Diagnosis and treatment of pure arterial malformation
Three case reports and literature review

Tian-Yi Liu, MD, Ning Xu, PhD, Zheng Wan, BSc, Zhe-Ming Zhang, BSc, Jian-Jun Xu, BSc, Hao Meng, PhD*, Hong-Lei Wang, PhD

Abstract
Rationale: The incidence of pure arterial malformations is relatively low, and few cases have been reported. Only 2 cases with pure arterial malformation have been reported to receive surgery or endovascular treatment.

Patient concerns: We report 3 cases and review the relevant literatures. The head examinations of the patients suggested the presence of high-density shadows in front of the pons and midbrain, the dilation of the supraclinoid segment of the right internal carotid artery, and moyamoya in the left brain with an aneurysm-like expansion located on the left posterior communicating artery respectively. After admission, head digital subtraction angiography (DSA) was performed.

Diagnoses: Digital subtraction angiography (DSA) for these 3 patients showed that the left posterior communicating artery, the supraclinoid segment of the right internal carotid artery, and the left posterior communicating artery appeared dilated, tortuous, and spirally elongated. In addition, the lesions in the latter 2 patients were accompanied with local aneurysmal changes.

Interventions: Two patients were given conservative treatment, and another patient was given endovascular treatment. A head DSA was reviewed 6 months after therapy.

Outcomes: The prognosis status of the 3 patients was good. Two patients in the conservative treatment group showed no changes in the lesions on head DSA examination. The DSA examination of the third patient indicated that the vascular remodeling of the diseased vessels was good, the blood vessels were unobstructed, and the aneurysms had disappeared.

Lessons: Pure arterial malformations mostly occur in young women and may involve any blood vessels in the brain. It can be accompanied with local aneurysms and calcification. The patients are often given conservative treatment but need to be reviewed regularly. However, it is beneficial to give endovascular treatment to the patients with local aneurysms.

Abbreviations: CT = computerized tomography, CTA = computed tomography angiography, DSA = digital subtraction angiography.

Keywords: aneurysm, arterial malformation, vascular disorders

1. Introduction

The incidence of pure arterial malformations is low; only few cases have been reported. In most patients, these are incidentally discovered during head computerized tomography (CT) or computed tomography angiography (CTA) examination performed for headache or other reasons. The concept of pure arterial malformation was first reported by McLaughlin et al, whose definition involves the presence of dilated, overlapping, and tortuous arteries forming a mass of arterial loops with a coil-like appearance in the absence of any venous components.[1] Due to its rarity, the optimal treatment remains controversial. Brinjikji et al proposed that patients with pure arterial malformations should receive conservative treatment.[2]

In this study, we reported 3 cases of pure arterial malformation, with 1 treated with endovascular intervention, and reviewed the related literatures.

2. Case presentation

2.1. Case 1

A 36-year-old female patient presented an intermittent headache. The head CT showed a high-density anterior to the pons and the midbrain. The head DSA revealed the left posterior communicating artery being dilated, overlapping, tortuous, and continuous to the P2 segment of the posterior cerebral artery. While other blood vessels were normal. We treated the patient with pain control and neurotrophic medications. The patient’s blood pressure was controlled within the normal range. Follow-up DSA 6 months after discharge did not show any morphological changes.

The authors have no funding and conflicts of interest to disclose.

Department of Neurosurgery, the First Hospital of Jilin University, Changchun, China.

* Correspondence: Hao Meng, Hong-Wei Wang, Department of Neurosurgery, the First Hospital of Jilin University, Changchun 130021, China (e-mails: menghao@jlu.edu.cn, wanghonglei@jlu.edu.cn).

Copyright © 2020 the Author(s). Published by Wolters Kluwer Health, Inc.

All data generated or analyzed during this study are included in this published article (and its supplementary information files).

This is an open access article distributed under the Creative Commons Attribution License 4.0 (CCBY), which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

How to cite this article: Liu TY, Xu N, Wan Z, Zhang ZM, Xu JJ, Meng H, Wang HL. Diagnosis and treatment of pure arterial malformation: three case reports and literature review. Medicine 2020;99:21(e20229).

Received: 30 May 2019 / Received in final form: 24 February 2020 / Accepted: 8 April 2020
http://db doi.org/10.1097/MD.0000000000020229
changes of the malformations, and the patient denied recurrence of headaches (Fig. 1).

### 2.2. Case 2

A 24-year-old female patient presented to our department with occasional dizziness. The head CTA revealed the dilation of the suprachinoid segment of the right internal carotid artery. Moreover, a local aneurysm-like change was found in the head DSA examination. The patient was given conservative treatment such as neurotrophic drugs and avoiding fatigue, and 6 months later, the head DSA examination showed no changes in the diseased blood vessels. The patient’s dizziness symptom was relieved (Fig. 2).

### 2.3. Case 3

A 53-year-old male complained of left limb numbness for 2 weeks. He had a history of hypertension for years and his blood pressure was not controlled well. The head MRI examination showed vascular emptying signals adjacent to the brainstem. The CTA examination revealed an aneurysmal expansion of the left posterior communicating artery. The DSA examination of the head indicated moyamoya disease on the left brain; the proximal portion of the left posterior communicating artery was obviously enlarged, and an aneurysm-like expansion was observed. Moreover, the left posterior communicating artery was distorted, prolonged, and continuous with the posterior cerebral artery. The patient was treated with stent-assisted-coil embolization. Six months later, the patient underwent a DSA review of the head. The vascular lumen remodeling was good, no dilatation was observed, and the aneurysm achieved Raymond I embolization (Fig. 3).

### 3. Ethic statement

Written informed consent was obtained from the patients and their parents for publication of these case reports. Patients have provided informed consent for publication of the cases.

### 4. Discussion

Pure arterial malformation is defined as the presence of dilated, overlapping, and tortuous arteries forming a mass of arterial loops with a coil-like appearance in the absence of any venous components. It may also be associated with local aneurysms or calcification. The diagnosis of the disease requires DSA and 3D imaging examinations. Pure arterial malformation needs to be differentiated from other vascular diseases such as arteriovenous
malformation, arterial dissection, dilatation and prolongation syndrome, and developmental arterial anomalies \(^{[1,11,23–28]}\) (Fig. 4). The abnormal arteries neither communicate with veins through the vascular network or vascular nest nor communicate with veins directly. This lack of communication with venous structures is the major characteristic differentiating from arteriovenous malformations and arteriovenous fistulas. Pure arterial malformation can involve any intracranial blood vessels, but it mostly occurs at the supraclinoid segment of the internal carotid artery, the posterior communicating artery, the anterior cerebral artery, and the M1 segment of the middle cerebral artery. The natural history of patients with pure arterial malformation is usually non-specific. On the other hand, the dilated arterial disease occurs mostly in the internal carotid and vertebral basilar artery.\(^ {\text{[6,29]}}\) Most patients with dilated arterial disease also have other risk factors, such as hypertension-related diseases, infectious diseases, immune diseases, or systemic macrovascular dilatation changes.\(^ {\text{[10,27,28,30–32]}}\) Developmental arterial anomalies often involve the distal branches of the intracranial vessels, and they are mainly characterized by small network-like dilated arterial clusters. Developmental arterial anomalies are also associated with embryonic dysplasia, and there may be cerebral cortical dysplasia around them.\(^ {\text{[5,6,23]}}\) In patients with pure arterial malformation, the abnormal vessel wall remains parallel, and the affected vessel segments are longer.\(^ {\text{[33]}}\) The form of the blood vessels and the vascular orientation can be seen clearly. The major feature differentiating pure arterial malformation from arterial dissection is that there is no hematoma in the vascular wall or double cavity sign on imaging examinations. However, theory exists pertaining to the relationship between arterial dissection and pure arterial malformation. Some scholars believe that repetitive healing could take place during the pathological process of arterial dissection by the formation of fusiform expansion, local stenosis, and complex odd shape aneurysms. Therefore, pure arterial malformations is a chronic healing of arterial dissection.\(^ {\text{[11,23,26,34]}}\) The etiology and pathogenesis of pure arterial malformation remain unclear. It may occur when there is a defect or injury on the vessel wall because of congenital dysplasia, they may also occur after acquired damage caused by bacterial or viral infection, or they may be a chronic healing process for dissection lesions.\(^ {\text{[14]}}\)

The optimal therapy for pure arterial malformation also remains controversial. A review of previous literatures showed that 94.4% of affected patients received conservative treatment and achieved a good prognosis. So far, only 2 cases treated with surgery or endovascular treatment have been reported. Brinzikji et al suggested that patients with pure arterial malformation should be treated conservatively. We proposed that conservative treatment is appropriate for the patients with a pure arterial malformation without a local aneurysm; however, patients with pure arterial malformation with local aneurysms can achieve a good prognosis when treated via intervention with endovascular treatment.\(^ {\text{[35,36]}}\) In the 3 cases reported here, the first 2 patients were young female patients with a good natural history. One of these patients had a left posterior communicating artery malformation, and the other patient had a pure arterial malformation in the supraclinoid segment of the right internal carotid artery with a local aneurysm (the shape of the aneurysm was regular). Considering that the patient’s main symptoms were not related to the diseased blood vessels and the patient was young, adopting endovascular treatment or surgical bypass therapy would result in a larger risk and may cause some problems of taking medicines post operation. Therefore, the patient was given conservative treatment. Moreover, the patient was regularly reviewed by imaging examination, and the risk factors for the disease were controlled. After 6 months, a review of a head DSA examination revealed no changes in the diseased blood vessels. The third patient had a history of hypertension for many years. Considering the distorted, dilated PCoA-PCA observed in this patient exhibited aneurysm-like changes, there was a risk of rupture and bleeding. Moreover, the patient had an embryonic posterior cerebral artery. The blood therefore
experienced turbulence in the posterior communicating artery of the aneurysm-like expansion, which is a risk factor for thrombosis. Once the thrombus falls off, it may cause infarction in the area which the posterior cerebral artery blood supplies, and may lead to serious consequences. Therefore, the patient was treated with stent-assisted-coil embolization. The stent was placed in the distorted and dilated posterior communicating artery. At the same time, we treated the local aneurysm with coils

Table 1

| Author                  | Sex, age (yr) | Lesion location | Symptom                                      | Therapy                  | Follow up | With aneurysm | Physical examination |
|-------------------------|---------------|-----------------|----------------------------------------------|--------------------------|-----------|---------------|---------------------|
| McLaughlin et al[17]    | F, 24         | Lt PICA&P2      | Headache dizziness                           | Conservative treatment   | Not mention | Yes            | Normal              |
| Abe et al[13]           | M, 32         | Lt MCA          | Seizures                                     | Conservative treatment   | Not mention | No             | Normal              |
| Sako et al[21]          | M, 35         | Lt PICA         | Vertigo                                      | Conservative treatment   | 6 mo       | No            | Normal              |
| Shankar et al[16]       | F, 26         | Distal rt PCA   | Incidental                                   | Conservative treatment   | Not mention | No             | Normal              |
| Uchino et al[21]        | F, 35         | Rt SCA          | Headache                                     | Conservative treatment   | 2 yr       | No            | Normal              |
| Hanakita et al[17]      | F, 43         | Rt distal ICA & proximal M1; PcA   | Dysphoria                                    | EC-IC bypass, wrapped    |            |               | Dysphoria            |
| Methy et al[21]         | F, 1          | Lt MCA & supraclinoid ICA | PHACE                                         | Conservative treatment   | Not mention | No             | Not mention          |
| Araki et al[14]         | F, 25         | Rt MCA, ACA&PCA | Rt hemiplegiaencephal                       | Conservative treatment   | Not mention | No             | Not mention          |
| Fekianova et al[21]     | M, 42         | Rt MCA          | Headache, nausea, vomiting, lt side weakness | Conservative treatment   | 1 yr       | Yes           | Lt side weakness    |
| Yamada et al[14]        | F, 17         | Lt supraclinoid, M1 & ACA | Nausea vomitting                             | Conservative treatment   | Not mention | No             | Normal              |
| Yamada et al[12]        | F, 40         | Lt supraclinoid, M1 &ACA | Rt side hemiparesis                          | Conservative treatment   | Not mention | No             | Rt side hemiparesis |
| Karamoto et al[11]      | M, 41         | Lt MCA          | Seizures                                     | Conservative treatment   | Not mention | No             | Normal              |
| Thompson et al[12]      | M, 39         | Distal lt ACA   | Seizures                                     | Conservative treatment   | 3 yr       | No             | Normal              |
| McLaughlin et al[14]    | F, 6          | Lt supraclinoid ICA & proximal M1 | Headache                                     | Conservative treatment   | Not mention | No             | Normal              |
| Lantena et al[12]       | M, 1          | Lt PICA&PCA     | Infarct from moyamoya                        | Conservative treatment   | Not mention | No             | Not mention          |
| Baccin et al[24]        | F, 4          | Lt supraclinoid ICA&PCA | PHACE syndrome, fever, & hypotonia          | Conservative treatment   | 16 mo      | No             | Rt-sided hemiparesis |
| Baccin et al[24]        | F, 1          | Lt supraclinoid ICA, PCA, P1; rt supraclinoid ICA | PHACE syndrome, fever, & hypotonia          | Conservative treatment   | Not mention | No             | Not mention          |
| Vanslambrock et al[8]   | M, 5          | Lt ICA, PCA, ACA, & Lt SCA | Minimal rt hemiparesis, & Lt SCA             | Conservative treatment   | Not mention | No             | Minimal rt hemiparesis, & Lt SCA |
| Wapler et al[24]        | M, 21         | Bilat pericallosal | Seizures, agniesis of corpus callosum        | Conservative treatment   | Not mention | No             | Normal              |
| Sack & Lindenburg[77]   | M, 2          | Bilat A2s       | Viral encephalitis                            | Conservative treatment   | Not mention | No             | Not mention          |
| Beringer & Aengst[16]   | M, 49         | Bilat pericallosal | Intermittent frontal headache                | Conservative treatment   | Several mo  | No             | Normal              |
| Krzywida et al[24]      | F, 72         | Bilat distal ACAs | Aphasia, r-sided hemiplegia                  | Conservative treatment   | Not mention | No             | Aphasia, r-sided hemiplegia |
| Tuskamoto et al[13]     | F, 37         | Bilat pericallosal | Mania                                         | Conservative treatment   | Not mention | No             | Normal              |
| Doran et al[17]         | F, 14         | Bilat per- & supraclinoid segments of ACAs | Partial complex Seizures                    | Conservative treatment   | Not mention | No             | Normal              |
| W. Brink et al[24]      | F, 10         | Lt supraclinoid, PCA, & ACA | Severe r-sided headache                      | Coil embolization         | 72 mo      | Yes            | Normal              |
| W. Brink et al[24]      | F, 19         | Lt MCA          | Incidental                                   | Conservative treatment   | 36 mo      | Yes            | Normal              |
| W. Brink et al[24]      | F, 27         | BA              | Headache, lt hemibody                        | Conservative treatment   | 5 mo       | No             | Normal              |
| W. Brink et al[24]      | F, 25         | Lt supraclinoid ICA & M1 | Headache                                     | Conservative treatment   | 60 mo      | Yes            | Normal              |
| W. Brink et al[24]      | F, 25         | Lt ACA          | Headache after minor trauma                  | Conservative treatment   | 30 mo      | No             | Normal              |
| W. Brink et al[24]      | F, 34         | Lt ACA          | Headache, left trauma                         | Conservative treatment   | 12 mo      | No             | Normal              |
| W. Brink et al[24]      | F, 38         | Lt PICA         | Transient hand numbness                      | Conservative treatment   | 2 mo       | No             | Normal              |
| W. Brink et al[24]      | M, 11         | Lt PICA         | Incidental                                   | Conservative treatment   | 1 mo       | No             | Normal              |
| W. Brink et al[24]      | M, 17         | Rt SCA          | Headache                                     | Conservative treatment   | 26 mo      | No             | Normal              |
| W. Brink et al[24]      | F, 47         | Rt ACA          | Prior thunderclap headache                   | Conservative treatment   | 27 mo      | No             | Normal              |
| W. Brink et al[24]      | F, 35         | Lt PCA&PCA      | Headache                                     | Conservative treatment   | 84 mo      | Yes            | Normal              |
| W. Brink et al[24]      | F, 20         | Rt ICA, P2 & P2 | Headache, trauma                             | Conservative treatment   | 1 mo       | No             | Normal              |

ACA = anterior cerebral artery, ICA = internal carotid artery, Lt = left, MCA = middle cerebral artery, PCA= posterior communicating artery, PICA= posterior inferior cerebellar artery, Rt= right, SCA = superior cerebellar artery.
to reduce the risk of bleeding and remodel the vascular lumen. The patient was given antiplatelet therapy after the operation. After 6 months, a head DSA examination showed that the aneurysm-like dilation had disappeared and the lumen remodeling was good.

We reviewed some individual and series of case reports of pure arterial malformations or similar pure arterial malformations reported in the past (Table 1). Among the 36 reviewed cases, 69.4% were female and 66.7% were young patients aged between 14 and 44 years old. The disease was usually observed in young women.[13] The clinical symptoms varied widely, with 38.9% of the patients presenting with headache, 13.9% with PHACE, and 8.3% diagnosed by chance, accompanied by congenital vascular dysplasia or systemic diseases, such as genetic diseases and infections, and they will therefore exhibit other symptoms at presentation.[6,29–31,37] Most elderly patients are associated with hypertension, diabetes, hyperlipidemia and other risk factors for damage to vessel walls. This disease can involve any part of any blood vessel in the brain, with 33% of patients having symptoms suggesting the simultaneous involvement of multiple blood vessels in the brain. Among the cases reviewed, the PCoA-PCA, ICA supraclinoid segment, and MCA-M1 were involved in 64.2%. A total of 6 patients with pure arterial malformations were accompanied with local aneurysm. The lesions which involved PCoA-PCA or MCA were more likely to have local aneurysms or calcification.[4,10,11,26] However, lesions involving the distal end of the ACA were mainly characterized by an expansion in the vascular diameter and was sometimes accompanied by peripheral cerebral cortical dysplasia.[15,17,19,24]

5. Conclusion

Among cerebrovascular diseases, pure arterial malformation is a relatively rare disease. Pure arterial malformation is the presence of dilated, overlapping, and tortuous arteries forming a mass of arterial loops with a coil-like appearance in the absence of any venous components. With the increasing awareness of pure arterial malformations, it is beneficial to give the patients conservative treatment, regular imaging examinations and to control risk factors. However, pure arterial malformations with local aneurysm may require surgery or endovascular treatment. The diagnosis and treatment of this disease remains a hot topic of discussion and are worth continued exploration.

Author contributions

Conceptualization: Hong Lei Wang, Ning Xu.
Data collection and writing of the original article: Tianyi Liu.
Data curation: Tianyi Liu.
Writing – original draft: Zheng Wan, Zheming Zhang, Tian-Yi Liu.
Writing – review & editing: Tian-Yi Liu.

References

[1] McLaughlin N, Raychev R, Duckwiler G, et al. Pure arterial malformation of the posterior cerebral artery: importance of its recognition. J Neurosurg 2013;119:653–60.
[2] Beijikji W, Cloft HJ, Flemming KD, et al. Pure arterial malformations. J Neurosurg 2018;129:91–9.
[3] Yamada K, Hayakawa T, Ushio Y, et al. Cerebral arterial dolichoectasia associated with moyamoya vessels. Surg Neurol 1985;23:19–24.
[4] VanSlambrouck K, Allegaert K, Goemans N, et al. Symptomatic unilateral dolicho-ectasia of the intracranial arteries in a child. Eur Radiol 2000;10:759–60.
[5] Abe T, Singer RJ, Marks MP, et al. Arterial vascular abnormality accompanying cerebral cortical dysplasia. AJNR Am J Neuroradiol 1997;18:144–6.
[6] Doran SE, Deveikis JP, Chandler WF. Dolichoectasia of the anterior cerebral arteries in an adolescent. AJNR Am J Neuroradiol 1995;16:1548–50.
[7] Lasjaunias PL. Segmental identity and vulnerability in cerebral arteries. Interv Neuroradiol 2000;6:113–24.
[8] Kanemoto Y, Hisanaga M, Kanoh M, et al. Congenital hemicerebral arterial ectasia complicating unilateral megalencephaly. Br J Radiol 1987;60:395–400.
[9] Wolpert SM, Carter BL, Fenns EJ. Lipomas of the corpus callosum. An angiographic analysis. Am J Roentgenol 1972;115:92–9.
[10] Hanakita J, Miyake H, Nagayasu S, et al. Surgically treated cerebral arterial ectasia with so-called moyamoya vessels. Neurosurgery 1986;19:271–3.
[11] Lanternier LA, Brembilla C, Grimi P. Letter to the editor. Pure arterial malformation of the posterior cerebral artery. J Neurosurg 2014;121:1007–8.
[12] Araki Y, Mori S, Kanoh M, et al. Congenital hemispheric arterial ectasia with so-called moyamoya vessels. Surg Neurol 2018;129:91–9.
[13] Tsukamoto Y, Nakata H, Suyama T, et al. Bilateral pericallosal arterial ectasia. Neuroradiology 1985;27:271–4.
[14] Baccin CE, Krings T, Alvarez H, et al. A report of two cases with dolichosegmental intracranial arteries as a new feature of PHACES syndrome. Childs nNerv Syst 2007;23:559–67.
[15] Sako T, Uchino A, Saito N. Pure arterial malformation of the posterior inferior cerebellar artery diagnosed by MR angiography. Neuroradiol J 2016;29:283–5.
[16] Beringer W, Aklenghat J. Pericallosal artery ectasia with associated stenosis. AJNR Am J Neuroradiol 2004;25:1197–8.
[17] Sacks JG, Lindenburg R. Dolicho-ectatic intracranial arteries: syndromatopathy and pathogenesis of arterial elongation and distention. Johns Hopkins Med J 1969;125:93–106.
[18] Hyogo T, Nakagawa J, Nakamura J, et al. Multiple segmental agenesis of the cerebral arteries: case report. Neuroradiology 1986;38:433–6.
[19] Thompson JR, Weinstein PR, Simmons CR. Cerebral arterial dolichoectasia with seizure. Case report. J Neurosurg 1976;44:509–12.
[20] Feliciano CE, Pamias-Portalatin E, Mendoza-Torres J, et al. Color-coded digital subtraction angiography in the management of a rare case of middle cerebral artery pure arterial malformation. A technical and case report. Interv Neuroradiol 2014;20:715–21.
[21] Metry DW, Dowd CF, Barkovich AJ, et al. The many faces of PHACE syndrome. J Pediatr 2001;139:117–23.
[22] Lee C, Pennington MA, Kenney CM3rd. MR evaluation of developmental venous anomalies: medullary venous anatomy of venous angiomata. AJNR Am J Neuroradiol 1996;17:61–70.
[23] Shankar JJ, Banerjee ST, Hogan M, et al. A rare case of cerebral cortical dysplasia with arterial vascular dysplasia. Can J Neurol Sci 2009;36:757–60.
[24] Kryst-Widgoswka T, Kozlowski P, Bienkiewicz M, et al. The angiographic and scintigraphic picture of dolichoectasia of the anterior cerebral artery. Eur J Nucl Med 1980;5:387–9.
[25] McLaughlin N, Duckwiler G, Martin NA. Pure arterial malformations of the posterior cerebral artery. Response. J Neurosurg 2014;120:575–7.
[26] Lanzino G, Burrows AM, Flemming KD, et al. Pure arterial malformations of the posterior cerebral artery. J Neurosurg 2014;120:575.
[27] Caplan LR. Dolichoectasia (dolicho-ectasia). What is known and not known. Ann Neurol 2005;57:469–71.
[28] Lou M, Caplan LR. Vertebrobasilar dolichoarteriopathy (dolichoectasia). Ann NY Acad Sci 2010;1184:121–33.
[29] Chao KH, Riina HA, Heier L, et al. Endovascular management of dolichoectasia of the posterior cerebral artery report. AJNR Am J Neuroradiol 2004;25:1790–1.
[30] Kwon HM, Lee YS. Dolichoectasia of the intracranial arteries. Curr Treat Options Cardiovasc Med 2011;13:261–7.
[31] Pico F, Jacob MP, Labreuche J, et al. Matrix metalloproteinase-3 and intracranial arterial dolichoectasia. Ann Neurol 2010;67:508–15.
[32] Pico F, Biron Y, Bousser MG, et al. Concurrent dolichoectasia of basilar and coronary arteries. Neurology 2005;65:1503–4.
[33] Duncan IC, Terblanche JM. Spontaneous isolated posterior communicating artery dissection in a young adult with hyperhomocysteinemia. AJNR Am J Neuroradiol 2005;26:2030–2.
[34] Krings T, Choi IS. The many faces of intracranial arterial dissections. Interv Neuroradiol 2010;16:151–60.
[35] Pierot L, Gawlitza M, Soze S. Unruptured intracranial aneurysms: management strategy and current endovascular treatment options. Expert Rev Neurother 2017;17:977–86.
[36] Raper DM, Webster Crowley R, Liu KC, et al. Endovascular techniques and devices for the treatment of intracranial aneurysms: a review of neurointerventional outcomes. J Neurosurg Sci 2016;60:104–15.
[37] Pico F, Labreuche J, Cohen A, et al. Intracranial arterial dolichoectasia is associated with enlarged descending thoracic aorta. Neurology 2004;63:2016–21.