Right Insular Atrophy in Neurocardiogenic Syncope: A Volumetric MRI Study

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*AJNR Am J Neuroradiol* 2014, 35 (1) 113-118
doi: https://doi.org/10.3174/ajnr.A3611
http://www.ajnr.org/content/35/1/113
ABSTRACT

BACKGROUND AND PURPOSE: Alterations in the central autonomic network are hypothesized to play a role in the pathophysiologic mechanism underlying neurocardiogenic syncope; however, few data are available regarding the structural changes of the brain in this condition. We used voxel-based morphometry and regional volumetry to identify possible neuroanatomic correlates.

MATERIALS AND METHODS: We prospectively studied 32 patients with neurocardiogenic syncope with a positive response to the head-up tilt test and 32 controls who had no history of syncope. We used voxel-based morphometry to compare GM volumes between patients and controls. In addition, regional volumes of structures known to be included in the central autonomic network were measured and compared between the groups. Correlation analyses were also performed between clinical variables and anatomic data.

RESULTS: Voxel-based morphometry showed a significant GM volume reduction in the right insular cortex in patients with neurocardiogenic syncope compared with controls (corrected $P = 0.033$). Regional volumetry showed a significant reduction of right insular volumes in patients compared with controls ($P = 0.002$, MANCOVA). Smaller right insular volumes in patients with neurocardiogenic syncope were related to larger drops in systolic ($P = 0.020$) and diastolic ($P = 0.003$) blood pressures during the head-up tilt test.

CONCLUSIONS: We observed a novel finding of right insular atrophy in patients with neurocardiogenic syncope with a positive response to the head-up tilt test, implicating the role of right insular dysfunction in the pathophysiologic mechanism underlying neurocardiogenic syncope. Our findings further support the hypothesis that right insular dysfunction may cause a decrease in sympathetic activity and a reciprocal increase in parasympathetic activity, leading to syncope.

ABBREVIATIONS: BP = blood pressure; HR = heart rate; HUT = head-up tilt test; NCS = neurocardiogenic syncope; VBM = voxel-based morphometry

N eurocardiogenic syncope (NCS) (simple fainting or vasovagal syncope) is the most frequent cause of syncope.1,2 It refers to a condition in which autonomic cardiovascular reflexes become inappropriate in response to triggers, leading to transient cerebral hypoperfusion and loss of consciousness with spontaneous recovery.3,5 Although the pathophysiologic mechanism of NCS remains incompletely understood, paradoxic compensatory reflex has been suggested as the most probable mechanism.3,6 Reduction of cardiac ventricular volume activates baroreceptor-mediated sympathetic outflow, which causes vigorous contraction of an empty cardiac ventricle. In turn, activation of mechanoreceptors in the ventricular wall might trigger an inhibitory response through increasing vagal tone, resulting in paradox bradycardia and hypotension.3,6 This widely accepted “ventricular theory” basically includes increased parasympathetic activity caused by alteration of autonomic controls in the CNS.2 Therefore, the understanding of CNS structures involved in the autonomic control system might be crucial for elucidating fundamental mechanisms of NCS.

To our knowledge, there is 1 voxel-based morphometry (VBM) study that investigated changes in brain structures in patients with NCS.8 VBM is an objective and fully validated method, which can detect regionally specific differences in GM structures on a voxelwise comparison between groups.8 In the present study, we applied VBM and regional volumetry to identify the structural GM changes associated with NCS. In addition, we investigated possible relationships between the structural changes and clinical variables such as disease duration, frequency of syncope attacks, and blood pressure (BP) and heart rate (HR) changes during head-up tilt test (HUT).
MATERIALS AND METHODS

Subjects
Consecutive patients with NCS were prospectively recruited from the outpatient clinic at the Neurology Department, Korea University Guro Hospital, from January 2010 to April 2012. Inclusion criteria were as follows: 1) unequivocal symptoms of syncope (defined as transient loss of consciousness and postural tone with spontaneous recovery) and presyncope (defined as a state of light-headedness, muscular weakness, and feeling faint without complete loss of consciousness); 2) orthostatic stress as the only trigger of syncope and a positive response to HUT (as described below); 3) no history of syncope relevant to situational syncope or carotid sinus hypersensitivity; 4) absence of any abnormal findings on electroencephalography; 5) no history of epileptic seizure; 6) no evidence of developmental and neurologic abnormalities; 7) neither abnormal nor unusual findings on conventional MR imaging; 8) no evidence of cardiogenic syncope based on electrocardiography and echocardiography; 9) no history of diabetes, Parkinsonism, and peripheral neuropathy; and 10) no history of significant head injury, alcohol or psychotropic drug abuse, or neurologic and psychiatric disorders. Demographic and clinical data such as age of onset, disease duration, and the number of syncope and presyncope attacks were obtained through interviews with the patients and their family members and reviews of medical records.

For group comparison, 32 right-handed healthy volunteers matched for age and sex were recruited to serve as control subjects. All control subjects underwent neurologic examination and a detailed interview to ensure that they did not have the following: 1) a history of syncope and presyncope; 2) a history of headache disorders, significant head injury, neurologic, psychiatric, or systemic disorders; and 3) a history of alcohol or psychotropic drug abuse. The local ethics committee approved the study protocol, and all participants gave written informed consent before study inclusion.

Head-Up Tilt Test
None of the patients were on any medications at the time of the HUT examination. Orthostatic BP recording was performed with a sphygmomanometer cuff over the brachial artery and a monitor console displaying systolic and diastolic BP. Simultaneous HR recording was performed as well. After a 20-minute rest in the supine position and measurement of baseline BP and HR, each subject was positioned at an angle of 70° from the supine position for as long as 45 minutes or until syncope occurred on a standard electrically driven tilt table with a footboard. Serial measurement of BP and HR was performed every 2 minutes during HUT. Pharmacologic provocation was not performed.10 The response to HUT was defined as positive when the development of symptoms recognized by the patients was similar to those of spontaneous syncope in association with hypotension, bradycardia, or both.4,11,12 As soon as syncope occurred, the patient was rapidly returned to the supine position. Changes in BP and HR were measured from the baseline to the onset of the syncope attack.

MR Imaging Acquisition
MR images were acquired on a Trio 3T scanner (Siemens, Erlangen, Germany) with a 12-channel phased array head coil. For volumetric analysis, a high-resolution 3D MPRAGE sequence was acquired with the following parameters: TR = 1780 ms, TE = 2.34 ms, matrix = 256 × 256, FOV = 256 × 256 mm, voxel dimension = 1 mm³. For identification of structural abnormalities, the following conventional MR images were acquired: axial T2-weighted and FLAIR images (4-mm thickness), oblique coronal T2-weighted and FLAIR images perpendicular to the long axis of the hippocampus (3-mm thickness), and gadolinium-enhanced axial T1-weighted images (5-mm thickness). The MR images of all participants were reviewed by a board-certified neuroradiologist (S.-i.S.) for any structural abnormalities, and findings were reported as normal.

Voxel-Based Morphometry
Data were processed and analyzed by using SPM8 (Wellcome Department of Imaging Neuroscience, London, UK), where we applied VBM implemented in the VBM8 toolbox (http://dbm.neuro.uni-jena.de/vbm.html) with default parameters. Images were bias-corrected, tissue-classified, and registered by using linear (12-parameter affine) and nonlinear transformations, within a unified segmentation model.13 Subsequently, analyses were performed on the volumes of the GM segments, which were multiplied by nonlinear components derived from the normalization matrix to compensate for possible volume changes during the nonlinear spatial normalization procedure. These modulated GM images were then smoothed with an 8-mm full width at half maximum isotropic Gaussian kernel. The analyses of modulated GM images can detect regional differences in GM volume.

Regionally specific differences in GM volume between patients with NCS and control subjects were assessed by using ANCOVA with total intracranial volume, age, and sex as nuisance variables. An absolute GM threshold of 0.2 (of a maximum value of 1) was used to avoid possible edge effects around the border between GM and WM. Two contrasts were defined to examine both increases and reductions in GM volume between the groups. Statistical significance was set at a height threshold of P < .001 and an extent threshold of cluster-level P < .05, corrected for multiple comparisons by using family-wise error.

Regional Volumetry
Cortical reconstruction and volumetric segmentation were performed by using FreeSurfer software (Version 5.1.0, http://surfer.nmr.mgh.harvard.edu). The technical details of the procedures are described elsewhere.14-17 Briefly, image preprocessing procedures included removal of nonbrain tissue,16 automated Talairach transformation, segmentation of the subcortical WM and deep GM volumetric structures,15 intensity normalization, tessellation of the GM/WM boundary, automated topology correction,18 and surface deformation to detect GM/WM and GM/CSF boundaries. Subsequent to cortical reconstruction, the cortex was subdivided into units on the basis of gyral and sulcal structures.19 This parcellation method was shown to be valid and reliable, with high intraclass correlation coefficients between the manual and automated procedures for volume measurement of 34 cortical regions of interest.19 A volume at each vertex was computed as the average area of the triangles around the vertex multiplied by the thickness at the vertex. The volume of a parcellated cortex was then computed as the sum of the volumes of the vertices in each parcellation.
The automated procedures for volumetric measurements of the subcortical GM structures were described in detail previously. Briefly, this procedure automatically provided segments and labels for up to 40 unique structures and assigned a neuroanatomic label to each voxel in an MR imaging volume on the basis of probabilistic information estimated automatically from a manually labeled training set. A Bayesian segmentation procedure was then performed, and the maximum a posteriori estimate of the labeling was computed. The segmentation uses 3 pieces of information to disambiguate labels: 1) the prior probability of a given tissue class occurring at a specific atlas location, 2) the likelihood of the image intensity given that tissue class, and 3) the probability of the local spatial configuration of labels given the tissue class. This technique has previously been shown to be comparable in accuracy with manual labeling. All segmentations were visually inspected for accuracy before inclusion in the analysis.

Among the automatically segmented regions, we focused on 3 cortical regions (rostral anterior cingulate, caudal anterior cingulate, and insula) and 1 subcortical region (amygdala) in each hemisphere, which are known to be parts of the central autonomic network. Volumes of the 8 regions were automatically measured and compared between patients and controls by using MANCOVA, adjusting for age, sex, and estimated total intracranial volume. Post hoc univariate tests with a Bonferroni correction were performed to follow the significant main effects yielded by the MANCOVA. Statistical significance was set at \( P < .00625 \) (.05/8). Correlation analyses were performed between the normalized volumes of 8 structures (each regional GM volume corrected for estimated total intracranial volume) and clinical parameters such as disease duration, total frequency of syncope and presyncope, and BP and HR changes during HUT (Spearman correlation analysis, \( P < .05 \)). Statistical analyses were performed by using the Statistical Package for Social Sciences (Version 19.0; IBM, Armonk, New York).

**RESULTS**

**Clinical Characteristics**

Thirty-two right-handed patients and 32 controls were enrolled in this study. Clinical characteristics and the results of HUT are summarized in the Table. There were no differences in age and sex between patients and controls. The mean age of onset was 20.8 ± 7.1 years (range, 11–39 years), the mean duration of NCS was 4.3 ± 4.7 years (range, 1–22 years), and the mean number of total syncope and presyncope attacks (mean number of attacks per year × disease duration in years) was 38.4 ± 74.4 (range, 2–300).

All patients showed positive responses to HUT. Mean baseline systolic BP was 109.5 ± 12.2 mm Hg (range, 87–148 mm Hg), mean baseline diastolic BP was 62.0 ± 9.6 mm Hg (range, 47–81 mm Hg), and mean baseline HR was 67.7 ± 10.0 beats/min (range, 52–86 beats/min). During HUT, mean change of systolic BP was −40.5 ± 15.3 mm Hg, mean change of diastolic BP was −18.3 ± 10.1 mm Hg, and mean change of HR was −33.5 ± 15.8 beats/min (range, −5 to −65 beats/min). According to the Vasovagal Syncope International Study (VASIS) classification, 31 of 32 patients had a type I mixed response and 1 patient had a type 3 vasodepressor response.

**Voxel-Based Morphometry and Regional Volumetry**

VBM showed a significant GM volume reduction in the right insular cortex in patients with NCS compared with control subjects (family-wise-error–corrected \( P = .033 \); cluster size = 1178 voxels [3976 mm³]; peak z score = 4.00; Montreal Neurological Institute coordinates of local maxima = 42, 3, −6; resolution element count = 1.5) (Fig 1). No region of a significant GM volume increase was found in patients compared with controls at the same threshold (family-wise-error–corrected \( P < .05 \)).

Regional volumes of 8 GM structures are presented in the Table. Volumes of the right insular cortex were significantly reduced in patients with NCS compared with controls (7323 ± 689.1 versus 8011 ± 932.5 mm³, \( P = .002 \)) (Fig 2). Volumes of the left rostral anterior cingulate cortex (\( P = .014 \) and left insular cortex (\( P = .018 \) were also reduced in patients compared with controls; however, the statistical significance of these 2 structures did not survive multiple comparisons by using a Bonferroni correction (\( P < .00625 \)).

Smaller right insular volumes in patients with NCS were reduced in patients compared with controls; however, the statistical significance of these 2 structures did not survive multiple comparisons by using a Bonferroni correction (\( P < .00625 \)).

**Table**

| VBM Score | Patients with NCS (n = 32) | Controls (n = 32) | P Value |
|-----------|---------------------------|------------------|---------|
| L insula  | 241 ± 6.9                 | 24.8 ± 5.2       | .655    |
| Male sex (No.) (%) | 11 (34.4)              | 12 (37.5)        | .794    |
| Age of onset (yr) | 20.8 ± 7.1               | 18.3 ± 10.1      | .018    |
| Disease duration (yr) | 4.3 ± 4.7               | −33.5 ± 15.8     | .003    |
| No. of total syncope and presyncope attacks | 38.4 ± 74.4            |                 |         |

**Head-up tilt test findings**

Baseline systolic BP (mm Hg) 109.5 ± 12.2
Baseline diastolic BP (mm Hg) 62.0 ± 9.6
Baseline mean HR (beats/min) 67.7 ± 10.0
Change in systolic BP (mm Hg) −40.5 ± 15.3
Change in diastolic BP (mm Hg) −18.3 ± 10.1
Change in HR (beats/min) −33.5 ± 15.8

**Volume of structure (mm³)**

| Structure | Patients with NCS (n = 32) | Controls (n = 32) | P Value |
|-----------|---------------------------|------------------|---------|
| R rostral anterior cingulate | 2288 ± 416.3          | 2461 ± 425.2     | .137    |
| R caudal anterior cingulate | 2386 ± 555.6          | 2488 ± 639.3     | .498    |
| R insula | 7323 ± 689.1              | 8011 ± 932.5     | .002    |
| R amygdala | 1755 ± 232.2           | 1829 ± 184.9     | .339    |
| L rostral anterior cingulate | 2791 ± 484.6          | 3141 ± 589.7     | .014    |
| L caudal anterior cingulate | 1819 ± 494.8          | 2142 ± 711.5     | .063    |
| L insula | 7324 ± 649.5             | 7657 ± 701.0     | .018    |
| L amygdala | 1770 ± 216.2          | 1816 ± 229.1     | .963    |

**Note:** R indicates right; L, left

*The values are presented as means. An independent t test and chi² test were used to compare between-group differences in age and sex, respectively. MANCOVA adjusting for age, sex, and estimated total intracranial volume was performed to compare regional volumes of each structure between the groups.*
correlations between right insular volumes and other clinical variables including HR changes, disease duration, and number of total syncope and presyncope attacks (all $P > .05$). Volumes of the left rostral anterior cingulate and left insular cortices were not correlated with any clinical variables (all $P > .05$).

**DISCUSSION**

This study attempted to explore cerebral structural changes associated with pathophysiologic mechanisms underlying NCS. Using VBM, we found that patients with NCS had regional atrophy in the right insular cortex. We also observed a significant reduction of whole right insular volume in patients with NCS compared with controls, consistent with the VBM finding. Moreover, smaller right insular volumes in patients with NCS were related to larger drops in both systolic and diastolic BP during HUT, suggesting an important involvement of the right insular cortex in NCS.

Among the regions belonging to the central autonomic network, the insular cortex has been recognized as playing a crucial role in cardiovascular modulation. An increase in nocturnal BP and occurrence of QT prolongation and cardiac arrhythmias were more frequently observed in patients with stroke with involvement of the insular cortex compared with those without insular involvement. Damage to the right insular cortex was associated with impairment in sympathovagal balance, supporting an important implication of the right insular cortex in autonomic control of cardiovascular activity.

Several lines of evidence indicate that there is a hemispheric lateralization of cardiovascular autonomic control of the insular cortex. A study investigating heart rate variability changes during intracarotid amobarbital injection in patients with temporal lobe epilepsy showed sympathetic activation with left hemisphere injection and parasympathetic activation with right hemisphere injection. An intraoperative study before temporal lobectomy in patients with epilepsy demonstrated that BP and HR decreased with left insular stimulation, whereas BP and HR increased with right insular stimulation. These findings strongly support a critical role of the left insula in parasympathetic cardiovascular control and of the right insula in sympathetic control.

There is a reciprocal interaction between sympathetic and parasympathetic control of cardiac autonomic functions. Damage confined to the left insular cortex by acute stroke could shift cardiovascular balance toward an increase in basal sympathetic tone and a decrease in parasympathetic tone. Likewise, our finding could be interpreted as right insular atrophy in NCS being associated with a decrease in sympathetic activity and a reciprocal increase in parasympathetic activity, resulting in a drop in BP and syncope. Our speculation

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**FIG 1.** Statistical parametric maps showing regional differences in GM volume between 32 patients with NCS and 32 controls. Orthogonal images show a reduction of GM volume in the right insula in patients with NCS compared with controls, superimposed on a standard template ($P < .05$, cluster-level-corrected for multiple comparisons using family-wise error). The color bar represents the $T$ values. The left side of each image is the left side of brain.

**FIG 2.** An illustration of segmentation and volume measurement of the right insula in a control subject (upper row, volume = 8119 mm$^3$) and a patient (lower row, volume = 7247 mm$^3$). The automatically segmented insular cortex is superimposed on axial (first column) and coronal (second column) GM images, and the third column illustrates a 3D reconstructed image of the right insula. The left side of each image is the right side of brain.
might be further supported by the findings of greater right insular volume reductions in relation to larger drops in both systolic and diastolic BP during HUT.

There are only a few studies evaluating hemodynamic and structural changes of the brain in patients with NCS. Studies by using SPECT showed decreases in regional cerebral blood flow in multiple cortical regions, yielding inconsistent results across the studies. The results of these SPECT studies might be interpreted as a consequence of cerebral hypoperfusion rather than a specific cause of NCS. To our knowledge, there is only 1 exploratory VBM study that showed a reduction of GM volume in the medulla in patients with NCS. The study suggested that NCS may be associated with structural changes of the medulla, which predisposes the patient to abnormal cardiovascular homeostasis and recurrent syncope. However, we did not observe any GM volume changes in the brain stem, even by using a lenient threshold of uncorrected P < .01. This difference could not be properly explained but might, in part, be ascribed to different sample sizes, MR scanners, and patient characteristics. Because a substantial number of conditions can mimic NCS, only patients with orthostatic stress and a positive response to HUT were included in our study. In their study, only 2 of 18 patients had orthostatic stress as a main trigger of syncope, and the remaining patients had variable triggers such as emotional distress, blood phobia, instrumentation, or pain. Because NCS comprises a heterogeneous group of conditions based on pathophysiologic classification of the principal triggers of syncope, the differences in the patient populations included might account for the inconsistency between the studies.

There are several limitations to our study. First, the current study is cross-sectional, and thus interpretation of our results with respect to a causal relationship is limited. Given that GM atrophy found in VBM could be a reflection of neuropathologic changes in many brain disorders such as temporal lobe epilepsy, Alzheimer disease, and schizophrenia, we hypothesized that right insular atrophy may represent a primary underlying pathology and therefore act as the specific cause that promotes NCS. Conversely, taken together with our finding of a trend toward significance for regional volume reductions of the left insula and anterior cingulate cortex in patients with NCS, it seems possible that right insular atrophy might be a consequence of repeated attacks of cerebral hypoperfusion. Further prospective study incorporating a longitudinal design would provide a hint of the causal relationship between NCS with a positive response to HUT, implicating a role of right insular dysfunction in the pathophysiologic mechanism underlying NCS. Together with previous investigations, our findings suggest that right insular atrophy may be associated with a decrease in sympathetic activity and a reciprocal increase in parasympathetic activity, resulting in cerebral hypoperfusion and syncope. Confirmation of our findings in larger, more homogeneous datasets of patients with NCS is warranted.

Disclosures: Ji Hyun Kim—RELATED: Grant: This work was supported in part by the National Research Foundation of Korea Grant funded by the Korean Government (Grant No. 20100004827, 20100005418) and a Korea University Grant.

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