Mirizzi syndrome from type I to Vb: a single center experience

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ABSTRACT

Objective: The present study describes a cohort of patients diagnosed with Mirizzi syndrome from type I to Vb, over a period of four years. It aimed to identify diagnostic and management pitfalls of Mirizzi syndrome, as well as their concomitant cholecystobiliary or cholecystoenteric fistulas.

Material and Methods: We retrospectively reviewed all electronic medical records of patients who underwent surgery for Mirizzi syndrome at a single institution.

Results: Twenty-two patients (0.6%) were diagnosed with Mirizzi syndrome. Most of the patients were females (n=19, 86.3%). Mean age was 43.8 years (range: 21-71 years). Ultrasound was performed in all (100%) patients. Six (27.2%) patients had a CT scan and six (27.2%) patients had endoscopic retrograde cholangiopancreatography. Overall preoperative diagnosis was achieved on 36.6% (n=8) of the patients. There were the same total and partial cholecystectomies, accounting for ten (45.5%) cases each, one hepaticojejunostomy with cholecystectomy (4.5%), and one enterolithotomy (4.5%). Laparoscopic cholecystectomy was attempted in 15 (68.1%) patients, with conversion to open surgery in 93.3% (n=14) of the patients. An open approach was made in five (22.7%) cases. Four (18.1%) patients were reported as MS type I, both types II and III each account for 22.7% (n=5) of the cases, there was only one (4.5%) patient with type IV, and seven (31.8%) patients with type V.

Conclusion: There are limited studies of patients with Mirizzi syndrome, including type V classification, and when this syndrome is suspected, a preoperative diagnosis should be made to avoid bile duct injuries or lesions to adjacent organs.

Keywords: Biliary disease, cholecystectomy, cholecystobiliary fistula, cholecystoenteric fistula, gallstone disease, mirizzi syndrome

INTRODUCTION

Gallstone disease is a common digestive disease with an estimated prevalence of 10-20% in adults in developed countries (1,2). This disorder occurs when there is an imbalance in the composition of the bile, resulting in precipitation of one or more of its components (3). Mirizzi syndrome (MS) is an uncommon phenomenon, with an incidence of 0.7-2.9% of all cholecystectomies (2,4,5). Pablo Luis Mirizzi first described this entity in 1948, defining it as an obstruction of the common hepatic duct (CHD) or common bile duct (CBD) by the compression of an impacted stone in the neck of the gallbladder or cystic duct, causing obstructive jaundice, and leading occasionally to fistulization to the bile duct or surrounding organs (6,7).

The treatment of MS is either by laparoscopy or by open approach, with high conversion rates with the former, consisting of partial or complete cholecystectomy with or without common bile duct exploration, and sometimes biliocystic anastomosis may be performed (8).

MS is a severe disease, and preoperative detection or intraoperative recognition of MS is essential for the surgeon to reduce the risk of operative complications, being the most common bile duct injury and residual stones (1).

The present study reviewed the experience of a single center with Mirizzi syndrome over a period of 4 years. It aimed to identify diagnostic and management pitfalls of Mirizzi syndrome, as well as their concomitant cholecystobiliary or cholecystoenteric fistulas.

This study was approved by IRB, Tecnologico de Monterrey, under the number 098.
MATERIAL and METHODS

From 2014 to 2018, 22 consecutive patients underwent surgery for MS at a single teaching institution. We searched the surgical database for all patients diagnosed postoperatively with MS, and retrospectively reviewed all electronic medical records. Patient demographics, clinical characteristics, diagnostic method, surgical procedures, outcomes, and follow-up were all documented and organized. All patients underwent physical examination, laboratory testing, and ultrasonographic (US) or computed tomography (CT) evaluation of the gallbladder. All the cases diagnosed were based on preoperative investigations or intraoperative findings. Beltran and Csendes modified classification (9), was followed to categorize patients as the diagnostic criteria. Laparoscopic cholecystectomy under general anesthesia with a standard four-port technique was attempted in most patients. Conversion to open surgery was left to the discretion of the surgeon. When dense adhesions to adjacent organs and impacted stones in Hartmann’s pouch rendered access to Calot’s triangle difficult, a fundus-first dissection technique was applied, and subtotal cholecystectomy was performed. All patients were seen in the outpatient clinic within the first month of the initial surgery, and all patients had a follow-up of up of at least six months after surgery. Patients were examined clinically, and liver function tests were evaluated for each follow-up visit.

This manuscript was approved by Tecnologico de Monterrey ethics committee and institutional review board (IRB) number 098, and was therefore performed in accordance with the ethical standards laid down in the 1964 Declaration of Helsinki and its later amendments. All patients provided informed consent to participate in the surveillance protocols.

Data were reported as mean, range, or percentages for continuous variables and frequencies and percentages for categorical variables. Data analysis was performed, with a combination of Excel 16.0 (Microsoft Corporation, United States) and R software 3.4 (R Core Team, New Zealand).

RESULTS

A total of 3556 cholecystectomies were performed during the study period. Twenty-two patients (0.6%) were diagnosed with Mirizzi syndrome. Most of the patients were females (n= 19, 86.3%), with only three (13.6%) male patients. Mean age was 43.8 years (range: 21-71 years), four (18.1%) patients had diabetes mellitus (DM) concomitantly with systemic arterial hypertension (SAH), two (9%) patients had only DM, and one (4.5%) patient had only SAH. One (4.5%) patient had hypothyroidism, and one (4.5%) patient was asthmatic. All patients had a history of abdominal pain of 10.9 months on average (range 1-76 months), with a mean time of exacerbation of abdominal pain before their surgical treatment of 6 days (range: 1-21 days). Thirteen (59%) patients were jaundiced at presentation. Liver function tests were altered in almost all patients (n= 18, 81.8%) (Table 1).

US was the initial imaging study performed in all patients. It revealed gallstones and features of acute cholecystitis in every case (100%), plus a single (4.6%) patient had choledocholithiasis. Mean size of gallbladder wall thickening was 4.86 mm (range: 1.2-10 mm), and mean diameter size of CBD was 8.21 mm (range: 3.17-14 mm). Ten (45.4%) had an impacted stone on the Hartman pouch or cystic duct, and in only six (27.2%) patients, a CT scan was ordered for suspicious of cholecystoenteric fistula, finding on 66% (n= 4) of the patients neumobilia and a sigmoid gallstone ileus on one (16.6%) patient. Endoscopic retrograde cholangiopancreatography (ERCP) was performed in six (27.2%) cases, performing sphincterotomy for bile duct decompression, placing bile duct stents at the end of the procedure. The diagnosis of MS was made in 50% of these patients by direct identification of a fistula.

Postoperatively, four (18.1%) patients were reported as type I, both types II and III each account for 22.7% (n= 5) of the cases, there was only one (4.5%) patient with type IV, treated with an open cholecystectomy and Roux-en-Y hepaticojejunostomy, and the remaining seven (31.8%) patients were found to have various types of cholecystoenteric fistulas (type V). We had the same number of patients with cholecystoduodenal and cholecystocolonic fistulas, accounting for three patients with each type of fistula. Of the three patients with cholecystoduodenal fistula, two were treated with a graham patch on the duodenum and primary closure of the CBD, and one was treated with choledochoplasty with a gallbladder flap and primary closure of the duodenum. Of the patients who presented with a cholecystocolonic fistula, a colostomy at the site of the fistula was performed on one patient, with primary closure of the CBD, another was treated with primary closure of the transverse colon and T tube insertion on the CBD, and the third patient was treated with enterolithotomy alone, for an intestinal obstruction due to a sigmoid gallstone ileus. The remaining patient with MS type V presented a gastrobiliary fistula on the lesser curvature.

<table>
<thead>
<tr>
<th>Table 1. Laboratories values at initial presentation</th>
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<td>Laboratories values</td>
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<tr>
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</tr>
<tr>
<td>WBC X 10^9/L</td>
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<tr>
<td>Total Bilirubin (mg/dL)</td>
</tr>
<tr>
<td>Direct Bilirubin (mg/dL)</td>
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<tr>
<td>Indirect Bilirubin (mg/dL)</td>
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<tr>
<td>AST (U/l)</td>
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<td>ALT (U/l)</td>
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<td>AP (U/l)</td>
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<td>GGT (U/l)</td>
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treated with primary closure, and an omental patch on the site of the fistula in the stomach, a T tube was placed with a reconstruction of the CBD (Figure 1).

Overall, there were equal number of total and partial cholecystectomies, accounting for ten (45.5%) cases each, one hepaticojejunoanostomy with cholecystectomy (4.5%), and one enterolithotomy (4.5%). Laparoscopic cholecystectomy was attempted in 15 (68.1%) patients, and because of difficult dissection and misleading anatomy in Calot’s triangle, conversion to open surgery was achieved in 93.3% (n= 14) of the patients. An open technique was used as the initial procedure in five (22.7%) cases. A trans-operative cholangiogram (TOC) was attempted in 76.1% (n= 16) of the cases where a cholecystectomy was performed (n= 21), being conclusive in 13 (81.25%) patients with evidence of either cholecystobiliary or cholecystoenteric fistula. In eight (38%) patients, a T-tube was placed on the common bile duct, and all patients had a tube drain left in the subhepatic space, which was removed within a mean of 5 days (range: 3-10 days) (Table 2).

Overall procedure-related morbidity was 13.6% (n= 3). Two patients developed bile leak, and one patient had a remaining stone. All were treated with ERCP and placement of biliary stents, removing them on postoperative week six, with an outstanding outcome. Mortality was accounted for 9% (n= 2) of the patients. In the remaining patients, mean length of hospital stay (LOS) was 8 days (range: 2-30 days). Final histopathology revealed chronic cholecystitis in all (100%) patients, except one 51-year-old patient, who presented associated adenocarcinoma of the gallbladder and treated with adjuvant therapy. All patients were followed up in the ambulatory clinic for at least 12 months, with a mean period of 23 months (range: 12-36 months), all symptoms free with normal liver function tests, except the patient treated with hepaticojejunoanostomy anastomosis, who had a persistent elevation of serum alkaline phosphatase and developed a mild episode of cholangitis treated successfully with antibiotics, with no further complications. The patient with gallbladder adenocarcinoma stage I did not need

Table 2. Type of surgical treatment

<table>
<thead>
<tr>
<th>Total (n= 22)</th>
<th>Type I (n= 4)</th>
<th>Type II (n= 5)</th>
<th>Type III (n= 5)</th>
<th>Type IV (n= 1)</th>
<th>Type Va (n= 6)</th>
<th>Type Vb (n= 1)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Open (n= 5)</td>
<td>-</td>
<td>2+</td>
<td>2</td>
<td>-</td>
<td>1</td>
<td>-</td>
</tr>
<tr>
<td>Laparoscopic + Conversion (n= 14)</td>
<td>3</td>
<td>3 (2*, 1*)</td>
<td>3*+</td>
<td>-</td>
<td>-</td>
<td>5 (2*, 3+)</td>
</tr>
<tr>
<td>Laparoscopic (n= 1)</td>
<td>1</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Hepaticojejunoanostomy (n= 1)</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>1</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Enterolithotomy (n= 1)</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>1</td>
</tr>
</tbody>
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*, T-tube; +: partial cholecystectomy.
any further surgery, and 12 months after his initial surgery, he is currently alive with oncology follow up.

**DISCUSSION**

The MS incidence of 0.6% in the present series correlates with the 0.7%-2.53% stated in the literature (10-12). This syndrome develops in patients with longstanding gallstone disease, with a female predominance, with this series supporting this majority. We had a mean age of 43.8 years, making it inferior to the ones reported in the literature that ranges from 48 to 61 years (9). The main confront in the management of MS is in accomplishing a precise preoperative diagnosis, with rates ranging from 8% to 62% of obtaining an accurate image identification of this condition, if this is not accomplished, the incidence of bile duct injuries could be as high as 17% (13,14). Several image studies can be used for achieving this; US is the most common modality used for the diagnosis of gallstones; however, it has limited sensitivity (48%). CT may show dilation of the biliary tree and the CBD, with low sensitivity for identifying stones at these sites, but it may exclude a malignancy in the porta hepatis area or the liver. Magnetic resonance cholangiopancreatography (MRCP), is a non-invasive imaging technique with a 50% diagnostic accuracy rate, having the advantage of avoiding the complications associated with ERCP, which is considered the gold standard for diagnosing this disorder with a mean sensitivity rate of 76.2%, with technical limitations in 5% to 10% of the cases, including inaccessibility to the bile ducts and incomplete filling of the ducts because of tight strictures, additionally it is not exempt from complications such as pancreatitis, cholangitis, and residual stones (4,14-16).

Since, we lack MRCP in our institution, a preoperative diagnosis was challenging to attain, and even though almost half of our patients had a US with an impacted stone, a direct compression to the hepatic or common bile duct was difficult to visualize, therefore in patients with high suspicious of MS, a ERCP or a CT scan was ordered, visualizing a fistula in three patients with the former study, and neumobilia in four patients with the latter. We achieved an overall preoperative diagnosis of 36.6% in our patients, with nearly the same rate as the series of Greiasov et al. (17) who accomplished a preoperative diagnosis rate of 27% in a large cohort of 284 patients.

An often-absent sign of identification of this disease and almost pathognomonic is neumobilia (18). This sign was seen in 66% of our patients who had CT (n=6), higher than the series from Li et al. (19) where this sign was seen in 33% of the CTs. We opted to treat our three patients with cholecystoduodenal fistula with primary closure and choledochoplasty in the CBD, performing primary closure of the duodenum with or without over sewing an omental patch, with satisfactory results. We had the same number of patients with cholecystoduodenal and cholecystocolonic fistulas, similar to the study of Pradeep et al. (20), who identified the duodenum and the colon, as their two most common organs involved. On the patients where the colon was involved, we performed on partial cholecystectomies. On one patient, we performed primary closure of the transverse colon and T tube insertion on the CBD, on a second patient, a colostomy was achieved with intestinal reconnection auspiciously implemented 20 weeks later, and on a third patient, a gallstone ileus on the sigmoid colon along with a cholecystocolonic fistula was faced, opting for a conservative management because of a low functional reserve, achieving only an enterolithotomy and antibiotic therapy with positive outcomes. This unusual site of impaction of the stone is exceptionally infrequent, occurring only in 8% of all patients with gallstone ileus (20-21).

There are two popular classifications well accepted for this condition. The first one is the one proposed in 1982 by McSherry and colleagues (22), categorizing MS into two types: type I, characterized by external compression of the adjacent common hepatic duct; and, type II, where a pressure necrosis of the common bile duct results in a cholecystocholedochal fistula. The second classification was developed seven years later by Csendes et al. (10) reclassifying MS in IV types, being type I lesions with external compression of the common bile duct, type II lesions where a cholecystocholedochal fistula is present with erosion of less than one-third of the circumference of the bile duct, distinguishing it from type III lesions where the fistula involves up to two-thirds of the duct circumference and finally type IV lesions where there is complete destruction of the bile duct. Almost two decades later, Csendes added an additional type to this classification, validated by Beltran (9), supplementing a type Va and Vb, corresponding with an MS with a cholecystoenteric fistula without and with a gallstone ileus respectively. Our most common type of MS was the type V, following equally by type II and III, and because MS type I does not involve a fistula, we think numerous MS type I passed undiagnosed, categorizing them as hydropic gallbladder or acute cholecystitis.

Beltran et al. (9) have supported that conventional surgery for patients with suspected MS is safer in most institutions of developing countries where access to diagnostic equipment, such as MRCP, ERCP, intraoperative ultrasonography, or cholecotography are not available. Laparoscopic management can be performed especially in MS type I with a visualized cystic duct, and only cholecystectomy either total or partial is needed. However, patients with the other four types of MS always require bile duct exploration with intraoperative cholangiogram and common bile duct reconstruction through simple closure of the fistula or T-tube insertion, leaving biloenteric anastomosis for patients with type IV (15). Of all surgeries performed at our hospital, only one patient with MS type I was completed by a laparoscopic approach; hence, we had a conversion rate of almost 100%. In patients with MS type II and III; a T tube was placed on 60% of them, with the remaining patients treated with primary closure or choledochoplasty. Almost all of these patients had adequate results, except for Turk J Surg 2020, 36 (4): 399-404
our two fatalities, being two elderly and diabetic patients, treated with T-tube placement and without T tube placement, MS type II and MS type III respectively, presenting bile leakage treated at first with ERCP and biliary stent placement, reinventing both cases with abdominal washout and drainage placement with no favorable evolution. The patients died 45 days and 80 days after the initial surgery due to abdominal sepsis. Moreover, a bilioenteric anastomosis was achieved for our patient with MS type IV with worthy results.

The clinical diagnosis of MS is challenging to attain since there are no pathognomonic symptoms for presentation (4). Thus, obtaining a preoperative diagnosis is deficient, because signs like jaundice, acholia, or choluria may be lacking, adding that 20% to 40% of the patients have normal serum bilirubin levels (23). In our series, 18.1% of our patient’s laboratory values were with standard parameters, and 59% of the patients presented with jaundice, differently from the series of Kwon et al. (23) where only 33% presented with this sign.

In their systematic review, Antoniou et al. (8) have stated that the present with this sign. Although signs and symptoms are not precise, the primary clinical manifestation is jaundice with a predominantly obstructive laboratory pattern, along with right upper quadrant pain, making it difficult to distinguish it from cholangitis. This is one of the few case series describing MS, including the type V classification, contributing with our experience to the existing literature regarding this unusual and controversial syndrome.

REFERENCES


Tip I'den Vb'ye Mirizzi Sendromu: Tek Merkez Deneyimi
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ÖZET

Giriş ve Amaç: Bu çalışma, dört yıllık bir süre boyunca tip I'den Vb'ye Mirizzi sendromu (MS) tanı konan bir hasta grubunu içermektedir. Mirizzi sendromunun tanısı ve yönetimindeki zorluklar ve eşlik eden kolesistobiliyer ve kolesistoenterik fistülleri tanımlayıcı amaçlarla incelemiştik.

Gereç ve Yöntem: Tek bir merkezden来的 Mirizzi sendromu nedeniyle ameliyat edilen hastaların tüm elektronik tıbbi kayıtlarını retrospektif olarak incelledik. Tek bir kurumda Mirizzi sendromu nedeniyle ameliyat edilen hastaların tüm elektronik tıbbi kayıtlarını retrospektif olarak incelledik.

Bulgular: Yirmi iki hastada (%0,6) Mirizzi sendromu tanısı konuldu. Hastaların çoğunun %9, %86,3, %21, %27 ve %11, %27 dişiye ait idi. Hastaların ortalaması 43,8 idi (aralıksız: 21-71 yaş). Tüm hastaların %100'sinde ultrasonografi tanısı konuldu. Alit hastada (%27,2), BT taraması, alit hastada (%27,2) endoskopi retrograd kolanjijiyopankreatografi uygulandı. Preoperatif tanı hastaların %36,6'sında (n=8) sağlandığı. Her birinde oran olsunun (%45,5) olduğu ancak parsiyel kolesistektomi, bir (%45) kolesistektomi ile hepatikojenüstomi ve bir de (%45) enterolitotomi uygulandı. %22,7'lik bir olgu شهرda mirizzi sendromu tanısı konuldu.

Sonuç: Mirizzi sendromlu hastaların tip V siniflandırılması dahi sinirli sayıda sıklıkla vardır ve bu sendromdan şuphelenildiğinde, safra kanalları yaralanmalarını veya komşu organlara lezyonları önlemek için ameliyat önerisi tanı konulmalıdır.

Anahtar Kelimeler: Safra hastalığı, kolesistektomi, kolesistobilyer fistül, kolesistoenterik fistül, safra taşı hastalığı, Mirizzi sendromu

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