INTRODUCTION

Infectious disease of the central nervous system (CNS), such as meningitis, is one of the most important categories of neurological conditions. Such diseases are challenging in terms of diagnosis, treatment, and prognosis. Meningitis occurs most often in young- and middle-aged patients, and the rate of misdiagnosis, especially during the early stages of the disease, is high. Correct diagnosis and treatment can save lives, but many factors, such as difficulty in medications crossing the blood-brain barrier (BBB) can cause difficulty in treatment.

ANALYSIS OF THE MISDIAGNOSIS RATE OF INFECTIOUS MENINGITIS

There are several reasons for the high rate of misdiagnosis in meningitis: (1) the development of this clinical sub-specialty is relatively recent. The professional setting and staffing for research into infectious CNS diseases fall behind those for research into cerebrovascular, demyelinating, neuromuscular, degenerative, and genetic diseases. At present, only a few hospitals in China, such as in Beijing, Shanghai, Xi’an, Shijiazhuang, and Yinchuan, have infection and cerebrospinal fluid cytology (CSFC) as a professional sub-specialty. (2) The content of current textbooks is out of date. It lacks data from large, double-blind, multicenter, case-control studies, as well as basic research data. The data on meningitis in current textbooks are about 40–50 years old, and some are from foreign studies, whereas the chapters on cerebrovascular disease are updated every 5–10 years. (3) The clinical manifestations of meningitis can be atypical. Meningitis has shown the greatest change in clinical manifestations over time compared with other nervous system diseases. “Atypical clinical manifestations” may, in fact, be the current “typical” features, but are different from the manifestations seen 50 years ago. The 50-year fight between the meningitis pathogen and the human immune system, the natural variation of the pathogenic organisms, the misuse of antibiotics and immune suppressants, and the prevalence of drug addiction and AIDS have all produced changes in the clinical manifestations of meningitis. (4) Obtaining samples of the pathogen is difficult. Taking a biopsy from the mater has technical limitations, and it may be difficult for the patient and family to accept. The positivity rate for pathogens in CSF is very low, except for Cryptococcus neoformans, which can be 99% in most references.[1,2] The textbooks state that the positivity rate for Mycobacterium tuberculosis by smear and culture of bacteria from CSF can be as high as 30%–40%, but clinical reports from most hospitals show a positive rate of below 10%. Of 167 patients with tubercular meningitis who were assessed during the period 1990–2010 in our hospital, only one had a positive result for M. tuberculosis by bacterial smear and culture from CSF samples.[3] Using the new acid-fast stain method of The Fourth Military Medical University, the positivity rate of CSF smear can be above 90%, enabling early diagnosis of tubercular meningitis.[4] Identifying the pathogens underlying cases of viral meningitis, weakly pathogenic bacterial meningitis, and parasitic meningitis is also difficult. For these reasons, the misdiagnosis rate in early meningitis is very high.

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DIFFICULTIES IN DIAGNOSIS AND TREATMENT OF INFECTIOUS MENINGITIS

The BBB protects the brain tissue, but is also the greatest obstacle to treatment of infectious meningitis. The focus is often on the effective permeation rate of treatment drugs across the BBB, instead of the sensitivity of the drug, which can influence the efficacy of treatment.\[^5\]

High doses and a long treatment course are needed for therapy of infectious meningitis. For example, an intensive course of treatment for tubercular meningitis requires a dose of isoniazid of 15 mg/kg per day, whereas the general dose in instruction is 0.6 g/day. The treatment course for intracranial tuberculosis is double that for extracranial tuberculosis, that is, 4–6 months of intensive treatment and 18–24 months for the whole course.\[^6\]\[^9\] Such doses and treatment courses pose challenges for both doctors and patients.

Because of the problems of pathogen isolation and difficulty in permeating the BBB, doctors need to perform experimental therapies and choose drugs that can effectively cross the BBB. That means doctors have to break the usage principle of antibiotics or the medical insurance regulations. This is also a great challenge.

From the three points above, we can see that the risk of treatment failure in infectious meningitis is higher than in other infectious diseases. Doctors are extremely concerned about the risk in specific countries and regions (tense physician – patient relationship) or of legal action.

DIAGNOSIS AND TREATMENT OF MENINGITIS

Differential diagnosis of infectious and noninfectious meningitis

A patient presenting with fever, headache, nausea, emesis, meningeal irritation, and abnormal CSF findings (high pressure, increased white cell count, and decreased glucose and chloride levels) is easily misdiagnosed as having meningitis. However, it is necessary to exclude noninfectious causes of meningitis such as chemicals, connective tissue diseases, and tumors. A patient with chemical meningitis usually has a clear history of intrathecal medicine injection such as cytarabine, methotrexate, or analgesics. Because chemical meningitis often occurs during a period of hospitalization or in patients with a clear history of using specific medicines, it is easier to exclude. Connective tissue disease-related meningitis is often ignored. In such cases, we need to clarify if there is a medical history of systemic lupus erythematosus, sarcoidosis, or rheumatoid arthritis; order the appropriate laboratory investigations; and request a rheumatology consultation. Tumor-related meningitis (immunity meningitis or cancerous meningeal disease) can present with fever or other symptoms of meningitis. Detection of tumor markers in blood and CSF, cytology testing of CSF, and scans (computed tomography, magnetic resonance imaging [MRI], and positron emission tomography) can be helpful in the differential diagnosis.

Differential diagnosis of possible infectious meningitis pathogens

Pathogens causing infectious meningitis include bacteria, fungi, and viruses. It is important to distinguish the species of pathogens with no result of CSF smear. Purulent meningitis is easier to identify by observing the CSF appearance, CSF cell number, and the percentage of multinucleate cells.

Viral, tuberculous, and \(C.\) \(neoformans\) meningitis are more difficult to distinguish. The disease course for tuberculous and \(C.\) \(neoformans\) meningitis is over 6 weeks, and may be as long as several months, but that for viral meningitis is often less than 3 weeks. The body temperature of a patient with viral or \(C.\) \(neoformans\) meningitis can be over 39°C, but a patient with tuberculous meningitis often has fever in the afternoon and the body temperature is below 39°C.

With regard to CSF examination, the differences between the various meningitis types are as follows:

1. **Pressure:** in \(C.\) \(neoformans\) meningitis, pressure is above 300 mmH\(_2\)O; in tuberculous meningitis, it is more often between 250 and 280 mmH\(_2\)O (rarely above 300 mmH\(_2\)O unless there is meninges adhesion); and in viral meningitis, it is normal or a little higher, rarely above 250 mmH\(_2\)O.\[^7\]\[^2\] (2) **Glucose and chloride levels:** in tuberculous meningitis, these are both decreased or at least glucose is decreased, sometimes below 1.0 mmol/L; in viral and \(C.\) \(neoformans\) meningitis, glucose is decreased or normal, often between 2.0 and 2.8 mmol/L, while chloride is generally normal, or if decreased, is often between 110 and 118 mmol/L. (3) **Protein levels:** in tuberculous meningitis, protein is increased; in viral meningitis, protein is normal or a little higher, rarely above 1.0 mmol/L,\[^8\]\[^3\] and may be over 10 g/L, but in viral and \(C.\) \(neoformans\) meningitis, it is rarely more than 1.0 g/L.

Using MRI with enhancement, we can see that the strengthened signals in the meninges are strongest for
tuberculous meningitis, and are sometimes accompanied by ring enhancement of tuberculoma. These signals are weakest for viral meningitis and may sometimes be absent.

**How to determine the diagnosis and treatment strategy without identifying the pathogen?**

It is difficult to determine the diagnosis and treatment strategy without having pathogen identification. On the basis of the stated above, experimental therapy can be carried out for 2–3 weeks if we have propensity diagnosis, and then the subsequent strategy can be determined based on the effect of treatment. If there is no tendency for diagnosis, my personal experience is as follows.

**Step 1:** we treat the condition as viral meningitis for 2–3 weeks. This treatment can continue if there is a positive effect on the clinical, CSF, or imaging findings; otherwise, we go on to the next step. **Step 2:** we treat it as tuberculosis meningitis for 2–3 weeks. This treatment can continue if there is a positive effect on the clinical, CSF, or imaging findings; otherwise, we go on to the next step. **Step 3:** we treat it as *C. neoformans* meningitis for 2–3 weeks, and then assess the effect.

During every treatment step, efforts must continue to identify the pathogen and then re-diagnose. If the pathogen is identified, targeted treatment can commence; otherwise, we can only perform experimental therapy based on the clinical, CSF, or imaging findings.

**REFERENCES**