Clinical Significance of Vertebral Artery Hypoplasia in Lateral Medullary Infarction

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**Objective:** The clinical significance of vertebral artery hypoplasia (VAH) and the possible pathomechanism of lateral medullary infarction in patients with VAH are still not completely clear.

**Methods:** Among 3,968 patients with acute ischemic stroke, we selected 102 patients with lateral medullary infarction (2.6% [102/3,968]; 67 men, 35 women; mean age, 66 years; range, 33 to 86 years) who underwent brain magnetic resonance imaging, contrast-enhanced magnetic resonance angiography, and transcranial Doppler (TCD) within 1 week of symptom onset. We compared the characteristics of VAH and non-VAH group.

**Results:** Of 102 patients with lateral medullary infarction, 34 (33.3%) had hypoplastic vertebral artery and 68 (66.7%) were non-VAH. The location of stroke in patients with right-sided VAH (n = 22) was predominantly ipsilateral and in the 12 patients with left-sided VAH, the stroke was predominantly ipsilateral. TCD parameters of Mean flow velocity and pulsatility index were significantly different on the affected side between VAH and non-VAH groups (P < 0.001). Patients having dominant vertebral artery is opposite to basilar artery curvature are 88.2% (30/34).

**Conclusion:** We concluded that VAH can be considered an additional risk factor for lateral medullary infarction. The influence of reduced blood flow in the hypoplastic vertebral artery is also important to consider, especially when other vascular risk factor are present. The presence of dolichoectasy of the basilar artery ipsilateral to the VAH causes reorganization of the blood flow around the vertebrobasilar junction.

**Keywords:** Vertebral artery; Hypoplasia; Infarction

INTRODUCTION

Vertebral artery hypoplasia (VAH) was described in the 19th century. Congenital variations of the caliber of the vertebral arteries or asymmetry of these vessels are not infrequent. VAH with caliber discrepancies of more than 1:1.7 was observed in up to 10% of normal individuals. There is a tendency for the contralateral vessel to compensate with an increased diameter of about 5 mm. There is no general agreement as to the definition of VAH. Operational definitions of VAH vary between diameters of less than 2 to less than 3 mm or an asymmetry ratio of equal or greater than 1:1.7. Additionally, reduced diastolic flow velocity associated with increased resistance index values indicates VAH [1,2].

Acquired morphological variations in the vertebral artery (VA), such as stenosis, occlusion, or dissection, are associated with different clinical symptoms. The clinical significance of VAH is not currently well understood. In the literature, a possible causal association between the presence of VAH and the incidence of posterior stroke has been shown. Sonographic finding of VAH has reported the presence of reduced flow velocities (compared to the contralateral side), and high resistance flow pattern in the Doppler spectrum. However, the exact pathomechanism of ischemic stroke in patients with VAH is still not completely clear [3-7].

Lateral medullary infarction (Wallenberg syndrome) is the clinical manifestation resulting from occlusion of the intracranial vertebral artery (ICVA) or of the posterior inferior cerebellar artery (PICA) or one of its branches, in which the lateral part of the medulla oblongata infarcts, resulting in a typical pattern. The most
commonly affected artery is the ICVA, followed by the PICA [8].

The aim of our study was to investigate clinical significance of VAH and the possible pathomechanism of lateral medullary infarction in patients with VAH.

**MATERIALS AND METHODS**

Participants were selected from 3,968 consecutive patients with acute ischemic stroke who were admitted to Soonchunhyang University Cheonan Hospital between September 2007 and August 2017. Among 3,968 patients, we selected 102 patients with lateral medullary infarction (2.6% [102/3,968]; 67 men, 35 women; mean age, 66 years; range, 33 to 86 years) who underwent brain magnetic resonance imaging, contrast-enhanced magnetic resonance angiography (MRA), and transcranial Doppler (TCD) within 1 week of symptom onset. The acute ischemic lesion on lateral medulla was assessed by high signal intensity on diffusion-weighted imaging with low signal intensity on apparent diffusion coefficient. MRA was analyzed blindly to the TCD results. VA was classified as hypoplasia and non-hypoplasia according to our MRA criteria.

VAH was defined to have a lumen diameter of less than 2–3 mm. Because there is no consensus on the value of VAH, we defined VAH as a diameter of ≤ 2.2 mm or an asymmetry ratio of equal or greater than 1:1.7. For defining the diameter of VAH, we examined the V2 of the largest VA by using contrast-enhanced MRA. The location of the VA was defined according to standard criteria (V1, prevertebral portion; V2, the portion within the vertebral column; V3, atlantoaxial portion; V4, intracranial portion) [8]. To assess reliability, the MRA of each patient was independently reviewed by two investigators (HG Oh, SC Lee), both of whom were blinded to all clinical information. The consensus for the presence of VAH was standardized according to neuro-radiological interpretation reports and was acceptable (κ = 0.85, P < 0.01). In non-VAH group, stenosis degree of the distal VA (V4) responsible for lateral medullary infarction was defined as occlusion, stenosis (> 50% reduction of arterial column width) and normal. Also we investigated the direction and degree of basilar artery (BA) curvature and relationship between VA dominance and BA dolichosis.

We assessed the risk factors in each patient for acute ischemic stroke, such as hypertension, diabetes, hyperlipidemia, current smoking, and heart disease. Cardiogenic embolic diseases (e.g., atrial fibrillation, left ventricular akinesia, recent myocardial infarction, or patent foramen ovale with right-to-left shunt) were routinely evaluated in all patients by electrocardiography and transthoracic echocardiography; further studies—such as transesophageal echocardiography or 24-hour holter monitoring—were performed if any abnormality was found on initial work-up. For each patient we attempted to determine the etiology of stroke.

TCD was performed with a 2-MHz pulsed Doppler instrument by an experienced sonographer. VA was evaluated between 40 and 80 mm depth through suboccipital window. Mean flow velocity (MFV), pulsatility index (PI), and flow velocity ratio to BA (VA/BA FVR) were compared between VAH and non-VAH group. Asymmetry index (AI) of VA calculated by $2\times100\times(\text{ipsilateral MFV}/\text{ipsilateral MFV}+\text{contralateral MFV})$ (%) was also analyzed.

Statistical analysis was done using SPSS ver. 17.0 (SPSS Inc., Chicago, IL, USA). Comparisons were performed using the Student t-test for continuous variables and Fisher’s exact test for nominal-level variables. P < 0.05 was considered as significant.

**RESULTS**

Of 102 patients with lateral medullary infarction, 34 (33.3%) had hypoplastic VA and 68 (66.7%) were non-VAH. We found right-sided VAH more frequently, that is, in 22 cases (64.7%). In 12 cases (35.3%), the VAH was located on the left side. None of non-VAH patients had experienced a VA dissection.

There was no substantial age difference between the non-VAH and the VAH groups. The difference in gender between VAH and non-VAH group was also not statistically significant. There was no correlation between the conventional vascular risk factors (hypertension, diabetes mellitus, hyperlipidemia, and smoking) and the presence or absence of VAH in patients with lateral medullary infarction (Table 1).

The location of stroke in patients with right-sided VAH (n = 22) was predominantly ipsilateral and in the 12 patients with left-sided VAH, the stroke was predominantly ipsilateral (Table 2). Patients having dominant VA is opposite to BA curvature are 88.2% (30/34). These findings were assigned to the blood flow reorganization mechanism based on the findings of Hong et al. [9] (Fig. 1).

TCD parameters of MFV and PI were significantly different on the affected side between VAH and non-VAH groups. On the non-affected side, MFV was significantly higher on VAH group. Also, AI was higher in VAH group, which was statistically significant (Table 3).
DISCUSSION

Recently, a cut-off point of < 2.2 mm was determined for VAH, based on significant hemodynamic changes, which is further sup-
ported by an increase in ipsilateral flow resistance (resistance index ≥ 0.75), contralateral diameter (side-to-side diameter difference ≥ 0.12 mm), and flow volume (side-to-side flow volume ratio ≥ 5) by color-coded duplex ultrasonography. The frequency of VAH (defined as VA diameter < 2.2 mm by duplex ultrasonography) was shown to be 11.6% (7.8% on the right and 3.8% on the left) in 447 subjects [1,2]. In Korean study, the frequencies of VAH were as high as 35.2% in 529 patients with ischemic stroke (3-dimension time-of-flight or contrast enhanced MRA) [4]. VAH is predominant on the right side, which is consistent with our finding.

VA hypoplasia as a congenital vessel variation was first mentioned in the 19th century. About a quarter of all ischemic strokes are localized in the vertebrobasilar circulation territory. Emerging evidences suggest that VA hypoplasia may contribute to posterior circulation ischemic events, especially when other risk factors coexist [4,5,10].

In patients with lateral medullary infarction, the most common vascular lesion is occlusion of the proximal or middle portion of the ICVA. Penetrating branches to the lateral medulla arise from the middle and distal 2/3 of the ICVAs and penetrate through the lateral medullary fossa to reach and supply the lateral medullary tegmentum. The medial branches of PICA supply only a small portion of the dorsal medullary tegmentum. The ICVA occlusive lesions decrease flow in these penetrators. Less often lateral medullary infarction is caused by occlusion of one of the small medullary branches [8].

It is assumed that the vessel diameter plays an important role in

Table 1. Basic demographic characteristics and stroke risk factors of patients with lateral medullary infarction between VAH group and non-VAH group (n = 102)

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>VAH group (n = 34)</th>
<th>Non-VAH group (n = 68)</th>
<th>P-valuea)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (yr)</td>
<td>65.5 ± 10.3</td>
<td>67.2 ± 9.4</td>
<td>NS</td>
</tr>
<tr>
<td>Male sex</td>
<td>24 (70.6)</td>
<td>43 (63.2)</td>
<td>NS</td>
</tr>
<tr>
<td>Hypertension</td>
<td>12 (35.3)</td>
<td>23 (33.8)</td>
<td>NS</td>
</tr>
<tr>
<td>Diabetic mellitus</td>
<td>3 (8.8)</td>
<td>7 (10.3)</td>
<td>NS</td>
</tr>
<tr>
<td>Hyperlipidemia</td>
<td>6 (17.6)</td>
<td>11 (16.2)</td>
<td>NS</td>
</tr>
<tr>
<td>Smoking</td>
<td>4 (11.8)</td>
<td>8 (11.8)</td>
<td>NS</td>
</tr>
<tr>
<td>Cardioembolism</td>
<td>4 (11.8)</td>
<td>10 (14.7)</td>
<td>NS</td>
</tr>
</tbody>
</table>

Values are presented as mean ± standard deviation or number (%)

Table 2. Laterality of VAH correlated to the location of ischemic lesion

<table>
<thead>
<tr>
<th>Location of stroke</th>
<th>Right sided VAH (n = 22)</th>
<th>Left sided VAH (n = 12)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Right lateral medulla</td>
<td>20</td>
<td>1</td>
</tr>
<tr>
<td>Left lateral medulla</td>
<td>2</td>
<td>11</td>
</tr>
</tbody>
</table>

VAH, vertebral artery hypoplasia.

Table 3. MFV, PI, FVR to BA, and AI of vertebral artery

<table>
<thead>
<tr>
<th>Variable</th>
<th>VAH group (n = 34)</th>
<th>Non-VAH group (n = 68)</th>
<th>P-valuea)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Affected site</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>MFV (cm/sec)</td>
<td>26.5 ± 13.1</td>
<td>33.0 ± 11.5</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>PI</td>
<td>0.91 ± 0.20</td>
<td>0.73 ± 0.15</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>VA/BA FVR</td>
<td>0.78 ± 0.04</td>
<td>0.77 ± 0.28</td>
<td>NS</td>
</tr>
<tr>
<td>Non-affected site</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>MFV (cm/sec)</td>
<td>38.5 ± 17.7</td>
<td>29.5 ± 12.5</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>PI</td>
<td>0.75 ± 0.18</td>
<td>0.74 ± 0.15</td>
<td>NS</td>
</tr>
<tr>
<td>VA/BA FVR</td>
<td>0.79 ± 0.48</td>
<td>0.78 ± 0.31</td>
<td>NS</td>
</tr>
<tr>
<td>AI of the VA (%)</td>
<td>24.5 ± 22.3</td>
<td>17.2 ± 13.7</td>
<td>&lt; 0.001</td>
</tr>
</tbody>
</table>

Values are presented as mean ± standard deviation. P < 0.05 is considered statistically significant.

MFV, mean flow velocity; PI, pulsatility index; FVR, flow velocity ratio; BA, basilar artery; AI, asymmetry index; VAH, vertebral artery hypoplasia; VA, vertebral artery; NS, non-significant.

a)Student t-test was used.
the ischemia mechanism. According to Poiseuille’s law, the flow rate is proportional to the fourth power of the vessel radius. Based on the hemodynamics in VAH, the reduced blood flow in a hypoplastic VA plays an important role in the mechanism of stroke in the vertebrobasilar territory. The VAH in coexistence with known risk factors for stroke may increase their negative clinical impact. Hong et al. [9] postulated that the unequally mechanical forces resulting from the asymmetric VA flow might influence the morphological deformation in the vertebrobasilar arterial system, for example, a lateral displacement or elongation (dolichoectasy) of the BA. The presence of dolichoectasy of the BA ipsilateral to the VAH causes reorganization of the blood flow around the vertebrobasilar junction. This BA deviation predisposes to increased frequency of stroke in the territory of the hypoplastic VA (Fig. 1) [9, 11,12].

Lateral medullary infarctions should be more frequent on the side of VAH. It is thought that luminal narrowing of the VAH might make it less feasible for cardiogenic emboli to pass through it. The VAH, which shows lower mean flow volume and decreased flow velocities, seems to be more susceptible to pro-thrombotic or atherosclerotic processes than normal or dominant vertebral arteries. Vertebral arterial thrombi can cause in situ strokes on lateral medulla and prone to cause distal embolization to PICA territory lateral medulla [12,13].

We concluded that VAH can be considered an additional risk factor for lateral medullary infarction when coexisting with conventional vascular risk factors. The influence of reduced blood flow in the hypoplastic VA is also important to consider, especially when other vascular risk factor are present.

Because of the relatively small sample size, further studies are necessary to confirm the association between VAH and lateral medullary infarction and to clarify the pathomechanism of stroke in the presence of VAH.

REFERENCES