Salmonella Enteritidis Empyema Preceding the Diagnosis of Non-Hodgkin’s Lymphoma and Subsequent Contralateral Chylothorax Treated with Radiolabeled Rituximab

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Abstract
Salmonella infection is common, but pleural involvement has rarely been reported. Only seven cases of Salmonella enteritidis pleural empyema have been reported; all had an associated preexisting underlying immunosuppression or malignancy. We report the case of an apparently healthy man who developed S. enteritidis empyema. On further follow-up and surveillance, he eventually presented with non-Hodgkin’s lymphoma and a contralateral recurrent chylothorax. The latter was successfully controlled with radiolabeled rituximab, which has never been described for the above purpose in literature before.

Keywords: Empyema, chylothorax, lymphoma, radiolabeled iodine, rituximab, Salmonella

INTRODUCTION
Salmonella infection is common; Salmonella enteritidis is among the most common cause of food poisoning in the United States, but presentation with focal pleuropulmonary diseases is uncommon (1, 2). Only seven cases of S. enteritidis pleural empyema have been reported; six had a known intra-abdominal infection, malignancy, or lupus. In a review of 1,543 non-typhoid Salmonellosis patients, focal infections were strongly associated with immunosuppression or existing underlying illnesses, including malignancies such as lymphoma.

We report the case of an apparently healthy man who developed S. enteritidis empyema. Kept under surveillance, he eventually presented with non-Hodgkin’s lymphoma and a contralateral chylothorax. The latter was successfully controlled with radiolabeled rituximab.

CASE PRESENTATION
A 57-year-old previously fit man presented with a three-day history of dyspnea. He had no risk factors for immunosuppression. He visited Bali one month prior and had diarrhea then. He was febrile and tachycardic at presentation. Peripheral blood showed neutrophilic leukocytosis 24.8 x 10^9/L, raised C-reactive protein (413 mg/L) level, and mildly deranged liver functions. His chest radiograph revealed a left-sided pleural effusion. Computerised tomography demonstrated a multi-loculated effusion, pleural thickening, and mild intrapulmonary and intra-abdominal lymphadenopathy. The drained fluid (1.9 L) was turbid and contained S. enteritidis; no malignant cells were found. Blood, stool, and urine cultures were sterile.

His empyema was successfully treated with intravenous piperacillin–tazobactam and intrapleural streptokinase. A repeat CT scan two months later showed resolution of his empyema and intrapulmonary lymphadenopathy but persistent mild retroperitoneal/mesenteric adenopathy and two tiny omental nodules. Gastroscopy and colonoscopy showed no malignancies. A provisional diagnosis of mesenteric panniculitis was made.

Eleven months after the initial empyema, he presented with breathlessness and a right-sided effusion (Figure 1). On pleuroscopy, 1.5 L of chylous, triglyceride-rich fluid was removed. The fluid and pleu-
ral biopsies both confirmed a low-grade B-cell non-Hodgkin’s lymphoma with kappa light chain restriction and no expression of CD5, CD10, or cyclin D1. Lymphoscintigraphy identified chyle leak at the level of the T6/7 vertebrae (Figure 2). Positron-emission tomography scan found no focal tracer uptake.

He remained asymptomatic and required no chemotherapy. His chylothorax gradually increased but completely resolved after a trial of treatment with 2.7GBq 131I-rituximab. He remained well with no recurrence of pleural effusions 12 months (Figure 1) after the lymphoma diagnosis.

Informed consent has been obtained from the patient for this case report and for the images.

Figure 1. a, b. Chest radiograph showing blunting of the right costophrenic angle and left-sided pleural thickening during initial presentation (a), chest radiograph on clinical follow-up showing resolution of the chylothorax effusion post rituximab treatment (b)

Figure 2. Lymphoscintigraphy using 99 Tm colloid at 3 h demonstrating persistent focal accumulation of the tracer at the expected location of the thoracic duct at the T6/7 level, which persisted after delayed imaging for 6 h, raising the possibility of a leak; however, there was no diffusion of the tracer into the effusion. The tracer appearance through major lymphatic channels of the legs to inguinal lymph nodes and subsequently pelvic and intra-abdominal retroperitoneal nodes is normal. The appearance in the liver also suggests that diffusion of the tracer into the systemic circulation was unaffected.

DISCUSSION
This is the first report of S. enteritidis empyema that preceded the diagnosis of malignancy and is also the first report of the successful control of lymphoma-associated chylothorax treated with radiolabeled rituximab.

Salmonella empyemas are rare. We found no other cases in an audit of 713 cases of culture-positive empyema in Western Australia (3). Although S. enteritidis is the most common serovar in causing enteric Salmonellosis, it only accounts for 7 of the 39 Salmonella empyema cases reported in literature (1, 2, 4-7). There are no clinical, imaging, or pleural biochemical features that clearly distinguish Salmonella empyema from empyemas of other microbial causes. The diagnosis of Salmonella empyema should be considered in patients with pleural infection and recent enteric symptoms.

The rarity of S. enteritidis empyema and the high rate of pre-existing diseases in reported series suggest that the lymphoma and empyema in our patient are causally related. Whether Salmonella empyema is an early manifestation of occult lymphoma in our patient or if it triggered the subsequent development of lymphoma is intriguing. Our case illustrates that such patients warrant thorough screening and longer-term surveillance for cancer.

The treatment of Salmonella empyema should be similar to that for other pleural infections, and it requires systemic antibiotics and prompt drainage of the infected pleural material (8). A wide choice of antibiotics has been quoted in literature, although drugs with beta-lactam activity and fluoroquinolones are often preferred regimens for Salmonella empyema (4). Surgery may be needed when conservative treatment fails. The contemporary use of combined
intrapleural tissue plasminogen activator and deoxyribonuclease therapy has significantly reduced the need for surgical drainage for empyema, but it has not been specifically tested in Salmonella empyema (9). The mortality of Salmonella empyema has been reported to be substantial, in cases of S. typhimurium infection in particular (4). It is unclear if this reflects the virulence of the organism or whether patients with significant comorbidity and risks of mortality are more susceptible to the organism.

Lymphoma-related chylothorax is often difficult to control, and a prolonged chyle leak has the potential to cause malnutrition and death. Compared with systemic intravenous use, single agent iodine-radiolabeled rituximab has been employed in cases of relapsed or refractory lymphoma with good responses and less myeloablative effects, thus preserving more aggressive retreatment options for future relapse (10). However, it has not been reported for the control of chylothorax. Our case suggests that this is an option worth exploring in future patients.

CONCLUSION
In patients with Salmonella empyema, further investigations for possible underlying occult malignancy are warranted. Our case suggested the need for extended surveillance and follow-up even if the initial screening is negative. In addition, radiolabeled rituximab may present a novel and minimally invasive option for recurrent chylothorax secondary to lymphoma.

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REFERENCES