was driven by a significantly positive association between left LC and global thickness in older adults (Table S1B) but not in younger adults (Table S1C).

An analogous set of tests did not indicate a significant interaction between right LC contrast and age group (Table S2A), although in older adults separately, right LC contrast had a significantly positive association with global thickness (Table S2B). There was no significant association between right LC contrast and global thickness in younger adults (Table S2C). These results indicate that both left and right LC contrast were associated with global cortical thickness in older adults, but a greater positive LC contrast-thickness association in older versus younger adults was only observed for left LC.

Table S1

Results of multiple linear regression analyses examining the association between left LC contrast and global cortical thickness

Predictor	β	SE	95% CI	t	p
A. Full sample, $F(3, 292) = 38.98, p < .001, R = 0.286, \text{adjusted } R^2 = 0.279$					
Left LC contrast	0.065	0.060	-0.053, 0.183	1.09	0.278
Age	-0.503	0.050	-0.600, -0.406	-10.2	<.001
Left LC contrast * Age group	0.120	0.060	0.002, 0.238	2.00	0.046
B. Older adults, $F(2, 226) = 8.84, p < .001, R^2 = 0.073, \text{adjusted } R^2 = 0.064$					
Left LC contrast	0.212	0.065	0.085, 0.340	3.28	0.001
Age	-0.139	0.065	-0.267, -0.012	-2.15	0.033
C. Younger adults, $F(2, 64) = 1.53, p = .225, R^2 = 0.046, \text{adjusted } R^2 = 0.016$					
Left LC contrast	-0.045	0.126	-0.296, 0.206	-0.36	0.723
Age	-0.199	0.126	-0.449, 0.052	-1.58	0.119

Note. Regression coefficients and standard errors are standardized values. Age group was coded as: older adults = 1, younger adults = -1. SE = standard error.

Table S2

Results of multiple linear regression analyses examining the association between left LC contrast and global cortical thickness

Predictor	β	SE	95% CI	t	p
A. Full sample, $F(3, 292) = 36.04, p < .001, R^2 = 0.270, \text{adjusted } R^2 = 0.263$					
Right LC contrast	0.020	0.056	-0.091, 0.131	0.35	0.724
Age	-0.510	0.050	-0.609, -0.412	-10.2	<.001
Right LC contrast * Age group	0.098	0.057	-0.013, 0.209	1.73	0.084
B. Older adults, $F(2, 226) = 5.42, p = .005, R^2 = 0.046, \text{adjusted } R^2 = 0.037$					
Right LC contrast	0.132	0.065	0.004, 0.260	2.03	0.044
Age	-0.173	0.065	-0.301, -0.045	-2.66	0.008
C. Younger adults, $F(2, 64) = 1.81, p = .173, R^2 = 0.053, \text{adjusted } R^2 = 0.024$					
Right LC contrast	-0.099	0.122	-0.342, 0.144	-0.81	0.420
Age	-0.211	0.122	-0.454, 0.032	-1.73	0.088

Note. Regression coefficients and standard errors are standardized values. Age group was coded as: older adults = 1, younger adults = -1. SE = standard error.

2.3.2. Vertex-wise analysis of left and right LC contrast and regional cortical thickness

Vertex-wise analyses were performed to determine cortical regions where each left and right LC contrast were associated with thickness, after accounting for the effect of chronological age. Cortical clusters where left and right LC contrast were associated with thickness after cluster-wise multiple comparison correction are presented in Tables S3 and S4, respectively. We identified two clusters where the association between left LC contrast and thickness differed in older versus younger adults (Table S3A); in each of these clusters, the association was more positive in older than younger adults. Further, we identified eleven clusters where left LC contrast was positively associated with thickness in older adults (Table S3B). These clusters corresponded to many of the clusters where overall and rostral LC contrast were associated with thickness in older adults (main text, Tables 5 and 6).

For right LC contrast, the association with thickness differed in older and younger adults in two cortical clusters (Table S4A); in this cluster, the association with right LC contrast was more positive in older versus younger adults. In older adults, right LC contrast was positively associated with thickness in one cortical cluster (Table S4B). There were no clusters where left or right LC contrast was associated

with thickness in younger adults. Together, these results indicate that in older adults, left LC was associated with thickness in a more widespread set of cortical regions than was right LC contrast.

Table S3

Cortical clusters where the association between left LC contrast and thickness was greater in older than in younger adults (A) and positive in older adults (B)

Region	Hemi-sphere	Size (mm ²)	Peak vertex*			CWP**
			X	Y	Z	
A. Left LC contrast-thickness association greater in OA vs. YA						
Pars triangularis	Left	900.35	-48.7	26.7	5.4	0.0230
Post central	Right	974.81	59.8	-13.0	30.7	0.0143
B. Left LC contrast positively associated with thickness in OA						
Precentral	Left	5075.26	-58.5	2.9	26.2	0.0001
Inferior parietal	Left	4654.32	-44.2	-64.5	42.0	0.0001
Pars triangularis	Left	3308.43	-47.9	35.6	-8.6	0.0001
Superior parietal	Left	1044.15	-23.2	-84.0	25.3	0.0098
Precentral	Right	5166.03	40.7	-10.3	59.3	0.0001
Inferior parietal	Right	3241.41	35.0	-74.2	40.0	0.0001
Rostral middle frontal	Right	1805.06	40.4	25.4	32.6	0.0001
Lateral occipital	Right	1779.82	39.4	-81.0	-12.8	0.0001
Supramarginal	Right	1359.55	59.9	-39.9	34.5	0.0007
Lingual	Right	1290.46	22.3	-72.4	-6.2	0.0014
Superior parietal	Right	1135.09	23.7	-85.8	30.5	0.0051

Note. OA = older adults; YA = younger adults.

*Talairach coordinates of vertex with peak CWP value.

**Cluster-wise probability (CWP) resulting from cluster-wise Monte Carlo correction for multiple comparisons, reflecting the probability of the cluster appearing by chance. Only clusters with CWP<.05 are listed.

Table S4

Cortical clusters where the association between right LC contrast and thickness was greater in older than in younger adults (A) and positive in older adults (B)

Region	Hemi- sphere	Size (mm ²)	Peak vertex*			CWP**
			X	Y	Z	
A. Right LC contrast-thickness association greater in OA vs. YA						
Fusiform	Right	1102.77	35.6	-41.9	-15.5	0.0060
Precentral	Right	994.52	40.1	3.6	13.7	0.0128
B. Right LC contrast positively associated with thickness in OA						
Postcentral	Right	801.08	41.6	-24.7	53.9	0.0498

Note. OA = older adults; YA = younger adults.

*Talairach coordinates of vertex with peak CWP value.

**Cluster-wise probability (CWP) resulting from cluster-wise Monte Carlo correction for multiple comparisons, reflecting the probability of the cluster appearing by chance. Only clusters with CWP < .05 are listed.

2.4. Analysis of associations between LC contrast and cortical thickness, controlling for sex

2.4.1. Analysis of LC contrast and global cortical thickness

Results of the analyses examining the association between overall LC contrast and global cortical thickness including sex as a predictor are presented in Table S5. Overall, these results indicate that accounting for sex did not qualitatively change the associations we observed between overall LC contrast and global thickness. Specifically, after accounting for the effects of sex and its interactions with age and overall LC contrast on global thickness, we found a significant interaction effect between overall LC contrast and age group on global thickness (Table S5A), indicating that the association between overall LC contrast and global thickness differed significantly between older and younger adults. In this analysis, sex and age were also significant predictors of global thickness, but we did not observe a significant interaction effect between overall LC contrast and sex or a significant interaction effect between overall LC contrast, age group, and sex. Thus, we performed post hoc regression analyses separately for each age group, including overall LC contrast, age, and sex as predictors of global thickness. These analyses indicated that overall LC contrast was positively associated with global thickness in older adults (Table S5B), even after accounting for effects of age and sex on global thickness. However, in younger adults, a similar model predicting global thickness from overall LC contrast, age, and sex fit the data poorly, with none of these three variables being significantly associated with global thickness (Table S5C).

Table S6 presents results of similar regression analyses examining the association between rostral LC contrast and global cortical thickness after accounting for age and sex. In this case, after taking sex into account, we did not observe a significant rostral LC contrast-by-age group interaction (Table S6A), which approached significance when sex was not included as a predictor (main text, Table 3). Consistent with analyses for overall LC contrast, we examined the association between rostral LC contrast and global thickness separately in each age group, including age and sex as predictors. These analyses demonstrated that in older adults, after accounting for age and sex effects on thickness, higher rostral LC contrast significantly predicted greater cortical thickness (Table S6B), whereas in younger adults, neither age, sex, nor rostral LC contrast were significant predictors of global thickness (Table S6C). These results indicate

that the positive association we observed between rostral LC contrast and global thickness in older adults persisted when also accounting for sex effects on thickness.

Results of analogous regression analyses examining the association between caudal LC contrast and global thickness including age and sex as predictors are presented in Table S7. As in main analyses, we observed a trend towards caudal LC contrast being negatively associated with global thickness; however, did not observe a significant interaction effect between caudal LC contrast and age group on global thickness (Table S7A), whereas this interaction effect approached significance when sex was not included as a predictor (main text, Table 4). Subsequent regression analyses in each age group separately indicated that, as was the case in main analyses, the effect of caudal LC contrast was driven by younger adults, with lower caudal LC contrast in younger adults being a significant predictor of greater global thickness (Table S7C). In contrast, caudal LC contrast was not significantly associated with global thickness in older adults after regressing out effects of age and sex on global thickness (Table S7B). These results indicate that the negative association we observed between caudal LC contrast and global thickness in younger adults persisted when also accounting for sex effects on thickness.

Table S5

Results of multiple linear regression analyses examining the association between overall LC contrast and global cortical thickness, including sex as a predictor

Predictor	β	SE	95% CI	t	p
A. Full sample, $F(7, 288)=20.26, p<.001, R^2=0.33, \text{adjusted } R^2=0.314$					
Overall LC contrast	0.043	0.053	-0.062, 0.147	0.80	.423
Age	-0.484	0.052	-0.588, -0.381	-9.24	<.001
Sex	0.127	0.063	0.003, 0.252	2.01	.045
Overall LC contrast * Age group	0.121	0.053	0.017, 0.225	2.29	.023
Overall LC contrast * Sex	0.086	0.053	-0.019, 0.190	1.62	.107
Age group * Sex	0.096	0.063	-0.027, 0.220	1.53	.126
Overall LC contrast * Age group * Sex	-0.016	0.053	-0.120, 0.088	-0.31	.760
B. Older adults, $F(3, 225)=11.67, p<.001, R^2=0.135, \text{adjusted } R^2=0.123$					
Overall LC contrast	0.151	0.064	0.024, 0.278	2.35	.020
Age	-0.162	0.062	-0.284, -0.039	-2.60	.010
Sex	0.263	0.067	0.132, 0.395	3.94	<.001
C. Younger adults, $F(3, 63)=1.16, p=.332, R^2=0.052, \text{adjusted } R^2=0.007$					
Overall LC contrast	-0.097	0.128	-0.352, 0.159	-0.75	.454
Age	-0.189	0.128	-0.444, 0.067	-1.47	.145
Sex	0.040	0.138	-0.235, 0.315	0.29	.773

Note. Regression coefficients and standard errors are standardized values. Age group was coded as: older adults = 1, younger adults = -1. Sex was coded as females = 1, males = -1. CI = confidence interval; LC = locus coeruleus; SE = standard error.

Table S6

Results of multiple linear regression analyses examining the association between rostral LC contrast and global cortical thickness, including sex as a predictor

Predictor	β	SE	95% CI	<i>t</i>	<i>p</i>
A. Full sample, $F(7, 288)=20.21, p<.001, R^2=0.329, \text{adjusted } R^2=0.313$					
Rostral LC contrast	0.064	0.059	-0.052, 0.180	1.09	.276
Age	-0.472	0.053	-0.577, -0.368	-8.89	<.001
Sex	0.109	0.064	-0.018, 0.235	1.69	.092
Rostral LC contrast * Age group	0.081	0.059	-0.035, 0.197	1.38	.168
Rostral LC contrast * Sex	0.090	0.059	-0.027, 0.206	1.51	.131
Age group * Sex	0.124	0.064	-0.001, 0.250	1.95	.052
Rostral LC contrast * Age group * Sex	-0.074	0.059	-0.190, 0.043	-1.24	.215
B. Older adults, $F(3, 225)=11.98, p<.001, R^2=0.138, \text{adjusted } R^2=0.126$					
Rostral LC contrast	0.160	0.063	0.035, 0.284	2.52	.013
Age	-0.168	0.062	-0.290, -0.046	-2.71	.007
Sex	0.268	0.066	0.138, 0.398	4.07	<.001
C. Younger adults, $F(3, 63)=0.96, p=.416, R^2=0.044, \text{adjusted } R^2=-0.002$					
Rostral LC contrast	-0.001	0.128	-0.258, 0.255	-0.01	.993
Age	-0.206	0.130	-0.465, 0.053	-1.59	.117
Sex	0.013	0.137	-0.261, 0.287	0.09	.925

Note. Regression coefficients and standard errors are standardized values. Age group was coded as: older adults = 1, younger adults = -1. Sex was coded as females = 1, males = -1. CI = confidence interval; LC = locus coeruleus; SE = standard error.

Table S7

Results of multiple linear regression analyses examining the association between caudal LC contrast and global cortical thickness, including sex as a predictor

Predictor	β	SE	95% CI	<i>t</i>	<i>p</i>
A. Full sample, $F(7, 288)=19.66$, $p<.001$, $R^2=0.323$, adjusted $R^2=0.307$					
Caudal LC contrast	-0.151	0.085	-0.319, 0.018	-1.76	.079
Age	-0.432	0.058	-0.545, -0.319	-7.51	<.001
Sex	0.172	0.070	0.035, 0.310	2.47	.014
Caudal LC contrast * Age group	0.101	0.086	-0.068, 0.271	1.18	.239
Caudal LC contrast * Sex	-0.029	0.086	-0.199, 0.141	-0.34	.735
Age group * Sex	0.111	0.069	-0.026, 0.247	1.59	.112
Caudal LC contrast * Age group * Sex	-0.046	0.087	-0.217, 0.124	-0.54	.592
B. Older adults, $F(3, 225)=9.69$, $p<.001$, $R^2=0.114$, adjusted $R^2=0.103$					
Caudal LC contrast	-0.032	0.063	-0.157, 0.093	-0.50	.616
Age	-0.174	0.063	-0.298, -0.050	-2.77	.006
Sex	0.308	0.066	0.178, 0.437	4.67	<.001
C. Younger adults, $F(3, 63)=2.83$, $p=.046$, $R^2=0.119$, adjusted $R^2=0.077$					
Caudal LC contrast	-0.276	0.119	-0.514, -0.037	-2.31	.024
Age	-0.236	0.122	-0.480, 0.008	-1.94	.057
Sex	0.022	0.128	-0.234, 0.278	0.17	.863

Notes. Regression coefficients and standard errors are standardized values. Age group was coded as: older adults = 1, younger adults = -1. Sex was coded as females = 1, males = -1. CI = confidence interval; LC = locus coeruleus; SE = standard error.

2.4.2. *Vertex-wise analysis of LC contrast and regional cortical thickness*

Vertex-wise analyses indicated that, after accounting for age and sex and correcting for multiple comparisons, the association between overall LC contrast and thickness was greater for older than younger adults in five cortical clusters. The size and details of these clusters which were located within frontal and parietal cortices are presented in Table S8A. Subsequent analysis of older adults only indicated that overall LC contrast was positively associated with cortical thickness in seven cortical clusters. These clusters were located in regions within frontal, parietal, and occipital cortices, and their location and size details are presented in Table S8B. Compared to results of main analyses in which sex was not considered, we observed similar associations in frontal, parietal, and occipital cortices, although the number of clusters observed was reduced when taking sex into account. In younger adults, we did not observe any cortical clusters where overall LC contrast was associated with cortical thickness after multiple comparison correction. Subsequently we examined where on the cortical surface rostral and caudal LC contrast were associated with thickness after controlling for age and sex. We found that rostral LC contrast was more positively associated with thickness in older than younger adults in one cluster in left superior parietal cortex (Table S9A); when examining older adults only, we identified ten clusters where rostral LC contrast was positively associated with thickness (Table S9B), many of which are similar to the regions observed in main results. In addition, there were 3 clusters where caudal LC contrast was negatively associated with thickness in older adults (Table S9C). After controlling for age and sex, there were no cortical clusters where the association between caudal LC contrast and thickness differed in older and younger adults or where caudal LC contrast was associated with thickness in younger adults.

Table S8

Cortical clusters where the association between overall LC contrast and thickness was more positive in older than in younger adults (A) and positive in older adults (B), after controlling for age and sex

Region	Hemi- sphere	Size (mm ²)	Peak vertex*			CWP**
			X	Y	Z	
A. Overall LC contrast-thickness association greater in OA vs. YA						
Superior parietal	Left	1002.58	-12.8	-88.5	26.4	0.0121
Rostral middle frontal	Left	922.37	-35.9	30.7	14.1	0.0208
Banks of superior temporal sulcus	Right	1320.74	45.6	-43.1	13.6	0.0015
Fusiform	Right	806.93	36.5	-42.9	-13.8	0.0453
Transverse temporal	Right	798.86	49.7	-14.5	2.6	0.0477
B. Overall LC contrast associated with thickness in OA						
Rostral middle frontal	Left	1521.81	-36.4	31.5	16.2	0.0003
Lateral occipital	Left	1074.83	-46.0	-71.1	8.6	0.0076
Superior parietal	Left	970.91	-15.2	-73.7	45.4	0.0153
Precuneus	Right	1469.78	15.3	-54.6	13.1	0.0002
Fusiform	Right	1403.35	37.4	-39.5	-14.9	0.0003
Precentral	Right	1222.49	40.4	-10.9	57.9	0.0020
Lateral occipital	Right	1158.57	36.7	-85.4	-13.8	0.0034

*Talairach coordinates of vertex with peak CWP value.

**Cluster-wise probability (CWP) resulting from cluster-wise Monte Carlo correction for multiple comparisons, reflecting the probability of the cluster appearing by chance. Only clusters with CWP<.05 are listed.

Table S9

Cortical clusters where the association between rostral LC contrast and thickness was greater in older than in younger adults (A) and positive in older adults (B) and where the association between caudal LC contrast and thickness was negative in older adults (C), after controlling for age and sex

Region	Hemi- sphere	Association with thickness	Size (mm ²)	Peak vertex*			CWP**
				X	Y	Z	
A. Rostral LC-thickness association greater in OA vs. YA							
Superior parietal	Left	Positive	859.78	-23.1	-73.0	29.4	0.0342
B. Rostral LC contrast associated with thickness in OA							
Pars opercularis	Left	Positive	1727.08	-49.9	25.4	12.0	0.0001
Superior parietal	Left	Positive	1215.68	-10.8	-86.7	28.7	0.0031
Caudal middle frontal	Left	Positive	830.72	-39.4	10.8	52.0	0.0337
Supramarginal	Left	Positive	830.63	-47.3	-25.5	33.4	0.0337
Supramarginal	Right	Positive	1559.34	57.4	-40.3	34.3	0.0003
Precuneus	Right	Positive	1475.78	6.1	-58.4	31.2	0.0003
Fusiform	Right	Positive	1459.01	34.9	-39.9	-17.5	0.0004
Precentral	Right	Positive	1048.24	49.7	-5.1	8.5	0.0089
Pars triangularis	Right	Positive	855.06	54.0	27.0	8.3	0.0325
Precentral	Right	Positive	803.61	40.8	-10.3	58.2	0.0461
B. Caudal LC contrast associated with thickness in OA							
Pars orbitalis	Left	Negative	2298.92	-36.3	48.5	-9.0	0.0001
Postcentral	Left	Negative	857.06	-23.6	-35.9	59.9	0.0338
Supramarginal	Left	Negative	835.16	-58.4	-21.1	25.0	0.0373

*Talairach coordinates of vertex with peak CWP value.

**Cluster-wise probability (CWP) resulting from cluster-wise Monte Carlo correction for multiple comparisons, reflecting the probability of the cluster appearing by chance. Only clusters with CWP<.05 are listed.

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