Subdural Empyema as a Complication of Sinusitis: A Diagnosis to Keep in Mind

Rosário Eça 1, Armando Graça 2, Rita Francisco 2, Jaime Pamplona 3

1. Internal Medicine, Centro Hospitalar de Lisboa Central, Lisboa, PRT
2. Critical Care, Centro Hospitalar de Lisboa Central, Lisboa, PRT
3. Radiology, Centro Hospitalar de Lisboa Central, Lisboa, PRT

Corresponding author: Rosário Eça, rosarinho.eca@gmail.com

Abstract

Subdural empyema (SDE) is a rare form of intracranial infection associated with a high morbidity and mortality rate. Infections of the middle ear and paranasal sinuses are the most common predisposing factors that can lead to bacterial proliferation in the subdural space, usually by direct extension in young patients. Clinicians must have a high level of suspicion for patients presenting with concomitant neurological deficits and signs of sinus pathology. Cross-sectional imaging is mandatory for the diagnosis, preferably contrast-enhanced magnetic resonance imaging. Treatment requires a prolonged course of intravenous antibiotic therapy and prompt neurosurgical drainage intervention.

Here, we present the case of a 20-year-old patient with long-term neurological sequelae following a left paranasal infection complicated by an SDE. This case report highlights the rapid progression and devastating consequences of SDE, an ominous neurosurgical emergency.

Categories: Internal Medicine, Emergency Medicine, Infectious Disease
Keywords: cross-sectional imaging, neurologic deficit, sinusitis, subdural empyema, intracranial infections

Introduction

Subdural empyema (SDE) is a rare acute neurological condition accounting for 20% of localized intracranial infections, mostly occurring as a life-threatening complication of sinusitis, otitis, mastoiditis, trauma, or surgical interventions [1]. It represents a loculated pyogenic infection, usually unilateral, located between the dura mater and the arachnoid that can potentially spread over the convexity of the brain and the midline to the contralateral hemisphere [2]. SDE is more common in the second decade of life and in males (male-to-female ratio of 3:1), although incidence has been decreasing over the last decades due to the use of effective antibiotics [2,3].

SDE is a challenging clinical diagnosis because of its non-specific presentation and rapid progression. The symptoms are reflective of increased intracranial pressure, meningeal irritation, and cerebritis, with the most common being fever, headache, vomiting, altered mental status, and focal neurological deficits [3,4]. Time is of the essence, with early treatment increasing the chances of a better outcome, usually requiring both medical and surgical treatment. Although early aggressive treatment of SDE empyema has reduced the mortality rate, it can be associated with significant long-term neurological morbidity [1,5].

Case Presentation

A 20-year-old male was admitted to our emergency department with a five-day history of frontal headache and fever, associated with nausea, vomiting, and photophobia since the previous day. He had a history of uncomplicated sinus disease, and his immunizations were up to date. He lived at home with his parents, none of whom were ill.

At admission, the physical examination revealed a lethargic but arousable patient, oriented to person but not to space or time. The patient had a fever (38.3°C), and his vital signs were stable. His neurologic examination was positive for mild right hemiparesis (left strength and reflexes were normal). The remainder of his physical examination was otherwise normal, and there were no signs of nuchal rigidity. Laboratory workup was remarkable for mild leukocytosis and elevated C-reactive protein. Urgent non-contrast head computed tomography (CT) was relevant for left pansinusitis with left frontal and parafalcine SDE causing a slight contralateral mass effect. A subsequent lumbar puncture showed an elevated white blood cell count of 214 per mm$^3$ (with 60% neutrophils) with no microorganisms isolated.

On the following day, his condition significantly worsened with gradually deteriorating mental status to a semi-comatose state and seizures. Afterward, he was admitted to the intensive care unit (ICU), and non-contrast-enhanced magnetic resonance imaging (MRI) confirmed the presence of an SDE with a slight mass effect (Figure 1).
After neurosurgical consultation, an emergent left frontotemporoparietal craniectomy was performed with the evacuation of a large amount of purulent material. Pus culture isolated a *Streptococcus intermedius* and *Enterococcus faecalis* infection and metronidazole was added to the already empirically introduced broad-spectrum antibiotics (ceftriaxone).

During the next two weeks in the ICU, the patient’s condition slowly recovered, and follow-up imaging showed a favorable progression. In the third week post-admission, the patient’s clinical condition relapsed with fever and rising inflammatory laboratory parameters; urgent contrast-enhanced MRI was consistent with relapse of the SDE with concerning findings for frontal bone osteomyelitis (Figure 2).
FIGURE 2: Contrast-enhanced cranial MRI three weeks post-craniecotomy.

T1 fat-saturated image after intravenous contrast showing a large midline extra-axial fluid collection consistent with relapsed subdural empyema.

MRI: magnetic resonance imaging

The patient was taken to the operating room for a repeated evacuation of the abscess. No changes were made to the antibiotic regimen being administrated (ceftriaxone plus metronidazole). Afterward, his condition improved, and two months later, he was transferred to the rehabilitation medical floor for continued care.

At the time of discharge (six months post-admission), the patient exhibited persistent mild right-sided hemiparesis and expressive aphasia, and the pre-discharge MRI revealed extensive left cortico-subcortical encephalomalacia with stable brain herniation through the craniectomy defect and passive left ventricle enlargement (Figure 3). There was no clinical improvement of the neurological sequelae at the clinical evaluation eight months post-discharge.
Discussion

Although most cases of sinusitis have a mild and uncomplicated natural history, rarely intracranial complications such as meningitis, empyema, brain abscess, or osteomyelitis can occur [3,4]. It is estimated that the risk of developing suppurative intracranial infection derived from sinusitis or otogenic infections is <0.1% [3]. SDE secondary to sinus pathology can present as an insidious process, with the frontal sinus most frequently implicated, whether by direct extension or retrograde thrombophlebitis via the valveless diploe veins [3,4]. A history of sinus disease is not mandatory as noted by Germiller et al. [6]. In a study of 25 children and adolescents with intracranial complications of sinus disease, only four had a significant history of sinus disease. Germiller et al. [6] also noted that most patients (59%) had a normal neurological examination at presentation and that neurologic deficits at presentation have prognostic significance [6].

Early symptoms of SDE are usually non-specific characterized by acute febrile illness, accentuated by acute neurological deterioration, and, if left untreated, coma and death. Laboratory workup is also non-specific but can show raised inflammation markers. Cross-sectional imaging has improved the rapid diagnosis of intracranial complications and is a cornerstone of the diagnosis of intracranial complications. Although MRI is more sensitive and accurate, head CT is usually done first because of its wide availability and rapid acquisition time [2]. The classic appearance of SDE on a CT scan is of a thin unilateral crescent fluid density collection over the cerebral convexity or along the falx with a rim of contrast enhancement. Of note, SDE exerts a mass effect that can be disproportionate to its volume due to accompanying cerebral edema and ischemia [1,3].

SDE secondary to paranasal sinus infection is commonly polymicrobial, more frequently by anaerobic and microaerophilic streptococci, in particular, those of the Streptococcus milleri group as in our patient (Streptococcus intermedius) [1,5,6].

Left untreated, SDE is invariably fatal. Since the introduction of antibiotics, the mortality rate has decreased but is still considerable, of up to 28% of patients [3,5]. Treatment of SDE includes broad-spectrum antibiotics, for example, third-generation cephalosporin, along with metronidazole and vancomycin to provide appropriate coverage against pathogens that commonly colonize the upper respiratory tract [3]. Appropriate antibiotic therapy should be tailored to positive culture results and continued for up to six weeks [3]. In the case of intracranial abscesses, emergent surgical drainage is needed to lower intracranial pressure, allowing for the collection of intraoperative cultures [1–3]. Relapse of the pus collection in the subdural space is a known complication, and CT should be repeated if unexplained neurological deterioration occurs [2,4]. Prognosis is mostly dependent on the preoperative mental status, presence of neurological deficits, and aggressiveness of the early treatment. An unfavorable prognosis is more common in older or comatose patients, and up to half of the patients can remain with permanent residual neurological deficits, particularly in older patients [2]. Neurological deficits were present at presentation in our patient, but, unfortunately, there was some initial mismanagement, which led to a potentially hazardous
lumbar puncture and a delayed neurosurgical consultation after imaging findings were consistent with a diagnosis of subdural collection.

**Conclusions**

Although paranasal sinusitis usually has a benign course, on rare occasions, it can lead to serious and potentially life-threatening complications such as SDE, which should be ruled out in patients with signs of sinusitis and neurological deficits. It is important to note that a history of previous sinus disease is not invariably present, especially in young patients. Early imaging is critical for accurate diagnosis, with successful treatment requiring a combination of aggressive medical and surgical intervention.

**Additional Information**

**Author Contributions**

All authors have reviewed the final version to be published and agreed to be accountable for all aspects of the work.

**Concept and design:** Rosário Eça

**Acquisition, analysis, or interpretation of data:** Rosário Eça, Armando Graça, Rita Francisco, Jaime Pamplona

**Drafting of the manuscript:** Rosário Eça

**Critical review of the manuscript for important intellectual content:** Armando Graça, Rita Francisco, Jaime Pamplona

**Supervision:** Armando Graça, Jaime Pamplona

**Disclosures**

**Human subjects:** Consent was obtained or waived by all participants in this study. **Conflicts of interest:** In compliance with the ICMJE uniform disclosure form, all authors declare the following: **Payment/services info:** All authors have declared that no financial support was received from any organization for the submitted work. **Financial relationships:** All authors have declared that they have no financial relationships at present or within the previous three years with any organizations that might have an interest in the submitted work. **Other relationships:** All authors have declared that there are no other relationships or activities that could appear to have influenced the submitted work.

**References**