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**complicated course of biliary inflammatory myofibroblastic tumor mimicking hilar cholangiocarcinoma: a case report and literature review**

Strainiene S *et al*. Complicated biliary IMT

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**Abstract**

BACKGROUND

The inflammatory myofibroblastic tumor (IMT) is a rare, idiopathic, usually benign, mass-forming disease with myofibroblastic proliferation and a varying amount of inflammatory cells. Although it can affect various organs, the biliary tract is a rare localization of primary IMT, clinically, endoscopically and radiologically imitating cholangiocarcinoma. The treatment options are based only on clinical practice experience.

CASE SUMMARY

A 70-year-old woman was referred to our center due to progressive fatigue, weight loss, abdominal pain, night sweats, and elevated liver enzymes. Magnetic resonance cholangiopancreatography and endoscopic retrograde cholangiopancreatography (ERCP) revealed proximal common hepatic duct and hilar biliary strictures extending bilaterally to lobular bile ducts. Although initial clinical, endoscopic and radiological signs were typical for hilar cholangiocarcinoma, histological examination showed no signs of malignancy. In total, 8 biopsies using different approaches were performed (several biopsies from dominant stricture during ERCP and direct cholangioscopy; ultrasound-guided liver biopsy; diagnostic laparoscopy with liver and lymph node biopsies). Histological examination revealed signs of IMT, and the final diagnosis of biliary IMT was stated. Although IMT is usually a benign disease, in our case, it was complicated. All pharmacological treatment measures were ineffective. The patient still needs permanent stenting, suffers from recurrent infections and mechanical jaundice. Despite that, the patient already survived 24 mo.

CONCLUSION

IMT presenting with hilar biliary strictures is a unique diagnostic and clinical challenge as it is indistinguishable from cholangiocarcinoma, and there are no evidence-based treatment options. Our goal is to increase the understanding of this rare disease and its possible course.

**Key Words:** Inflammatory myofibroblastic tumor; Hilar cholangiocarcinoma; Biliary strictures; Recurrent cholangitis; Case report; Literature review

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**Core Tip:** Biliary inflammatory myofibroblastic tumor (IMT) is a rare, idiopathic, usually benign, mass-forming disease with myofibroblastic proliferation and the varying amount of inflammatory cells that can occur in almost every organ. IMTs of the biliary ducts are an uncommon cause of obstructive jaundice. The clinical and radiological presentation usually mimics cholangiocarcinoma (Klatskin tumor). However, histological examination shows no malignancy. We present a rare, difficult to diagnose and treat biliary IMT, which was unresponsive to the pharmacological treatment and complicated by recurrent infections and disease progression. The global experience towards diagnosing and treating this disease is limited and based mostly on clinical practice experience.

**INTRODUCTION**

An inflammatory myofibroblastic tumor (IMT) is a rare, usually benign mass or multiple masses forming disease with myofibroblastic proliferation along with the varying amount of inflammatory infiltrate. These lesions can be found in every organ throughout the body[1–3]. Although their appearance clinically and radiologically mimics various aggressive malignancies, IMTs are usually considered as benign neoplasms. However, these tumors might also have a diverse spectrum of biological behavior[4]. According to the World Health Organization (WHO) classification, IMT is an intermediate-grade tumor[3]. Several terms have been applied to this condition, such as inflammatory pseudotumor (IPT), fibrous xanthoma, plasma cell granuloma, pseudosarcoma, lymphoid hamartoma, myxoid hamartoma, inflammatory myofibrohistiocytic proliferation, benign myofibrosblastoma, and most recently, IMT. The diverse nomenclature is mainly descriptive and reflects the uncertainty regarding the true biologic nature of these lesions[5,6].

The biliary tract is rarely involved by IMT[1,2]. It should be emphasized that since the first description of liver IMT by Pack and Baker[7] in 1953, over 300 Liver IMT cases have been reported, and very few IMTs were located in a biliary tract[8]. Biliary IMT is a great challenge for clinicians to diagnose and treat, as there are no confirmed recommendations or systemic trials towards treating this condition. We present a unique case of unknown origin, difficult to diagnose and treat biliary IMT presenting with significant biliary strictures, clinically, endoscopically, and radiologically imitating perihilar cholangiocarcinoma, Bismuth Corlette type IV.

**CASE PRESENTATION**

***Chief complaints***

A 70-year-old woman was referred to our tertiary Center of Hepatology, Gastroenterology and Dietetics due to progressive fatigue, weight loss (about 10 kg per year without special diets), right upper abdominal quadrant pain, heatwaves, and night sweats.

***History of present illness***

The patient was observed by a family physician due to hepatosteatosis and slightly elevated liver enzymes since 2017. Two years later, during the routine yearly check-up, laboratory tests showed a more significant elevation of liver enzymes (aspartate aminotransferase (AST) by 76 U/L, alanine aminotransferase (ALT)) by 80 U/L, alkaline phosphatase by 296 U/L, and gamma-glutamyltransferase by 455 U/L)). She was then further investigated for possible liver diseases. Viral hepatitis B and C, autoimmune hepatic diseases were excluded: the patient was negative for viral hepatitis B and C markers, antimitochondrial (AMA) and antinuclear (ANA) antibody titers were also negative. Abdominal ultrasound showed dilatation of intrahepatic ducts in the right lobe and segment IV, without apparent perihilar tumor signs. There were no gallstones in the gallbladder and common bile duct. The patient was referred to our center for further investigation.

***History of past illness***

The patient was diagnosed with dyslipidemia, hepatosteatosis, arterial hypertension, nontoxic multinodular goiter. There was no history of jaundice, previous hepatobiliary diseases, infections, or abdominal operations.

***Personal and family history***

The patient denied alcohol consumption, allergies to food or medicines.

***Physical examination***

On admission, the patient was asthenic (normal body mass index 22 kg/m2), afebrile (body temperature 36.8ºC) with normal vital signs (blood pressure 120/70 mmHg, pulse 88 bpm). There was no visible jaundice and no palpable abdominal pain. The patients’ liver, spleen and superficial lymph nodes were not enlarged.

***Laboratory examinations***

Laboratory tests revealed signs of inflammation, elevated liver enzymes, mild hyperbilirubinemia, and slightly elevated carbohydrate antigen (CA) 19-9 of 111.98 kU/L (Table 1).

***Imaging examinations***

Abdominal ultrasound showed dilated intrahepatic ducts with a normal common bile duct. Contrast-enhanced magnetic resonance cholangiopancreatography (MRCP) was performed to clarify the diagnosis, revealing a radiological image resembling perihilar cholangiocarcinoma, Bismuth-Corlette type IV, also known as Klatskin tumor (Figure 1).

Further, we proceeded with endoscopic retrograde cholangiopancreatography (ERCP). ERCP revealed tight stricture in the proximal part of the common hepatic duct (CHD) and the confluence extending to the left and right hepatic ducts (Figure 2A). Right and left hepatic ducts were not visualized due to complete obstruction. The radiological image was typical for cholangiocarcinoma, Bismuth Corlette IV. The biopsy was taken from the dominant CHD stricture, and a 7 Fr 12 cm pigtail stent was placed into the left hepatic duct to manage cholestasis. The preventive pancreatic 5 Fr 5 cm stent was placed in the pancreatic duct during the first ERCP procedure after nonintentional cannulation of the pancreatic duct.

Histological examination of the first biopsy revealed no signs of malignant disease. Therefore, direct cholangioscopy with biopsies was performed (Figure 2B). However, histological examination repeatedly showed no atypical changes.

***Further diagnostic work-up***

Based on clinical, radiological and endoscopic findings, the patient was diagnosed with perihilar cholangiocarcinoma during the multidisciplinary team meeting (MDT). The treatment with re-stenting every 3 mo and stereotaxic radiotherapy was recommended. However, radiotherapy was postponed until the pathological confirmation of the tumor. Therefore, ultrasound-guided liver and lymph node biopsies were performed. Histological examination revealed signs of the IMT/IPT. IMT could have been the primary tumor or similar chronic active inflammatory infiltrate (IPT) due to cholangitis. As the patient did not experience any signs of cholangitis before, it was more likely to be IMT.

The second MRCP 3 mo later showed a periductal mass in the liver hilum obstructing left and right main hepatic and segmental ducts to a similar extent as on the previous MRCP with a cluster of rim enhancing lesions on segments VIII and VII subsistent with small abscesses due to cholangitis. Enlarged cardiophrenic lymph nodes were also observed due to tumor spread or long-lasting inflammation of the liver (Figure 3).

During the second MDT meeting, it was decided to perform diagnostic laparoscopy to specify the lesions and finally withdraw cholangiocarcinoma. The radical surgical treatment was contraindicated as the mass was extending into both hepatic and some segmental ducts. Liver transplantation was not an option because of the patient‘s age. Histological examination of the CBD, liver mass and lymph node biopsies was compatible with the previous dominant stricture biopsies. Histopathology of the lesions showed no evidence of malignancy, atypical mitosis or necrosis. The non-specific changes were noted: widespread myofibroblastic and fibroblastic spindle cells with inflammatory infiltrate of lymphocytes, plasma cells, eosinophils, histiocytes. The immunohistochemical reaction for anaplastic lymphoma kinase (ALK)-1 and ALK (D5F3) was negative (Figure 4). Histologically, the tumor proved an IMT arising from the bile duct epithelium and confirmed the diagnosis of biliary IMT*.*

To specify the possible etiology of the tumor, several additional laboratory tests were performed. The patient was tested negative for cytomegalovirus, Epstein-Barr virus, HIV 1 and 2, and immunoglobulin (Ig) G4. Tuberculosis was also withdrawn. The biochemical markers for common oncological diseases were also negative (Table 1). The patient did not have a history of cholangitis before the first ERCP.

**FINAL DIAGNOSIS**

Although imaging examinations showed signs of perihilar cholangiocarcinoma, the pathological investigations revealed only inflammatory changes, and the other possible diseases were excluded. Accordingly, the patient was diagnosed with biliary IMT and dominant CHD stricture.

**TREATMENT**

The primary symptomatic treatment with permanent CBD stenting every 3 mo was applied and continued throughout patients’ treatment and follow-up in our hospital. After confirming the diagnosis of biliary IMT, it was decided to start the treatment with a 1-month course of steroids (40 mg/d of prednisolone tapering by 5 mg) and non-steroidal anti-inflammatory drugs (NSAIDs) (celecoxib 100 mg/d) based on the cohort study by Casanova *et al*[5]. However, prednisolone was discontinued 10 d after due to significantly increased C-reactive protein (to 100 mg/L).

We continued restenting every 3 mo. However, the majority of procedures have been followed by cholangitis and septic complications. Therefore, the culture from the bile was taken after elective ERCP. Six pathogens were cultured: *Pseudomonas aeruginosa, Klebsiella pneumoniae, Escherichia coli, Enterococcus avium, Enterococcus gallinarum and* *Enterococcus faecium.* The patient was treated with antibiotics according to the antibioticogram. However, this approach did not affect the stricture as control MRCP 9 mo after the initial diagnosis showed no significant changes in stricture length. The concentration of carbohydrate antigen 19-9 significantly increased from 112.15 to 1184.1 kU/L and continued to rise (Figure 5).

Further, the patient was consulted by the oncochemotherapist. As steroids and celecoxib therapy failed due to infectious complications, it was decided to start the second-line treatment with vinblastine and methotrexate. MRCP after 3 cycles of chemotherapy (18 mo since diagnosed) showed no positive results. There were no liver abscesses seen on subsequent MRI. However, the stricture length of the bile ducts remained the same, and cardiophrenic and liver hilum lymph nodes did increase by few millimeters in size. It is hard to attribute lymph node enlargement for tumor progression as they can also enlarge due to recurrent infections. The patient’s clinical condition and well-being did improve at that time, but this is most likely due to treated infectious complications.

**OUTCOME AND FOLLOW-UP**

The patient continued chemotherapy (5 cycles). Unfortunately, this treatment was discontinued, as the patient was repeatedly hospitalized due to recurrent cholangitis and sepsis. What is more, she gained a COVID-19 infection. After fighting COVID-19 disease, the patient’s condition significantly worsened as she developed cholecystitis (*Pseudomonas aeurginosa*) – related to cystic duct blockade with the ductal stents. The whole body computer tomography (CT) images after biliary stenting showed the same biliary stricture in the liver hilum with a large gallbladder and a new solid mass at the site of the gallbladder fundus. Enlarged lymph nodes in the right lung were also observed (Figure 6). A percutaneous cholecystostomy was performed, and the biopsy was taken from the newly developed mass in the gallbladder fundus. Histological examination showed fibrotic changes without any signs of a malignant tumor. Although CA 19-9 rose to 7884 kU/L, which could have been a reaction to constant infections, disease progression, and COVID-19 disease. Improvement in infectious status was observed after cholecystostomy. The patient‘s survival time is already 24 mo, and her general condition is satisfactory due to timely management of recurrent infections (Figure 7). Although the stricture site remains stable, the patient might have a poor prognosis due to constant infectious complications.

**DISCUSSION**

IMTs are rare, usually benign mass forming lesions of unknown origin. It encompasses a spectrum of myofibroblastic proliferation and varying amounts of polymorphous inflammatory cell infiltrate consisting of spindle cells, plasma cells, lymphocytes, eosinophils and macrophages. Also, there are variable amounts of fibrosis, necrosis, granulomatous reactions[3,9]. These lesions can be found in every organ throughout the body, most commonly in the lung, orbit, peritoneum, omentum and mesentery, followed by soft tissue, mediastinum, gastrointestinal tract, pancreas, genitourinary tract, oral cavity, skin, breast, nerve, bone, and central nervous system[1–3].

IMTs of the biliary ducts are an uncommon cause of obstructive jaundice. The cases found during the literature search using PubMed are presented in Table 2(used keywords: biliary IMT, inflammatory myofibroblastic tumor of the biliary tract, perihilar IMT, hepatic bifurcation IMT).

***Etiopathogenesis***

The exact origin and pathogenesis of the disease are poorly understood. The tumor can affect both sexes and all ethnic groups equally, and can occur in any age group[10,11]. The most popular and widely accepted theories describe IMTs as either primary reactive and inflammatory processes or low-intermediate grade neoplastic processes with secondary inflammation. Infection, minor trauma, previous abdominal surgery or procedure, radiotherapy, chemotherapy, and steroid use - all might play a role in the pathogenesis of the heightened inflammatory reaction and result in IMT/IPT. An immune-autoimmune mechanism has also been implicated[9]. The unusual inflammatory or immune responses such as sclerosing cholangitis, phlebitis, and retroperitoneal fibrosis also can be found in association with IPT.

Some organisms like parasitic fragments, *Escherichia coli*, gram-positive cocci, *Klebsiella pneumonia*, *Epstein-Barr* virus were found in splenic and nodal pseudotumors. *Actinomycetes* and *nocardiae* were found in hepatic and pulmonary pseudotumors, respectively; mycobacteria associated with spindle cell tumor and mycoplasma found in pulmonary pseudotumors[9,12]. There have been reports of IMTs/IPTs related to *Mycobacterium avium–intracellulare* complex, *Escherichia coli, Klebsiella, Bacillus sphaericus, Pseudomonas, Helicobacter pylori,* *Corynebacterium equi,* and *Coxiella burnetti*[9,12–14]. However, in most cases, no organisms were cultured or identified in tissue sections, and nothing was cultured from the specimens. There are some cases where liver IPT has been reported in patients with a history of cholangitis and intra- and extrahepatic bile duct dilatation[15-17]. The sequence in these cases indicates that cholangitis may play a role in the development of hepatic IPT.

In our case, the etiology of the lesion remained unclear. The patient didn’t have any predisposing infections, no history of cholangitis or any abdominal surgery procedures. Recurrent cholangitis manifested only after the first ERCP and stenting. The microabscesses in the liver could have been a consequence of repeated invasive procedures, maintaining the inflammation in the biliary tract.

***Clinical manifestation***

Painless obstructive jaundice, abdominal pain, weight loss and fever are the most common clinical findings in patients with biliary IMTs. Their clinical presentation and imaging features are non-specific and are indistinguishable from cholangiocarcinoma, especially when hepatic hilum or bile ducts are involved[18].

Laboratory tests usually show leukocytosis, elevated C-reactive protein (CRP), anemia, thrombocytosis, polyclonal hypergammaglobulinemia, and slightly elevated liver enzymes. Tumor markers, such as carcinoembryonic antigen (CEA) and serum alpha-fetoprotein (AFP), are usually normal, although in some patients, marginally elevated CA 19-9 can be found[8,19].

Highly increased CA 19-9 is characteristic of cholangiocarcinoma and other malignancies[20]. Moreover, elevated CA 19-9 can also be observed in patients with benign hepatobiliary diseases. For instance, CA 19-9 elevation above 1000 U/mL can be observed in obstructive jaundice, acute liver injury or alcoholic liver injury and benign hepatic strictures[19,21]. In our case, CA 19-9 value was slightly increased on the onset and further increased to 1184 U/mL before starting chemotherapy. We found only one case in the literature regarding IPT with CA 19-9 elevation above 1000 U/mL[22]. CA 19-9 marker elevation may also be induced by chemotherapy or severe inflammatory process[23]. The latest CA 19-9 elevation to 7884 kU/L could have been a reaction to constant infections, disease progression, and COVID-19 disease. Considering this, the importance of CA 19-9 as a negative predictive factor of IMT and a prognostic mark of the disease progression is questionable as there were no changes in the stenosis site.

Imaging findings of IMT are not specific, and their radiological identification is not always possible. Typical radiological features of the biliary IMT are infiltrating hilar lesions and intrahepatic ductal dilatation[24].Tublin *et al*[18] studied biliary pseudotumors in seven patients and concluded that there were no radiological signs to differentiate them from cholangiocarcinoma. Thus, the main method to diagnose IMT is the histological examination of the biopsy or postoperative material[2,24].

Immunohistochemical studies of T and B-cell subpopulations may help to distinguish IMT from lymphoma. IMTs usually contain both T cells and B cells, whereas in lymphoma, a clonal B- or T-cell population predominates. Also, the heterogeneity of the inflammatory cell population in IMT tends to exclude lymphoma. Most IMT cases can be considered as IgG4 related based on their histopathologic features and immunohistochemistry criteria[25,26].

About half of recently analyzed cases of IMT harbors an anaplastic lymphoma kinase (ALK-1) gene rearrangement locus on chromosome 2p23, causing aberrant ALK-1 expression[26]. Although there is no statistically significant correlation between ALK-1 expression, tumor type, recurrence and metastasis, it is a useful diagnostic aid, based on which decision on chemotherapeutic treatment might be made[27]. It has been suggested that predominating histiocytic cells in IPT are associated with infection, whereas myofibroblastic cells characterize the lesions more likely to be considered true neoplasms[9,12]. Even if the biopsy of the mass shows signs of IMT, there is still a concern regarding the biopsy’s adequacy because only the inflammatory changes surrounding cancer may have been sampled. These fears often lead to an aggressive surgical approach as the initial treatment of choice.

***Treatment***

Although poorly understood, the etiology of IMT seems to relate closely to an altered immune response to injury. Infection, trauma, previous abdominal surgery, radiotherapy, chemotherapy, and steroid use - all have been thought to play a role in the pathogenesis of the heightened inflammatory reaction. Therefore, it seemed logical to treat this disease with an anti-inflammatory agent[28]. Some studies described the complete reduction and disappearance of the pseudotumor with conservative treatment with NSAIDs and steroids (such as prednisolone)[29–32].

Conservative management of IMT includes NSAIDs, corticosteroids and chemotherapeutic agents (like vinorelbine, methotrexate, azathioprine, cyclosporine, cyclophosphamide)[5,33-35]. It should also be emphasized that this conservative treatment could not be started in many cases due to the lack of definitive diagnosis of the mass preoperatively. Therefore, we performed diagnostic laparoscopy for a definite diagnosis.

However, in our case, treatment with NSAIDs was not effective, and steroids were discontinued due to increased inflammatory markers. Therefore, an alternative treatment plan was applied. Cytotoxic chemotherapy with methotrexate and/or vinorelbine/vinblastine (MTX-V) has been shown to be very effective, and long-term disease control could be achieved[34]. Casanova *et al*[5]reported the clinical findings and treatment results in the cohort of patients with IMT managed according to the European pediatric Soft Tissue Sarcoma Study Group protocol from 2005 to 2016. However, data is inconclusive on long-term remission, as studies have also shown recurrence of IMT combining surgery and chemotherapy[35]. Possible treatment options for IMT are summarised in Table 3.

Despite reported successful treatment, complete reduction and tumor regression using a conservative approach (corticosteroids, NSAIDs, antibiotics, chemotherapy), tumors identified as IMTs by histopathology are locally progressive and often need surgical resection, especially if medical therapy is not effective[36,37]. Although there are no approved recommendations for operative hepatobiliary IMTs treatment, the literature suggests that patients with resectable IMTs should be managed with radical surgical resection when it is anatomically and physiologically feasible[5,35-37]. The selection of liver resection candidates must be based on the patient's condition, tumor location, size and extension, the functional reserve capacity, and a sufficient liver remnant assessed by clinical and biochemical measures and by hepatic volume in cases of major hepatectomy[38,39]. Complete resection of hilar tumors requires a partial hepatectomy or extended hemi-hepatectomy[39,40]. There have also been some reports of a successful liver transplant, pancreaticoduodenectomy and combined liver transplant with pancreaticoduodenectomy in patients with hilar IMTs (Table 3)[1,37].

Cases with hilar tumors are always challenging (particularly those in perihilar cholangiocarcinoma) as anatomically these tumors in the hepatic hilum are in intimate relation with the portal vein and hepatic artery. Only about 1 in 5 patients with perihilar cholangiocarcinoma is eligible for surgery at the time of presentation[41]. The biliary extent of the tumor towards the segmental bile ducts is often more extensive than seemed on preoperative imaging[40].

The outcome of liver surgery during the past few decades improved. However, postoperative liver failure remains the leading cause of postoperative mortality. The assessment of hepatic functional reserve remains one of the most important issues in liver surgery[42]. The limit for "safe" liver resection is leaving a future liver remnant of at least 25% of the preoperative liver volume in patients with normal liver parenchyma or at least 30% to 40% in livers that are compromised by steatosis, chronic cholestasis, cirrhosis or chemotherapy[40,43,44]. The potential risks and benefits of the surgery must be considered, especially when there are no established recommendations in the surgical treatment of biliary IMT.

Contraindications for liver resection are similar to those of cholangiocarcinoma and include: (1) patient factors (medically unfit for operation, cirrhosis/portal hypertension, malnutrition); (2) local factors (bilateral involvement of secondary biliary radicles, encasement or occlusion of the main portal vein, atrophy of one lobe with encasement of contralateral portal vein branch, atrophy of one lobe with contralateral involvement of secondary biliary radicles); and (3) distant metastases[40,42].

Usually, IMTs are considered as benign neoplasms. According to the WHO classification, IMT is an intermediate-grade tumor with the potential for recurrence and rare metastasis[3]. In rare cases, IMT takes a malignant course and is hard to treat. There are no definite histopathologic, molecular, or cytogenetic features to predict malignant transformation, recurrence or metastasis[5]. Some IMTs are classified as neoplastic lesions, or reactive lesions, that have undergone a malignant transformation because a subset of IMT grows aggressively and shows a malignant behavior, sometimes with the formation of metastases. Whether a pseudotumor is a neoplastic or reactive process remains a debatable question. Some features, such as progressive growth, local recurrence, the development of multifocal masses, the destruction of liver substance and bile ducts, and the vascular (portal-venous) invasion, deny the purely reactive/inflammatory nature of these lesions. However, there are no signs of a neoplastic process in the histologic pattern[9,27,33].

***Diagnostic and clinical challenges of the case***

The patient’s clinical course of the disease and instrumental examinations were compatible with malignant disease, most likely perihilar cholangiocarcinoma. Therefore, multiple biopsies were taken using different approaches (from the site of strictures during ERCP and direct cholangioscopy; ultrasound-guided liver biopsy; diagnostic laparoscopy; biopsy from new mass near the gallbladder) and reviewed by several pathologists. However, none of them confirmed perihilar carcinoma or any other malignancy and were compatible with the diagnosis of IMT, which was negative for ALK-1. What is more, despite ineffective treatment and constant infections, the patient already survived 24 mo, and her general condition is satisfactory, which is not characteristic of cholangiocarcinoma. The median survival time of inoperable patients with Klatskin tumor is 6 to 12 mo[45].

This case’s management included repeated biliary tract stenting, glucocorticoids, NSAIDs, and, lastly, chemotherapy. Radical surgical treatment was impossible as the mass was locally advanced - extending into both hepatic and some segmental ducts. Liver transplantation was also not an option due to the patient's age and recurrent infections. The infectious complications almost after each ERCP and re-stenting procedure limited the ability to prescribe and continue steroids. Based on the literature, chemotherapy is one of the possible treatment methods for IMT as it is defined as intermediate-grade sarcoma. Therefore, after rigorous discussion with oncochemotherapists, we decided to start cytotoxic chemotherapy, as other treatment methods showed no effect. Although the patient‘s overall condition improved after the MTX-V scheme, it is probably misleading as the stricture length remained almost the same, and the MRCP images were performed during the time with and without cholangitis. Later on, it became impossible to continue chemotherapy due to constant infections. The patient developed COVID-19 disease, after which cholangitis became more severe: the patient developed cholecystitis, and some more solid masses were observed in the CT, indicating disease progression.

The reported outcomes of IMTs is ranging from completely benign to malignant and even fatal outcomes. In our case, IMT took a complicated course with constant infections leaving us with no effective treatment options.

**CONCLUSION**

IMTs can occur in various sites of the body. However biliary tract as the tumor’s primary location is reported rarely. Differentiation between benign and malignant strictures is a challenge for radiologists, endoscopists and gastroenterologists, as lesions are often indistinguishable from cholangiocarcinoma. Although IMTs are generally benign, these lesions can lead to hardly controlled recurrent infections or malignant course. Data about disease characteristics and effective treatment methods is lacking. Therefore, we report a case of an unusual localization, non-operable hard to diagnose and treat IMT of the biliary tract, and we hope this article will contribute to increasing the understanding of this rare condition.

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**Footnotes**

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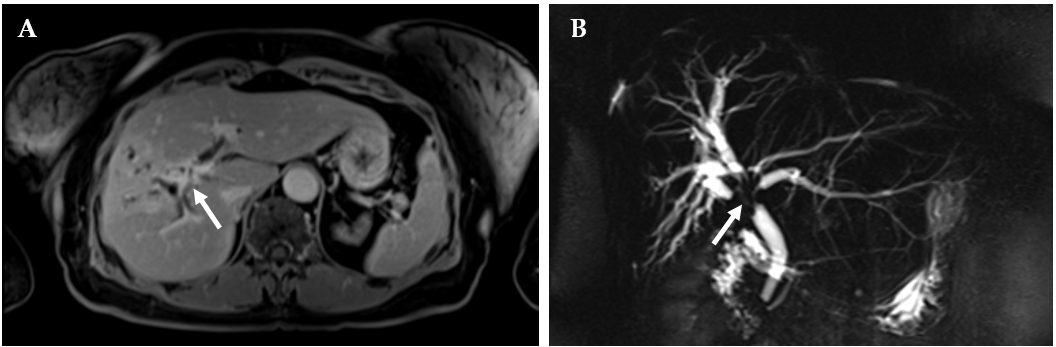
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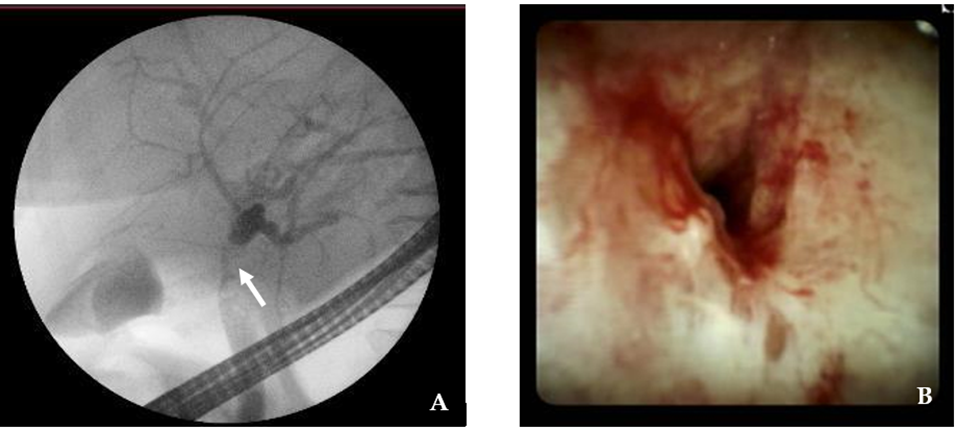
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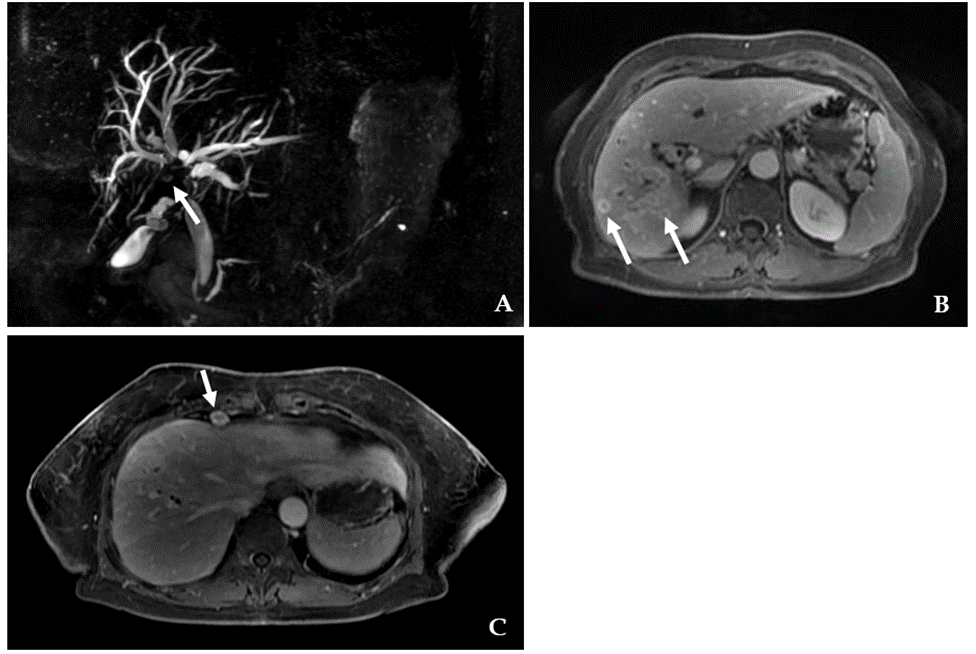
**Figure Legends**

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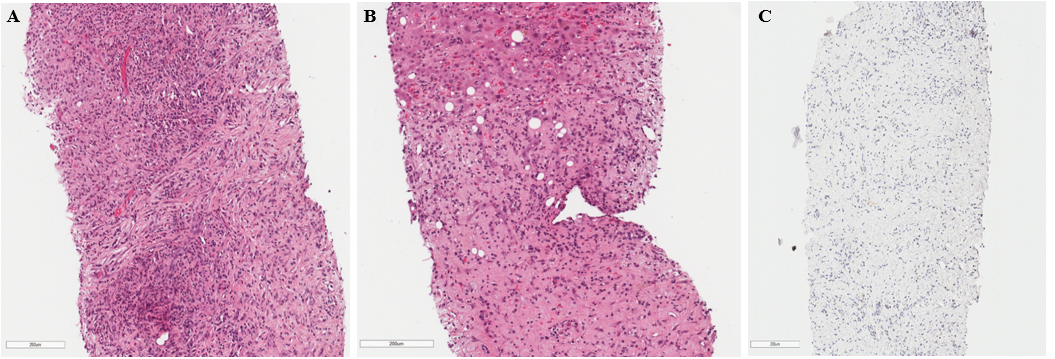
**Figure 1 magnetic resonance cholangiopancreatography images at presentation.** A: T1 FS late phase contrast-enhanced image showing soft tissue mass with gradual enhancement in the late phase around the wall of the common hepatic duct; B: Three-dimensional magnetic resonance cholangiopancreatography images showing a mass extending to segmental intrahepatic ducts on both lobes with dilatation of peripheral ducts.



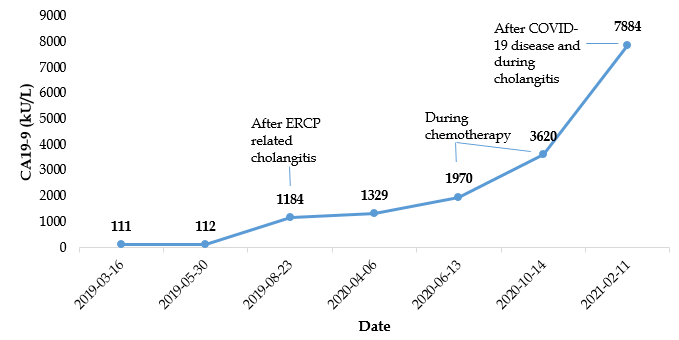
**Figure 2 endoscopic retrograde cholangiopancreatography and direct cholangioscopy images.** A:endoscopic retrograde cholangiopancreatography image showing proximal common hepatic duct (CHD) stricture (arrow) involving hepatic confluence with complete obstruction of the right hepatic duct; B: Direct cholangioscopy endoscopic image showing stricture of the proximal CHD.

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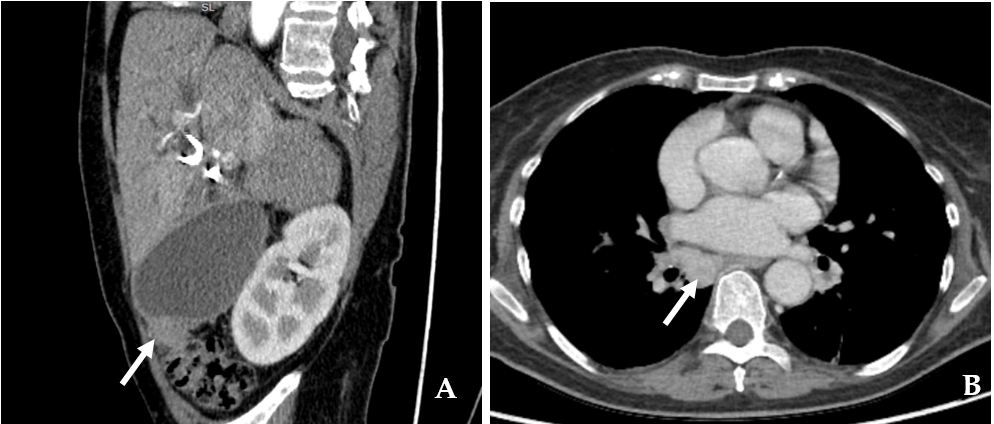
**Figure 3****magnetic resonance cholangiopancreatography images 3 mo later.**A: Three-dimensional magnetic resonance cholangiopancreatography (MRCP) images showing a common hepatic duct stricture (arrow) extending to intrahepatic segmental ducts extent to similar extent as on the previous MRCP with peripheral duct dilatation; B: FS contrast-enhanced images showing a cluster of rim enhancing lesions most likely small abscesses due to cholangitis (arrows); C: Enlarged cardiophrenic lymph nodes (arrow).



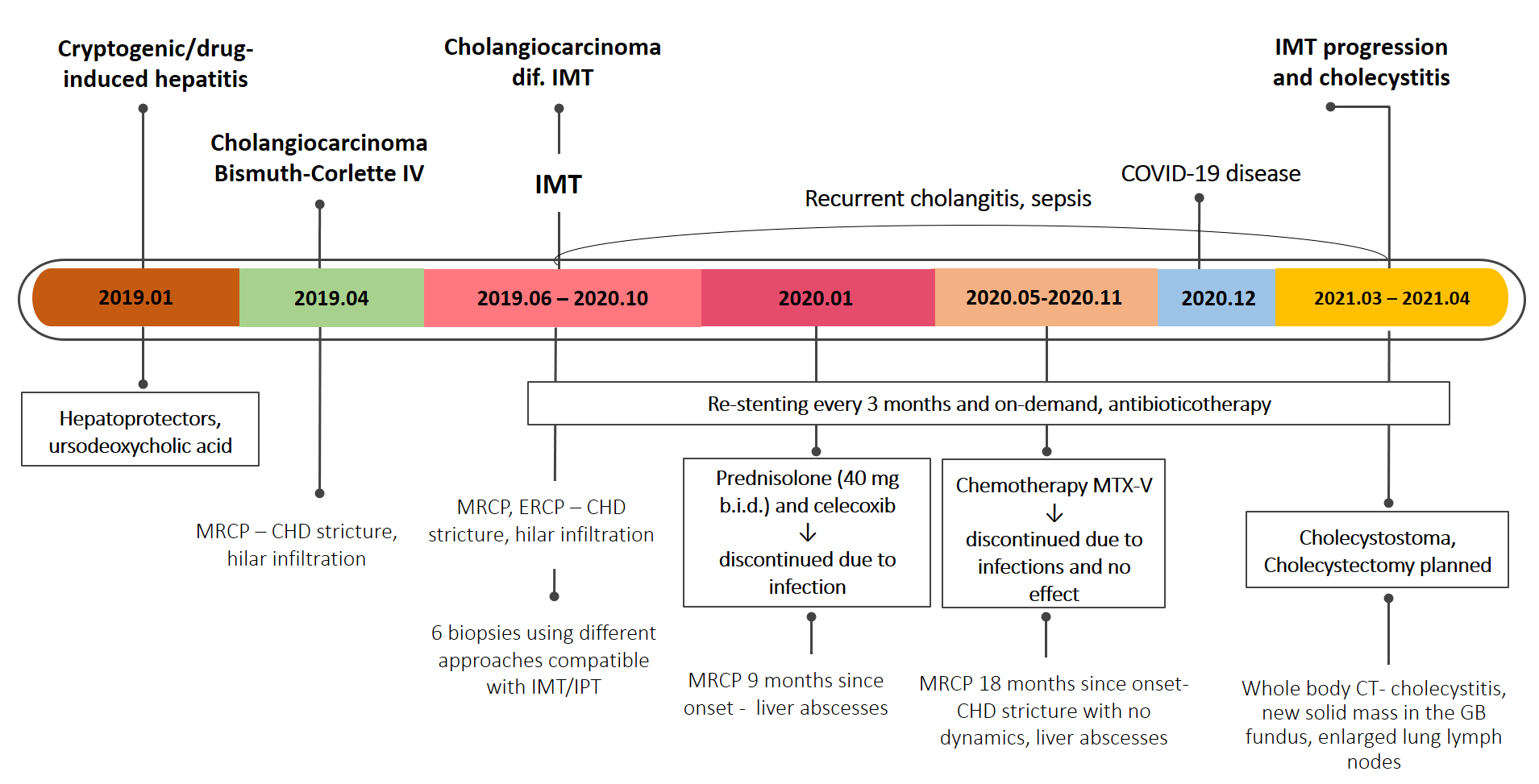
**Figure 4 Histological images.** A, B:The tumor is formed of disordered myofibroblasts and fibroblasts. Focal abundant polymorphonuclear infiltration can be seen (top and bottom left) (HE x 200); C: Negative ALK-1 immunohistochemical reaction.



**Figure 5 carbohydrate antigen 19-9 dynamics during the treatment.** CA 19-9: carbohydrate antigen 19-9;ERCP: Endoscopic retrograde cholangiopancreatography.



**Figure 6 computed tomography images 22 mo after the initial diagnosis.**A: Reformatted two-dimensional computed tomography (CT) images in the portal venous phase showing a contrast-enhancing soft tissue mass (arrow) arising from the gall bladder fundus and extending into colon mesenterium; B: Axial CT portal venous phase images showing a large lymph node (arrow) in the right lung hilum.



**Figure 7 Case history timeline.** MRCP: magnetic resonance cholangiopancreatography; CHD: common hepatic duct; IMT: inflammatory myofibroblastic tumor; ERCP: endoscopic retrograde cholangiopancreatography; IPT: inflammatory pseudotumor; MTX-V: methotrexate-vinorelbine; CT: computed tomography; GB: gallbladder.

**Table 1** **Main laboratory findings**

|  |  |  |
| --- | --- | --- |
|  | **Value** | **Normal range** |
| White blood cell count | **10.04** | 4.0 x 109/L-9.8 x 109/L |
| Hemoglobin | **125** | 128-160 g/L |
| Platelet count | 328 | 130 x 109/L-400 x 109/L |
| C reactive protein | **55,4** | ≤ 5 mg/L |
| Aspartate aminotransferase | **76** | < 40 U/L |
| Alanine aminotransferase | **135** | < 40 U/L |
| γ- glutamyl transferase | **344** | ≤ 36 U/L |
| Alkaline phosphatase | **376** | 40-150 U/L |
| Total bilirubin | **30** | < 21 μmol |
| Albumin | 36.2 | 36-52 g/L |
| K+ | 4.3 | 3.8-5.3 mmol/L |
| Na+ | 132 | 134-145 mmol/L |
| SPA | 95 | 70%-130% |
| INR by Owren | 1.02 | 0.90-1.19 |
| Creatinine | 51 | 62-11 µmol/L |
| Urea | 3.7 | 2.5-7.5 mmol/L |
| CA 19-9 | **111.98** | < 37 kU/L |
| CEA | 1.0 | < 5 mkg/L |
| AFP | 1.30 | 0.5-5.5 kU/L |
| HBsAg | 0.11 negative | negative s/co |
| Anti-HCV | 0.11 negative | negative |
| AMA: M2 | Negative | negative |
| ANA | negative (1:40) | < 1:40 |
| Immunoglobulin G4 | 0.653 | 0.08-1.40 g/L |

INR: international normalized ratio; SPA: Stago prothrombin assay; CA 19-9: carbohydrate antigen 19-9; CEA: carcinoembryonic antigen; AFP: alpha-fetoprotein; HBsAg: hepatitis B surface antigen; HCV: hepatitis C virus; AMA: antimitochondrial; ANA: antinuclear.

**Table 2 Cases of biliary inflammatory myofibroblastic tumors**

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
| **Ref.** | **Age /gender** | **Location/extension** | **Treatment** | **Follow- up** | **Outcome** |
| Haith *et al*[46] | 6/M | Distal CBD | PD and celecoxib | 5 mo | NR |
| Stamatakis *et al*[47] | 13/F | Proximal CBD, cystic duct, CHD | Extrahepatic BD excision | 21 mo | NR |
| Ikeda *et al*[48] | 43/M | Proximal CBD/ intrahepatic ducts, CHD, gall bladder, lymph nodes | Surgery | 7 mo | Lung metastasis |
| Fukushima *et al*[49] | 58/F | Mid-CBD/pancreas and lymph nodes | PD | - | NR |
| Walsh *et al*[50] | 50/M | Proximal CBD | PD | 19 yr | Metastasis |
| Sobesky *et al*[51] | 51/F | Distal CBD | PD | 2 yr | NR |
| Venkataraman *et al*[52] | 17–52/3 M, 1 F | 1 liver mass  1 liver mass and periportal infiltration  2 periportal infiltration | - | - | - |
| Büyükyavuz *et al*[53] | 8/F | Hepatic hilar | Surgery, antibiotics | - | - |
| Lopez-Tomassetti Fernandez *et al*[54] | 55/F | Distal CBD | PD | 4 yr | R |
| Martín Malagón *et al*[55] | 51/F | Distal CBD | PD | - | - |
| Kim *et al*[56] | 63/F | Hilar bile duct | Hepatectomy, caudate lobectomy | 5 mo | R |
| Sekaran *et al*[57] | 17/F | Left hepatic, CHD, proximal CBD | Left hepatectomy | 6 wk | NR |
| Abu-Wasel *et al*[58] | 55/F | Distal CBD | Extrahepatic BD excision | 14 mo | NR |
| Vasiliadis *et al*[2] | 70/F | Mid-CBD | Extrahepatic BD excision | 8 mo | NR |
| D’Cunha *et al*[59] | 12/F | Distal CBD | Debulking, corticosteroids | - | NR |
| Verma *et al*[1] | 24/F | Mid-CBD | Extrahepatic BD excision | 12 mo | NR |
| Karimi *et al*[36] | 12/F | Hepatic duct bifurcation | CBD resection | - | R |
| Present case | 70/F | Proximal CBD, CHD | Corticosteroids, chemotherapy | 24 mo | Disease progression |

M: male; F: female; CBD: common bile duct; PD: pancreatoduodenectomy; NR: no recurrence; CHD: common hepatic duct; BD: bile duct; US: ultrasound; R: recurrence.

**Table 3 Possible treatment options for inflammatory myofibroblastic tumors**

|  |
| --- |
| **Possible treatment options for IMTs** |
| High-dose steroids  Low dose steroids  Non-steroidal anti-inflammatory drugs (*e.g.*, celecoxib)  Vinblastine and methotrexate  Anaplastic lymphoma kinase inhibitors  Ifosfamide-based chemotherapy  Vinorelbine and low-dose cyclophosphamide  Vincristine and actinomycin-D  Cyclosporine, azathioprine  Radical surgical treatment (when anatomically and physiologically feasible) |

IMTs**:** inflammatory myofibroblastic tumors.