Spontaneous Obliteration of a Dissecting Aneurysm of Recurrent Artery of Heubner Monitored by Serial Magnetic Resonance Vessel Wall Imaging

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Abstract

Aneurysms of the recurrent artery of Heubner (RAH) are known to be one of the uncommon cerebral aneurysms, predominantly presenting with bleeding symptoms. Previously, nine cases of the RAH aneurysms have been reported, all of which were treated surgically or endovascularly and most cases developed postoperative cerebral infarct in the ipsilateral caudate nucleus. Herein, we report a man presenting with transient ischemic attack due to diffuse cerebral vasospasm from a minor non-disabling subarachnoid hemorrhage (SAH) from an RAH aneurysm. He visited our hospital 7 days after the first experience of a thunderclap headache complaining with transient unilateral motor weakness and thin SAH in the right sylvian fissure. Diagnostic catheter angiography revealed a dissecting fusiform aneurysm (8 mm in size) originating from the left RAH contralateral to the thin SAH. Contrast-enhanced magnetic resonance vessel wall imaging (MR-VWI) helped to identify the ruptured nature of the RAH aneurysm. Owing to his delayed ischemic condition after minor SAH, he was conservatively treated with serial MR-VWI monitoring. The aneurysm was spontaneously obliterated with an asymptomatic lacunar infarct in the ipsilateral caudate nucleus in a month. Together, this case was considered as the dissecting aneurysm of RAH with a favorable outcome after the conservative management. Although long-term follow-up is mandatory because the disappearance of the vessel wall enhancement does not necessarily secure the permanent cure of the lesion, serial MR-VWI is helpful to diagnose the ruptured nature and monitor its chronological change in combination with conventional radiological imaging techniques.

Keywords
► MRI
► recurrent artery of Heubner
► vascular disorder
► vessel wall imaging

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Key Messages

Spontaneous obliteration of a dissecting aneurysm of recurrent artery of Heubner could be monitored by serial magnetic resonance vessel wall imaging as a clear contrast enhancement of the affected vessel.

Introduction

Aneurysms of the recurrent artery of Heubner (RAH) are known as uncommon cerebral aneurysms located nearby the horizontal portion of the anterior cerebral artery at the base of the brain. To the best of our knowledge, only nine cases of the RAH aneurysms have been reported, all of which were treated surgically or endovascularly. In most cases, hemorrhagic presentation was the clinical symptom and clinically silent, postoperative cerebral infarcts were observed in the ipsilateral caudate nucleus, except for two cases with unfavorable outcome.\(^1\)\(^-\)\(^7\) Previous reports mostly claimed that RAH aneurysms can be treated surgically because RAH occlusion might only cause clinically silent cerebral infarct, which is debatable based on the two above-mentioned unexpected cases. Herein, we report a man with a dissecting fusiform aneurysm of RAH, monitored by repeated magnetic resonance (MR) imaging of the vessel wall, which was helpful in identifying the ruptured nature of this aneurysm as a clear contrast enhancement of the affected vessel.

Case History

A 58-year-old man presented to the emergency room of our hospital after transient weakness in his left extremities. He had a thunderclap headache a week ago without any neurological deficit. He had essential thrombocytosis and was on once-daily low-dose aspirin, while his past brain health check-up screening showed no abnormalities in his brain and cerebral arteries. A brain computed tomography (CT) scan showed no subarachnoid hemorrhage. Transient ischemic attack (TIA) due to reversible cerebral vasoconstriction syndrome (RCVS) was suspected at this time. One week later,
however, he complained of sustained headache without repeated TIA symptoms (i.e., transient weakness in his left extremities). Brain magnetic resonance (MR) images demonstrated thin subarachnoid hemorrhage (SAH) in the right distal Sylvian fissure (►Fig. 1A) and MR angiography demonstrated diffuse cerebral vasospasm and a fusiform aneurysm above the horizontal portion of the left anterior cerebral artery (►Fig. 1B). Notably, MR vessel wall imaging with an improved motion-sensitized driven equilibrium (iMSDE) technique showed marked enhancement of the aneurysm and adjacent brain tissue (►Fig. 1C–E). Catheter carotid angiography revealed a fusiform aneurysm (8 mm in size) originating from the left RAH, which arose from the junction of the horizontal portion of the left anterior cerebral artery and the anterior communicating artery (►Fig. 2). There was no evidence of moyamoya-like vessels, vasculitis, or tumor stains in the four vessel observations. Given the “string and pearl” appearance of the aneurysm along with the RAH (►Fig. 2C) and the clear enhancement of the aneurysm noted during vessel wall MR imaging (►Fig. 1C–E), a ruptured nature was strongly suspected. Thus, we considered this patient first developed minor subarachnoid bleeding from the dissecting aneurysm of the RAH (presenting with thunderclap headache), followed by TIA as a delayed neurological deficit (DIND) due to diffuse cerebral vasospasm from the thin SAH. Based on these considerations, we decided to start with conservative treatment for cerebral vasospasm with strict blood pressure maintenance and close observation for the rebleeding from the ruptured aneurysm of the RAH. Aspirin therapy was discontinued. After a month, his headache disappeared, and TIA did not recur. Notably, repeated catheter angiography and MR vessel wall imaging revealed obliteration of the aneurysm and RAH without any neurological sequelae (►Fig. 3A–D). As expected, MR images showed a clinically silent cerebral infarct in the ipsilateral caudate nucleus (►Fig. 3E). His clinical outcome was excellent (modified Rankin scale 0 at 3 months after the onset), and daily oral low-dose aspirin was re-started for the essential thrombocytosis. We still continue his follow-up to monitor recurrence of the arterial lesion.

Discussion

We present the first case of spontaneous obliteration of a ruptured dissecting aneurysm of the RAH after a month of conservative treatment. This case showed thunderclap headache as an initial symptom with no evidence of SAH on CT. Consequently, TIA occurred and MR imaging showed thin SAH, suggesting either RCVS or DIND due to diffuse cerebral vasospasm secondary to SAH. Repeat diagnostic work-up revealed a fusiform cerebral aneurysm on the horizontal part of the anterior cerebral artery and diffuse cerebral vasospasm, indicating the patient developed aneurysmal SAH but not RCVS. Catheter angiography clearly demonstrated a fusiform aneurysm located along with the RAH. Moreover, clear contrast enhancement of the aneurysm by the MR vessel wall imaging suggested the ruptured nature of the dissecting aneurysm of the RAH.

There were nine previously reported RAH aneurysms presenting mostly with SAH (seven of nine cases). All were

![Fig. 2](image-url) Catheter angiography showing a recurrent artery of Heubner aneurysm. The anteroposterior view (A) and three-dimensional reconstruction of conventional and rotational angiography (B) of the left carotid artery demonstrates a recurrent artery of Heubner (RAH) fusiform aneurysm (arrow heads) on the left A1. The oblique view of the right carotid artery angiogram (C) shows the fusiform aneurysm (white arrows) more clearly along with the left RAH (black arrows).
treated with open or endovascular surgery, as summarized in Table 1. 

Previous reports all concluded that the RAH aneurysm can be surgically treated with favorable outcome. The postoperative course was uneventful, even if RAH was obliterated after the surgery. However, Ogata et al reported an RAH aneurysm presenting with non-severe SAH, which was resected after trapping of the RAH with postoperative neurological deficit and cerebral infarct in the ipsilateral caudate nucleus. The deficit was fully recovered; the authors stated that preservation of the RAH might be difficult in surgical treatment of the aneurysm. Mansfield et al reported a dissecting RAH aneurysm in a patient with osteogenesis imperfecta who ultimately died from postoperative severe cerebral vasospasm. The patient presented with severe SAH (Glasgow coma scale 5); therefore, surgical treatment of the RAH aneurysm was unavoidable and performed successfully with the expected clinically silent cerebral infarct. Our patient presented with TIA due to cerebral vasospasm from non-severe SAH. Presumably, the aneurysmal SAH occurred a week before without any sign of rebleeding. Due to the ischemic symptoms of symptomatic diffuse cerebral vasospasm, we decided to perform conservative management. If rebleeding from the aneurysm occurred or was anticipated, surgical clipping was a treatment option as previously reported.

Our diagnostic effort by catheter angiography and contrast-enhanced vessel wall MR imaging clearly demonstrated the aneurysm of the RAH, suggesting a dissecting nature as the bleeding source of SAH. As mentioned above, RCVS and aneurysmal SAH were suspected first in this case. Khoo et al previously reported a case of RAH aneurysm masquerading as an anterior cerebral artery (A1) aneurysm radiologically and concluded that careful interpretation of the cerebral vasculature shown by angiography was important in this clinical setting. Although catheter angiography clearly demonstrated the “string and pearl” appearance along with the RAH, we still suspected the possibility of simultaneous RCVS, causing thin SAH and vasospasm, and unruptured RAH aneurysm. Notably, the vessel wall MR imaging was helpful in diagnosing the ruptured nature of this aneurysm, which showed clear contrast enhancement of the aneurysm and adjacent brain tissue. As demonstrated by Omodaka et al previously, the circumferential enhancement of the saccular cerebral aneurysm by vessel wall MR imaging could be useful in differentiating ruptured from unruptured intracranial aneurysms. Given the fusiform, “string and pearl” appearance and clear contrast-enhancement of the aneurysm, we could diagnose this aneurysm as a dissecting aneurysm as the bleeding source of the SAH in this patient.
Table 1  Summary of past reports and present case for the aneurysm of recurrent artery of Heubner

<table>
<thead>
<tr>
<th>Author, year</th>
<th>Country</th>
<th>Age</th>
<th>Sex</th>
<th>Clinical Presentation</th>
<th>Underlying condition</th>
<th>DSA findings</th>
<th>Clinical course</th>
<th>Outcome</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bechan, 2014</td>
<td>Netherlands</td>
<td>24</td>
<td>F</td>
<td>SAH</td>
<td>Moyamoya angiopathy</td>
<td>Ruptured right RAH aneurysm</td>
<td>Obliterated (Coiling)</td>
<td>No infarct</td>
</tr>
<tr>
<td>Mansfield, 2015</td>
<td>USA</td>
<td>50</td>
<td>F</td>
<td>SAH</td>
<td>Osteogenesis imperfecta</td>
<td>Fusiform left RAH 4mm-aneurysm</td>
<td>Obliterated (Clipping)</td>
<td>Infarct (caudate nucleus/Clinically silent)</td>
</tr>
<tr>
<td>Ogata, 2017</td>
<td>Japan</td>
<td>31</td>
<td>M</td>
<td>SAH</td>
<td>None</td>
<td>Fusiform right RAH &lt;5 mm aneurysm</td>
<td>Trapped (Clipping)</td>
<td>Infarct (caudate nucleus/Clinically silent)</td>
</tr>
<tr>
<td>Vellore, 2014</td>
<td>Australia</td>
<td>58</td>
<td>F</td>
<td>SAH</td>
<td>Hypertension</td>
<td>Saccular right RAH 2.7 mm aneurysm</td>
<td>Obliterated (Clipping)</td>
<td>No infarct</td>
</tr>
<tr>
<td>Wanibuchi, 2001</td>
<td>Japan</td>
<td>56</td>
<td>M</td>
<td>SAH</td>
<td>None</td>
<td>Saccular right RAH 5 mm aneurysm</td>
<td>Obliterated (Clipping)</td>
<td>N.D.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>66</td>
<td>M</td>
<td>Incidental</td>
<td>None</td>
<td>Saccular left RAH 2.5 mm aneurysm</td>
<td>Obliterated (Clipping)</td>
<td>N.D.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>65</td>
<td>M</td>
<td>Incidental</td>
<td>None</td>
<td>Saccular left RAH 2.7 mm aneurysm</td>
<td>Obliterated (Clipping)</td>
<td>N.D.</td>
</tr>
<tr>
<td>Khoo, 2019</td>
<td>Australia</td>
<td>71</td>
<td>F</td>
<td>SAH</td>
<td>None</td>
<td>Left RAH 5 mm aneurysm</td>
<td>Obliterated (Clipping)</td>
<td>Infarct (caudate nucleus/Clinically silent)</td>
</tr>
<tr>
<td>Hong, 2019</td>
<td>Korea</td>
<td>53</td>
<td>F</td>
<td>SAH</td>
<td>None</td>
<td>Left RAH 8 mm aneurysm</td>
<td>Obliterated (Clipping)</td>
<td>Infarct (caudate nucleus/Clinically silent)</td>
</tr>
<tr>
<td>Nakazaki, 2021 (present case)</td>
<td>Japan</td>
<td>58</td>
<td>M</td>
<td>Vasospasm from SAH</td>
<td>Essential thrombocytopenia</td>
<td>Fusiform left RAH 8 mm aneurysm</td>
<td>Obliteration (Spontaneous)</td>
<td>Infarct (caudate nucleus/Clinically silent)</td>
</tr>
</tbody>
</table>
Conclusion

In conclusion, we experienced a case of RAH aneurysm presenting with delayed cerebral ischemia due to diffuse cerebral vasospasm with minor SAH who was conservatively treated. Serial MR vessel wall imaging with contrast enhancement is helpful to diagnose the ruptured nature and monitor its chronological change in combination with conventional radiological imaging techniques. Long-term follow-up is mandatory because the disappearance of the vessel wall enhancement does not necessarily secure the permanent cure of the lesion.

Note
This paper was presented at Mt. Fuji workshop on CVD held at Sendai, Japan, on August 28, 2021.

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None.

Conflict of Interest
None declared.

References