Acute Acalculous Cholecystitis in Neurological Patients; Clinical Review, Risk Factors, and Possible Mechanism

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INTRODUCTION

Acute cholecystitis is known to be caused by the pathologic mechanism of inflammation and ischemia in the gallbladder (GB) wall, stagnant pooling of bile juice, and the lithogenicity of bile juice in a complex manner. The incidence of acute cholecystitis is
especially high after a major surgery or serious illness. Acute cholecystitis may occur in the presence of a GB stone, however acute acalculous cholecystitis (AAC) is reported to occur more often 1.

Hypoperfusion and bile stasis are the key pathogenesis of AAC, which could be aggravated by hypotension, dehydration, and the usage of vasoactive drugs. Bile stasis is caused by the use of opioids, fasting with total parenteral nutrition (TPN), and mechanical ventilation with positive end-expiratory pressure (PEEP) 2.

AAC is reportedly a complication of a disease with high mortality unless treatment, long-term and short-term mortality may reach 59% and 35%, respectively 3. The incidence of AAC is rare in patients with severe neurological conditions with alterations in consciousness, such as cerebrovascular disease (CVD) or severe brain trauma. However, diagnosing and treating AAC are important in avoiding high mortality. Therefore, it is important to recognize the symptoms and signs of AAC, make an accurate diagnosis, and provide expedient treatment. Neurological patients are accompanied by severe neurological symptoms and signs, such as loss of consciousness, paraplegia, and aphasia, which can increase the severity of the symptoms of AAC.

Nevertheless, abdominal evaluations are not routinely done in patients with CVD or brain trauma and there are no established clinical characteristics of AAC in the intensive care unit (ICU). The aim of this study is to investigate the clinical features of AAC and the possible mechanisms of AAC in neurological patients.

**METHODS**

Among the patients with neurological conditions admitted to our hospital between March 2007 and September 2018, the medical records of 52 patients who underwent cholecystectomy after being diagnosed with cholecystitis were reviewed retrospectively. Data regarding age, sex, neurological conditions, mental status on admission, past medical history, time from admission to the onset of AAC, initial symptoms or signs of AAC, and fasting period were investigated.

AAC was diagnosed through abdominal sonography or computed tomography (CT), which showed no gallstones. The date of diagnosis was defined as the day in which AAC was diagnosed through the abdominal imaging.

**RESULTS**

The clinical characteristics of total 52 patients who underwent cholecystectomy is summarized in Table 1. The average age was 67.1 years with broad age spectrum spanning from 17 to 90 years-old, only one 17 years-old, no one in the 20s, 17 in the 60s, and 19 in the 70s with the age group in their 70s showing the highest prevalence. There were 36 males and 16 females with the males showing 1.89 times higher incidence than the females. For the underlying disease, there were HTN in 23 patients, diabetics mellitus (DM) in 10 patients. The mean Glasgow Coma Scale (GCS) was 10 which implies neurological deterioration with drowsy to stuporous mentality being the most prevalent. Also, Intracerebral hemorrhage was the most common neurological diagnosis among brain insults with subarachnoid hemorrhage, subdural hemorrhage, and epidural hemorrhage in order of descent.

The clinical factors related to AAC are summarized in Table 2. The mean time from admission to onset of AAC was 22.5 days and length of hospital stay showed an average of 77.9 days. The mean fasting period was 8.1 days. The initial symptoms and signs of AAC were elevation of liver function test (LFT), which was also the most common symptoms and signs in 27 patients; there were right upper quadrant (RUQ) pain or tenderness in 4 patients, fever, in 2 patients, C-reactive protein (CRP) elevation in 5 patients, and incidentally diagnosed in 10 patients. The 10 incidentally diagnosis of AAC was done through abdominal sonography or CT after suspicions of acute kidney injury, cystitis, pneumonia, and pleural effusion. The asymptomatic 10 patients received cholecystectomy

| Table 1. Clinical characteristics of 52 patients |
|-----------------------------------------------|
| Characteristics                              |
| Sex (number)                                 |
| Male                                         | 36 |
| Female                                       | 16 |
| Age (mean)                                   | 67.1 |
| Past medical history                         |
| Hypertension                                 | 23 |
| Diabetes mellitus                            | 10 |
| Cerebrovascular disease                      | 2 |
| Heart disease                                | 5 |
| Alcoholics                                   | 4 |
| Glasgow Coma Scale on admission              |
| ≥ 13                                         | 26 |
| 7-12                                        | 4 |
| < 6                                         | 22 |
| Neurological conditions                      |
| Subarachnoid hemorrhage                      | 12 |
| Intracerebral hemorrhage                     | 19 |
| Subdural hematoma                            | 11 |
| Epidural hematoma                            | 3 |
| Intraventricular hemorrhage                  | 8 |
| Cerebral infarction                          | 3 |
| Hypoxic brain damage                         | 1 |
| Cerebral abscess                             | 2 |
after discussion among board certified gastroenterologists or general surgeons. Of the patients who were treated for neurological diseases, there were 35 patients who have received neurosurgical operative treatment and 17 patients who have received conservative management. Of which the group received neurosurgical operative treatment had a higher prevalence of AAC.

The patients who have received cholecystectomy after being diagnosed with AAC were divided into the neurosurgical treatment group and the conservative management group (Table 3). The mean time from admission to onset of AAC in the neurosurgical treatment group was 27.3 days, which was 14.7 days longer than the conservative management group (12.6 days). The mean fasting period in the neurosurgical treatment group was 9.1 days and the that in the conservative management group was 6.1 days. Mean GCS in the neurosurgical treatment and conservative management group were 9.4 and 12.1, respectively.

### DISCUSSION

The prevalence of cholecystitis in GB stone patients after surgery is reported to be similar between male and female. However, over 80% of AAC after other operations unrelated to trauma are male patients aged 50 years old and over. This indicates that emergent operation or an elective operation complicated by hypotension and blood loss is pronounced in male patients. The incidence of AAC in this study showed a 1.89 times higher rate of incidence in males than in females as noted. This could be explained that the rate of comorbidities is higher in the male patients, however sex alone cannot be considered an independent factor for cholecystectomy in neurological patients. Nevertheless, special attention could be paid to the occurrence of AAC in male patients considering what is reported in previous studies and this study.

Most of the neurological deteriorated patients admitted to the ICU requires PEEP, which can be associated with bile stasis. 7-10cm ventilation in H2O PEEP can raise hepatic venous pressure, which causes a decrease in portal perfusion, in turn can cause bile stasis and increases the possibility of gallbladder mucosal injury. There was no statistically significant difference among the patients in this study as 4-5cm ventilation of H2O PEEP was applied, but it is recommended to always keep in mind the possibility of bile stasis in patients requiring 7-10cm high PEEP.

The average duration in ICU stay was 13.3 days with mean fasting period 7.3 days. Yang et al. reported that successive fasting time as a risk factor of acute cholecystitis in aneurysmal subarachnoid hemorrhage patients. Successive fasting time reported was

### Table 2. Clinical factors associated with acute acalculous cholecystitis

| Clinical factors                                      | Mean time from admission to onset of AAC (day) | Hospital stay (day) | Mean fasting period (day) | Initial symptoms or signs (number) |
|-------------------------------------------------------|-----------------------------------------------|--------------------|--------------------------|----------------------------------|
| Mean time from admission to onset of AAC (day)        | 22.5                                          | 77.9               | 8.1                      | RUQ pain or tenderness: 8         |
| Hospital stay (day)                                   |                                               |                    |                          | Fever: 2                         |
| Mean fasting period (day)                             |                                               |                    |                          | Elevation of LFT: 27             |
| Initial symptoms or signs (number)                    |                                               |                    |                          | Elevation of CRP: 5              |
| RUQ pain or tenderness                                |                                               |                    |                          | Diagnosis incidentally: 10       |
| Fever                                                 |                                               |                    |                          | Neurological treatment:          |
| Elevation of LFT                                      |                                               |                    |                          | Operative: 35                    |
| Elevation of CRP                                      |                                               |                    |                          | Conservative: 17                 |
| Diagnosis incidentally                                |                                               |                    |                          |                                  |
| Neurological treatment                                |                                               |                    |                          |                                  |
| Operative                                             |                                               |                    |                          |                                  |
| Conservative                                          |                                               |                    |                          |                                  |

AAC: acute acalculous cholecystitis; CRP: c-reactive protein; LFT: liver function test; RUQ: right upper quadrant.

### Table 3. Clinical factors associated with acute acalculous cholecystitis

| Variables                                      | Neurosurgical treatment (n=35) | Conservative management (n=17) |
|------------------------------------------------|------------------------------|--------------------------------|
| Mean time from admission to onset of AAC (day) | 27.3                         | 12.6                           |
| Mean fasting period (day)                      | 9.1                          | 6.1                            |
| Mean GCS on admission                         | 9.4                          | 12.1                           |
| Neurological conditions                       |                              |                                |
| Subarachnoid hemorrhage                       | 10                           | 2                              |
| Intracerebral hemorrhage                      | 11                           | 8                              |
| Subdural hematoma                             | 9                            | 2                              |
| Epidural hematoma                             | 1                            | 2                              |
| Intraventricular hemorrhage                   | 7                            | 1                              |
| Cerebral infarction                           | 1                            | 2                              |
| Hypoxic brain damage                          | -                            | 1                              |
| Cerebral abscess                              | 2                            |                                |

AAC: acute acalculous cholecystitis; GCS: Glasgow coma scale.
5.38 ± 2.78 days and the pathogenesis of acute cholecystitis included bile stasis as a result of fasting time as its mechanism. Loss of enteral stimulation during parenteral nutrition impairs the gallbladder motor function and causes GB “sludge” formation. Long-term TPN can increase the incidence of AAC by up to 30%5, GB “sludge” formation can occur in about 50% of patients treated with TPN for 4 weeks and in 100% of patients after 6 weeks9. Autonomic nerve dysfunction in neurological patients may cause bile stasis. Sevastos et al.10 was proved to cause abnormalities in the flow control of bile and pancreatic juice through increase in vagal paresis and catecholamine secretion following the damage of the brain center with the cessation of oral feeding.

Gallbladder ischemia is pathogenesis of AAC. Orlando et al.11 reported that gallbladder ischemia and bile stasis leading to hypoperfusion. Intraluminal pressure is increased by bile stasis, which results in a decrease in gallbladder perfusion pressure. In this hypothesis, bacterial invasion of ischemic tissue is a secondary phenomenon, acute bacterial and viral diseases, such as hepatitis, infection of upper respiratory tract, are important precipitating factors11-14. Gallbladder ischemia increases mucosal phospholipase A2 and superoxide dismutase activities, and mucosal lipid peroxide content. Whereas gallstone-related disease is associated with arterial dilation and extensive venous filling, but AAC is associated with arterial occlusion and minimal to absent venous filling15.

Another mechanism closely related to the pathogenesis of AAC is systemic inflammatory response. The response to systemic injury and subsequent release of inflammatory mediator, such as factor XII, platelet-activating factor, induce the inflammatory response of the gallbladder16. The neurological patients admitted to the ICU is prone to lung problems, including pneumonia, atelectasis, pleural effusion, catheter-related infections, operations-related infection, drug fever, inflammation due to hemorrhage without infectious causes, pseudomembranous colitis, bacteremia, and urinary tract infection (UTI), which could be related to the systemic inflammatory response. Pneumonia and UTI took up more than half of the infections in the ICU. Systemic inflammatory response can cause cholestasis, gallbladder ischemia, therefore could affect the incidence of AAC in neurological patients2).

CONCLUSION

AAC may be a significant complication in patients with neurological conditions. It is important to identify symptoms and signs of AAC, accurately diagnose the AAC, particularly in long-term stay at ICU, a long period of fasting, state of hypotension or hypoperfusion, PEEP ventilation.

NOTES

Conflict of interest

No potential conflict of interest relevant to this article was reported.

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