Emphysematous cholecystitis presenting as gas-forming liver abscess and pneumoperitoneum in a dialysis patient: a case report and review of the literature

Abstract

Background: Emphysematous cholecystitis is a rare variant of acute cholecystitis with a high mortality rate. The combination of emphysematous cholecystitis, liver abscess and pneumoperitoneum are even rarer. Herein we present a case of emphysematous cholecystitis in a senile diabetic lady who had worsening hemodynamics while undergoing hemodialysis.

Case presentation: A 64-year-old woman with history of type 2 diabetes mellitus and end stage renal disease with regular hemodialysis presented to the emergency department with a 1-day history of sudden onset of lassitude and hypotension during hemodialysis. The result of a computed tomography (CT)-scan revealed air encircling the gallbladder, liver parenchymal and minimal pneumoperitoneal and liver abscess with no cholelithiasis. The patient had received empirical antibiotics with piperacillin-tazobactam 2.25 g intravenous route every 6 h for 14 days and cholecystectomy with surgical debridement and lead an uneventful postoperative hospital course. Escherichia coli was demonstrated as well as blood culture and peritoneal fluid culture.

Conclusion: In a senile diabetic and dialysis patient, we should take emphysematous cholecystitis into consideration once vague abdominal pain occurs. Empirical antibiotic therapy and adequate surgical intervention should take place as soon as possible.

Keywords: Emphysematous cholecystitis, Gas-forming liver abscess, Pneumoperitoneum, Dialysis

Background

Emphysematous cholecystitis (EC) is a rare life-threatening form of acute cholecystitis representing between 1 and 3 % of acute cholecystitis presenting mainly in male patients aged 50–70 years, and mostly occurring in patients with diabetes mellitus, immunosuppressed and peripheral vascular disease [1].

EC has been characterized clinically by the imaging with gas in the gallbladder lumen, the gallbladder wall and adjacent structure, and elsewhere in the biliary tracts in the absence of an abnormal communication with the gastrointestinal tract.

The gas may disseminate to subcutaneous tissue, as well as to the peritoneal and retroperitoneal cavity. The combination of emphysematous cholecystitis, liver abscess and pneumoperitoneum are rarely seen. Dialysis patients rarely develop such complications according to the review of the literature from Pubmed [2].

Subhepatic abscess involved associated with emphysematous cholecystitis is rare [3–5]. Emphysematous cholecystitis occurring in association with a pneumoperitoneum is relatively rare [6, 7]. A review of the literature from pubmed revealed 18 other
cases of this combination and this is the first reported case occurring during dialysis.

Herein we present a dialysis case with clinical image composed of emphysematous cholecystitis, liver abscess and pneumoperitoneum.

**Case presentation**

A 64-year-old woman with history of type 2 diabetes mellitus and end stage renal disease with regular hemodialysis presented to the emergency department with a 1-day history of sudden onset of lassitude and hypotension during hemodialysis. She complained of fluctuating and persistent dull pain over the epigastric area. The painful sensation could not be relieved by lying down or adopting the decubitus position. She did not have nausea, vomiting, tea color urine, clay-like stool, muscle spasm or focal neurologic signs. She denied contact with animals or travel to foreign countries in recent days. On physical examination, the patient was actually ill and had a body temperature of 37.3 °C, pulse rate of 110 beats per minute, respiratory rate of 40 times per minute and blood pressure of 87/65 mmHg. The abdomen revealed right upper quadrant tenderness with Murphy's sign and muscular defense of the upper abdomen. In reviewing of the system, no diarrhea, no melena or hematochezia, no dysuria, no hematuria, no flank pain, no periumbilical and flank ecchymosis/petechiae been found. In addition, the laboratory examinations revealed leukocytosis (12800/μL) with a left shift (90.4 % neutrophil), elevated C-reactive protein (44.92 mg/dL), liver function impairment (aspartate aminotransferase (AST) of 237 U/L and alanine aminotransferase (ALT) of 232 U/L) and mild jaundice (total bilirubin:1.48 mg/dL). A plain radiography of the chest with the patient in a supine position suggested the presence of a dilated gallbladder with air in the lumen and wall (Fig. 1). The result of a computed tomography (CT)-scan revealed air encircling the gallbladder, liver parenchymal and minimal pneumoperitoneal and liver abscess (Fig. 2a, b, c) with no cholelithiasis. The patient had received empirical antibiotics with piperacillin-tazobactam 2.25 g intravenous route every 6 h for 14 days. The blood

![Fig. 1 Plain-film radiography showing air in the lumen and wall of the enlarged gallbladder of a 64-year-old woman with abdomen pain and shock while undergoing hemodialysis (arrows)](image-url)
A general surgeon was consulted and cholecystectomy and surgical debridement performed. The postoperative course went smoothly without any complications. The gallbladder was found to be necrotic. The culture of the bile collected during the operation and the peritoneal fluid collected from the pneumoperitoneum were the same as the blood culture yielded *Escherichia coli*. Pathologic analysis of the resected gallbladder disclosed empyema with extensive transmural necrosis and neutrophils infiltration of the whole organ (Fig. 3). The patient had developed acute delirium status with response to antipsychotic medication and active upper gastrointestinal tract bleeding with response to proton pump inhibitor therapy during the latter hospital course. She successfully recovered without any sequelae after adequate antibiotic treatment.

**Discussion**

EC, also known as acute gaseous cholecystitis is pathophysiologically different from acute or chronic cholecystitis. Obstruction of the gallbladder neck secondary to cholelithiasis induces acute and chronic cholecystitis. EC frequently results from thrombosis or occlusion of the cystic artery with ischemia necrosis of the gallbladder wall with subsequent gallbladder necrosis and secondary infection by gas-forming organisms. EC can be subclassified into 3 different variants including gas in the gallbladder lumen, gallbladder wall and pericholecystic tissues.

The mortality rate in EC is as high as 15 % compared with 4 % in acute cholecystitis [8–11]. The most common symptoms in EC are right upper quadrant abdominal pain. Fever, nausea and vomiting are also the main clinical symptoms of emphysematous cholecystitis.

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**Fig. 2** a Computed tomography revealed air encircled the gall-bladder lumen as well as intramural and pericholecystic air pockets, with findings pathognomonic for emphysematous cholecystitis. b Liver parenchymal destruction by air and partial liver abscess denoted. c Pneumoperitoneum as denoted by white arrow.

**Fig. 3** Pathologic analysis of the resected gallbladder disclosed empyema with extensive transmural necrosis and neutrophils infiltration of the whole organ. Mucosa slough with bacteria colonies been observed (arrow).
However, the presenting symptoms of EC are sometimes very vague and initially indistinguishable from those of uncomplicated acute cholecystitis, frequently causing a diagnostic dilemma, as in our patient initially masquerading as hypotension while performing hemodialysis. The symptoms may be even trivial in patients with diabetes mellitus and end-stage renal disease [2, 12].

We had conduct a systemic review from the series literatures of Pubmed with the linkage between “emphysematous cholecystitis”, “diabetes mellitus” and “hemodialysis” which enclosed 23 papers, which include 29 cases of EC, of which 21 cases have the cormobilities of diabetes, one with regular CAPD and the other one received temporary hemodialysis due to acute kidney injury with anuria resulted from hemolytic uremia syndrome. (Table 1) To the best of the author’s knowledge, concurrent EC, liver abscess and pneumoperitoneum in a dialysis diabetic patient has not been previously reported (Table 2).

Diabetes usually provide an environment for submucosa thrombosis of biliary tract and predispose patients to fulminant infections. The reason for delay in diagnosis probably involves diabetes neuropathy, which sometimes masks the symptoms of acute abdomen. Hyperglycemia and ischemia environments in diabetic patients can lead to reduced mobility of phagocytes in the areas of infection and further reduce antimicrobial activity, making EC possible. Appropriate control of blood sugar levels can lower the chance of bacterial overgrowth and associated severity of the disease [11].

Chen et al. reported that end-stage renal disease was an independent risk factor for acute cholecystitis. The independent risk factors were older age, higher Charlson’s score, atrial fibrillation, severe liver disease, diabetes, and dialysis modality. Haemodialysis patients had a higher risk of acute cholecystitis than PD patients [12].

Another possible postulated mechanism in EC is the fluctuating hemodynamic change during haemodialysis compared with peritoneal dialysis [13]. Hypotension in dialysis patient results from rapid reduction of blood volume owing to ultrafiltration and decrease in extracellular osmolarity during the dialysis session especially in older and diabetic patient with coexisting illnesses, such as cardiovascular diseases, which might contribute to systemic hypoperfusion and further compromise visceral circulation such as the cystic artery which lead to gallbladder ischemia and facilitates the proliferation of gas-forming organisms and bacterial translocation in the devitalized tissue with low oxygen saturation. The hypotension episode in our patient can be overlooked due to underlying bacteremia related septic shock may mimic the presentation of dialysis process. Besides, the inflammation and oxidants produced after ischaemia/reperfusion also impair the emptying of the gallbladder, increase the residual volume, and reduce smooth muscle

Table 1 Review of the literature (1955–2015) of emphysematous cholecystitis with association without diabetes mellitus

| Number | Age | Gender | Comorbidities | Diabetes | GB stone | Operation/survival | Bacteria source/Concurrent disease/CAusative agents other than bacteria | Author (year) |
|-------|-----|--------|----------------|---------|----------|-----------------|---------------------------------------------------------------------------------|--------------|
| 1     | 54  | M      | N/A            | Yes     | –        | Yes (Ce)/Yes    | B/C: C. baratii/Liver abscess/-                                                                                                        | Huang et al. (2012) [5] |
| 2     | 80  | F      | –              | –       | PTGBD/-  | –               | Bi/C & B/C & abd soft tissue: Clostridium difficile/myonecrosis/-                                                                      | Safioleas et al. (2007) [26] |
| 3     | 47  | M      | Alcoholism     | –       | –        | Yes (Ce)/Yes    | Bi/C: Escherichia Coli & Enterobacter Femoral tissue culture: Escherichia Coli, Bacteroids & Enterobacter/myonecrosis/-                  | Safioleas et al. (2007) [26] |
| 4     | 72  | M      | N/A            | –       | –        | Yes (Li)/Yes    | N/A                                                                                                                                  | Ise et al. (2002) [36] |
| 5     | 67  | F      | –              | –       | –        | Yes (Ce)/Yes    | B/C, Bi/C negative/Serum: antibodies against Escherichia coli O157,adult-onset HUS; liver abscess/-                                 | Yoshida et al. (1998) [37] |
| 6     | 41  | M      | ESRD secondary to Fabry’s with regular CAPD; status post two living related donor transplants; Abdomen vessel calcifications | –       | –        | Yes (Ce)/Yes    | S/C: Clostridium difficile; P/C: Clostridium perfringens/recent massive UGIB/-                                                           | Mirza et al. (1997) [2] |
| 7     | 64  | M      | –              | –       | –        | Yes (Ce)/Yes    | Bi/C: Streptococcus group D/-                                                                                                           | Carvalho et al. (2007) [26] |
| 8     | 63  | M      | –              | N/A     | –        | /Yes           | N/A                                                                                                                                  | Tooms et al. (1955) [39] |

M male, F female, CA cancer, PAD peripheral arterial disease, CAD coronary artery disease, N/A unknown, -- none, SAH subarachnoidal hemorrhage, ESRD end-stage renal disease, HUS hemolytic-uremic syndrome, OP operation, CAPD continuous ambulatory peritoneal dialysis, HG hyperglycemia, FBG fasting blood glucose, GB gallbladder, Ce cholecystectomy, Co cholecystostomy, Cd cholecodochotomy, PTGBD percutaneous transhepatic gallbladder drainage, Lo laparotomy, Lc laparoscopy, Bi/C bile culture, B/C blood culture, S/C stool culture, P/C peritoneal fluid culture, APN acute pyelonephritis, UGIB upper gastrointestinal tract bleeding
| Number | Age | Gender | Comorbilities | Diabetes | GB stone | Operation/survival | Bacteria source/Concurrent disease/Causative agents other than bacteria | Author (year) |
|--------|-----|--------|---------------|----------|----------|-------------------|---------------------------------------------------------------------|---------------|
| 1      | 65  | F      | CAD           | Yes      | –        | PTGBD & liver abscess drainage/Yes | B/C: Clostridium perfringens/Liver abscess, hemolysis/- | Cochrane. et al. (2015) [4] |
| 2      | 85  | M      | CAD           | Yes      | –        | Yes/Yes           | B/C: Clostridium perfringens/- - | Mirrakhimov et al. (2014) [20] |
| 3      | 77  | F      | Gastric CA    | Yes      | –        | Yes/Yes           | -/-/Chemotherapeutic agents | Kuroda et al. (2013) [21] |
| 4      | 73  | M      | Nephropathy   | Yes      | –        | Yes/Yes           | B/C: negative/Escherichia Coli related APN/- | Ogawa et al. (2012) [22] |
| 5      | 11  | M      | Obesity       | Yes (type1) 1D | –        | Yes (Lc)/Yes      | B/C: Enterococcus P/C: Escherichia Coli/Secondary appendicitis/- | Pal et al. (2011) [23] |
| 6      | 82  | F      | SAH           | Yes (type1) 1D | –        | Yes/Yes           | B/C: Clostridium species/Subarachnoidal hemorrhage/ - | Uesaka et al. (2009) [24] |
| 7      | 65  | M      | Hypertension  | Yes      | multiple GB stones | Yes (subtotal Ce & Co) /Yes | B/C/toxin A of Clostridium difficile and Escherichia Coli/- - | Theodossis et al. (2008) [25] |
| 8      | 87  | F      | Bedridden state | Yes      | –        | PTGBD/-          | B/C & B/C & abdomen soft tissue culture: Clostridium difficile/ Renal failure; myonecrosis/- | Safioleas et al. (2007) [26] |
| 9      | 32  | M      | Yes           | Yes      | –        | Yes (Ce)/No      | B/C: Enterococci & Clostridium Welchii; Femoral tissue culture: Escherichia Coli/Myonecrosis- | Safioleas et al. (2007) [26] |
| 10     | 70  | M      | Heart disease | Yes      | Multiple small GB stones | Yes (Ce)/Yes | B/C: Clostridium perfringens/- - | Shresth et al. (2007) [27] |
| 11     | 68  | M      | Hypertension  | Type 1   | –        | Yes (Ce)/Yes      | B/C: Clostridium perfringens and Corynebacteria/- - | Bernstein et al. (2007) [28] |
| 12     | 64  | F      | Hypertension  | Yes      | –        | Yes (Ce)/Yes      | B/C: Salmonella derby B/C: Negative/- - | Moanna et al. (2006) [11] |
| 13     | 62  | M      | Alcoholism    | Yes      | –        | Yes (Ce)/Yes      | B/C: Klebsiella pneumonia/- - | Prieto Fernández et al. (2004) [29] |
| 14     | 62  | M      | Alcoholism    | Yes      | –        | Yes (Ce)/Yes      | B/C: Klebsiella pneumonia/- - | Prieto Fernández et al. (2004) [29] |
| 15     | 42  | M      | Recurrent UTI | Yes      | –        | Yes (Ce)/Yes      | U/C & B/C : Negative /Emphysematous pyelonephritis/- - | Bhansali et al. (2004) [30] |
| 16     | 55  | M      | Hypertension  | Yes      | –/Yes    | N/A               | | Chiu et al. (2004) [31] |
Table 2: Review of the literature (1955–2015) of emphysematous cholecystitis with association with diabetes mellitus or dialysis (Continued)

|   |   |   |   |   |   |   |
|---|---|---|---|---|---|---|
| 17 | 70 | M | N/A | Yes | Yes (Mirizzi syndrome) | Yes (Ce)/Yes | N/A | Ozkan et al. (2003) [32] |
| 18 | 66 | F | Gastric CA post OP | Yes | – | Yes (PTGBD; Ce and Cd) | B/C: Clostridium perfringens & E. coli/ pneumomobilia/- | Ohtani et al. (1996) [33] |
| 19 | 77 | M | Breast CA post OP | – | Yes | – | N/A | Matsura et al. (1995) [34] |
| 20 | 66 | M | – | Yes | Yes | Yes (Ce)/Yes | N/A/- | Carvalho et al. (1965) [35] |
| 21 | 68 | M | – | Yes | N/A | – | N/A/- | Carvalho et al. (1965) [35] |

M: male, F: female, CA: cancer, PAD: peripheral arterial disease, CAD: coronary artery disease, N/A: unknown, –: none, SAH: subarachnoidal hemorrhage, ESRD: end-stage renal disease, HUS: hemolytic-uremic syndrome, OP: operation, CAPD: continuous ambulatory peritoneal dialysis, HG: hyperglycemia, FBG: fasting blood glucose, GB: gallbladder, Ce: cholecystectomy, Co: cholecystostomy, Cd: cholecdothomy, PTGBD: percutaneous transhepatic gallbladder drainage, Lo: laparotomy, Lc: laparoscopy, B/C: bile culture, B/C: blood culture, S/C: stool culture, P/C: peritoneal fluid culture, APN: acute pyelonephritis, UGB: upper gastrointestinal tract bleeding
compared with percutaneous cholecystostomy due to cystitis, cholecystectomy may be a better initial choice under-went hemodiaysis with concurrent acute cholecystitis [17]. Gunay et al. had proposed that in a patient treated ESRD patients compared with the normal popula-
tion, morbidity, mortality, and hospital stay were achieved by applying laparoscopic cholecystectomy to treat ESRD patients compared with the normal popula-
tions [17]. Gunay et al. had proposed that in a patient underwent hemodialysis with concurrent acute cholecys-titis, cholecystectomy may be a better initial choice compared with percutaneous cholecystostomy due to higher success rate and lower morbidity and mortality rate [18]. Gumus et al. suggested that in the manage-ment of acute cholecystitis patients with chronic hemodialysis states especially in poor surgical candidate, percutaneous cholecystostomy may be the alternative choice [19].

Conclusions
In conclusion, emphysematous cholecystitis is a rare form of cholecystitis especially in dialysis patients which could be fatal if delayed in diagnosis and progress to pneumoperitoneal and liver abscess. In a senile diabetic and dialysis patient, we should take emphysematous cholecystitis into consideration once vague abdominal pain and hypotension occurs in a patient underwent hemodialysis. Empirical antibiotic therapy and adequate surgical intervention should take place as soon as possible.

Consent
Written informed consent was obtained from the patient for publication of this Case report and any accompany-ing images. A copy of the written consent is available for review by the Editor of this journal.

Abbreviations
ALT: alanine aminotransferase; AST: aspartate aminotransferase; CT: computed tomography; EC: emphysematous cholecystitis.

Competing interests
The authors declare that they have no competing interests.

Authors’ contributions
CYL participated in writing the manuscript. HEW participated in revision of the manuscript. MKT carried out the study and is the original physician of the patient. RJB participated in infectious survey and interpretate findings of the culture results. WHK participated in providing knowledge of the disease etiology and provide possible differential diagnosis. HCL, CCL, and KJS collected information of patient and wrote the contents of Clinical course of the patient. CAM helped draft the manuscript and revise it. IHC was in charge of the imaging examinations. CCT followed the patient. STC help provide the pathology of the emphysematous gallbladder. All authors read and approved the final manuscript.

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Author details
1Department of Medicine, Kaohsiung Armed Forces General Hospital, No.2, Zhongzheng 1st Rd, Lingya Dist, Kaohsiung City 802, Taiwan R.O.C.
2Department of Pathology, Kaohsiung Armed Forces General Hospital, Kaohsiung, Taiwan R.O.C.
3Department of Internal medicine, Division of Nephrology, Tri-service general hospital, National defense Medical center, No.325, Section 2, Cheng-Kung Road, Neihu 114, Taipei, Taiwan R.O.C.

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