Locus Coeruleus magnetic resonance imaging: a comparison between native-space and template-space approach

F. S. Giorgi1 · N. Martini2 · F. Lombardo3 · A. Galgani4 · L. Bastiani5 · D. Della Latta2 · H. Hlavata3 · C. L. Busceti6 · F. Biagioni6 · S. Puglisi-Allegra6 · N. Pavese7,8 · F. Fornai1,6

Received: 16 February 2022 / Accepted: 2 March 2022 / Published online: 20 March 2022 © The Author(s) 2022, corrected publication 2022

Abstract
Locus Coeruleus (LC) is the main noradrenergic nucleus of the brain, which is involved in many physiological functions including cognition; its impairment may be crucial in the neurobiology of a variety of brain diseases. Locus Coeruleus-Magnetic Resonance Imaging (LC-MRI) allows to identify in vivo LC in humans. Thus, a variety of research teams have been using LC-MRI to estimate LC integrity in normal aging and in patients affected by neurodegenerative disorders, where LC integrity may work as a biomarker. A number of variations between LC-MRI studies exist, concerning post-acquisition analysis and whether this had been performed within MRI native space or in ad hoc-built MRI template space. Moreover, the reproducibility and reliability of this tool is still to be explored. Therefore, in the present study, we analyzed a group of neurologically healthy, cognitively intact elderly subjects, using both a native space- and a template space-based LC-MRI analysis. We found a good inter-method agreement, particularly considering the LC Contrast Ratio. The template space-based approach provided a higher spatial resolution, lower operator-dependency, and allowed the analysis of LC topography. Our ad hoc-developed LC template showed LC morphological data that were in line with templates published very recently. Remarkably, present data significantly overlapped with a recently published LC “metaMask”, that had been obtained by averaging the results of a variety of previous LC-MRI studies. Thus, such a template space-based approach may pave the way to a standardized LC-MRI analysis and to be used in future clinic-anatomical correlations.

Keywords Locus Coeruleus · Magnetic resonance imaging · Aging · Noradrenaline · Biomarker

Introduction
Locus Coeruleus (LC) is the main noradrenergic (NA) nucleus of the brain and provides the NA innervation for the whole cortical mantle and subcortical structures (Counts and Mufson 2012). LC belongs to the brainstem isodendritic core and is a key element of the so-called...
reticular ascending activating system (Moruzzi and Magoun 1949; Theofili et al. 2015). LC contributes to a variety of brain functions, mainly by regulating neuronal homeostasis and modulating neural network activity (Poe et al. 2020). In particular, the LC-NA system plays a pivotal role in neurovascular coupling, blood–brain barrier homeostasis, and microglia modulation (Giorgi et al. 2020a, b); moreover, NA is crucial in wake/sleep cycle, attention and alert systems, and strongly contributes to learning and memory (Poe et al. 2020). This is due to a strong effect on modulating synaptic plasticity (e.g., Giorgi et al. 2006, 2008; Hansen 2017).

In the last decade, several studies have addressed the potential involvement of LC in the pathogenesis of neurodegenerative disorders (Gesi et al. 2000; Kelly et al. 2017). In particular, the development of MRI-based approaches aimed to visualize LC has allowed evaluating its integrity in vivo (see the reviews by Galgani et al. 2020; Beardmore et al. 2021). LC Magnetic Resonance Imaging (LC-MRI) has been already used to explore LC involvement in healthy aging (Dahl et al. 2019; Liu et al. 2020; Giorgi et al. 2021), Alzheimer’s Disease (Betts et al. 2019a; Jacobs et al. 2021), Parkinson’s Disease (Sommerauer et al. 2018; Li et al. 2019) and other pathological conditions (reviewed in Galgani et al. 2020). Thus, LC-MRI might add to the large variety of promising novel biomarkers that have been proposed in the last decade to improve diagnostic accuracy in neurodegenerative disorders (Frisoni et al. 2017; Baldacci et al. 2020).

Post-acquisition analysis often represents a major difference among LC-MRI studies; in particular, some authors have profited from data analysis performed in native-space (NS) (Olivieri et al. 2019; Jacobs et al. 2021), while others took advantage of an MRI template-space (TS) ad hoc built from their study sample itself (Dahl et al. 2019; Liu et al. 2020).

In a very recent study, Dahl et al. (2021) used a TS-based approach (which requires the construction of an LC mask—see Methods paragraph—identifying the region of interest of the LC in the MRI images), and developed an LC “MetaMask” taking into account a number of previously published LC masks (cited in Dahl et al. 2021). This MetaMask was made freely available, thus giving the opportunity also to other groups to test the reliability of their study-specific LC mask and of the MetaMask itself.

In a previous study, we explored the association of LC-MRI features with aging in a group of healthy and cognitively intact elderly subjects, using an NS-based post-processing protocol (Giorgi et al. 2021). Then, in line with the most recent studies on LC-MRI (Dahl et al. 2019), we specifically developed a post-acquisition analysis, building a study-specific LC template, by profiting of those MRI scans acquired in the context of the previous study (Giorgi et al. 2021).

In the present paper, we aimed to assess and discuss the degree of inter-method agreement between the NS- and the TS-based approach. Finally, the LC mask we developed in the present study was compared with the above-described LC metaMask (Dahl et al. 2021).

**Methods**

**Characteristics of the study sample**

The demographic characteristics and the recruitment protocol of the study sample are reported in detail in Giorgi et al. (2021). Briefly, 53 healthy and cognitively intact elderly subjects (mean age 71.70 ± 4.69 years, 20 males; mean Mini Mental State Examination 27.04 ± 1.27) were recruited at the Pisa University Hospital and Fondazione “G. Monasterio”- CNR/Tuscany Region. Exclusion criteria were: severe medical/cardiological and/or psychiatric comorbidities; neurological disease potentially associated with cognitive decline; history of drugs/alcohol abuse; MRI signs of moderate-severe chronic vascular encephalopathy, according to Fazekas et al. (1987), or other significant alterations. Cognitive and neurological integrity had been evaluated at baseline (T0) and confirmed after 1-year follow-up (T1). Brain MRI was performed within 30 days from T0.

**LC-MRI protocols**

MRI scans were performed using a 3T MR-Unit (GE Excite HDx, GE, USA) with an 8-channel phased-array head coil. The LC-sensitive sequence was acquired along the oblique axial plane (Fig. 1A), perpendicular to the fourth ventricle floor, covering an area from the inferior border of the pons to the posterior commissure. We used a 2D-FSE T1-weighted sequence with the following parameters: TR 600 ms; TE 14 ms; flip angle 90°; echo train length 2; NEX 5; matrix size 512×384; FOV 200×200 mm; pixel size 0.39×0.52 mm; 12 contiguous slices, slice thickness 2.2 mm, slice gap 0; acquisition time 14.29 min.

The complete Brain MRI protocol also included 2D-FLAIR, T2* GRE, Spin EchoT1- and FSE T2-weighted with fat saturation and diffusion-weighted imaging sequences. In addition, 3D-Fast-SPGR T1-weighted images were obtained: TR 10.7 ms; TE 4.9 ms; FOV 256×256 mm; matrix size 256×256; isotropic voxel 1 mm; NEX 1; acquisition time: 5.50 min.

The post-processing imaging analysis provided the quantification of two LC-MRI parameters: LC Contrast Ratio (LC_Cr), which represents the signal intensity of the LC, and the total amount of voxels considered as belonging to the LC (LC_Vox), which was considered as an indirect estimation of LC volume.
Native space-based approach

The semi-automatic NS protocol was already described in Giorgi et al. (2021). Briefly, the analysis was performed by two independent trained operators, using a software developed in-house in Java language. LC region of interest (ROI) was manually placed at the level of the fourth ventricle floor, from the inferior border of the pons to the posterior commissure. In panel B, it is reported the probabilistic map of LC-MRI; voxels with a probability higher than 10% to carry LC signal among included subjects are distributed in consecutive MRI slices with a pattern strongly suggestive of LC anatomical counterpart.

Template space-based approach

Our templates for the 3D and the LC-MRI scans were built according to the same procedure proposed by other authors (Dahl et al. 2019), with minor modifications. The workflow steps are detailed below.

Whole brain 3D template creation  As the first step, a common space for the 3D anatomical MRI (template3D) was created using MRI images of the recruited cognitively intact healthy subjects. These were interpolated to an isotropic resolution of 0.5 mm and a nonuniform intensity correction was applied (N4 bias field correction) and then submitted to multi-resolution iterative registration of 3D scans (antsMultivariateTemplateConstruction2 of the Advanced Normalization Tools v.2.3.4 software (Avants et al. 2011). Registration parameters used were: number of iterations 30 × 90 × 20, cross-correlation similarity metric, Greedy SyN transformation model. Once the registration process had been completed, the aligned 3D volumes were averaged in order to obtain the group whole-brain template3D. We also saved the transformation matrices and warping fields that warped the 3D from the individual subject’s space to the template3D space (see below).

Brainstem template creation  The second step was the creation of a common brainstem space for the acquisition of LC-sensitive sequence. Using the transformations calculated during the whole-brain template3D creation procedure, each scan obtained with LC-sensitive sequence was warped from the native 2D space to the space of the template3D itself. A second multi-resolution iterative registration was run with the same parameters (iterations, similarity metric, transformation model) described above. This spatial registration took into account the anatomical variability between subjects in the LC acquisitions, as well as the possible intra-subject misalignment between the LC and the 3D scans. As a result, we obtained the brainstem template, whose isotropic resolution was 0.5 mm.

LC mask creation  In the brainstem template, we found increased intensity and higher signal-to-noise ratio com-
pared to the ventral pons, in a symmetrical bilateral region below the floor of the fourth ventricle. Considering the anatomical location of the signal, the LC/sub-coeruleus complex was considered the main signal source candidate (Bettss et al. 2019a; Dahl et al. 2019). To identify those voxels more likely belonging to the LC and, thus, to create the LC mask, a semi-automatic thresholding procedure was performed. An expert operator placed two square 8.5 mm × 8.5 mm reference ROIs bilaterally in the ventrals pons in each slice, using ITKsnap (www.itksnap.org) (Yushkevich et al. 2006). The threshold was computed following this formula: μROI + 4*σROI, where μROI and σROI are the mean value and the standard deviation in the reference ROIs, respectively. Then, a custom program written in Python selected all the voxels exceeding the threshold in the brainstem template, for both hemispheres. These represent the candidate LC-belonging voxels of the brainstem template. The segmentation mask was further refined by an expert neuroradiologist (FL) to exclude voxels that were anatomically incongruent with the LC (e.g., within the fourth ventricle).

**LC-MRI parameters extraction** Once the LC mask was obtained, LC-sensitive scans of each subject were warped from the native space to the brainstem template space for data analysis, and the parameters LC_{CR} developed by TS-based approach (LC_{CR}-TS) and LC_{VOX} developed by TS-based approach (LC_{VOX}-TS) were extracted. The parameter LC_{CR}-TS was calculated using the following formula: LC_{CR}=[max(LC)—max(Ref)] / max(Ref), where max(Ref) and max(LC) are the maximum signal intensities in the left and right reference regions, and LC regions derived from the LC template segmentation, respectively.

LC_{VOX}-TS was calculated as follows. First, the LC mask in the brainstem template was considered as the search space of the LC voxels. Then, reference ROIs were drawn on the LC scan warped to the brainstem template, and a subject-specific threshold was calculated using the same formula used for the LC mask. The number of voxels in the search space that exceed the subject-specific threshold constituted LC_{VOX}-TS parameter value.

For each subject, the two LC-MRI parameters (LC_{CR} and LC_{VOX}) were calculated for left and right hemisphere separately (“left LC” and “right LC”, respectively), or for left and right LC combined (“combined LC”).

In order to maximally reduce any potential side-related intensity artifacts which might affect the results, we applied to LC scans a field correction algorithm (Tustison et al. 2010), and LC-related parameters were standardized using bilateral ROIs as reference regions, placed symmetrically in the left and ventral pons (see above).

**Comparison with previously published metaMask**

Recently, an LC mask obtained by pooling a variety of study-specific LC masks (“metaMask”), was released and made freely available (https://osf.io/sf2ky/) (Dahl et al. 2021). To assess the compatibility between our TS-based approach and the published metaMask, we evaluated the overlap in the MNI space between our LC mask and the metaMask. To warp our LC mask into the MNI space, we first co-registered the brainstem template to the 0.5-mm iso-voxel MNI template using linear (rigid, then affine), followed by nonlinear (SyN) registration (Avants et al. 2011). Then, these transformations were applied to the LC mask using a nearest-neighbor interpolation to warp the LC mask into the MNI space. The degree of compatibility between our LC mask and the metaMask was expressed using the same parameters that Dahl et al. themselves described in their study, namely specificity and sensitivity (Dahl et al. 2021). These were calculated as the ratio between the number of voxels the study-specific LC mask and the LC metaMask have in common and the total number of voxels of the former and the latter, respectively. We calculated also the “accuracy” which was defined as the mean between sensitivity and specificity of the mask (Dahl et al. 2021). For comparison, we also warped the metaMask into the brainstem template through inverse transformation, and we additionally calculated the LC parameters using the metaMask in the TS approach.

**Statistical analysis**

Data normality was assessed through Shapiro–Wilk test and visual checking of heteroskedasticity. Since they did not show a Gaussian distribution, variables were Z-standardized. Pearson’s correlation was used to evaluate inter-method agreement (van Stralen et al. 2008) and age effect on TS-based LC parameters. A Paired-sample t-test was used to assess side-related differences within TS-based LC variables. Multiple comparisons were adjusted with False Discovery Rate (FDR) correction to rule out type I error. The level of significance was set at p < 0.05. All statistical analyses were performed using SPSS Version 25. GraphPad Prism 8.0 was used for graphical representations.

**Results**

**Template space-based LC reconstruction**

The spatial co-registration of each LC-sensitive scan to the brainstem template was carefully visually inspected. The obtained brainstem template showed marked hyperintensities bilaterally in the LC region, while the region of the ventral pons showed a rather homogeneous distribution. The
final LC mask included a total number of 234 voxels across 17 axial slices. Figure 1B shows the 3D visualization of the LC probabilistic map, reflecting the coordinates of the maximum value of the left and right LC for each slice.

**Template space-based LC features**

We did not find any effect of age on TS-based LC parameters, neither considering the combined LC (LC_{CR-TS}, $r = -0.291$, $p = 0.07$, LC_{VOX-TS}, $r = -0.254$, $p = 0.066$), nor analyzing the two LC separately (LC_{CR-TS}, $r = -0.213$, $p = 0.167$, LC_{VOX-TS}, $r = -0.273$, $p = 0.096$ for the left LC; LC_{CR-TS}, $r = -0.322$, $p = 0.076$, LC_{VOX-TS}, $r = -0.254$, $p = 0.152$, for the right LC).

A side-related difference was observed between the two LCs. LC_{CR-TS} and LC_{VOX-TS} values of the left LC were higher than those of the right LC ($t = 5.556$, df = 52, $p < 0.001$; $t = 4.434$, df = 52, $p < 0.001$, respectively) (Fig. 2).

Remarkably, there was no significant difference between left and right reference regions intensity values ($t = 0.518$, df = 52, $p = 0.607$).

**Inter-method agreement**

There was a good agreement between the signal intensity measures obtained by the two methods. LC_{CR-NS} showed a direct correlation with LC_{CR-TS} values obtained from two LC combined ($r = 0.413$, $p = 0.004$) (Fig. 3A). It was
evaluated also the correlation of left and right LC values, separately assessed with the TS-based approach, with the single (i.e., including the values from left and right LC) parameters extracted with the NS-based approach, in order to assess whether the latter ones were more related to the LC signal of one side or the other one. We found that the same trend described above for the combined LC was observed when left and right LC were analyzed separately ($r = 0.383$, $p = 0.010$ and $r = 0.371$, $p = 0.006$, respectively) (Fig. 3B and C).

The NS- and TS-based methods showed a significant agreement with each other also concerning volume-related measures (although the inter-method agreement was lower compared to the $LC_{CR}$ ones), when considering either the combined LC ($r = 0.340$, $p = 0.013$) (Fig. 3D) or the two LC separately ($r = 0.330$, $p = 0.032$ for the left $LC_{VOX}$, $r = 0.305$, $p = 0.026$ for the right $LC_{VOX}$) (Fig. 3E and F).

**Compatibility of the template-based approach with LC metaMask**

We found a high degree of compatibility between our LC mask and the LC metaMask (specificity 69.8%, sensitivity 49.5% and accuracy 59.6%; Fig. 4). Moreover, the LC-MRI parameters calculated using our LC mask showed very strong direct correlation coefficients with those calculated using the LC metaMask (Supplementary Table 1).

**Discussion**

*In vivo* LC analysis by MRI has been performed by several authors and it will likely be a matter of interest also in future years, as LC degeneration represents a promising marker in a number of neurodegenerative disorders (Betts et al. 2019a; Galgani et al. 2020). In this study, we presented the results of an LC-MRI analysis performed using both an NS- and a TS-based approach in a group of cognitively healthy subjects. We successfully built an LC-MRI template and used it to extract LC imaging parameters. Then, we compared the results obtained with both approaches, and we showed a good inter-method agreement for $LC_{VOX}$, and mostly for $LC_{CR}$.

$LC_{CR}$ appears to be more directly related to LC features, since it simply represents the ratio between the MRI signal intensity of LC and the reference regions. This might explain why $LC_{CR}$ showed a higher correlation coefficient between the two methods. Nonetheless, there was not a complete concordance between NS- and TS-based approaches regarding this parameter. This may depend on the higher spatial resolution of the TS-based approach, which allows the identification of many voxels that were potentially excluded from the NS-based analysis, thereby affecting $LC_{CR}$ values.

Such a higher spatial resolution may also explain the lower concordance between NS- and TS-based approaches concerning the parameter $LC_{VOX}$. Thus, $LC_{VOX}$ may be useful for LC-MRI analysis studies, since it provides the estimation of authentic LC volume. Even though we have already tried to extrapolate this parameter also using the NS-based approach (Giorgi et al. 2021), $LC_{VOX}$ computed with the TS-based method might be more appropriate to calculate the LC anatomical volume. In fact, as shown in present paper, and in line with what already shown by the groups that used a similar approach (e.g., Betts et al. 2017; Dahl et al. 2019), the spatial distribution of LC voxels closely followed LC anatomy. Concerning this aspect, it is also worth emphasizing that the distribution of $LC_{VOX}$ showed a higher value in the middle part of the LC mask and a lower one in its rostral and caudal parts; this is in line with anatomical data, which repeatedly showed that LC neurons are more abundant and densely packed in the middle part of the nucleus, while they are scattered towards LC rostral and caudal poles (Schwarz and Luo 2015). Also, the length and the volume of the nucleus estimated from the LC mask were close to what measured in histological studies (Fernandes et al. 2012) (Fig. 1A).

In a previous paper, in which we profited from the NS-based approach only (Giorgi et al. 2021), we did not find any association between LC features and age in this group of subjects. Here, we confirmed this observation also with the TS-based approach, both when merging data from right and left LC, and when considering the left and right LC...
separately. These findings are in line with recent post-mortem data, which, by profiting from unbiased stereology, did not show any significant LC involvement during physiological aging (Theofilas et al. 2017).

The left/right side-related difference we observed in the distribution of LC-MRI parameters deserves a special emphasis. Left LC showed higher $LCC_{CR}$ and $LCC_{VOX}$ values compared to the right one. To our knowledge, this has never been specifically addressed in post-mortem studies of LC. MRI studies of LC carried out by other authors (Betts et al. 2017; Liu et al. 2019) are compatible with the present finding in spite of different MRI scanners and LC post-acquisition methods. In the present study, all efforts were made to avoid any potential machine-related influence on LC signal (as witnessed also by the lack of differences between left and right ventral pons reference ROIs intensity), nonetheless side-related difference may still be partly due to uncommon artifacts (see also Betts et al. 2017). Such an asymmetry needs to be validated and, when confirmed, it should be specifically analyzed to better understand the role of LC in brain activities and brain disorders.

So far, all research teams that profited from a TS-based approach developed specific LC mask based on their own subjects’ sample (e.g., Betts et al. 2017; Dahl et al. 2019). However, in a very recent study Dahl et al. (2021) developed a so-called LC metaMask. This was built-up by pooling together the LC masks developed by various research groups. Concordance of the metaMask with each specific study-dependent mask was assessed in the same study (Dahl et al. 2021). The fact that our LC mask remarkably overlaps with Dahl’s group’s metaMask, even though it was not used to build it, lends further substance to the reliability of the TS-based approach for LC imaging studies.

In conclusion, our study showed a good agreement between NS- and TS-based approaches, with the latter method offering more advantages in terms of spatial resolution and regional analysis of LC. Furthermore, since the TS-based approach drastically reduces operator-dependency, it would pave the way to a standardized methodological protocol for LC-MRI analysis and, thus, to its possible clinical application. As already mentioned, LC has been receiving growing attention in recent years for its possible role both in the pathogenesis and pathophysiology of Alzheimer’s Disease (Rorabaugh et al. 2017; Weinshenker et al., 2018), and thus it might even become a target of potential disease-modifying drugs. In such a context, LC-MRI may be used in the future to identify the Alzheimer’s Disease patient phenotype with the highest burden of LC pathology and more likely to benefit from a LC-targeted therapy.

Supplementary Information The online version contains supplementary material available at https://doi.org/10.1007/s00702-022-02486-5.

Author contributions FSG: conceptualized the study, received specific fundings, coordinated the research team, participated in including the subjects, contributed to data interpretation, to critical review of the literature, reviewed and finalized the manuscript. NM: developed the software for LC post-acquisition analysis (both the native-space and the template-space ones), performed imaging analysis, re-evaluated all the MRIs, concurred to critical review of the literature on LC and MRI, and to reviewing the manuscript. FL: performed all MRI acquisitions, concurred to MRI analysis development and application, to critical review of the literature on LC and MRI, and to reviewing the manuscript. AG: participated in including the subjects, contributed to data analysis and interpretation, to critical review of the literature, contributed to the statistical analyses, tables/figures generation, and drafted the paper. LB: performed statistical analysis and reviewed the paper. DDL developed the native-space approach. HH: concurred to MRI acquisition and analysis. CLB, FB, SP-A: contributed to finalizing the article. NP and FF: contributed to discussing the data, to critically reviewing and finalizing the article.

Funding Open access funding provided by Università di Pisa within the CRUI-CARE Agreement. This work was funded by a grant of the Italian Ministry of Health to FSG [grant number “Ricerca Finalizzata PE-2013–02359574—In vivo assessment of the role of Locus Coeruleus in the development of Alzheimer’s Disease and other types of Dementia” (PI: FSG)]. FF is funded by Ministero della Salute—Ricerca Corrente.

Declarations

Conflict of interest DDL is presently an employee of Terarecon Inc; his collaboration to the present study occurred before its present affiliation, his contribution to this article reflects entirely and only his own expertise on the matter, and he declares no competing financial interests related to the present article. All the other authors do not have competing interests to disclose concerning the present manuscript.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit http://creativecommons.org/licenses/by/4.0/.

References

Avants BB, Tustison NJ, Song G et al (2011) A reproducible evaluation of ANTs similarity metric performance in brain image registration. Neuroimage 54:2033–2044. https://doi.org/10.1016/J.NEUROIMAGE.2010.09.025

Baldacci F, Mazzucchi S, dellaVecchia A et al (2020) The path to biomarker-based diagnostic criteria for the spectrum of neurodegenerative diseases. Expert Rev Mol Diagn 20(4):421–441

Beardmore R, Hou R, Darekar A et al (2021) The locus coeruleus in aging and Alzheimer’s disease: a postmortem and brain imaging review. J Alzheimers Dis 83:5–22. https://doi.org/10.3233/JAD-210191

DDL is presently an employee of Terarecon Inc; his collaboration to the present study occurred before its present affiliation, his contribution to this article reflects entirely and only his own expertise on the matter, and he declares no competing financial interests related to the present article. All the other authors do not have competing interests to disclose concerning the present manuscript.
Betts MJ, Cardenas-Blanco A, Kanowski M et al (2017) In vivo MRI assessment of the human locus coeruleus along its rostrocaudal extent in young and older adults. NeuroImage 163:150–159. https://doi.org/10.1016/j.neuroimage.2017.09.042

Betts MJ, Cardenas-Blanco A, Kanowski M et al (2019a) Locus coeruleus MRI contrast is reduced in Alzheimer’s disease dementia and correlates with CSF Ap levels. Alzheimers Dement 11:281–285. https://doi.org/10.1016/j.dadm.2019.02.001

Betts MJ, Kirilina E, Otaduy MCG et al (2019b) Locus coeruleus imaging as a biomarker for noradrenergic dysfunction in neurodegenerative diseases. Brain 142:2558–2571. https://doi.org/10.1093/brain/awz193

Counts SE, Mufson EJ (2012) Locus Coeruleus. In: Paxinos G (ed) The Human Nervous System, Third Edition, pp 427–440

Dahl MJ, Mather M, Düzsel S et al (2019) Rostral locus coeruleus integrity is associated with better memory performance in older adults. Nat Hum Behav 3:1203–1214. https://doi.org/10.1038/s41562-019-0715-2

Dahl MJ, Mather M, Werkle-Bergner M et al (2021) Locus coeruleus integrity is related to tau burden and memory loss in autosomal-dominant Alzheimer’s disease. Neurobiol Aging 112:39–54. https://doi.org/10.1016/j.neurobiaging.2021.11.006

Fazekas F, Chawluk JB, Alavi A et al (1987) MR signal abnormalities at 1.5 T in Alzheimer’s dementia and normal aging. Am J Roentgenol 149:351–356. https://doi.org/10.2214/ajr.149.2.351

Fernandes P, Regala J, Correia F, Gonçalves-Ferreira AJ (2012) The human locus coeruleus 3-D stereotactic anatomy. Surg Radiol Anat 34:879–885. https://doi.org/10.1007/s00276-012-0979-y

Frisoni GB, Boccardi M, Barkhof F et al (2017) Strategic roadmap for an early diagnosis of Alzheimer’s disease based on biomarkers. Lancet Neurol 16(8):661–676. https://doi.org/10.1016/S1474-4422(17)30159-X

Galganì A, Lombardo F, dellaLatta D et al (2019) The role of the locus coeruleus in phenotypic variants of Alzheimer’s disease. Ann Clini Trans Neurol 6(7):1345–1351. https://doi.org/10.1002/acn3.50818

Gesi M, Soldani P, Giorgi FS et al (2000) The role of the locus coeruleus in the development of Parkinson’s disease. Neurosci Biobehav Rev 24:655–668. https://doi.org/10.1016/S0149-7634(00)00028-2

Giorgi FS, Mauceli G, Blandini F et al (2006) Locus coeruleus and neuromelanin plasticity in a model of focal limbic epilepsy. Epilepsia 47(Suppl 5):21–25. https://doi.org/10.1111/j.1528-1167.2006.01235.x

Giorgi FS, Blandini F, Cantafora E et al (2008) Activation of brain metabolism and fos during limbic seizures: the role of locus coeruleus. Neurobiol Dis 30:388–399. https://doi.org/10.1016/j.nbd.2008.02.008

Giorgi FS, Biagioni F, Galganì A et al (2020a) Locus coeruleus modulates neuroinflammation in parkinsonism and dementia. Int J Mol Sci 21(22):8630. https://doi.org/10.3390/ijms21228630

Giorgi FS, Galganì A, Puglisi-Allegra S et al (2020b) Locus coeruleus and neurovascular unit: from its role in physiology to its potential role in Alzheimer’s disease pathogenesis. J Neurosci Res 98(12):2406–2434. https://doi.org/10.1002/jnr.24718

Giorgi FS, Lombardo F, Galganì A et al (2021) Locus coeruleus magnetic resonance imaging in cognitively intact elderly subjects. Brain Imaging Behav. https://doi.org/10.1007/s11682-021-00562-0

Hansen N (2017) The longevity of hippocampus-dependent memory is orchestrated by the locus coeruleus-noradrenergic system. Neural Plast 2017:2727602. https://doi.org/10.1155/2017/2727602

Jacobs HL, Becker JA, Kwong K et al (2021) In vivo and neuro-pathology data support locus coeruleus integrity as indicator of Alzheimer’s disease pathology and cognitive decline. Sci Trans Med 13(612):eabj2511. https://doi.org/10.1126/scitranslmed.abj2511

Kelly SC, He B, Perez SE et al (2017) Locus coeruleus cellular and molecular pathology during the progression of Alzheimer’s disease. Acta Neuropathol Commun 5:8. https://doi.org/10.1186/s40478-017-0411-2

Li Y, Wang C, Wang J et al (2019) Mild cognitive impairment in de novo Parkinson’s disease: a neuromelanin MRI study in locus coeruleus. Mov Disord 34(6):884–892. https://doi.org/10.1002/mds.27682

Liu KY, Acosta-Cabronero J, Cardenas-Blanco A et al (2019) In vivo visualization of age-related differences in the locus coeruleus. Neurobiol Aging 74:101–111. https://doi.org/10.1016/j.neurobiaging.2018.10.014

Liu KY, Kievit RA, Tsvetanov KA et al (2020) Noradrenergic-dependent functions are associated with age-related locus coeruleus signal intensity differences. Nat Commun 11:1712. https://doi.org/10.1038/s41467-020-15410-w

Moruzzi G, Magoun HW (1949) Brain stem reticular formation and activation of the EEG. Electroencephalogr Clin Neurophysiol 1:455–473. https://doi.org/10.1016/0013-4694(49)90219-9

Oliviieri P, Lagarde J, Lehericy S et al (2019) Early alteration of the locus coeruleus in phenotypic variants of Alzheimer’s disease. Ann Clini Trans Neurol 6(7):1345–1351. https://doi.org/10.1002/acn3.50818

Poe GR, Foote S, Eschenko O et al (2020) Locus coeruleus: a new look at the blue spot. Nat Rev Neurosci 21(11):644–659. https://doi.org/10.1038/s41583-020-0360-9

Rorabaugh JM, Chalermpalanupap T, Botz-Zapp CA et al (2017) Chemogenetic locus coeruleus activation restores reversal learning in a rat model of Alzheimer’s disease. Brain 140(11):3023–3038. https://doi.org/10.1093/brain/awx232

Schwarz LA, Luo L (2015) Organization of the locus coeruleus-norepinephrine system. Curr Biol 25(21):R1051–R1056. https://doi.org/10.1016/j.cub.2015.09.039

Sommerauer M, Fedora TD, Hansen AK et al (2018) Evaluation of the noradrenergic system in Parkinson’s disease: an 11 C-MeNER PET and neuromelanin MRI study. Brain 141(2):496–504. https://doi.org/10.1093/brain/awx348

Theofilas P, Dunlop S, Heinzen H, Grinberg LT (2015) Turning on the light within: subcortical nuclei of the isodentritic core and their role in Alzheimer’s disease pathogenesis. J Alzheimers Dis 46(17–18):33–43. https://doi.org/10.3233/JAD-142682

Theofilas P, Ehrenberg AJ, Dunlop S et al (2017) Locus coeruleus volume and cell population changes during Alzheimer’s disease progression: a stereological study in human postmortem brains with potential implication for early-stage biomarker discovery. Alzheimers Dement 13:236–246. https://doi.org/10.1016/j.jalz.2016.06.2362

Tustison NJ, Avants BB, Cook PA et al (2010) N4ITK: improved N3 bias correction. IEEE Trans Med Imaging 29:1310. https://doi.org/10.1109/TMI.2010.2046008

van Stralen KJ, Jager KJ, Zoccali C, Dekker FW (2008) Agreement between methods. Kidney Int 74:1116–1120. https://doi.org/10.1038/KI.2008.306

Weinshenker D (2018) Long road to ruin: noradrenergic dysfunction in neurodegenerative disease. Trends Neurosci 41(4):211–223. https://doi.org/10.1016/j.tins.2018.01.010

Yushkevich PA, Piven J, Hazlett HC et al (2006) User-guided 3D active contour segmentation of anatomical structures: significantly improved efficiency and reliability. Neuroimage 31:1116–1128. https://doi.org/10.1016/j.neuroimage.2006.01.015

Publisher’s Note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.