Case Report: Idiopathic Traumatic Neuroma of the Gallbladder Without Previous Surgery

Tianyu Lin¹, Abdul Saad Bissessur², Yujie Zhu¹, Teruko Fukuyama³, Guoping Ding¹ and Liping Cao*¹

¹Department of Hepatobiliary and Pancreatic Surgery, Sir Run Run Shaw Hospital, School of Medicine, Zhejiang University, China, ²Department of Surgical Oncology, Sir Run Run Shaw Hospital, School of Medicine, Zhejiang University, China, ³School of Medicine, Zhejiang University, Hangzhou, China

INTRODUCTION

Neuroma arising in the biliary system is a very rare benign lesion and can be classified into two types: primary and traumatic neuroma (1). Pathologies, such as infection, inflammation, trauma, gallstones, or surgery, involving the biliary tree can lead to traumatic neuroma (2). Traumatic neuroma is a tumor-like proliferation formed at the proximal end of damaged nerves after surgery or trauma, usually presenting as a firm, oval, slowly growing, and painful nodule (3). Traumatic neuroma, also known as amputation neuroma, arising in the gallbladder without prior surgery and cholelithiasis is extremely rare (4).

Diagnosis of a gallbladder neuroma is ascertained based on pathological findings of spindle-shaped cells and immunohistochemistry findings of the S-100 protein of postoperative or biopsy samples (5).
Traumatic neuroma usually follows an asymptomatic course (6). Surgical excision remains the mainstay treatment (3). However, traumatic neuroma frequently mimics cholangiocarcinoma and extensive radical surgery is performed (7).

Due to the lack of specific clinical symptoms and imaging findings, it is most commonly misdiagnosed as malignancy. Herein, we report one case of an idiopathic traumatic neuroma of the gallbladder treated at our hospital and preoperatively misdiagnosed as cholangiocarcinoma.

**CASE PRESENTATION**

A 60-year-old woman presented at our hospital with abnormal liver function tests discovered 2 weeks earlier on physical examination. Enhanced abdominal computed tomography (CT) scans revealed a space-occupying lesion at the hepatic hilum that was presumably considered hilar cholangiocarcinoma. The patient was admitted to our hospital on February 21, 2019. The patient denied any prior abdominal surgery or trauma. No abnormalities were found on physical examination.

Laboratory investigations included liver function enzymes and tumor markers: alanine aminotransferase (ALT) 187 U/L (normal: 7–55 U/L), aspartate aminotransferase (AST) 111 U/L (normal: 8–48 U/L), alkaline phosphatase (ALP) 737 U/L (normal: 40–129 U/L), gamma-glutamyltransferase (γ-GT) 1,074 U/L (normal: 8–61 U/L), total bilirubin (TBil) 28.5 μmol/L (normal: 4–19 μmol/L), direct bilirubin (DBil) 13.7 μmol/L (normal: 0–5 μmol/L), albumin 44.5 g/L (normal: 30–50 g/L), CA19-9 50.01 U/ml (normal: 0–37 U/ml), CA125 12.18 U/ml (normal: 0–35 U/ml), alpha fetoprotein 8.63 ng/ml (normal: <20 ng/mL), and CEA 1.93 ng/ml (normal: 0–2.5 ng/mL).

Abdominal enhanced CT scan (Figure 1A) revealed a 10 × 17 mm mass in the hepatic hilum with an unclear boundary. In the arterial phase (Figure 1B), the mass showed obvious enhancement. The intrahepatic bile duct showed obvious enhancement.

![Figure 1](https://example.com/figure1)
dilation, and the mass surrounded the cystic duct. The wall of the gallbladder appeared thickened, but there was no significant enlargement. No enlarged lymph nodes were noted in the retroperitoneum. Furthermore, 3D reconstruction revealed a space-occupying lesion at the hilar bile duct, invading the right branch of the portal vein. Furthermore, the postoperative left residual liver volume (RLV) was estimated to be only 27.6% (Figure 1C).

After a multidisciplinary team (MDT) discussion, the location of the tumor was considered unordinary, and, since a biopsy would have been highly risky, the pathological nature of the tumor was difficult to establish preoperatively. Thus, preoperative diagnosis was considered to be hilar cholangiocarcinoma based on enhanced CT scans. However, direct surgical intervention (right hepatectomy) would be inappropriate due to insufficient residual liver volume. The patient thus underwent preoperative percutaneous transhepatic cholangiodrainage (PTCD) on March 1, 2019, followed by percutaneous transhepatic portal vein embolization (PVE) on March 4, 2019, to improve residual liver volume.

One month later, the patient was admitted again for surgery. Preoperative laboratory investigations revealed elevated liver function enzymes (ALT 131 U/L, AST 256 U/L, ALP 188 U/L, γ-GT 209 U/L), while bilirubin and tumor markers were all within the normal range. A contrast-enhanced CT (Figure 2A) of the abdomen revealed improvement in the dilation of the hepatic duct, while the volume of the left hepatic liver had increased by 55.57% (Figure 2B). Magnetic
resonance cholangiopancreatography (MRCP) (Figure 2C) indicated an irregular shadow at the porta hepatitis with a vague boundary and involving the cystic duct. The gallbladder was not enlarged, but the walls were significantly thickened. On April 11, 2019, the patient underwent surgery. Intraoperatively, the gallbladder was severely attached to the liver. The gallbladder had atrophied with a hardened texture (Figure 3A). The tumor invaded the hilar bile duct and the

![Figure 3](image-url)

**FIGURE 3** | (A) Intraoperative findings revealing an atrophied gallbladder, with a hardened texture (the red arrow points out the tumor). (B) Grayish-white nodule (pointed by the red arrow) seen at the neck of the gallbladder (represented by the red circle). (C) Low-magnification and (D) high-magnification pathological analysis showing spindle cells. (E) Immunohistochemical positivity of S-100.
right branch of the portal vein. Lymph node enlargement at the hepatoduodenal ligament was observed. Based on the intraoperative findings, gallbladder carcinoma was considered. The surgical intervention included laparoscopic cholecystectomy, laparoscopic right extended hepatectomy, and laparoscopic hepatoduodenal ligament lymph node dissection. A cholangiojejunostomy was then performed. Intraoperative freezing biopsy showed no cancer cells at the level of the resected left bile duct and cystic duct.

Postoperative pathological examination revealed that the mucosa of the gallbladder was grayish-white with a hard texture. Fibrous tissue hyperplasia of the gallbladder wall with hyaline degeneration was seen together with chronic interstitial inflammatory cell infiltration and focal multinucleated giant cell aggregation. A nodule, 1.4 × 1.0 cm in size, with a grayish-white cross-section (Figure 3B), was seen at the neck of the gallbladder. Pathological analysis and immunohistochemistry validated the diagnosis of traumatic neuroma through spindle cells (Figure 3C) and S-100 positivity (Figure 3D). Four resected lymph nodes showed reactive hyperplasia. The final validated diagnosis was traumatic neumia of the gallbladder, chronic cholecystitis, fibrosis of the gallbladder wall, hyaline degeneration, and gallstones. The patient recovered well after the operation, without complications, and was discharged from the hospital. There was no obvious discomfort in the postoperative follow-up.

**DISCUSSION**

Benign neoplasms of the biliary system are most commonly adenomas or papillomas and account for only 6% of all biliary tumors (8). However, the actual incidence of gallbladder neumia remains uncertain as the patients most commonly follow an asymptomatic course. According to an autopsy study of post-cholecystectomy patients, 11 out of 40 patients had histological findings consistent with traumatic neuromas, revealing that 25% of cholecystectomy patients develop traumatic neuromas after surgery (9).

Traumatic neumia is a non-neoplastic overgrowth of tissues and disorderly proliferation of nerve fibers and most frequently arises after radical neck dissection, orthopedic or limb amputation surgeries (3, 10). Traumatic neumia of the biliary system emerges from the sympathetic and parasympathetic nerve endings surrounding the bile ducts and gallbladder (2). As the common bile duct is a rich innervated area (11), traumatic neuroma commonly occurs after cholecystectomy or even bile duct exploration (12). The tumor commonly arises at the cystic duct stump or remnant after cholecystectomy (13). Traumatic neumia can occur as early as 1 month and as late as 56 years post-cholecystectomy (14). However, traumatic neumia arising in the gallbladder is a rare occurrence (15). Moreover, due to sustained injuries, traumatic neumia can occur after orthotopic liver transplantation (16, 17). Rarely, traumatic neumia might arise in the gallbladder post-vagotomy (18).

Even though most patients follow an asymptomatic course, some patients present with abnormal liver function, jaundice, pruritus, abdominal pain, weight loss, abnormal urine color, or clay-colored stools (1, 7). Our patient presented with abnormal liver function tests. Surgical excision remains the mainstay treatment of traumatic neumia, and there is no response to chemotherapy or radiotherapy (3). Due to its rarity and the limitations of specific imaging findings, traumatic neumia of the biliary tract is most commonly misdiagnosed as cholangiocarcinoma (7), as happened in our case. Neumia is composed of disordered nerve fibers and its signal density is the same as soft tissue and the pancreatic head, thus making it difficult to identify by CT and MRI examinations (19). Just as in our case, the elevation of tumor marker CA19-9 (7) and liver function enzymes (1) makes correct preoperative diagnosis a real challenge.

A case report by Shimura et al. (6) proved that intraductal ultrasonography (IDUS) could demarcate traumatic neumia from other malignancies as it provided accurate details. The IDUS showed that the lesion demonstrated a smooth homogeneously hypoechoic mass with a distinct margin and was able to determine whether the surrounding structures such as the portal vein, hepatic artery, and pancreas were invaded. These characteristics pointed toward traumatic neumia, which meant that extensive surgery could be avoided. Furthermore, two post-cholecystectomy cases of traumatic neumia were diagnosed by biopsies through the combined use of a cholangioscopy and an endoscopic ultrasound (5, 14). In our case, however, the tumor was close to the hepatic hilum, making a preoperative biopsy difficult.

In this case, the traumatic neumia was located in the gallbladder. Erratically, our patient had no prior history of

**TABLE 1** | Case reports of idiopathic traumatic neumia of gallbladder published in the past 60 years.

| Study          | Age | Sex | Symptoms                           | Gallstones  | Pre-operative diagnosis | Surgical scope                   |
|----------------|-----|-----|------------------------------------|-------------|-------------------------|----------------------------------|
| Janes et al.   | 66  | M   | Epigastric pain, jaundice          | Present     | Gallstones              | LC converted to OC, repair of fistula b/w the gallbladder and colon |
| Matsuoka et al.| 74  | M   | None ( incidental finding)         | Absent      | Cholangiocarcinoma      | LC converted to OC, partial liver resection |
| Peison et al.  | 88  | M   | RUQ abdominal pain, abdominal distension, loss of appetite | Present     | Gallstones              | Cholecystectomy                  |
| Yoshida et al. | 68  | M   | None ( incidental finding)         | Absent      | Cholangiocarcinoma      | Cholecystectomy + hepatic bed resection |

M, male; LC, laparoscopic cholecystectomy; OC, open cholecystectomy; b/w, between; RUQ, right upper quadrant.
abdominal surgery, gallstones, or polyps. Since there are no typical causes, such as surgery and physical trauma, we delineate such a finding as “idiopathic traumatic neuroma of the gallbladder.” PubMed, Embase, CNKI, and other websites were thoroughly searched for relevant publications, and only four cases of traumatic neuroma without previous surgery have been reported in the last 60 years (Table 1). All the patients were elderly males, of which two patients presented with abdominal pain and gallstones (4, 20–22). In contrast, our patient was an elderly female presenting with abnormal liver function enzymes. In one of the cases (22), reported in the Japanese literature, physical stimuli to surrounding nerves from a pedunculated polyp were speculated to be the causative factor of the neuroma in the gallbladder.

It is worth mentioning that long-term chronic inflammation was confirmed pathologically in our case and the above-mentioned cases. Physical trauma from surgery or gallstones or polyps may not be the only triggering factors for traumatic neuroma formation in the gallbladder; chronic inflammation may also spawn traumatic neuroma (23). Long-term continuous inflammation and destruction are required to cause a large number of abnormal proliferation of nerves in the gallbladder wall. Hence, chronic cholecystitis could be the inductive factor in our case, thus the reason why we delineate it as “idiopathic traumatic neuroma of the gallbladder.” The rarity of traumatic neuroma could be accounted for by the fact that the nerves near the mucosal surface are too thin to produce colossal proliferation enough to lead to the formation of a neuroma (20). Moreover, chronic inflammation can also cause severe local adhesion, gallbladder rupture, and fistula of the gallbladder with the colon, thereby forming an illusion similar to tumor infiltration, resulting in incorrect intraoperative diagnosis and expanding the scope of surgery. In this case, the gallbladder closely adhered to surrounding tissue.

In patients presenting with a history of abdominal surgery such as a cholecystectomy or liver transplant, traumatic neuroma should be included in the differential diagnosis, especially if the tumor arises at the remnant stump of the bile duct. However, even without prior surgery, inflammation triggered by gallstones or polyps or idiopathic inflammation can be an etiology of traumatic neuroma. While imaging modalities are not specific, liver function enzymes and tumor markers are elevated, and an intraoperative frozen biopsy should be incorporated to avoid extensive radical surgery, thus improving patients’ quality of life. Although idiopathic traumatic neuroma of the gallbladder is very rare, it should be included in the differential diagnosis in patients diagnosed with hilar cholangiocarcinoma.

DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article/Supplementary Material, further inquiries can be directed to the corresponding author/s.

ETHICS STATEMENT

Written informed consent was obtained from the individual(s) for their participation in this case report. Written informed consent was obtained from the individual(s) for the publication of any potentially identifiable images or data included in this article.

AUTHOR CONTRIBUTIONS

LC and GD designed the study. TL and ASB researched the literature and drafted the manuscript. YZ, TF, GD, and LC edited the manuscript. All authors contributed to the article and approved the submitted version.

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