A new ENG mutation in a Japanese family with hereditary hemorrhagic telangiectasia and pulmonary arteriovenous malformations

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ABSTRACT

We present a case series of four siblings with hereditary hemorrhagic telangiectasia (HHT) and pulmonary arteriovenous malformations (PAVM). The patients' mother has HHT. Case 1: A 22-year-old man developed dyspnea and epistaxis. CT revealed a large PAVM, treated by segmentectomy. Case 2: A 27-year-old woman developed epistaxis and dyspnea. CT revealed three PAVMs, treated by partial resection. Case 3: A 20-year-old woman developed dyspnea. CT revealed multiple PAVMs, treated with endovascular occlusion of the largest one. Case 4: A 12-year-old woman developed epistaxis. CT revealed multiple PAVMs, observed without treatment. Genetic testing identified a new mutation, ENG c.1517T > C (p.Leu506Pro), in all patients and their mother. We suspect that HHT in these patients may be associated with this ENG mutation.

1. Introduction

Hereditary hemorrhagic telangiectasia (HHT) or Osler-Weber-Rendu disease, is an autosomal dominant disease characterized by mucocutaneous telangiectasias, epistaxis and visceral arteriovenous malformations (AVM) most commonly found in the lungs, liver, and brain [1,2]. In Japan, the incidence of HHT is estimated to be 1:5000 to 1:8000 [3], which is similar to reports from other countries [4–6]. HHT is clinically diagnosed based on the Curaçao criteria: 1) an affected first-degree family member, 2) recurrent epistaxis, 3) multiple telangiectasia along the mucocutaneous surface, and 4) arteriovenous malformations in major organs. The diagnosis is considered “confirmed” in an individual with at least three, and “suspected” with two of the above features [4]. These features progress with age and pulmonary arteriovenous malformation (PAVM) development is thought to be complete by the end of puberty [7,8].

Recently, genetic research has demonstrated that heterozygous mutations including ENG, ACVR1I, and rarely SMAD4 are causative genes of HHT. There are at least two other unidentified genes that can cause HHT [7,8]. The majority of HHT patients have mutations in ENG, encoding endoglin, or ACVR1I, encoding activin receptor like kinase. These genes are associated with the transforming growth factor (TGF)-β superfamily signaling pathway, which is important for maintaining vascular integrity [5–7]. HHT with ENG mutation is characterized by a high incidence of PAVMs and cerebral AVMs [9], whereas HHT with ACVR1I is associated with hepatic AVMs. It is thought that clinical profiles have some correlation with the genotype.

Here we report four patients with familial HHT with PAVM associated with a new ENG mutation.

2. Case reports

In August 2015, a 48-year-old woman with HHT visited Sapporo Medical University Hospital for clinical genetic counseling with her four children, a son (Case 1) and three daughters (Case 2, 3, and 4). She had undergone left lower lobe lobectomy of a PAVM in the past.
A 22-year-old man complained of dyspnea and recurrent epistaxis. He had a history of bronchial asthma, attention deficit hyperactivity disorder, learning disabilities and dysgraphia. He had telangiectasias in his fingers and lips, clubbing, and expiratory wheezing. A vascular murmur (bruit) was auscultated in his right lower back. Arterial blood gas showed a partial pressure of arterial oxygen (PaO2) of 51.4 mmHg and a SpO2 of 83% on room air. Gas exchange showed a PaO2 of 93.8 mmHg and a SpO2 of 95% on room air. Chest CT revealed multi PAVMs in the right S4, the right S6 and the left S4 (Fig. 4) and they were 5 mm, 7 mm and 5 mm in size, respectively. They were of the simple type and supplied by feeding arteries of 3–4 mm in diameter. A brain MRI, an abdominal CT and a gastrointestinal endoscopy showed no abnormal finding. Percutaneous embolization of the feeding artery was thought to be difficult because the PAVM was very large and both the feeding artery and the draining vein were thick. There was a higher risk of paradoxical embolization such as migration of occlusive devices or blood clots. Therefore, we treated the PAVM by segmentectomy with video-assisted thoracoscopic surgery (VATS). After the surgery, the shunt fraction decreased to 14.2% and SpO2 rose to 94% on room air.

Chest CT taken 16 months post-operatively revealed a new asymptomatic small PAVM, 3 mm in diameter, in the right upper lobe. As the feeding artery was less than 3 mm in diameter and the patient had no symptoms, we recommended careful observation.

2.2. Case 2

A 27-year-old woman complained of recurrent epistaxis and mild dyspnea on exertion. She had history of bronchial asthma and irritable bowel syndrome. She had telangiectasias in fingers and lips. Auscultation was normal. Arterial blood gas analysis showed a PaO2 of 95.5 mmHg and SpO2 of 98% on room air. Chest CT revealed multiple PAVMs in the right S4, the right S6 and the left S4 (Fig. 4) and they were 5 mm, 7 mm and 5 mm in size, respectively. They were of the simple type and supplied by feeding arteries of 3–4 mm in diameter. A brain MRI, an abdominal CT and a gastrointestinal endoscopy showed no abnormal finding. Percutaneous embolization of the feeding artery was not necessary for transcatheter embolization and due to the risk of anaphylaxis and cardio-pulmonary failure, we performed surgical removal of three PAVMs via VATS instead. Moreover, three additional PAVMs were discovered on the surface of the visceral pleura of the right S3, S8 and S10 by direct observation during the operation. These PAVMs on the pleural surface were excised by electroablation using the SOFT COAG electrosurgical output system.

2.3. Case 3

A 20-year-old woman complained of dyspnea on exertion. She had history of idiopathic thrombocytopenic purpura and bronchial asthma as well as autistic spectrum disorder, depression, and intellectual impairment. She had given birth twice. Mucosal and skin examinations showed no abnormality. Auscultation was normal. Arterial blood gas demonstrated a PaO2 of 93.8 mmHg and a SpO2 of 95% on room air. Chest radiograph showed well-defined nodular lesions in the right lower lung field. Chest CT revealed multiple PAVMs in both lungs (Fig. 5). The largest one was 18 mm in diameter, it was located in right S10, and had two feeding pulmonary arteries of 3 and 5 mm in diameter. Percutaneous embolization of the feeding artery was not necessary for transcatheter embolization and due to the risk of anaphylaxis and cardio-pulmonary failure, we performed surgical removal of three PAVMs via VATS instead. Moreover, three additional PAVMs were discovered on the surface of the visceral pleura of the right S3, S8 and S10 by direct observation during the operation. These PAVMs on the pleural surface were excised by electroablation using the SOFT COAG electrosurgical output system.

2.4. Case 4

A 12-year-old woman complained of recurrent epistaxis. She had history of mental and adjustment disorders. Physical examination revealed telangiectasias in fingers and lips. Chest CT revealed small diffuse PAVMs with a feeding artery of less than 3 mm in diameter in both lungs (Fig. 6). Enhanced abdominal CT showed no abnormal finding. Due of her psychological instability, she was unable to undergo brain MRI. Since the size of the PAVMs was small and the diameter of the blood vessel was 3 mm or less, we managed her conservatively. Laser ablation treatment for nasal mucosal telangiectasia reduced the
episodes of epistaxis.

3. Discussion

We report a family with HHT with PAVMs associated with a new ENG mutation. Although more than a hundred mutations in the ENG gene have been reported related to HHT [6], there is no report of HHT caused by the ENG c. 1517T > C (p. Leu506Pro). To our knowledge, this is the first reported case. We would have been able to confirm whether this mutation was causative if there was an unaffected sibling without the mutation. Further research is needed in order to determine precisely if this mutation can cause HHT. In addition, their clinical profiles showed all of our patients had PAVM and three had psychiatric or developmental disorders. Recently, it was reported that pediatric patients with HHT had a relatively high prevalence of malformations of cortical development [10]. There may be a relationship between the observed psychiatric and developmental disorders and this new ENG mutation.

Causative genes of HHT including ENG, ACVRL1 and SMAD4 are involved in the TGF-β pathway [11,12]. The TGF-β superfamily signaling pathway has a crucial role in regulating proliferation, differentiation, and migration in angiogenesis. HHT gene mutations may negatively influence some forms of angiogenesis, with specific effects on the stability of newly formed vascular sprouts [7]. Endoglin, a membrane protein expressed in the endothelial cells, works as an endothelial specific co-receptor for multiple receptor complexes of the TGF-β superfamily. It has been hypothesized that the mutated proteins may be misfolded and unstable, and fail to reach the cell surface, rendering them unstable to form heterodimers with normal endoglin [6]. ENG mutations causing endoglin haploinsufficiency are thought to impair recruitment of mural cells to vessels [7].

Currently, transcatheter embolization with vascular coils and occluders (Amplatzer™) are the standard treatment for PAVMs [12]. Treatment of PAVM with transcatheter embolization of the feeding vessels significantly decreases right-to-left shunting, leading to

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Table 1
Summary of four cases.

| Case | Age | Sex | Criteria | Symptom | PAVM Type | Location | Size (mm) | Treatment | Past history/Complications |
|------|-----|-----|----------|---------|-----------|----------|-----------|-----------|---------------------------|
| 1    | 22  | M   | confirmed| Dyspnea | Simple    | Right S10| 60        | Surgery   | BA, ADHD, LD, Dysgraphia |
| 2    | 27  | F   | confirmed| Dyspnea | Diffuse   | Right S4, S6, Left S4 | 5, 7, 5   | Surgery   | BA, Irritable bowel syndrome |
| 3    | 20  | F   | suggested| Dyspnea | Diffuse   | Both lungs | 2-18      | TE        | BA, ITP, ASD, Mental retardation, Depression, Mental intellectual disorder |
| 4    | 12  | F   | confirmed| Epistaxis| Diffuse   | Both lungs | 2.8-6     | none      | Adjustment disorder, Mental disorder |

BA: bronchial asthma, ADHD: attention deficit hyperactivity disorder, LD: learning disabilities, ITP: idiopathic thrombocytopenic purpura, ASD: autistic spectrum disorder, TE: transcatheter embolization.

Fig. 3. Case 1. Three-dimensional computed tomography (3D-CT) angiography reveals a large pulmonary arteriovenous malformation (PAVM, arrow) located in the subpleural area of the right segment (S) 10. The size is 60 mm in diameter and feeding pulmonary artery is 10mm in diameter.

Fig. 4. Case 2. 3D-CT angiography reveals PAVMs (arrows) located in right middle and lower lobe, and left upper lobe. They are located in right S4, S6, and left S4. The sizes of them are 5 mm, 7 mm and 5 mm in diameter, respectively. They are simple type and supplied by a feeding artery of 3-4 mm in diameter.
improved oxygenation. In our cases, we performed this procedure for Case 1 and Case 2, and embolotherapy for Case 3. In Case 1, because of a large PAVM with thick feeding artery and draining vein which had presumably high flow of blood within the PAVM, endovascular treatment of the large PAVM was thought to have a higher risk of complications such as device embolism or thrombus passing through the PAVM. Akiyama et al. [13] reported a surgically treated case of PAVM who had turbulent thrombi in one of the PAVMs, which was capable of inducing embolic strokes such as symptomatic paradoxical thrombosis. Therefore, segmentectomy was thought to be preferable to transcatheter embolization in Case 1 for preventing severe complications with embolotherapy [13,14]. In Case 2, surgery was performed because embolotherapy was not feasible due to allergy to contrast medium. On the other hand, there is no agreement regarding the treatment of PAVM in pediatric patients [15]. A significant concern in treating growing lungs is that treated PAVMs may be at increased risk for reperfusion via pulmonary collaterals. Therefore, we determined to follow up Case 4 who was a 12-year-old child with asymptomatic multiple small PAVM. On the other hand, Case 1 showed recurrence and progression of PAVMs after treatment. The manifestations of progressing HHT and PAVM may take various forms with increasing age. Long-term follow-up is necessary for HHT patients even after AVM treatment.

Genetic testing is not always essential to diagnose of HHT, however, this knowledge might be useful for a patient with suggestive clinical features of HHT. In addition, when the affected parent has the mutation, the child without the mutation need not to undergo further screening test for AVM. In our cases, Case 3 was thought to be a pre-symptomatic mutation carrier. In this report, we presented 4 cases of HHT in a single family, in which the clinical diagnosis was confirmed using genetic testing, providing an opportunity for early detection of complications and management of the HHT.

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Fig. 5. Case 3. 3D-CT angiography reveals multiple PAVMs located in bilateral peripheral lung. The largest PAVM (arrow) with complex type is located in right S10.

Fig. 6. Case 4. Chest CT shows three PAVMs (arrows) in right upper lobe (A), lower lobe (B) and left lower lobe.

Conflicts of interest
The authors state that they have no conflict of interest.

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