Recurrent stroke from paradoxical embolism in a case with hereditary hemorrhagic telangiectasia

Rendu Osler Weber or Hereditary Hemorrhagic Telangiectesia (HHT) is an autosomal dominant genetic disorder causing abnormal development of vasculature characterized by telangiectases and arteriovenous malformations of specific organs. The most common presentation of HHT is epistaxis; however, ischemic stroke occasionally occurs due to paradoxical shunt such as pulmonary arteriovenous malformation (PAVM). We reported a 37-year-old woman presenting with recurrent ischemic stroke as a result of paradoxical emboli from PAVMs. The patient was brought to emergency department and the examination revealed left facial palsy and left hemiplegia. The CT angiography demonstrated complete occlusion of the distal right M1 MCA. Transcranial Doppler showed positive microbubble examination with suspected extracardiac shunts corresponding with positive bubble test from transesophageal echocardiogram. Pulmonary angiography demonstrated 3 PAVMs and the shunts were successfully embolized. The patient was disabling due to large volume of the right MCA infarction. (J Thai Stroke Soc. 2017; 16 (3): 49-54.)

Keywords: Rendu Osler Weber, hereditary hemorrhagic telangiectasia, pulmonary arteriovenous malformation, recurrent ischemic stroke
Rendu Osler Weber or Hereditary Hemorrhagic Telangiectesia (HHT) is a rare autosomal dominant genetic disorder causing multiple vascular dysplasia such as telangiectasia on mucosal area. Patients may suffer from several manifestations resulting from vascular defects including nose bleeding (epistaxis) which is the most common symptom, and gastrointestinal bleeding.\(^1\) Ischemic stroke is one of vascular manifestations accounting of 15%\(^2\) caused by paradoxical emboli via shunting from extracardiac arteriovenous malformation in lungs, liver, or pancreas.\(^3\) Clots in the venous system are able to bypass the pulmonary capillary system through those shunts and result in embolism into terminal organs such as cerebral hemispheres.

Extracardiac or intracardiac shunts can be effectively detected by contrast-enhanced transcranial Doppler study (TCD), and the TCD gains as high as 100% sensitivity comparable with transthoracic contrast echocardiography (TTCE)\(^4,5\). Microbubbles in agitated saline are frequently used as contrast media and those microemboli give rise to distinctive characterized Doppler signals in both examinations.

**Case Report**

A 37-year-old Thai woman presented with sudden left hemiparesis while engaging leisure activity. She had two episodes of non-disabling ischemic stroke. In the first attack, 6 year earlier, the patient presented with acute dysarthria and weakness on her right arm. Diagnostic work up for young onset ischemic stroke was done including transthoracic echocardiogram, Electrocardiography, VDRL, TPHA, Anti-HIV, thrombophilia, and inflammatory vasculitis work up. All of the test results appeared normal. The MRI/MRA scan demonstrated a small segmental caliber of the basilar artery with bilateral thalamic infarction. The diameter of the basilar artery was later recovered, therefore reversible cerebral vasoconstriction syndrome was initially suspected. After the ischemic attack, she was then given an antiplatelet for secondary prophylaxis. 4 years later, she experienced acute right facial palsy with a complaint of right sided hemiparesthesia. She was diagnosed with recurrent ischemic stroke and her deficits eventually improved to normal state within a few months.

One day earlier, the patient had bleeding per nose in the morning concurrently with bilateral tensed headache.

On this admission, the patient abruptly experienced dysarthria and left sided hemiparesis with facial palsy approximately 40 minutes earlier. At emergency department, she became drowsy. Physical examination revealed bleeding per nose with suspected telangiectasia at nasal septum, tongue and both lips. Left hemiplegia was observed and her NIHSS score was 13. A large right middle cerebral artery (MCA) infarction was preliminary suspected. The CT scan showed a dense MCA sign on the right MCA and old infarctions on both thalami owing to previous stroke.

![Image](https://example.com/image.png)

**Figure 1:** Axial non contrast CT brain: dense MCA signs (cord sign) on the right MCA (arrow).
Intravenous thrombolysis was not given because of active recurrent epistaxis within one month. Following CT angiography demonstrated an obvious distal right M1 MCA occlusion.

Mechanical thrombectomy was undergone and the procedure achieved recanalization. The decompressive craniectomy was done to treat severe cerebral swelling and she later received bed-side rehabilitation. Causes of stroke in the young were then re-evaluated. Results from all blood tests for hypercoagulable states and inflammatory vasculopathy were normal.

**Figure 2:** DSA: left image showed a total occlusion of the distal right M1 MCA and right image showed recanalization after an endovascular therapy.

Transcranial Doppler was done and the study revealed positive bubble test at a period of time longer than 10 cardiac cycles after agitated saline was administered.

**Figure 3:** TCD study demonstrated time of bubble injection (yellow arrow), time of valsava initiation (red arrow) and embolic signals appeared at a time longer than 10 cardiac cycles after injection.
Regarding previous study by using echocardiography in detecting right-to-left shunt in HTT, if the bubbles contrast were detected within 3 cardiac cycles after agitated saline injection, intracardiac shunt should be considered. On the other hand, if the bubbles were detected following 8 cardiac cycles or longer time after the contrasted injection, extracardiac shunt should be suspected.\(^5\)

Transoesophageal echocardiogram was done and similarly confirmed a positive bubble test. Extracardiac shunt was then highly suspected.

*Figure 4:* transesophageal echocardiogram revealed numerous microbubbles in the left ventricle.

The interventionist was consulted for angiographic study of the thoracic. The pulmonary angiography revealed 3 pulmonary arteriovenous malformations at left lower lung in which were later successfully embolized by the interventionist.

*The patient symptoms remained stable.* She was disabling due to large left MCA infarction. The antiplatelet was continued and she has not been suffering another recurrent episode until present.

*Figure 5:* Left, CT angiogram of chest demonstrated abnormal dilated vascular structure in the left basal lung. Right: Angiographic study showed arteriovenous malformations in the left lung.
Discussion

Diagnosis of hereditary hemorrhagic telangiectasia is challenging and essential. Our patient had experienced acute ischemic events prior to this visit. She had not been diagnosed of HHT until current visit which eventually resulted in more severe disability. Routine worked up for routine unusual cause in young onset of stroke should be strictly monitored, for example, bubble or contrasted echocardiography or bubble transcranial Doppler study. To diagnose Osler Weber Rendu disease, we use Curacao clinical criteria which are as follows:

1. Epistaxis
2. Telangiectasias—characteristic sites include lips, oral cavity, fingers and nose;
3. Visceral lesions including gastrointestinal, hepatic, PAVMs, cerebral AVMs (CAVMs) or spinal AVMs
4. Family history with a first-degree relative with HHT

Presence of three or more clinical criteria makes a definite diagnosis while two criteria is a possible diagnosis.

Our patient had three out of four criteria: recurrent epistaxis, mucosal telangiectasia and established pulmonary AVMs, whereas family history could not be confirmed. Though, epistaxis is the most common presentation affecting about 85–90% of patients. History of bleeding nose had not been documented during the first 2 visits of ischemic events in our patient. In study concerning delayed in diagnosis of HHT, an ischemic stroke was found as the onset presentation of disease in only 3/233 (1.3%) while epistaxis alone occurred 202/233 (86.7%).

Consideration is given to screening for PAVMs in all patients with HHT who present with neurological manifestation. Therefore, Bubble study in echocardiogram or transcranial Doppler ultrasound should be performed to detect right to left shunting in such patient.

Conclusion

Transcranial Doppler with bubble study is useful in detecting right to left shunting and the examination helps distinguish intracardiac shunt and extracardiac shunt with high sensitivity comparable to bubble echocardiography.

Reference


บทคัดย่อ
Rendu Osler Weber หรือ Hereditary Hemorrhagic Telangiectesia (HHT) เป็นโรคทางพันธุกรรมที่ส่งต่อแบบยีนเด่น ทำให้เกิดความผิดปกติในการพัฒนาหลอดเลือดและทำให้เกิดหลอดเลือดโป่ง และการสร้างการเชื่อมต่ออัตติกติของหลอดเลือดแดงและดำในอวัยวะต่างๆ อาทิเช่น ปอด ตับ หรือตัวอ่อน อาการแรกที่พบได้บ่อยที่สุดคือเลือดก่าพาแตก การที่มีการเชื่อมต่อของหลอดเลือดแดง และการเกิดกังวลโดยเฉพาะในปอด ทำให้เกิดกังวลที่ผ่านเข้าไปยังหัวใจและทำให้เกิดหลอดเลือดสมองอุดตัน โดยหากไม่ได้รับการแก้ไขทำให้เกิดหลอดเลือดสมองอุดตันกลับเป็นซ้

ผู้นิพนธ์รายงานผู้ป่วยหญิง 37 ปี ที่เป็นโรคหลอดเลือดสมองแบบกลับเป็นซ้ ในครั้งที่สามอาการ เลือดก่าพาแตก 1 เดือนเป็นๆหายๆ ร่วมกับอาการอ่อนแรงแขนขาข้างซ้าย เลือดผิด และอุดตันขวา จากผลตรวจทางรังสีการอุดตันส่วนปลายของหลอดเลือดสมอง MCA ข้างขวาและได้รับ การรักษาด้วยการหยุดหลอดเลือด การตรวจหลอดเลือดสมองด้วยคลื่นเสียงความถี่สูงร่วมกับภาพอากาศ ช่วยวินิจฉัยสาเหตุการเชื่อมต่อของเส้นเลือดแดงและด้านปลายน้ำอาจเกิดขึ้น ผลการตรวจหลอด เลือดทางรังสีการเชื่อมต่อของเส้นเลือดแดงและด้านปลายน้ำอาจเกิดขึ้นได้ เวลาที่มีการอุดเส้นเลือดสมองกลับเป็นซ้

คำสำคัญ: Rendu Osler Weber, hereditary hemorrhagic telangiectasia, pulmonary arteriovenous malformation, recurrent ischemic stroke, TCD bubble test, right to left shunt