Predictive Role of External Carotid Artery Vasospasm on Cerebral Ischemia After Subarachnoid Hemorrhage: Experimental Study

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ABSTRACT

AIM: Cerebral vasospasm after subarachnoid hemorrhage (SAH) may lead to a devastating neurological outcome by inducing cerebral ischemia. However, the role of external carotid artery (ECA) vasospasm has been rarely reported in the literature. The aim of this study was to elucidate the effect of ECA vasospasm on cerebral ischemia related neurodegeneration in the cerebral cortex after SAH.

MATERIAL and METHODS: This study was performed on 23 rabbits, divided into three groups: control (n=5), sham (n=5), and SAH (n=13). Experimental SAH was performed by injecting 0.75 mL auricular arterial homologous blood into the cisterna magna. After three weeks, the animals were decapitated and the common carotid arteries with their external and internal branches and the brains were examined histopathologically. Vasospasm indexes (VSI) of ECAs and internal carotid arteries (ICAs) and degenerated glial cell numbers of temporal cortices (n/mm³) were estimated stereologically and the results were compared statistically.

RESULTS: Temporal cortex glial cell density was estimated as 136.950±9.257/mm³ in normal rabbits, 131.324±7.987/mm³ in sham, 112.320±6.112/mm³ in light, and 97.543±5.432/mm³ in severe ECA vasospasm. The mean VSI values of ECA of all groups were 1.95±0.21, 2.15±0.29, 2.95±0.65 and 3.12±0.276 respectively. Statistical differences between the VSI values of ECA and degenerated neuron densities in temporal cortices were significant (p < 0.005).

CONCLUSION: ECA vasospasm was observed to have a more important predictive role on the serious cerebral ischemia and neuronal degeneration after SAH. The mechanism may be related to ischemia of the parasympathetic ganglia of the lower cranial nerves and dorsal root ganglion.

KEYWORDS: Cerebral ischemia, Cerebral vasospasm, External carotid artery vasospasm, Subarachnoid hemorrhage

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ubrarachnoid hemorrhage (SAH) accounts for 5-10% of all cerebrovascular accidents. Cerebral vasospasm is the vasoconstriction of the arteries of the circle of Willis leading to secondary cerebral ischemia. Symptomatic vasospasm occurs 4-12 days after SAH with a peak incidence between 6 and 10 days. The predictors of cerebral vasospasm are the volume, density and prolonged presence of subarachnoid blood. The impairment of cerebral autoregulation and decreased blood flow results in delayed ischemic injury. Angiographic vasospasm is identified in 70% of the patients while 40% have clinical vasospasm, and delayed ischemic injury is seen in 30% of the patients (21, 48).

The pathogenesis of the cerebral vasospasm is still not fully understood. Vasoconstriction of the pial vessels with smooth muscle and myofibroblast proliferation with cellular necrosis and remodeling, intimal hyperplasia, collagen deposition and fibrosis as well as intraluminal platelet aggregation with microthrombus formation, arterial thrombosis and distal embolization have been described (21). Impairment of the nitric oxide pathway and neuronal nitric oxide synthase containing neurons in the adventitia is one of the mechanisms (29). Research has shown the inflammatory role of the cytokines, cell adhesion molecules, and leukocytes on the development of cerebral vasospasm (16).

SAH causes cerebral vasospasm through various neurohumoral mechanisms. Cerebral arteries are innervated by several systems contributing to the autonomic control of cerebral blood flow. Parasympathetic fibers have vasodilatory effects, and sympathetic fibers are vasospastic on cerebral arteries. The neuron density of peripheral nerve ganglia plays a regulatory role in these functions (9, 25).

Cerebrovascular sensory nerves are mainly from the trigeminocerebrovascular system. The trigeminal sensory nerves project to the ipsilateral internal carotid artery (ICA), middle cerebral artery (MCA), anterior cerebral artery (ACA), the rostral part of the basilar artery, the posterior cerebral artery (PCA), and the posterior communicating artery (11, 35).

Cerebral ischemia and infarction lead to poor outcome and death. The use of nimodipine, hemodynamic treatment, endovascular procedures, and the recent use of endothelin antagonist for prevention of vasospasm have improved the prospects for these patients but outcomes remain poor (47).

External carotid artery (ECA) vasospasm has not been much defined in the literature. In case of cerebral vasospasm, the collateral system helps to save the brain and initiates its self-treating process. However, external carotid spasm together with cerebral vasospasm impairs this collateral salvation network and leads to neurodegeneration of the temporal area supplied by both the internal and external carotid system.

This study investigates the pathogenesis of the cerebral vasospasm with an emphasis on ECA vasospasm and the relationship with decreased parasympathetic activity in order to help develop new therapeutic methods.

MATERIAL and METHODS

This study has been conducted on 23 rabbits. The animals were divided into 3 groups: SAH (n=13), sham (n=5) and control (n=5) group. The animal protocols were approved by the Ethics Committee of Erzurum Ataturk University, Medical Faculty. The care of the animals and the experiments themselves were conducted according to the guidelines set forth by the same ethics committee. A balanced, injectable anesthetic was used in order reduce pain and mortality. After anesthesia was induced with isoflurane given by a facemask, 0.2 mL/kg of the anesthetic combination (Ketamine HCL, 150 mg/1.5 mL; Xylazine HCL, 30 mg/1.5 mL; and distilled water, 1 mL) was subcutaneously injected before surgery. During the procedure, a dose of 0.1 mL/kg of the anesthetic combination was used when required. Autologous blood (0.75 mL) was taken from the auricular artery and injected using a 22-gauge needle into the cisterna magna of animals in the SAH group over the course of 1 minute. In the sham group, 1 mL of physiological saline was injected into the cisterna magna. Electrocardiography and respiratory rhythm parameters were monitored once/day and animals were decapitated after 3 weeks. Their common carotid arteries and brains were extracted bilaterally and fixed with 10% formalin solution for four days. Sections of paraffin embedded brains at the levels of superior temporal gyri and common carotid arteries at the just post-bifurcation levels were taken and stained with hematoxylin & eosin (H&E) and terminal deoxynucleotidyl transferase dUTP nick end labeling (TUNEL) for SAH-related apoptotic damage detection and examined stereologically. In order to detect hypertrophy, vasospasm, endothelial apoptosis in the ECA and ICA, vasospasm index (VSI) was used. Vasospasm was observed by the development of smooth muscle cells hypertrophy, arterial wall enlargement, and luminal narrowing. The hypertrophied muscle cells and intimal edema cause thickening of the vessel wall surface. The proportion of ICA or ECA wall section surface values (wall ring) to lumen surface values was accepted as VSI. This index was calculated with the following formula: VSI = (πR²−πr²)/πr² = (R²−r²)/r². Low luminal surface values and high VSI index was accepted as prominent vasospasm.

The differences of VSI values of ECA and the temporal cortex cell number data were analyzed using a commercially available statistics software package (SPSS® for Windows v. 12.0, Chicago, USA). The Kruskal-Wallis and Mann-Whitney U tests were used for data analysis. Differences were considered to be significant at p<0.05.

RESULTS

One animal in the sham group and two animals in the SAH group died in the first postoperative week and the remaining animals were followed for 21 days. Fever, meningeal irritation, consciousness, convulsions, apnea, cardiac arrhythmia, and breathing disturbances were observed frequently in the five of the surviving animals of the study group. Angiographic appearance of normal caroticovertebral arteries and histopathological appearance of a normal ECA-ICA (Figure 1) and of the ECA-ICA after SAH-without vasospasm were seen (Figure 2). There were no inflammatory changes in the arteries but intimal
derangement was observed in a rabbit with slight ECA vasospasm (Figure 3). Minimally convoluted inner elastic membrane, swelling endothelial and contracted smooth muscle cells, and thickened vessel walls were seen. Among animals with severe vasospasm, we observed intimal hypertrophy and prominent luminal narrowing in the ECA and ICA of a rabbit with severe temporal lobe neurodegeneration (Figure 4). A convoluted inner elastic membrane (IEM) and swelling of the endothelial and contracted smooth muscle cells (SMC), endothelial cell loss, luminal narrowing, endothelial basal lamina rupture, and thickened vessel wall were seen. Prominent vascular hypertrophy, endothelial and muscular and also adventitial proliferation were observed. The stereological neuron estimation method is summarized in Figure 5, and the VSI calculation method is shown in Figure 6 A, B. Figure 7 shows histopathological appearance of a normal temporal cortex, and histopathological appearance of temporal cortex with normal and apoptotic neurons with severe ECA vasospasm.

The cervical carotid artery and its branches were measured in 23 consecutive rabbits. The mean±SD size of the common carotid artery was 1.95±0.25 mm, the main trunk of the ECA 1.10±0.26 mm, and the ICA was 1.01±0.19 mm. Normal temporal cortex glial cells density was estimated as 136,950±9.257/mm³ in normal rabbits, 131,324±7.987/mm³ in sham, 112,320±6.112/mm³ in slight ECA vasospasm and 97,543±5.432/mm³ in severe ECA vasospasm. The mean VSI values of ECA were estimated as 1.95±0.21; 2.15±0.29; 2.95±0.65 and 3.12±0.276 respectively (Table I). Statistical differences between the VSI values of ECA and degenerated...
neuron densities in temporal cortices were significant (p < 0.005). The difference was more significant for the severe vasospasm group (p < 0.0001).

**DISCUSSION**

Cerebral vasospasm is a frequent complication after SAH contributing to overall morbidity and mortality. Various neuronal, humoral, and chemical factors are involved in cerebral vessel innervation (12, 14, 23).

Nitric oxide (NO) is a vasodilatory substance. However, NO may have a detrimental effect on the vascular system by reactive induction of inducible nitric oxide synthase (iNOS) in SAH (67). The peroxidation of membrane proteins by the nitric oxide metabolite may contribute to the morphological damage in chronic vasospasm (50). Calcitonin gene related peptide (CGRP) is a neurotransmitter with tachykinins in sensory fibers in the cerebral vasculature with possible vasodilating properties. The trigeminal cerebrovascular system may be involved in SAH (33). Neuropeptide Y-like immunoreactivity (NPY-L1) levels were increased compared to controls in patients with hemodynamic changes. CGRP-L1 levels were also increased in connection with vasospasm (32-34).

Parasympathetic fibers have vasodilation effect, and sympathetic fibers have vasoconstrictor effect on cerebral arteries (69). Dysfunction of neuronal signal processing and transmission occurs after SAH and contributes to the high morbidity and mortality. The underlying mechanism may be neuronal cell death. Direct influence of subarachnoid blood metabolites on neuronal signaling should also be considered (69).

Autonomic nerve fibers provide neural innervation. The post-ganglionic fibers of the ciliary ganglion of the third cranial

![Figure 5: VSI value estimation methods are seen in external carotid arteries (ECA) (H&E, x4) both normal and severe vasospasm developed animals following SAH.](image)

![Figure 6 A, B: Stereologic cell counting of the superior temporal lobe in a rabbit. A, B were taken from two parallel, adjacent thin sections separated by a distance of 5 µm. The upper and right lines represent the inclusion lines, and the lower and left lines are exclusion lines. The number of neurons from the two dissectors occurs in a volume given by the product of the counting frame area and the distance between the sections. The numerical density of the neurons is calculated as Nv=GN –ΣQ·GN/txA. The nucleoli marked with ‘2, 3, 4, 5, 7’ are dissector particles in A. Section B shows them as they disappeared. The nucleoli marked with ‘2, 3, 7, 8’ are not a dissector particle in A. Section B shows ‘1, 6, 8’ as it disappeared (TUNEL, x10).](image)
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Table I: Vasospasm Index (VSI) of External Carotid Artery (ECA) in Control, Sham, Slight and Severe ECA Vasospasm Group; and Temporal Cortex Glial Cell Number in These Groups Respectively (p < 0.005)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Control</th>
<th>Sham</th>
<th>Slight</th>
<th>Severe</th>
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<tbody>
<tr>
<td>VSI of ECA</td>
<td>1.95±0.21</td>
<td>2.15±0.29</td>
<td>2.95±0.65</td>
<td>3.12±0.276</td>
</tr>
<tr>
<td>Temporal cortex glial cells (/mm³)</td>
<td>136.950±9.257/mm³</td>
<td>131.324±7.987/mm³</td>
<td>112.320±6.112/mm³</td>
<td>97.543±5.432/mm³</td>
</tr>
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Figure 7: Histological appearance of a normal temporal cortex (LM, Tunnel, x4/Base), and histopathological appearance of temporal cortex with normal and apoptotic neurons with severe ECA vasospasm (LM, Tunnel, x10).

Cerebral vasospasm causes neuronal and astrocytic apoptosis after SAH. SAH causes ischemic insults to the vagal nerve ganglia and ischemic neurodegeneration in the petrosal ganglia resulting in hypertension by decreasing the number of neurons in these ganglia (52).

The cerebral vasculature, in particular the pial vessels, are densely supplied with noradrenergic sympathetic nerve fibers mainly originating in the superior cervical ganglion, accompanying the carotid artery, and projecting into the ipsilateral hemisphere. Ischemic injury of the upper cervical ganglia after SAH may cause vasospasm of the anterior spinal arteries (63).

Intracerebral hemorrhage causes descending neurodegeneration from the cortex to the dorsal root ganglion (DRG) (55). The low neuron density of the C3 dorsal root ganglion was reported to be related with severe vasospasm of anterior spinal artery (35). Retrograde neuronal death is well established in dorsal root ganglia after peripheral nerve injury or spinal cord trauma (4, 5). Upper cervical ganglia innervate anterior spinal arteries (ASA). Ischemic injuries after SAH may cause vasospasm at the ASA as a result of vasospasm of the arteries supplying DRG, leading to ischemic degeneration in the DRG. SAH results in neurodegeneration in DRG and C4 dorsal root ganglia (63). Low neuron density of the C4 dorsal root ganglia may be considered as an important factor in the pathogenesis of severe respiratory disturbances in SAH (63). Parasympathetic ischemia of the lower cranial nerve ganglia and DRG may have the major effect on the vasospasm of the external carotid artery. Spinal SAH should also be considered in the development of ECA vasospasm since the ASA vasospasm is related with the DRG ischemia (35).

High neuron density of the stellate ganglion can be considered as sympathetic overactivity that causes basilar vasospasm and neovascularisation, whereas low neuron density of stellate ganglion can be considered as sympathetic hypoactivity that prevents basilar vasospasm and causes vascular wall thinning after bilateral common carotid artery ligation (39).

In fatal SAH, respiratory muscles are paralyzed due to ischemia of vagal nerve and other upper and lower cervical nerve injuries. The parasympathetic system plays a major role in the continuation of spontaneous respiration. In SAH, vagal nerve roots are affected by vasospasm (6, 7, 10, 15).
A brief review of the literature discloses the importance of the neurohumoral mechanisms and autonomic nervous system on vasospasm after SAH but vasospasm of the extracranial arteries are rarely reported and discussed.

Clinically recurrent strokes due to transient vasospastic occlusions of the ICA have been observed, whereas recurrent vasospasms of the extracranial ICA are rare. ECA vasospasm after SAH has been reported even less commonly. Migrainous vasospasms of the extracranial ICA have been asserted to be associated with sympathetic vasomotor innervation and adrenoreceptor blockade was proposed as a treatment option (31, 68). In the angiography of a patient with posterior inferior cerebellar artery (PICA) aneurysm, fibromuscular dysplasia was observed in the external carotid and vertebral arteries (27).

After the rupture of an intracranial aneurysm, vasospasm of the extracranial as well as intracranial arteries was reported. Digital subtraction angiography revealed severe vasospasm of the external carotid arteries (46). Superior cervical ganglionectomy and perivascular sympathectomy of cervical ICA improved the symptoms. In 23 angiographically investigated patients, ECA vasospasm was observed in 50%. Pathogenesis of the vasospasm of the external carotid arteries was reported to be related with the sympathetic nerves (24).

Collateral system plays an important role in case of occlusion of the arteries. In cerebral vasospasm, this system is expected to be on stage. The suboccipital network consists of anastomoses of the cervical, vertebral, occipital, and carotid arteries. This arterial network has a salvaging effect in arterial occlusions. The suboccipital knot enlarges to provide collateral blood flow from subclavian to external carotid artery giving extravascular support to the brain by way of intraorbital ophthalmic anastomoses (2, 8). These important anastomotic vascular channels between the extracranial and intracranial arteries are located in three regions along the skull base: the orbital region via the ophthalmic artery between the internal maxillary artery (IMA) and ICA; the petrocavernous region via the inferolateral trunk, the petrous branches of the internal carotid artery, the meningoypophyseal trunk to the carotid artery; and the upper cervical region via the ascending pharyngeal, the occipital, and the ascending and deep cervical arteries to the vertebral artery (17, 28, 44).

In SAH patients, cerebral catheter angiography showed segmental vasoconstriction involving bilateral internal carotid, posterior cerebral and external carotid branches. External carotid artery branch involvement in reversible cerebral vasoconstriction syndrome was reported (58). Trapping of the ICA in the acute stage of the subarachnoid hemorrhage may result in ischemia secondary to hemodynamic hypoperfusion or occlusion of the perforating artery with delayed vasospasm. Evaluation of the ECA-ICA bypass is important (43).

Since the temporal lobe receives blood from both the carotid and the vertebrobasilar systems (41), vasospasm of both internal and external carotid arteries increases the morbidity impairing the collateral salvation system. Temporal lobe degeneration leads to psychiatric pathologies (20, 37). Bilateral damage to the medial temporal lobe has been reported to cause severe and lasting impairment in declarative memory. The affected structures are hippocampal region with the CA fields, the dentate gyrus, and the subicular complex and the adjacent perirhinal, entorhinal, and parahippocampal gyrus (45). Studies have shown psychiatric co-morbidities in temporal lobe epilepsy and temporal lobe atrophy in relation to Alzheimer’s disease (13, 22, 36, 51, 66).

In our study, ECA vasospasm together with ICA vasospasm has led to temporal lobe degeneration due to lack of collateral supply. Diagnosis and treatment of vasospasm is important at the early stage before the development of cerebral ischemia.

Our study has also shown that ECA vasospasm together with the intracerebral vasospasm have more deleterious effect on the brain, leading to neurodegeneration. We hypothesize that the ischemic degeneration of the lower cranial nerve ganglions and DRG have a major influence on the external vasospasm of the arteries. The sympathetic overdischarge during this phase with the depression of the parasympathetic activity should be also taken into consideration.

Medical treatment is the initial step when clinical vasospasm is suspected. Papaverine, Amrinone, Milrinone, Verapamil, Nimodipine, Nicardipine and Fasudil hydrochloride are pharmacological agents used in the treatment. Intrathecal nicardipine has been reported to be effective for the prevention and treatment of vasospasm (21). Memantine can prevent SAH by restoring endothelial nitric oxide synthase (29).

The vagal system has been proposed to modulate the inflammatory system through acetylcholine receptors. The neuroinflammatory reflex is vagally mediated that may relate to parasympathetic nervous system activation that suppresses inflammation.

Parasympathetic dysfunction coupled with catecholamine release may play a role in unchecked inflammation leading to myocardial dysfunction and cell death (49) and in the inflammatory cascade leading to vasospasm after SAH (16, 49).

At the early stage of SAH parasympathetic activity is enhanced (65). The acute activation was supposed to contribute to cardiac injury. Subcutaneous vagal nerve blockade at the beginning of SAH, whereas vagal nerve stimulation or sympathetic nerve blocks was proposed as a treatment method (1, 49).

ICA is innervated mainly by the superior cervical sympathetic nerve and the stimulation of this nerve results in cerebral vasoconstriction. Several reports have proposed cervical sympathectomy as a successful treatment method. Blood flow increase is observed especially in spastic vessels (24).

ECA may be spared during the acute stage of SAH, before the ischemia of the lower cranial parasympathetic and dorsal root ganglia with an early intervention.

Early endovascular treatment of cerebral vasospasm by mechanical dilatation with balloon angioplasty and pharmacological dilatation with intraarterial drug infusion or combination therapy is the treatment of choice (21).
Flow augmentation strategies are used as a novel and alternative therapeutic option for SAH-induced cerebral vasospasm (42). A dual balloon catheter designed to increase cerebral blood flow (CBF) via partial occlusion of the aorta above and below the origin of the renal arteries, intraaortic balloon pump counter pulsation in patients refractory to triple-H therapy are reported as new methods of treatment (21).

Transcutaneous electrical nerve stimulation (TENS) and spinal cord stimulation have been shown to influence peripheral and CBF through a sympathetic pathway (61, 62,64).

Sphenopalatine ganglion stimulation activates perivascular vasodilatory nerves in the ipsilateral anterior circle of Willis. Sphenopalatine ganglion stimulation was reported to decrease vasospasm and increase CBF after SAH in monkeys and it was associated with opening of the blood-brain barrier (59).

Cervical sympathetic block can relieve cerebral vasospasm after SAH and increase NO content and NOS activity in plasma and cerebrospinal fluid to promote neural functional recovery (19).

The neuron number of stellate ganglia may play an important role in the regulation of basilar artery volume. Stellate ganglia ablation may prevent severe vasospasm after SAH (39).

Stimulation of the ophthalmic branch of the trigeminal ganglion may be a therapeutic option in the prevention of severe vasospasm in SAH (54).

Radial artery graft bypass with ICA sacrifice was considered as a treatment method in the acute phase of SAH (40). ECA-radial artery-M2 segment of the MCA bypass was used for the treatment of large or giant ICA aneurysms (30). Revascularization technique with direct and indirect bypass surgery has been used for the treatment of ischemic neurological symptoms (38).

Extracranial–intracranial bypass is performed in patients with atherosclerotic MCA occlusion, ischemic symptoms or poor cerebral hemodynamics unresponsive to medical treatment (18, 30). The ischemic area surrounding the cerebral infarction in the eloquent area was reported to be saved by superficial temporal artery (STA)-MCA bypass recently (26). STA-MCA anastomosis is a direct bypass considered to be effective for the acute improvement of cerebral hemodynamics (70). The rapid increase of the CBF due to direct bypass in the acute stage of cerebral infarction was thought to cause cerebral edema, hemorrhage, and reperfusion injuries. However, these complications were not observed in the recently reported studies (70).

Vasospasm of venous grafts in extra-intracranial bypass was observed, though the patients were treated successfully with calcium antagonists as well as hypertensive and hypervolemic medication (57).

STA–MCA and STA–ACA double anastomosis with pansynangiosis was reported to have excellent outcome. Double bypass with indirect bypass or single bypass in the treatment of acute stage of ischemia may prevent the neurological deterioration (71).

ECT stenting with STA–MCA anastomosis is regarded as a good therapeutic option with symptomatic ICA occlusion and severe stenosis of the ipsilateral ECA (53).

External carotid spasm together with intracranial spasm has a more severe neurodegenerative effect than cerebral vasospasm alone. Medical and invasive methods have been proposed in the treatment of vasospasm after SAH. The important point is to intervene as early as possible in the acute stage of SAH, when the parasympathetic activity is dominant and yet collaterals are not impaired.

The non-invasive, functional neurosurgical treatment methods increasing the parasympathetic activity or endovascular and bypass techniques may be life-saving in the early period before the ischemia develops and infarct settles.

■ CONCLUSION

ECA vasospasm is an important predictive factor on the serious cerebral ischemia and neuronal degeneration after SAH that has not been extensively mentioned in the literature. We theorized that decreased vasodilatory functions of cervical dorsal root ganglia, trigeminal-glossopharyngeal-vagal nerves and/or increased vasospastic effects of cervical sympathetic ganglia may be responsible for this phenomenon.

■ REFERENCES


