A Patient with Cavernous Sinus Dural Arteriovenous Fistula Complicating Klippel-Trenaunay Syndrome

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Objective: We report a case of cavernous sinus dural arteriovenous fistula (CSdAVF) associated with Klippel-Trenaunay syndrome (KTS).

Case Presentation: A 58-year-old woman was diagnosed with KTS with port-wine stain, overgrowth of tissues and bones, and venous malformation on the left upper limb. She was admitted to our hospital with the primary complain of ptosis and diplopia due to left oculomotor palsy, and her cerebral angiography revealed CSdAVF with retrograde leptomeningeal venous drainage. The shunt point was located at the posteromedial part of the left cavernous sinus (CS) on the angiogram. An enlarged subclavian vein and giant varix was detected in the left upper limb, and abnormality of the coagulation-fibrinolysis system caused by localized intravascular coagulation was confirmed. We performed transvenous coil embolization, and the symptoms improved after a few weeks with no new neurological deficits. However, the activation of coagulation-fibrinolysis system continued even after the surgery. Conclusion: The formation of dAVF occurrence in this case is unclear. If genetic abnormalities that cause angiogenesis are involved in KTS, follow-up is important in the future.

Keywords ▶ Klippel-Trenaunay syndrome, dural arteriovenous fistula, transvenous embolization, venous malformation, coagulopathy

Introduction

Dural arteriovenous fistula (dAVF) is a disease with acquired shunt formation in the dura, and the annual incidence is reported to be 0.16/100000 people overseas1) and 0.29/100000 people in Japan.2) Inducers of dAVF include head injury and infection, past medical history of craniotomy, venous thrombosis due to abnormal coagulation, and venous hypertension, but the cause is unclear in many cases.

In the neurosurgery field, hemangioma on the body surface and in soft tissue and vascular malformation are rarely encountered. Multiple or very large vascular malformations are resistant to various treatments and life-long treatment is necessary due to pain, ulcer formation, abnormal growth of the affected limb, and problems with the appearance. We encountered a patient with dAVF complicating Klippel-Trenaunay syndrome (KTS), which is classified as vascular malformation-related syndrome.

Case Presentation

Chief complaints: Left ptosis and diplopia.
Past medical history: Port-wine stains in the left upper limb and left precordium have been pointed out since birth, a varix was noted in the left upper limb as a juvenile, and the patient was diagnosed with KTS. In addition, the left pointing finger was amputated due to persistent hemorrhage, although the details were unclear. The patient regularly visited the hospital for venous hemorrhage from peripheral ulcer in the left upper limb and received sclerotherapy from 1 year earlier, but she had no previous medical history.
of head injury and malignant disease or treatment with an oral antithrombotic or anticoagulant.

History of present illness: The disease suddenly manifested as left ptosis and diplopia. The patient visited a physician and was referred to our hospital, suspecting an intracranial lesion. On the first examination, consciousness was clear, left conjunctival hyperemia, left oculomotor paresis, and left pulsatile tinnitus were observed, the affected left upper limb was entirely swollen, and a dilated varix was present (Fig. 1). On close examination, Borden type II cavernous sinus (CS) dAVF was observed, in which blood flows from the left middle meningeal artery, accessory meningeal artery, artery of foramen rotundum, and ascending pharyngeal artery into the medial posterior part of the left CS and flowed out into the left superior ophthalmic vein (SOV), superficial middle cerebral vein (SMCV), inferior petrosal sinus (IPS), and pterygoid plexus, and it was accompanied by cortical venous reflux. In addition, the shunt also flowed into the lateral CS. In the delayed phase of contrast-enhanced CT of the trunk, a giant varix was present in the left upper limb and vascular dilatation expanded to the left subclavian vein, but the left internal jugular vein was not dilated (Fig. 2). Many phleboliths were present in the left upper limb, but no filling defect suggesting thrombus was present in the varix (Fig. 3). On
blood testing, the platelet count was 10.6 × 10^4/μL; fibrinogen, 104.6 mg/dL; fibrin degradation product (FDP), 39.1 μg/mL; D-dimer, 20.5 μg/mL; showing abnormality of the coagulation-fibrinolysis system (Table 1). For treatment, transvenous coil embolization (TVE) with heparinization only during treatment without pre- or postoperative anti-thrombotic therapy was planned. Treatment course: The right common femoral artery was punctured and a catheter for diagnosis was placed in the left external carotid artery. The right femoral vein was punctured and 6Fr Launcher (Medtronic, Minneapolis, MN, USA) was placed in the left internal jugular vein and advanced to the CS through the IPS using 4Fr BHW (Katecs, Aichi, Japan) and Radifocus Guidewire 0.035 (Terumo, Tokyo, Japan). Using this as an index, SL-10 (Stryker, Kalamazoo, MI, USA) was guided to the CS through the IPS as described above. Reflux to the SOV had resolved but reflux to the SMCV had aggravated due to re-canalization of the region embolized in the first treatment. The region of the SMCV embolized in the first treatment over the proximal region was embolized, followed by additional embolization of the postomedial compartment of the CS, and the shunt disappeared (Fig. 5). Pulsatile tinnitus disappeared immediately after treatment, and oculomotor paralysis completely resolved 3 weeks after the second treatment. No recurrence has occurred for 3 years thereafter.

### Discussion

KTS is a syndrome of congenital abnormal vasculogenesis developing port-wine stains in the four limbs, varix, and bone soft tissue overgrowth as the three main symptoms, and it develops in the unilateral lower limb in 75% of cases. In the classification established by the International Society for the Study of Vascular Anomalies (ISSVA) classification, simple vascular malformation is classified into capillary blood vessel malformation, lymph vessel malformation, venous malformation, arteriovenous malformation (AVM), and arteriovenous fistula, and in the mixed type, several vascular malformations are present. KTS is complicated by capillary blood vessel malformation, lymph vessel malformation, and venous malformation in many cases and enlargement of bone soft tissue accompanies as a concomitant symptom so that it is classified as a related syndrome in the ISSVA classification. It is considered that KTS, in which the flow rate is low, should be distinguished from Parkes Weber syndrome (PWS) complicated with high-flow-rate arteriovenous fistula, hyperplasia in the affected limb, and heart failure in severe cases. However, both syndromes were collectively reported as Klippel-Trenaunay Weber Syndrome (KTWS) in many previous studies.

KTS-associated complication by vascular lesions in the central nervous system has been reported. In all, 12 cases...
of KTS or KTWS complicated by cerebral aneurysm have been reported and one or more giant cerebral aneurysms were present in three cases of them.4–6) KTS or KTWS was complicated by spinal A VM or fistula in 33 cases7–11) and it tended to develop in the lower thoracic over the lumbar spinal cord. In addition, involvement of the Angiogenic factor with G-patch and FHA domain 1 (AGGF1) and phosphoinositide-3-kinase, catalytic, alpha polypeptide (PIK3CA) genes in KTS has been reported.12,13) Overexpression of the AAGF1 gene promotes expression of flt4, dab2, and EPH receptor B4 (EPHB4), which are venous markers, and it activates protein kinase B (AKT) playing
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an important role in cell proliferation, survival, and metabolism, through which venous angiogenesis occurs and leads to vascular malformation. Regarding the PIK3CA gene, when activating mutation occurs in the developmental process, mammalian target of rapamycin having an influence on vascular growth and proliferation is excessively activated and causes angiogenesis and tissue overgrowth. Reduction of vascular lesions and tissue overgrowth in PIK3CA-related overgrowth syndrome by administration of a molecular target drug inhibiting this PIK3CA gene has been reported.14) Sgubin et al.9) reported that KTS may be complicated by spinal AVM due to the influence of angiogenic factor. Angiogenesis due to these gene abnormalities may have contributed to CSDaVF formation in the present patient. On the other hand, Alomari et al.15) denied the association between KTS and spinal AVM. They reviewed cases diagnosed as KTS in previous reports on the association with spinal AVM and found that the cases were actually other similar diseases, such as Congenital lipomatous overgrowth, vascular malformations, and epidermal nevi (CLOVE) syndrome and capillary malformation-AVM (CM-AVM), suggesting difficulty in definitely diagnosing KTS. Gene abnormality causing vascular malformation has been confirmed in several diseases and the association of PWS, which is a KTS-related disease, with RAS p21 protein activator 1 (RASA1) gene abnormality has been reported,16 and involvement of RASA1 and EPHB4 mutations in CM-AVM has also been reported.17) CM-AVM was complicated by AVM/fistula in the brain, face, and four limbs in 30% and when a genetic test was performed in five patients with CM-AVM complicated by spinal AVM/fistula, RASA1 gene abnormality was detected in all five patients in a study.18)

To our knowledge, this was the initial case report of KTS with intracranial dAVF. In this patient, the left subclavian
vein dilated to three times the thickness of the vein on the healthy side accompanying the very large venous malformation in the left upper limb, and phleboliths were also present. Phlebolith is calcified thrombus formed in a vascular lumen with a slow flow velocity and it is considered a characteristic finding of venous malformation. Abnormal coagulation occurs when the volume of the lesion is large and when multiple phleboliths are present. First, we discuss the relationship between dilatation of the left subclavian vein and development of CsdAVF. dAVF development after diagnosing sinus occlusion has been reported, but in our patient, although stenosis of the left IPS was observed at the time of diagnosis, no sinus occlusion inducing dAVF or preceding symptom suggesting its presence was noted. It has been reported that in CsdAVF, the sinus is gradually thrombosed by shunt blood flow, and the posterior outflow path and then anterior outflow path is occluded with progression of the disease stage. Applying this to our patient, the disease stage was still early so that the condition was still in the process leading to occlusion of the posterior outflow path. Congestion of the left internal jugular vein was also absent on angiography and no phlebolith suggesting thrombosing was noted. Therefore, it is less likely that dAVF was induced by perfusion injury caused by dilatation of the left subclavian vein. Second, we discuss the relationship between abnormal coagulation and CsdAVF formation in the present patient. The blood test findings before and after treatment are shown in Table 1. A mild decrease in platelets and increases in FDP and D-dimer were noted 1 year before the onset of CsdAVF. FDP and D-dimer markedly increased and fibrinogen decreased immediately after the first TVE and these may have reflected coil-induced thrombosing. The values of the coagulation-fibrinolysis system including TAT and PIC were still abnormal at 3 years after treatment. This abnormal coagulation in this patient was localized intravascular coagulation (LIC) caused by massive consumption of coagulation factor in lesions due to chronic retention of blood in the varix of the affected limb, and this has to be distinguished from Kasabach-Merritt syndrome complicating infantile and pediatric Kaposiform hemangioendothelioma and tufted hemangioma. In addition, local abnormal coagulation may spread to other regions due to sclerotherapy, surgical treatment, fracture, and menstruation and LIC may progress to disseminated intravascular coagulation (DIC). In our patient, thrombosing did not progress in CsdAVF and no finding of DIC was noted. Therefore, it is less likely that abnormal coagulation in pre-existing LIC-induced CsdAVF. The efficacy of perioperative administration of low-molecular-weight heparin to reduce LIC-induced local pain and prevent aggravation to DIC in surgery and sclerotherapy for venous malformation complicated by LIC has been reported. Catheter treatment-induced progression of LIC to DIC has not been reported, but it would have been good to consider perioperative administration of low-molecular-weight heparin although there is a problem with coverage of TVE for CsdAVF with national health insurance.

No gene search was performed in this patient and the cause of CsdAVF formation was unclear. Since involvement of gene abnormality inducing angiogenesis in KTS has been reported, its influence on dAVF formation cannot be ruled out. Three years have passed without recurrence after treatment, but continuation of careful follow-up is necessary.

### Conclusion

We performed endovascular treatment in a rare case of CsdAVF complicating KTS and achieved a favorable outcome. Many points are unclear with regard to the cause of dAVF, for which further elucidation of the pathology is expected.

### Disclosure Statement

None of the first and coauthors has conflicts of interest.

### References


