Survival of a case of *Bacillus cereus* meningitis with brain abscess presenting as immune reconstitution syndrome after febrile neutropenia – a case report and literature review-

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**Abstract**

**Background:** *Bacillus cereus* sometimes causes central nervous system infection, especially in compromised hosts. In cases of meningitis arising during neutropenia, CSF abnormalities tend to be subtle and can be easily overlooked, and mortality rate is high. We report a survived case of *B. cereus* meningitis/brain abscess in severe neutropenia, presenting as immune reconstitution syndrome.

**Case presentation:** A 54-year-old Japanese female with acute myelogenous leukemia developed *B. cereus* bacteremia and meningitis during consolidation chemotherapy. At the onset, she presented with mild meningism. She had marked leukocytopenia (WBC <100/μL, neutrophils 0/μL) and lumbar puncture yielded only mild pleocytosis. She was transferred to intensive care unit, and meropenem, linezolid and vancomycin was started. With intensive therapy, she recovered and once became afebrile. On day 19, however, her fever, meningism and consciousness level dramatically worsened despite recovery of bone marrow function. The antimicrobial chemotherapy was continued and finally she was cured with no complications.

**Conclusions:** With early diagnosis and prompt initiation of antibiotics, the case was successfully treated without any sequelae. It is important to remember that, even under optimal antimicrobial therapy, bone marrow recovery can cause transient reaggravation of the disease. In such cases, timely and appropriate evaluation should be done to make the clinical decision to change, continue, or intensify treatment.

**Keywords:** *Bacillus cereus*, Meningitis, Bacteremia, Febrile neutropenia, Immune reconstitution syndrome

**Background**

*Bacillus cereus* is a Gram positive, aerobic facultative, spore-forming, rod-shaped bacterium. It is motile, catalase positive and ubiquitous, found in air, soil, water, and food [1]. Because of the ubiquitous distribution of *B. cereus* in food products, the bacterium is ingested in small numbers and becomes part of the transitory human intestinal flora [2]. Traditionally, the detection of *Bacillus* species in sterile samples has been regarded as contamination. However, the organism has the potential to cause severe infectious diseases in immunocompromised hosts [1]. Herein, we report a case of *B. cereus* meningitis with brain abscess, arising in the setting of neutropenia, and presenting as immune reconstitution syndrome after bone marrow recovery.

**Case presentation**

A 54-year-old Japanese female was diagnosed with acute myeloid leukemia (AML). Complete remission was
achieved after induction chemotherapy, but 14 months later the disease relapsed. During the re-induction chemotherapy (FLAGM; fludarabine, cytarabine, granulocyte colony stimulating factor, and mitoxantrone), she experienced severe septic shock caused by *Klebsiella pneumoniae* bacteremia requiring intensive care. During the consolidation therapy (high dose cytarabine), oral polymyxin B, sulfamethoxazole/trimethoprim, voriconazole, and acyclovir were administered as prophylaxis.

On day 18 of consolidation chemotherapy (day 1 of infection), she complained of fever and chills. Meropenem 3 g/day was started after blood culture sampling. On day 2, headache and nausea appeared. Diarrhea was absent, and the vital signs were as follows: blood pressure 124/67 mmHg, heart rate 86 bpm, and body temperature 39.5 °C. At this time, she was conscious and lucid. Neurologically, no abnormal focal signs, including nuchal rigidity, were noted. Laboratory tests showed marked leukocytopenia (WBC < 100/μL, neutrophils 0/μL) with slight C-reactive protein elevation (1.32 mg/dL). The blood culture detected a Gram-positive rod-shaped bacterium on day 2, and the pathogen was identified as *B. cereus* on day 4. The strain was resistant to penicillin and quinolones, but sensitive to carbapenems, vancomycin, and linezolid.

On day 3, disorientation, slurred speech, and confusion appeared. Slight nuchal rigidity was observed, and the neck flexion test was positive. Although meningitis was suggested, lumbar puncture was not performed because of severe thrombocytopenia (21,000/μL). Instead, linezolid (1200 mg/day), vancomycin (targeted trough of 20 mg/dL), and acyclovir (30 mg/kg/day) were administered in addition to high dose meropenem (6 g/day).

On day 5, she was transferred to the intensive care unit because of shock. At this time, lumbar puncture yielded the following results: opening pressure: 18 cmH₂O; cells: 43/mm³ (mononuclear cells dominant, with no abnormal cells); glucose: 31 mg/dL; protein: 155 mg/dL; negative for HSV-DNA, VZV-DNA, CMV-DNA, HHV-6 DNA, Cryptococcus-Antigen and WT-1 (Wilms Tumor-1). Cerebrospinal fluid (CSF) culture was negative, but *B. cereus* PCR was positive, leading to the diagnosis as *B. cereus* meningitis.

With inotropic agents, deep sedation, and antimicrobials, she recovered and became afebrile. On day 19, however, her fever, meningism and consciousness level dramatically worsened despite recovery of bone marrow function. The CSF findings revealed markedly increased cells (2040 cells/mm³), and decreased glucose level (12 mg/dL). The CSF culture and PCR for *B. cereus* were both negative, and other diseases were excluded by the aforementioned examinations. Therefore, in concert with the increased leukocytes in peripheral blood and CSF, it was suggestive of immune reconstitution syndrome after *B. cereus* meningitis. The gadolinium enhanced magnetic resonance imaging (MRI) performed on day 15 had already detected meningeal thickening and several ring enhancement lesions. Glucocorticoid administration was not used to treat immune reconstitution syndrome. Meropenem, vancomycin, and linezolid were continued for 26, 40, and 45 days, respectively. They were stopped after CSF and MRI findings were confirmed to be normalized. On day 90, she was discharged to her home without any sequela, and her AML remains in remission for 5 years.

### Discussion and conclusions
Central nervous system infections caused by *B. cereus* can be classified as three diseases: 1) postsurgical meningitis, ventriculitis or brain abscess, which can occur several days after a surgical procedure and often affects devices [3]; 2) neonatal meningitis or encephalitis occurring in premature neonates at 7 days of age (median, range 1–49 days) [4]; 3) necrotizing meningitis or

| Table 1 Three Types of Central Nervous System *Bacillus cereus* Infection |
|-------------------------------|------------------|------------------|------------------|
| **Risk Population** | Ommaya reservoir | Premature birth: 32 (27–36) weeks | WBC count <100/μL |
| **Disease Onset** | Several days after procedures | 7 (1–49) days after birth | Around 10 days of neutropenia |
| **Mode of Infection** | Meningitis | Meningoencephalitis | Necrotizing meningoencephalitis |
| | Ventriculitis | | Cerebral hemorrhage |
| | Brain abscess | | Subarachnoid hemorrhage |
| **Clinical Course and Prognosis** | Responds well to antimicrobial therapy | Progresses within hours | Progresses within hours |
| | | Leads to death within days | Leads to death within days |
| **Cerebrospinal Fluid Findings** | Up to 10⁵ cells /mm³ | Around 10²–³ cells /mm³ | Around 10⁵ cells /mm³ |
| **Mortality rate** | Low | 75% (12/14 cases) | 79% (11/14 cases) |
| **References** | 3 | 3, 4 | 3, 6–14 |
## Table 2: Clinical Characteristics of Neutropenia-Related *Bacillus cereus* Meningitis

| Case | Year | Age | Underlying Diseases | WBC Count at Onset | Fever | Headache | Vomiting | Diarrhea | Abdominal Pain | Neurological Abnormalities | CNS Disease | Treatment | Outcome | Reference |
|------|------|-----|---------------------|--------------------|-------|----------|----------|----------|---------------|---------------------------|-------------|-----------|---------|-----------|
| 1    | 1973 | 63  | AML                | 400*               | +     |          |          |          |               | Coma                      | Brain abscess | GM, OXA, CBC | Dead    | 5         |
| 2    | 1981 | 19  | AML                | <100               | +     |          |          |          |               | Meningitis                 | GM, PCG      | Dead      | 6       |           |
| 3    | 1988 | 67  | MDS/AML            | 100*               | +     | +        |          |          |               | Meningoencephalitis/SAH    | GM, LM, PIPC | Dead      | 7       |           |
| 4    | 1989 | 3   | ALL                | 20                 | +     |          |          |          |               | Lethargy                  | Brain abscess | CP, VCM, GM, RFP | Alive   | 8         |
| 5    | 1995 | 26  | AML                | 20                 | +     |          |          |          |               | Visual disturbance          | Meningoencephalitis/SAH | CAZ        | Dead    | 9       |           |
| 6    | 1997 | 64  | AML                | 300                | +     | +        |          |          |               | Speech disturbance, Sensory disturbance | Meningoencephalitis/SAH | PIPC, GM, CPZ, CTX, ABPC | Dead    | 10        |
| 7    | 1997 | 20  | ALL                | 0                  | +     | +        | +        |          |               | Speech disturbance, Sensory disturbance | Cerebral infarction (basal ganglia) | Dead       | 11       |           |
| 8    | 1999 | 13  | ALL                | 0                  | +     |          |          |          |               | Meningoencephalitis/hydrocephalus | CAZ, AMK     | Alive, Severe sequelae |           |           |
| 9    | 1999 | 15  | ALL                | 0                  | +     |          |          |          |               | Meningoencephalitis         | CAZ, AMK     | Dead      | 11      |           |
| 10   | 1999 | 30  | AML                | <100               | +     |          |          |          |               | Delirium, hyperventilation | Necrotic meningoencephalitis | CAZ, AMK & others | Dead    | 12        |
| 11   | 1999 | 43  | AML                | <100               | +     | +        | +        |          |               | Delirium, decerebrate rigidity | Meningoencephalitis | VCM, GM & others | Dead    | 12        |
| 12   | 1999 | 14  | ALL                | <100               | +     |          |          |          |               | Delirium, epilepsy, coma | Meningoencephalitis | CAZ, AMK & others | Dead    | 12        |
| 13   | 2003 | 16  | AA                 | 90                 | +     |          |          |          |               | Nuchal rigidity, epilepsy, consciousness disturbance | Meningoencephalitis | CAZ, IPM/CS | Dead    | 13        |
| 14   | 2005 | 19  | HD                 | 0                  | +     |          |          |          |               | Confusion, epilepsy, hemiparesis | Meningoencephalitis | ABPC, AMK, CPFX, TEIC, CLDM | Alive   | 14        |
| Our case | 54  | AML  | <100               | +                  | +     |          |          |          |               | disorientation, slurred speech, and confusion | Meningoencephalitis/hydrocephalus brain abscess | MEPM, LZD, VCM | Alive   |           |

*denotes the number of neutrophils

**Abbreviations:** AA Aplastic anemia, ABPC Ampicillin, ALL Acute lymphoblastic leukemia, AMK Amikacin, AML Acute myelogenous leukemia, CAZ ceftazidime, CBC, carbenicillin, CLDM Clindamycin, CNS Central nervous system, CP Chloramphenicol, CPFX Ciprofloxacin, CPZ Cefoperazone, CTX Cefotaxime, GM Gentamicin, HD Hodgkin disease, IPM/CS Imipenem/cilastatin, LM Lincomycin, LZD Linezolid, MDS Myelodysplastic syndrome, MEPM Meropenem, OXA Oxacillin, PCG Penicillin G, PIPC Piperacillin, RFP Rifampin, VCM Vancomycin
hemorrhage in cases of severe neutropenia, predominantly affecting patients with leukemia (Table 1) [3–14].

While postsurgical infections typically resolve in spite of marked CSF pleocytosis, in the order of $10^5$ cells/mm$^3$, the latter two infection types have very poor prognoses with mortality rate greater than 75%. Notably, in many cases of these two malicious diseases, CSF cell counts are usually less than $10^2$ cells/mm$^3$.

Fourteen cases of neutropenia-associated $B.~cereus$ meningitis have been reported so far (Table 2). All but one case had hematological malignancies (AML: 6, acute lymphoblastic leukemia: 7, myelodysplastic syndrome: 1, aplastic anemia: 1) and all had severe neutropenia of less than 500/$\mu$L at the onset of disease. Moreover, 79% had less than 100/$\mu$L blood leukocytes, suggesting a role for severe neutropenia in the pathogenesis of this disease. The cases demonstrated mild or absent CSF pleocytosis, which is considered to be a result of both the severely immunodeficient state of the patient and the character of $B.~cereus$ infection in itself, which causes little inflammatory reaction [15].

Our case shared common features with previous reports such as severe neutropenia at the onset of meningitis and the complication of brain abscesses. While those are typical findings of neutropenia associated $B.~cereus$ meningitis, there are two particular features of this report. First, with an early diagnosis and administration of a contemporary antimicrobial regimen, we were able to successfully treat the infection. The clinical course of the patient is shown in Fig. 1.
able to cure the disease without any complications. The regimen contained antibiotics with good penetration, including linezolid, high dose meropenem, and a well-controlled dose of vancomycin that achieved a serum trough level of 20 μg/mL. This well-structured regimen was started immediately after the onset of meningitis, though the physical findings were subtle in the early phase, and CSF findings at day 5 showed only a moderate level of pleocytosis. This early diagnosis is crucial for the rapid initiation of intensive antimicrobial therapy.

Second, we were able to closely track the CSF and MRI findings with the recovery of hematopoiesis. The patient's meningitis symptoms were improving until day 19, when neutrophils rapidly increased. With the recovery of neutrophil count, a dramatic worsening of fever, consciousness and meningism was demonstrated. This could be considered as a type of immune reconstitution syndrome, though the term originally denoted the recovery of cellular immunity from AIDS or other immunosuppressive states. As shown in Fig. 1, we serially observed parameters of cellular immunity as well as neutrophil count. Interestingly, the severity of meningitis symptoms clearly synchronized with the blood neutrophil count, and the correlation is even more dramatic for CSF cells, CSF soluble interleukin-2 receptor (sIL-2R), and CSF adenosine deaminase (ADA). On day 19, the vast majority of CSF cells were polymorphonuclear cells (neutrophils). Later, CSF, sIL-2R and ADA increased, indicating that a typical inflammation cascade was ongoing. As differential diagnosis, other infections or an intracranial recurrence of AML were considered, but negative culture, cytology and clinical course were indicative of *B. cereus* meningitis. Therefore, we continued the antibiotics and the symptoms gradually improved.

In conclusion, we experienced a case of *B. cereus* meningitis with brain abscess during severe neutropenia from which the patient survived. With serial measurement of inflammatory / immunological markers, our report described the phenomenon resembling immune reconstitution syndrome, and we also proved that contemporary antimicrobial regimens can cure this severe disease. In cases of meningitis arising during neutropenia, CSF abnormalities tend to be subtle and can be easily overlooked. Because immediate diagnosis is crucial for survival, clinicians should be able to carefully recognize the symptoms and interpret the clinical data in order to rapidly initiate intensive treatment. Furthermore, it is important to remember that bone marrow recovery can cause transient reaggravation of the disease, even under optimal antimicrobial therapy. In such cases, timely and appropriate evaluation should be done to make the clinical decision to change, continue, or intensify treatment.

**Abbreviations**

ADA: Adenosine deaminase; AIDS: Acquired immunodeficiency syndrome; AML: Acute myeloid leukemia; CMV: Cytomegalovirus; CSF: Cerebrospinal fluid; FLAGMT: Fludarabine, cytarabine, granulocyte colony stimulating factor, and mitoxantrone combined therapy; HHV-6: Human herpes virus-6; HSV: Herpes simplex virus; LZD: Linezolid; MEPM: Meropenem; MNC: Mononuclear cells; PNC: Polymorphonuclear cells; sIL-2R: Soluble interleukin-2 receptor; VCM: Vancomycin; VZV: Varicella zoster virus; WBC: White blood cells

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**Authors’ contributions**

YK wrote this paper. YK, TO, HM, and KH contributed to diagnosis and clinical decision for treatment. HM and AA contributed in giving advice from the point of Microbiology, and Immunology, respectively. All authors reviewed the manuscript. All Authors read and approved the final version of this manuscript.

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**Availability of data and materials**

The authors are ready to provide the data in response to the need.

**Ethics approval and consent to participate**

Informed consent was obtained from the patient after verbal and written information provision. All the procedures have been performed in accordance with the ethical standards laid down in the 1964 Declaration of Helsinki and its later amendments.

**Consent for publication**

We have confirmed that the participant gave written consent for her personal or clinical details along with any identifying images to be published in this study.

**Competing interests**

The authors declare that they have no competing interests.

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