Fulminant Superior Ophthalmic Vein and Cavernous Sinus Thrombophlebitis with Intracranial Extensions: Case Reports

Cavernous sinus thrombophlebitis (CST) is a rare and life-threatening disease with- out prompt diagnosis and treatment. Two cases of fulminant superior ophthalmic vein (SOV) and CST caused by maxillary periodontitis and sphenoid sinusitis are described. A 65-year-old woman presented with right proptosis, headache, and fever. A 74-year-old woman presented with left periorbital swelling. In both patients, MRI with gadolinium showed expansion of the bilateral cavernous sinus and diffuse dilatation of the SOV with non-enhancement of central thrombus, which indicated CST. The condition was complicated by brain abscess, meningitis, and ischemic stroke. These conditions were improved by antibiotic treatment, but one patient underwent exen- teration of the orbit due to orbital rupture during hospitalization.

CASE REPORT

A 65-year-old female was hospitalized after a 2-week history of right-sided proptosis, headache, and fever. The patient was suffering from toothache since several weeks. Physical examination revealed right-eye proptosis with limitation of eyeball movement. The leukocyte count was 15600/mm³ and the C-reactive protein level was 21.3 mg/dL. Orthopantomography revealed a periapical abscess of the left secondary maxillary molar.
MR imaging was performed. The contrast-enhanced three-dimensional fast spoiled gradient-echo images showed multifocal, irregular filling defects within an expanded, enhanced CS (Fig. 1A). The left SOV was dilated in the absence of enhancement (Fig. 1B). Both the CS and the left SOV exhibited restricted diffusion (Fig. 1C). We suspected that the periapical abscess was the primary source of infection of the SOV and CST, and commenced parenteral broad-spectrum antibiotics after tooth extraction. *Parvimonas micra* was identified on blood culture. Follow-up contrast-enhanced T1-weighted spin echo images [repetition time (TR)/echo time (TE), 600/9.3] showed that the CST was controlled but brain abscesses developed on the left anterior temporal convexity at 7 days after admission (Fig. 1D). The patient continued to take antibiotics for 1 month and was then discharged.

A 74-year-old female presented with periorbital swelling and
purulent discharge from the left eye. She was suffering from diabetes mellitus since 20 years. Her body temperature was 37.2°C and periorbital edema was evident. The leukocyte count was 18880/mm³ and the C-reactive protein level was 16.6 mg/dL.

A post-contrast enhanced CT scan revealed left-sided proptosis, a gas-containing abscess in the preseptal region, and left sphenoid sinusitis (Fig. 2A). MR imaging was performed. The contrast-enhanced T1-weighted fat-suppressed spin-echo image (TR/TE, 816.6/9) showed diffuse dilatation of the left SOV and a non-enhanced central thrombus (Fig. 2B). Multiple abscesses were evident in the preseptal region and the intraconal space. CST was noted, along with thick enhancement of the dura along the left frontotemporal cerebral convexity.

Antibiotic therapy was commenced and patient underwent exenteration of the left orbit due to orbital rupture on the third hospital day. Corneal culture yielded Actinomyces. Aspergillus was cultured from a fungal ball from the left sphenoid sinus, and from the specimen obtained after exenteration. Therefore, antifungal agents were added. However, patient experienced a sudden episode of dysarthria and right-sided weakness at 15 days after admission. Multifocal acute infarctions in the left basal ganglia and periventricular white matter were identified on diffusion-weighted image (Fig. 2C). Combination treatment with antiplatelet and antimicrobial agents was continued for 6 weeks, and the patient recovered and was discharged.

DISCUSSION

The CSs are heavily trabeculated venous sinuses located along the sides of the sella turcica, extending from the anterior superior orbital fissure posteriorly to the petrous portion of the temporal bone. The CSs have extensive vascular connections that are valveless, rendering the CSs prone to septic thrombosis after infection (2). Paranasal sinusitis has become the most frequent cause because complicated facial infections have been effectively reduced by antibiotic treatment (3, 4). Two-thirds of cases in the antibiotic-era develop chronic residual neurological deficits including ocular palsy, blindness, hemiparesis, or pituitary insufficiency (4). Because of the risk of such poor outcomes, if CST is clinically suspected, it is imperative that prompt treatment should be given and careful evaluation including radiographic studies should be considered.

When paranasal sinusitis causes CST, as in one of our patients, sphenoid sinusitis is the common cause. This occurs, first, via direct spread of an infection through communicating veins, facilitated by osteomyelitis of the intervening diploic bone or by mucosal breach caused by bony defects (5). Second, indirect spread occurs via the ophthalmic veins after penetration of the lamina papyracea, as in our cases. The latter route of infection is more common (6). Recently, oral and dental infections have been described as uncommon causes of CST. Infections can extend from the maxillary molar to enter the orbit through the inferior orbital fissure, eventually spreading to the CSs (1).

The organism most commonly isolated from CST patients is Staphylococcus aureus (69%) (1). Also, Rhinocerebral mucormycosis and Aspergillus can cause CST in immunocompromised and diabetic patients (7). Mixed infections involving beta-hemolytic streptococci, other streptococci, and anaerobes, are common in patients with oral and dental infections (1). Actinomycosis rarely extends to the CSs, either direct extension from the ear or sinuses or hematogenous spread may play a role (8).

The most common presentations of CST patients include fever, proptosis, and external ophthalmoplegia triggered by secondary dysfunction of cranial nerves III, IV, and VI (4). Moreover, intracranial extension of infections can develop, triggering pituitary insufficiency, meningitis, cortical vein thrombophlebitis with subsequent hemorrhagic infarction, subdural empyema, or development of brain abscesses (3).

CT and MRI are the primary imaging modalities to confirm the suspected origin of the infectious process and to promptly detect fatal intracranial complications of CST. Multiple filling defects within the expansive contours of enhanced CSs, caused by a thrombus, are direct signs. Filling defects in CST tend to be larger in size and show non-fat density compared with intracavernous fat deposits. Also, the thrombus is irregular in shape, whereas neural structures in CSs are either longitudinal or round to oval and confined to the lateral wall (9).

Indirect signs are related to concomitant venous obstruction, including dilatation of the SOVs, exophthalmos, and abnormal enhancement of the dura along the lateral margins of the CSs (7, 8). SOV thrombophlebitis is one of the well-known imaging findings of CST, which was present in all of our cases. Brain abscesses developed on the left anterior temporal convexity in the first case. According to Mahapatra (10), if CST is complicated by
cerebral abscesses, the common locations of such abscesses are the frontal and temporal lobes.

Management of CST patients includes prompt and aggressive administration of parenteral antibiotics targeting the most common pathogens associated with primary infection, and such a therapy should be maintained for at least 2 weeks after clinical resolution (1). Surgical treatment is reserved for patients with primary sinusitis, dental infection, or complicating brain or orbital abscesses. The use of corticosteroids to treat cranial nerve dysfunction has been reported (4), and adjuvant anticoagulation to prevent thrombus propagation can be helpful (1).

In conclusion, although CST has become a rare disease, it remains potentially lethal, triggering life-threatening complications, long-term sequelae, and significant morbidity. Rapid establishment of the diagnosis of CST using contrast-enhanced CT scans and MR studies with commencement of prompt medical and surgical treatment are crucial.

REFERENCES

두개 내 합병증을 동반한 전격성 상안와 정맥, 해면정맥동 혈전성 정맥염: 증례 보고

황지선 1·홍현숙 1·박지상 1·이아름 1·추은주 2·장기현 1

해면정맥동 혈전성 정맥염은 현재 드물게 발생하는 질환이지만 즉각적인 진단과 치료가 시행되지 않을 경우 생명을 위협할 수 있다. 저자는 상악 치주염과 접형동 부비동염에서 발생한 상안와 정맥과 해면정맥동의 혈전성 정맥염 2예를 보고하고자 한다. 우안의 안과출혈증과 두통, 발열을 주소로 내원한 65세 여자 환자와 좌안의 안과 주위 부종을 주소로 내원한 74세 여자 환자의 자기공명영상에서 양측 해면정맥동과 상안와 정맥이 폐쇄되고 내부에 조영증강 되지 않는 저신호강도를 보여 해면정맥동 혈전증이 진단되었다. 환자들은 합병증으로 뇌농양, 뇌막염, 허혈성 경색이 발생하였다. 이는 항생제 치료 후 호전되었으나 한 환자는 안과내로 인해 안과내용물제거술을 시행받았다.

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