Perimedullary arteriovenous fistulas of the craniovertebral junction: A systematic review

ABSTRACT

Perimedullary arteriovenous fistulas (PMAVFs) are uncommon vascular malformations, and they rarely occur at the level of the craniovertebral junction (CVJ). The therapeutic management is challenging and can include observation alone, endovascular occlusion, or surgical exclusion, depending on both patient and malformation characteristics. A systematic literature search was conducted using MEDLINE, Scopus, and Google Scholar databases, searching for the following combined MeSH terms: (perimedullary arteriovenous fistula OR dural arteriovenous shunt) AND (cranio cervical junction OR craniovertebral junction). We also present an emblematic case of PMAVF at the level of the craniovertebral junction associated to a venous pseudoaneurysm. A total of 31 published studies were identified; 10 were rejected from our review because they did not match our inclusion criteria. Our case was not included in the systematic review. We selected 21 studies for this systematic review with a total of 58 patients, including 20 females (34.5%) and 38 males (65.5%), with a female/male ratio of 1:1.9. Thirty-nine out of 58 patients underwent surgical treatment (67.2%), 15 out of 58 patients were treated with endovascular approach (25.8%), 3 out of 58 patients underwent combined treatment (5.2%), and only 1 patient was managed conservatively (1.7%). An improved outcome was reported in 94.8% of cases (55 out of 58 patients), whereas 3 out of 58 patients (5.2%) were moderately disabled after surgery and endovascular treatment. In literature, hemorrhagic presentation is reported as the most common onset (subarachnoid hemorrhage in 63% and intramedullary hemorrhage in 10%), frequently caused either by venous dilation, due to an ascending drainage pathway into an intracranial vein, or by the higher venous flow rates that can be associated with intracranial drainage. Hiramatsu and Sato stated that arterial feeders from the anterior spinal artery (ASA) and aneurysmal dilations are associated with hemorrhagic presentation. In agreement with the classification by Hiramatsu, we defined the PMAVF of the CVJ as a vascular lesion fed by the radiculomeningeal arteries from the vertebral artery and the spinal pial arteries from the ASA and/or lateral spinal artery. Considering the anatomical characteristics, we referred to our patient as affected by PMAVF, even if it was difficult to precisely localize the arteriovenous shunts because of the complex angioarchitecture of the fine feeding arteries and draining veins, but we presumed that the shunt was located in the point of major difference in vessel size between the feeding arteries and draining veins. PMAVFs of CVJ are rare pathologies of challenging management. The best diagnostic workup and treatment are still controversial: more studies are needed to compare different therapeutic strategies concerning both long-term occlusion rates and outcomes.

Keywords: Aneurysm, angiography, arteriovenous fistula, craniovertebral junction, perimedullary, subarachnoid hemorrhage

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INTRODUCTION

Perimedullary arteriovenous fistulas (PMAVFs) of the craniovertebral junction (CVJ) represent <1% of all cranial or spinal arteriovenous fistulas (AVFs)\(^1-3\) and are classified as type 5 AVFs according to Hiramatsu et al.\(^4\) PMAVFs may present clinically either with acute neurological deficits due to subarachnoid or intramedullary hemorrhage, or with progressive myelopathy caused by venous congestion;\(^5,6\) a differential diagnosis must be made with spinal thoracolumbar AVFs, whose main presentation is myelopathy;\(^7,8\) Therapeutic management includes options such as observation alone, endovascular occlusion, surgical exclusion, or a combination of these two last techniques, depending on patient-specific and lesion-specific characteristics. Observation carries the risk of rebleeding, surgical treatment presents challenges related to the operative approach, and endovascular treatment presents several technical limitations due to vessels’ size and risk of medullary ischemia. We present a rare case of a patient affected by ruptured PMAVF of CVJ associated to a venous pseudoaneurysm, providing new insights into the diagnosis and management of this rare entity.

MATERIALS AND METHODS

Study selection

A systematic review of the literature was conducted using MEDLINE, Cochrane, and EBSCO databases using the following combined MeSH terms: (perimedullary arteriovenous fistula OR dural arteriovenous shunt) and (craniovertebral junction OR craniovertebral junction). This search was limited to studies published in English. The PRISMA flow diagram of our search is available in Figure 1. Eligibility criteria were limited by the nature of the existing literature on PMAVFs, which consists of case series, case reports, and only one multicentric study. A definition of PMAVF has been reported only in 2017,\(^9\) all previous papers were meticulously analyzed to define the anatomical features of every AVF. All papers reporting PMAVFs not located at the CVJ were not included in this review. The reference list of all the articles included in this review was examined searching for additional papers.

Data extraction

All included papers were meticulously reviewed and analyzed for their study design, methodology, and patient characteristics. All patients’ data were recorded when available, including age, sex, clinical presentation, interval between onset of symptoms and diagnosis, treatment options (surgery or endovascular), and outcome.

Descriptive statistics are reported, including means and proportions. No formal statistical comparisons were performed due to small sample size and insufficient power to detect differences between groups.

RESULTS

A total of 31 published studies from 1996 to 2019 were identified through our search and additional reference evaluation. After detailed examination, ten were rejected from our review, as they were written in other languages or AVFs were not at the level of the CVJ. Our case was not included in the systematic review. Patients data are reported in Table 1.\(^4,9-28\) We selected 21 studies for this systematic review, with a total of 58 patients, including 20 females (34.5%) and 38 males (65.5%), with a female/male ratio of 1:1.9. Patients presented a mean age of 55.9 years, ranging from 38 to 82 years. The presentation with subarachnoid hemorrhage (SAH) occurred in 45 out of 58 patients (77.6%), whereas 11 out of 58 patients (18.9%) presented with sensory and/or motor dysfunction; 2 out of 58 cases (3.4%) were incidental findings. In 35 out of 58 patients (60.3%), the onset was acute. In 9 out of 58 patients (15.5%), the mean interval between the presentation of symptoms and diagnosis was 12.3 months, ranging between 0.5 and 48 months; in 13 out of 58 patients (22.4%), this time interval was not reported. Thirty-nine out of 58 patients underwent surgical treatment (67.2%), whereas 15 out of 58 patients were treated with the endovascular procedure only (25.8%); 3 out of 58 patients were treated with combined surgical and endovascular approach (5.2%); only one patient was managed conservatively (1.7%). An improved outcome was reported in 94.8% of cases (55 out of 58 patients), whereas 3 out of 58 patients (5.2%) were moderately disabled after treatment.

Case presentation

Medical history and clinical examination

A 73-year-old woman with a history of osteoporosis presented to our institution with sudden headache, vomiting, and rigor nucalis associated to dysesthesia of the upper limbs.

Imaging

Brain angio-computed tomography (CT) scan demonstrated diffuse SAH (Fisher grade 3) [Figure 2], as well as magnetic resonance angiography revealed the presence of a PMAVF associated to a venous pseudoaneurysm at C1–C2 level [Figure 3]. Subsequent digital subtraction angiography (DSA) confirmed a PMAVF of the CVJ, responsible for the recent SAH; the PMAVF was located in the right paramedian position, posterior to the middle third of the odontoid process with caudal extension up to C2–C3, with maximum dimensions on the sagittal plane of about 1.2 cm [Figure 4]. Cranially to the PMAVF, there was evidence of venous pseudoaneurysm. The PMAVF had arterial feeders from both hypertrophic anterior spinal arteries,
originating from the intracranial portion and from cervical radicular branches – which stemmed at the right C3–C4 and left C2 level, of greater caliber and flow to the left – of both vertebral arteries (VAs); the main venous drainage, dilated, was at the level of the basal venous plexus, with drainage in both inferior petrous sinuses and at the level of the left posterior cervical paravertebral plexus, also markedly dilated.

Management
After a multidisciplinary discussion, we decided to perform surgical treatment of the PMAVF because no navigable afferent vessels allowed the exclusion of the shunt point during the endovascular procedure. However, the patient refused the treatment, probably due both to progressive clinical improvement and to the high surgical risks explained during informed consent. She improved to complete recovery, CT scan revealing progressive reabsorption of the SAH. After 9-month follow-up, the PMAVF is stable in seriated DSA and angio-CT scans, and the patient is in good clinical conditions.

DISCUSSION
Historically, PMAVFs were considered congenital.\textsuperscript{[17]} Anteriorly located shunts represent an atypical feature.\textsuperscript{[29]} PMAVFs present direct shunts between the vessels at the level of the pia mater, superficially to the spinal cord, and they can be located anteriorly or anterolaterally to the spinal cord.\textsuperscript{[15,30,29]} Lawton \textit{et al.}\textsuperscript{[31]} suggested that angiogenesis could be induced by venous hypertension, responsible for the development of AVFs in experimental models. Sato \textit{et al.}\textsuperscript{[17]} hypothesized that venous hypertension can cause venous hypoxia, which induces additional pial shunts in patients with AVFs of the craniovertebral junction. According to Hassler \textit{et al.},\textsuperscript{[12]} venous drainage close to the nerve root sleeve could worsen a sump effect. Moreover, blood supply to the radiculomedullary arteries may be near the origin of the
### Table 1: Systematic review of demographic, symptoms, interval time to diagnosis, treatment, and outcome data of patients affected by perimedullary arteriovenous fistulas of the craniovertebral junction reported in the literature to date

| Authors and year | Number of patients | Age (years) | Sex | Symptoms | Time to diagnosis | Treatment | Outcome |
|------------------|--------------------|-------------|-----|----------|-------------------|-----------|---------|
| Hiramatsu et al. 2018<sup>[4]</sup> | 6                  | Mean 67     | 2 males, 2 females | SAH symptoms | Not reported | 4 endovascular treatments, 2 surgeries | Improved |
| Mascalchi et al. 1996<sup>[2]</sup> | 2                  | 69, 53      | Male | Lower limb paresthesias, urinary retention, progressive gait disturbance | 4 years, 2 years | Surgery (clipped and excised); endovascular treatment (injection of glue) | Both improved |
| Kinouchi et al. 1998<sup>[3]</sup> | 10                 | 63, 65, 44, 52, 70, 68, 50, 61, 60, 45 | 7 males, 3 females | 2 incidental, 1 back pain, bowel and bladder disturbance, sensory disturbances, trigeminal nerve disturbance, quadriparesis, and respiratory disturbance, 7 SAH symptoms | Not reported, 1 year, 7 acute onset | Surgery | Improved |
| Kowayama et al. 1999<sup>[4]</sup> | 3                  | 73, 52, 52  | 2 males, 1 female | SAH symptoms | Acute onset | 1 conservative, 1 endovascular treatment, 1 surgery | Moderately disabled |
| Yoshida et al. 1999<sup>[9]</sup> | 1                  | 68          | Male | Lower limb motor and sensory disturbances | 6 months | Surgery | Improved |
| Asakawa et al. 2002<sup>[10]</sup> | 1                  | 64          | Male | Lower limb weakness, dyspnea | 2 weeks | Embolization + surgical exclusion | Improved |
| Lv et al. 2011<sup>[11]</sup> | 4                  | 73, 66, 51, 40 | 3 males, 1 female | SAH symptoms | Acute onset | Endovascular treatment | Improved |
| Ohba et al. 2011<sup>[12]</sup> | 1                  | 55          | Male | SAH symptoms | Acute onset | Surgery | Improved |
| Kulvin et al. 2012<sup>[13]</sup> | 2                  | 44, 54      | Female | Altered mental status, right hemiparesis and diplopia | Acute onset | Surgery | Improved |
| Sato et al. 2013<sup>[14]</sup> | 9                  | 59, 82, 78, 62, 64, 62, 60, 64, 66 | 5 males, 4 females | SAH symptoms | Acute onset | Surgery | Improved |
| Gilard et al. 2013<sup>[15]</sup> | 1                  | 59          | Female | SAH symptoms | Acute onset | Surgery | Improved |
| Kasliwal et al. 2015<sup>[16]</sup> | 1                  | 59          | Female | SAH symptoms | Acute onset | Surgery | Improved |
| Jeon et al. 2015<sup>[17]</sup> | 1                  | 73          | Male | Progressive weakness of both lower extremities | 10 months | Endovascular treatment + surgery | Improved |
| Pop et al. 2015<sup>[18]</sup> | 1                  | 38          | Male | Paraparesis and acute urinary retention, generalized tonic-clonic seizures | 28 days | Endovascular treatment | Improved |
| Ogita et al. 2016<sup>[19]</sup> | 1                  | 58          | Male | SAH symptoms | 9 months | Surgery | Improved |
| Kawasaki et al. 2016<sup>[20]</sup> | 1                  | 73          | Female | Headache and left hemiparesis | Acute onset | Conservative | Improved |
| Enokizono et al. 2017<sup>[21]</sup> | 1                  | 50          | Female | Neck pain and urination difficulty, paralysis, and numbness of all four extremities | 1 month | Clipping | Improved |
| Fujimoto et al. 2018<sup>[22]</sup> | 6                  | 69, 79, 76, 64, 68, 46 | 4 males, 2 females | SAH symptoms | Not reported | 5 surgeries, 1 endovascular treatment | Improved |
| Kanemaru et al. 2019<sup>[23]</sup> | 4                  | 67, 76, 45, 53 | 3 males, 1 female | SAH symptoms | Acute onset | Surgery | Improved |
| Sato et al. 2019<sup>[24]</sup> | 1                  | 69          | Male | SAH symptoms | Acute onset | Endovascular treatment + surgery | Improved |
| Yoshida et al. 2019<sup>[25]</sup> | 1                  | 65          | Male | SAH symptoms | Acute onset | Endovascular treatment | Improved |

SAH - Subarachnoid hemorrhage

In literature, hemorrhagic presentation is reported as the most common onset (SAH in 63% and intramedullary hemorrhage in 10%)<sup>[1,6,34]</sup> frequently caused either by venous dilation, due to an ascending drainage pathway into an intracranial vein<sup>[6,10,25‑37]</sup> or by the higher venous flow rates that can be associated with intracranial drainage<sup>[10,38]</sup>. For Hiramatsu et al.<sup>[3]</sup> and Sato et al.<sup>[17]</sup> arterial feeders from the anterior spinal artery (ASA) and aneurysmal dilations are associated with hemorrhagic presentation.<sup>[1,17]</sup> However, there is controversy about the definition of PMAVFs.
CONCLUSIONS

We presented one emblematic patient, affected by rare PMAVF of the CVJ that refused treatment, but serial radiological follow-up demonstrated the stability of the lesion with good clinical performance. Thus, the present paper focused on a peculiar subtype of AVF of the CVJ that presented controversies both for radiological diagnosis and treatment indications. Therefore, we suggest a combined endovascular and neurosurgical approach to treat this type of patients, as it is possible to obtain the best postoperative outcome.

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Conflicts of interest
There are no conflicts of interest.

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