Vertebral artery dissection in term pregnancy after cervical spine manipulation: a case report and review the literature

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Abstract

Background: Vertebral artery dissection is an uncommon, but potentially fatal, vascular event. This case aimed to describe the pathogenesis and clinical presentation of vertebral artery dissection in a term pregnant patient. Moreover, we focused on the differential diagnosis, reviewing the available evidence.

Case presentation: A 39-year-old Caucasian woman presented at 38 + 4 weeks of gestation with a short-term history of vertigo, nausea, and vomiting. Symptoms appeared a few days after cervical spine manipulation by an osteopathic specialist. Urgent magnetic resonance imaging of the head was obtained and revealed an ischemic lesion of the right posterolateral portion of the brain bulb. A subsequent computed tomography angiographic scan of the head and neck showed a right vertebral artery dissection. Based on the correlation of the neurological manifestations and imaging findings, a diagnosis of vertebral artery dissection was established. The patient started low-dose acetylsalicylic acid and prophylactic enoxaparin following an urgent cesarean section.

Conclusion: Vertebral artery dissection is a rare but potential cause of neurologic impairments in pregnancy and during the postpartum period. It should be considered in the differential diagnosis for women who present with headache and/or vertigo. Women with a history of migraines, hypertension, or autoimmune disorders in pregnancy are at higher risk, as well as following cervical spine manipulations. Prompt diagnosis and management of vertebral artery dissection are essential to ensure favorable outcomes.

Keywords: Vertebral artery dissection, Pregnancy, Vertebrobasilar ischemia, Cervical spine manipulation, Osteopathy

Background

Vertebral arterial dissection (VAD) is a rare complication of pregnancy and puerperium. A data registry reported that 2.4% of symptomatic, spontaneous VADs occurred in the postpartum period [1]. Aortic, coronary, and cervical/vertebral artery dissection was reported to be associated with preeclampsia in the antenatal setting [1]. On the other hand, VAD incidence in hypertensive disorders of pregnancy is unknown, due to the paucity of reports documenting only adverse outcomes [1]. Hormonal and mechanical factors might increase the risk of VAD during pregnancy and puerperium [2]. Indeed, identified predisposing factors of VAD include intimal injury related to Valsalva maneuvers during labor, and alterations in arterial wall integrity due to hormonal or vasoactive substances, in addition to an overall state of hypercoagulability [3]. Another possible condition that can lead to VAD is cervical spine manipulation [4]. It is known that any type of trauma can cause a dissection such as cervical manipulation. Therefore, in nonpregnant patients, it is...
not uncommon, but most patients are asymptomatic, and this serious accident after manipulation has an underestimated incidence. Nevertheless, the risk related to the development of VAD is decidedly low if the manipulation maneuvers are carried out according to good clinical practice [4]. Several vascular and connective tissue disorders have also been associated with dissection; in particular, migraines, fibromuscular hyperplasia, and vascular Ehlers–Danlos syndrome [5]. As previously stated, the etiology of VAD is complex and often multifactorial, especially when trivial trauma and manipulations are involved. Other risk factors or conditions, such as fibromuscular dysplasia, Marfan’s syndrome, migraines, use of oral contraceptives, recent infections, and mild hyperhomo-cysteinemia, should be considered in any given case [6]. There exists uncertainty of how to counsel women with a previous VAD, regarding the risk of recurrence during pregnancy [2].

We aim to describe a 39-year-old female who presented with vertigo, nausea, and vomiting and was found to have a VAD. We discuss the presentation, differential diagnosis, and pathogenesis of this uncommon, but clinically significant, vascular event. Finally, we briefly review other described VAD cases.

Case presentation
A 39-year-old pregnant Caucasian woman presented to the Obstetric Emergency Room reporting vertigo, vomiting, nystagmus, dizziness, and hindrance in the execution of fine movements of the right arm. The maternal parameters on admission are regular: pulse 98 beats per minute, pressure 110/68 mmHg, and temperature of 36.2 °C. She had an obstetric history of a first-trimester spontaneous abortion and a medical history of tension headache. She is married and graduated. She has a high socioeconomic status and is employed as an engineer.

The ongoing pregnancy coursed physiologically until that moment. The fetus was screened for aneuploidy with a noninvasive prenatal test (NIPT), while second- and third-trimester ultrasounds for the study of malformations were both normal. The oral glucose tolerance test during pregnancy [2].

The neonate showed good adaptation to postnatal life with an Apgar score of 9 at 1', 10 at 5', and 10 at 10'; blood gas analysis was regular both in the artery and vein. Normal anthropometric parameters were present: 3250 g (52nd centile), length of 49 cm, and cranial circumference of 36 cm.
On day 4, for better study suspected dissection on small vertebral vessels, the patient underwent CT angiography of the neck, which showed a focal dissection at the V2 distal segment of the right vertebral artery (Fig. 1C–D). The puerperium course was normal, and the midwife helped the patient during breastfeeding because of the difficulty of standing up and walking due to the diplopic symptom. Psychological support was offered during the hospitalization, with daily physiotherapy rehabilitation and orthoptic evaluation. Congenital and acquired
thrombophilia tested negative. After 12 days of rehabilitation, the patient was discharged with continued complaints of diplopia and a walker for mobility. After the VAD diagnosis and for the entire length of hospitalization, the patient was treated with Cardioaspirin 100 mg/day and prophylactic enoxaparin 4000 UI/day subcutaneous injection for 60 days. At the subsequent neurological evaluation, during the follow-up of 2 and 4 months, the patient showed persistence of vertical diplopia and a circumscript and wide gait, and life-long ASA was prescribed. A follow-up MRI was scheduled for 6 months after the stroke, which confirmed the signs of the previous ischemic lesion on the posterolateral right medulla oblongata. The remaining findings are unchanged.

Ethical approval was obtained, and the patient gave written informed consent to publish this case and any accompanying images.

Discussion and conclusion
To our knowledge, this is one of the rare reports of an ischemic lesion due to VAD in low-risk pregnancy secondary to cervical spine manipulation. Cervical artery dissection (CAD), including VAD, is a rare complication of pregnancy; however, Salehi Omran et al. recently demonstrated that the incidence in pregnancy is twice as common as in the rest of the female population [7].

VAD has typically been associated with hypertensive disorder of pregnancy (HDP) [1, 8], autoimmune disease [9], and migraines [10]. A recent nationwide American cohort study on pregnancy-associated arterial dissection showed that VAD is the fourth most common dissection after or prior birth and found a significant association with older maternal age, chronic hypertension, dyslipidemia, tobacco use, alcohol use, obesity, heart failure, chronic liver disease, arthritis, depression, Marfan syndrome, and Ehlers–Danlos syndrome [11]. However, one recent meta-analysis demonstrated that nearly 50% of cases occur in the absence of such risk factors [12, 13]. To date, the etiology of VAD is not well established. Borelli et al. proposed a dual mechanism of pathogenesis occurring in the postpartum period: (1) advanced age, causing increased arterial stiffness, and (2) hormone fluctuations, inducing structural vascular changes [14], which may also happen at the end of pregnancy in our patient. McKinney et al. also suggested that endothelial damage may occur due to the release of vasoactive or angiogenic substances during pregnancy [15].

VAD should be considered in the differential diagnosis of women who present with nonspecific symptoms, such as headache, vomiting, and/or vertigo, particularly in the context of HDP [1]. Women older than 35 years and those with a history of HDP or autoimmune disease (that is, systemic lupus erythematosus, anti-phospholipid syndrome) are particularly at high risk. Other predisposing factors of arterial dissection in the peripartum period include intimal injury related to Valsalva maneuvers during labor, alterations in arterial wall integrity due to pregnancy-related hormonal or vasoactive substances [9], and reactive thrombocytosis (subsequent to postpartum hemorrhage), which may all play potential roles in this process and require further investigation [16]. Moreover, prompt diagnosis and management of VAD are essential to ensure favorable patient outcomes.

One recent report discussed the contributing factors in a case of VAD following chiropractic treatment in a pregnant woman with systemic lupus erythematosus [4]. Migraine disorder was shown to be associated with a twofold increased risk of VAD in a recent meta-analysis [9] and has been frequently reported in several case series [17–19]. Stuber et al. recently published a review of the literature regarding adverse effects of spinal manipulation in the pregnant and postpartum periods [20], identifying adverse events in five pregnant women and two postpartum women.

Table 1 summarizes all cases of VAD reported both prior and after delivery, with 24 cases distributed with a prevalence during the postpartum period (19 of the 24 cases). The clinical presentation is varied, with a higher frequency of headaches, vertigo, and diplopia, and the risk factors most represented are hypertension and migraines.

The association between cervical spine manipulation and neurovascular complications is still strongly debated [21, 22]. CAD is thought to occur spontaneously, but neck trauma, especially in hyperextension and rotation, has been reported as a trigger [23]. A population-based, case–control study found no evidence of excess risk of vertebrobasilar stroke associated with chiropractic care compared with controls [24]. A recent retrospective case–control study, however, found a significantly increased risk of VAD in individuals less than 55 years of age with recent neck manual therapy [25]. A recent multivariable regression analysis of a retrospective study assessed the risk factors and clinical outcomes associated with CAD-related strokes. Patients with CAD were younger and more likely to have a history of migraines and recent neck manipulation [26]. A systematic review and meta-analysis of chiropractic care and CAD concluded that the quality of the published data was very low, and the authors showed a small association between chiropractic neck manipulation and CAD [27, 28].

Finally, the future of these women is somewhat debated. They should be advised about their increased risk of developing a new stroke, so these patients will need to continue Cardioaspirin prophylaxis for life [2]. Regarding reproductive future, a recent observational
| Cases                  | Age | Presentation                                                                                                                                         | Risk factors                      | VAD affected | Mode of delivery | Time to and from delivery (days) |
|------------------------|-----|------------------------------------------------------------------------------------------------------------------------------------------------------|-----------------------------------|--------------|-----------------|----------------------------------|
| Current report         | 39  | Vertigo, vomiting, nystagmus, dizziness, and hindrance in the execution of fine movements with the right arm                                              | Migraine                          | Right        | Cesarean section | Antepartum, 39 w                 |
| Gasecki et al. (1999)  | 34  | Neck pain, headache, 1 week later: right facial numbness, left-sided weakness, and vertigo, right. Horner’s syndrome, right-side ataxia.          | Healthy                           | Right        | Vaginal delivery | 14 days postpartum               |
| McKinney et al. (2002) | 41  | Severe headache, blurred vision, HTN                                                                                                                   | Preeclampsia                      | Left         | Cesarean section | 5 days postpartum                |
| Tuluc (2006)           | 39  | Headache preceding loss of consciousness                                                                                                               | HTN                               | Right        | Cesarean section | Antepartum                       |
| Arnold et al. (2008)   | 41  | Ipsilateral neck pain, thundertap headache                                                                                                             | Migraine, hyperlipidemia          | Left         | Vaginal delivery | 18 days postpartum               |
|                        | 27  | Ipsilateral neck pain, thundertap headache                                                                                                             | Migraine, HTN, hyperlipidemia     | Right        | Vaginal delivery | 11 days postpartum               |
|                        | 38  | Thundertap headache                                                                                                                                     | Migraine, hyperlipidemia          | Bilateral    | Vaginal delivery | 7 days postpartum                |
|                        | 34  | Ipsilateral neck pain, headache                                                                                                                       | Chiropractor neck manipulation    | Right        | Vaginal delivery | 7 days postpartum                |
| Sharma et al. (2010)   | 28  | Non-exertional, intermittent, substernal, sharp chest pain, and left arm numbness, intermittent bifrontal headache                                      | Atherosclerotic risk factors      | Left         | Vaginal delivery | 10 days postpartum               |
| Cenkowski et al. (2012) | 35  | Sudden-onset retrosternal chest pain radiating to the jaw, nausea, vomiting (7 months postpartum), diplopia, numbness to left arm and face (8 months postpartum) | None                              | Right        | Vaginal delivery | 7 months postpartum              |
| Drazin et al. (2012)   | 37  | Thundertap headache, neck pain                                                                                                                        | Migraine, high pressure during labor | Bilateral    | Vaginal delivery | 3 days postpartum                |
| Morton A. (2012)       | 38  | Occipital headache, severe right-sided anterior neck pain, ipsilateral Horner’s syndrome (after chiropractic treatment; spinal manipulation)       | Migraine, SLE, HTN, heterozygous for prothrombin gene mutation | Right        | 4 days after the onset of neurological symptoms intrauterine fetal demise | Antepartum 16 weeks was noted |
Table 1 (continued)

| Cases                  | Age | Presentation                                                | Risk factors                      | VAD affected | Mode of delivery | Time to and from delivery (days) |
|------------------------|-----|-------------------------------------------------------------|-----------------------------------|--------------|-----------------|----------------------------------|
| Kelly et al. (2014)    | 39  | Thunderclap headache, ipsilateral neck pain, blurred vision, and horizontal diplopia | HTN, hyperlipidemia               | Bilateral    | Vaginal delivery | 24 days postpartum               |
|                        | 39  | Right eyelid ptosis, headache, and bilateral neck pain     | Healthy                           | Right        | Vaginal delivery | 11 days postpartum               |
|                        | 29  | Right-sided weakness, sensory loss, and expressive aphasia, followed by severe headache and right-sided hemiplegia | Migraine                          | Left         | Vaginal delivery | 53 days postpartum               |
|                        | 32  | Severe headache and neck pain, followed by left-sided facial droop and left arm weakness | Migraine                          | Right        | Vaginal delivery | 0 days postpartum                |
|                        | 28  | Severe headache, neck pain, bilateral leg weakness        | HTN                               | Left         | Cesarean section | 4 days postpartum                |
| Finley et al. (2015)   | 35  | Thunderclap headache, intractable vertigo                 | Migraine                          | Right        | Vaginal delivery | 21 days postpartum               |
| Nishimura et al. (2015)| 35  | Thunderclap headache                                      | Eclampsia, PRES                   | Right        | Cesarean section | 8 days postpartum                |
| Shanmugalingam et al. (2016)| 32  | Left-sided neck pain                                      | Preeclampsia/eclampsia            | Left         | Cesarean section | Antepartum, 38 + 2 weeks         |
|                        | 33  | Right-sided neck pain                                     | Preeclampsia                      | Right        | Cesarean section | Antepartum, 36 weeks             |
|                        | 30  | Headache with left-sided neck pain                         | NSAID-induced postpartum HTN, migraines, obesity | Right        | Vaginal delivery | 3 days postpartum                |
|                        | 30  | Left-sided neck pain                                      | Previous IUGR and postpartum hemorrhage with DIC | Left         | Vaginal delivery | 6 days postpartum,               |
| Manasewitsch et al. (2020)| 31  | Frontal headache, vertigo, nausea, vomiting               | Preeclampsia, smoking             | Left         | Cesarean section | 10 days postpartum               |

HTN hypertension, SLE systemic lupus erythematosus, PRES posterior reversible encephalopathy syndrome, NSAID nonsteroidal antiinflammatory drug, IUGR intrauterine growth restriction, DIC disseminated intravasal coagulation
German study concluded that the risk of recurrent VAD may not be significantly increased with pregnancies, starting at least 12 months after the event, in women without connective tissue disease, such as our patient [2].

Despite the absence of hypertension and autoimmune diseases in our patient, previous chiropractic treatment, pregnancy hormonal condition, and advanced age (39 years) may have contributed to vessel fragility. The risk of VAD was also increased due to the history of tension headache/migraine. Osteopathy practitioners should be aware of the possible complications of neck manipulation in pregnancy and the postpartum period, particularly in mothers with underlying medical disorders that may predispose to vessel fragility and VAD.

In conclusion, we recommend that obstetric professionals carefully consider VAD as a differential diagnosis when evaluating women with dizziness, headache, and neck pain with or without a recent history of spinal manipulation, both in pregnancy and in the postpartum period. Moreover, they should consider with caution the risks and benefits of any cervical osteopathy practice in pregnant women with risk factors for VAD (hypertension and autoimmune diseases, history of tension headache/migraine).

Abbreviations
VAD: Vertebral artery dissection; CAD: Cervical artery dissection; NIPT: Noninvasive prenatal test; ORL: Otolaryngologist; CT: Computed tomography; MRI: Magnetic resonance imaging; ASA: Acetylsalicylic acid; CS: Cesarean section; HDP: Hypertensive disorder of pregnancy; DWI: Diffusion-weighted imaging; ADC: Apparent diffusion coefficient; PICA: Posteroinferior cerebellar artery.

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Authors’ contributions
FM and SB contributed to the clinical management of the patient and the literature review, and wrote the initial draft of this manuscript and subsequent revisions. MGI, FF, IN, FC, MG, AG, and FC contributed to the clinical management and manuscript preparation. All authors have read and approved the final manuscript.

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Availability of data and materials
This is a case report of a single patient. To protect privacy and respect confidentiality, none of the raw data has been made available in any public repository. The original reports, laboratory studies, imaging studies, and outpatient clinic records are retained, as per normal procedure, within the medical records of our institution.

Declarations
Ethics approval and consent to participate
No ethics approval was required, as this is a case report. Written and verbal consent was obtained from the patient discussed in this case report.

Consent for publication
Written informed consent was obtained from the patient for publication of this case report and any accompanying images. A copy of the written consent is available for review by the Editor-in-Chief of this journal.

Competing interests
The authors declare they have no financial or other conflicts of interest concerning this research and its publication.

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