sinus stenosis as a possible mechanism. However, when evaluated with an MRV, over 2/3 of these patients demonstrated asymmetric venous sinus or frank stenosis. The referral of such patients to neurointervention was rare. Our study highlights the overall under-consideration of transverse venous sinus asymmetry/stenosis as a treatable entity and stresses the importance of obtaining an MRV and referring patients for neurointerventional evaluation.

Disclosures P. Ramakrishnan: 3; C; Cerenovus. C. Berry: None. S. Leinfelder: None. W. Leech: 3; C; Penumbra Inc. 4; C; Cerebrotech Inc. 6; C; Stryker, Medtronic, Terumo. J. Sanderson: None.

E-062 SUBDURAL CONTRAST EFFUSION DURING ENDOVASCULAR THERAPY

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Background Accumulation of contrast medium in the subdural space after diagnostic intraarterial and intravenous contrast administrations is a rare observation. The authors report the case of a subdural contrast effusion (SCE) mimicking an acute subdural hematoma (SDH) presenting during embolization of an intracranial dural arteriovenous fistula (DAVF).

Clinical history A 52-year-old woman was admitted to our department with a four months history of left-sided numbness of the upper lip, cheek, tongue and forehead. Digital subtraction angiography showed a right tentorial DAVF with a venous ectasia. The DAVF was mainly supplied by the right middle meningeal artery, occipital artery and the artery of Ber- nasconi-Cassinari and drained into the Galenic venous system.

Procedure The patient underwent endovascular treatment with triaxial catheterization of the right occipital artery. Transarterial embolization using PHIL was performed and monitored by control runs made with an intermediate (distal access) catheter. After several injections, increased accumulation of contrast medium along the cerebellar tentorium and the walls of both transverse sinuses was noted. An immediately performed Dyna CT showed extensive contrast medium in the subdural space of the posterior cranial fossa and foramen magnum suspicious for an acute SDH. The catheters were removed, and the procedure was terminated. The patient woke up presenting no new symptoms and had an uneventful postoperative course. A 24-hour follow-up Dyna CT was completely normal suggesting the diagnosis of an asymptomatic SCE. We believe that repeat high pressure contrast injections via a large bore intermediate catheter into the territory of a (even partly) occluded DAVF may have induced leakage of contrast medium into the extravascular subdural space thereby causing a SCE.

Summary In conclusion we present the unique case of an asymptomatic SCE, which developed during transarterial embolization of a DAVF. SCE can occur during endovascular therapy and may mimic an acute SDH. Differentiation between the two by computed tomography or Dyna CT and early neurological examination can be crucial for patient management.

Disclosures R. Dahl: None. V. Eskesen: None. G. Benndorf: None.

E-063 RECURRENT SYNCOPE CAUSED BY A DURAL ARTERIOVENOUS FISTULA: A CASE REPORT

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Introduction Dural arteriovenous fistulas (DAVFs) are pathologic vascular connections that shunt dural arterial flow directly to dural venous drainage. DAVFs constitute 10–15% of all intracranial arteriovenous malformations and are most commonly located in transverse, sigmoid, and cavernous sinuses. Symptoms from DAVFs vary based on lesion location and correlate with their respective patterns of venous drainage. Common presenting symptoms include pulsatile tinnitus, chronic headache, vision difficulties, cranial nerve abnormalities, cerebellar symptoms, focal neurologic deficits, and seizures. Only a few isolated case reports describe syncope on presentation, all in combination with other symptoms. In this article, we report a rare case of DAVF causing recurrent syncope.

Methods A 29-year-old female presented with a 9-year history of progressive syncopal episodes exacerbated by positional changes, strenuous activity and emotional stressors.

Results The patient was referred to us after symptoms of dizziness and syncope persisted despite treatment by multiple cardiologists, endocrinologists, and psychiatrists. Symptoms occurred upon waking and lasted for 2–3 hours before she was able to regain functionality. Furthermore, she found that her symptoms would remit if she was to lay in the Trendelenburg position immediately after symptom onset. Physical exam revealed no abnormalities. MRI of the brain showed no irregularities. MRA revealed abnormal serpiginous structures in the left jugular foramen which communicated with the ascending pharyngeal branch of the left external carotid artery. Cerebral angiogram disclosed a left jugular foramen dural AVF (Borden/Cognard type 1) supplied by the left ascending pharyngeal and left occipital arteries. There was no retrograde flow or cortical venous reflux. The DAVF was successfully managed by progressive endovascular embolization with coils and Onyx 34 via transvenous route. The final cerebral angiogram demonstrated complete obliteration of the fistula. On clinical follow-up evaluation, the patient had no further episodes of dizziness or syncope.

Conclusion We present an atypical case of DAVF in a patient presenting with recurrent syncope. Only 3 cases of DAVF causing syncope have been reported, all in combination with other neurologic symptoms. In comparison, we report a unique case of DAVF presenting solely with recurrent syncope, a previously undocumented finding in the literature. Our case adds to other reports of nonspecific DAVF presentations and highlights the importance of considering this etiology.

Disclosures D. Sheinberg: None. E. Luther: None. D. McCarthy: None. R. Starke: None.