Case Report

Immunoglobulin (IgG4) Sclerosing Cholecystitis-Camouflage for Gall Bladder Cancer- Case report and Review of Literature

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Received Date: 31 May, 2022; Accepted Date: 08 June, 2022; Published Date: 10 June, 2022

Abstract

IgG4 related disease is a rare systemic disorder supposed to have an underlying autoimmune phenomenon. Majority of these disorders are associated with autoimmune pancreatitis component. These disorders are difficult to diagnose as these disorders mimics inflammatory as well malignancy and poses a real mammoth task to diagnose and treat. Our case report includes a gall bladder mass without any pancreas involvement suspected to be a gall bladder cancer on clinically as well radiologically basis. Patient managed surgically and final histopathology revealed immunoglobulin G4 (IgG4) related cholecystitis. This disease could be managed conservatively by giving oral steroids, if it has been picked up preoperatively and major surgical intervention have been avoided. No defined blood test or tumour markers are currently available to diagnose this entity except serum immunoglobulin G4 that is costly and not feasible in developing nations like India.

Keywords: Immunoglobulin G4; Sclerosing cholecystitis; Gall bladder mass, Gall bladder cancer

Abbreviations: IgG4: Immunoglobulin G4; IgG4-RD: Immunoglobulin G4 Related Disorder

Introduction

IgG4 related disease is a rare systemic disorder, which can involve any organ of our body. To club these different organs involvement by this rare similar autoimmune phenomenon, Kamisawa, et al. [1] proposed a new common term of systemic IgG4-related autoimmune diseases in 2003 as IgG4 related disorder (IgG4-RD). These disorders are difficult to diagnose clinically as well radiologically due to the overlapping of clinical and radiological imaging features. These disorders mimics inflammatory as well malignancy and poses a real difficulty in diagnosis, treatment and predicting the natural course of disease. These disorders are immune mediated and mainly affects pancreatico biliary tree i.e. pancreas, gall bladder and biliary tree (extrahepatic as well intrahepatic). Apart from these organs, it can involve head and neck region, retroperitoneal organs, genitourinary organs, salivary glands and thorax [2,3]. However, the incidence of involvement of all these organs/sites is less as compared to pancreas. Autoimmune pancreatitis is a well-studied entity and along with it involvement of gall bladder bilary tree i.e. cholecystitis and cholangitis respectively is common. Approximately 20-30% of gall bladder involvement is there in autoimmune pancreatitis [4,5].

All these immune related entities are covered under one umbrella and termed as-IgG4 related diseases (IgG-RD) [6]. Isolated gall bladder involvement and its clinical manifestations are rare and still evolving entity and literature also has paucity of
It is imperative to differentiate this autoimmune entity of gall bladder IgG 4 sclerosing choleystitis from other causes especially malignancy of gall bladder in order to avoid overtreatment as both of these have different treatment and completely different prognosis. Diagnosis of this entity is really a head scratching task as it has a totally similar presentation and most of the time it is misdiagnosed and wrongly treated.

Isolated IgG4 cholecystitis mimicking as cancer is not much reported in literature and here we are reporting a case report of IgG4 cholecystitis masquerading as gall bladder mass/cancer. Most of the case report of IgG4 cholecystitis are associated with autoimmune pancreatitis (AIP) and reporting of isolated involvement of IgG4 cholecystitis are very few. Our case report will help in suspecting IgG4 cholecystitis whenever we encounter gall bladder mass on imaging and adds up to the existing literature of isolated involvement of gall bladder in IgG4 cholecystitis.

**Case Report**

64-year lady, post-menopausal, non-smoker and non-alcoholic residing in rural area of the state with known history of diabetes mellitus with poor glycemic control as patient was not taking oral hypoglycemics in a regular manner. She was evaluated for pain in the right upper abdomen along with dyspeptic symptoms lasting for one year. Patient was referred to us in a tertiary care hospital. On examination, her vitals were stable and performance status was ¼ as per ECOG (Eastern cooperative oncology group) and her GPE (general physical examination) was within normal limits and abdomen was tender at right hypochondriac and epigastric region.

Laboratory investigations: total leucocyte count 9800/mm³, total bilirubin 1.07 mg/dL, aspartate aminotransferase 21 U/L, alanine aminotransferase 27 U/L, alkaline phosphatase 169 U/L, albumin 4.2 g/dL and PT-11.8, INR 0.64. HbA1c was 7.3 Carbohydrate antigen 19–9 (CA 19–9) levels were 70 U/mL and serum carcinoembryonic antigen (CEA) level was within normal range.

Ultrasound abdomen showed focal gall bladder thickening at fundal approximately 5-6 mm along with gall bladder stones. Patient was planned for triple phase CECT scan abdomen, which showed (Figure 1) contracted gall bladder with irregular asymmetrical thickening of 6 mm along with calcification in fundal region of gall bladder. Fat planes with adjacent liver was ill defined. Subcentimetric lymphadenopathy in inter-aortocaval region. Pancreas and bile duct and rest of the organ were normal.

First possibility of gall bladder cancer was kept under the background of porcelain gall bladder and irregular thickness of 6 mm. Due to suspicion of gall bladder cancer, patient underwent ultrasound guided fine needle aspiration cytology (FNAC) from gall bladder which was descriptive. In view of abnormal thickening and porcelain gall bladder, patient planned for surgery. Patient underwent extended cholecystectomy after 3 weeks of acute attack of pain and good glycemic control was obtained. On opening the abdomen, there was no ascites and metastasis. Gall bladder was thickened at fundal region along with multiple stones in lumen. Centimetric lymph nodes were found along hepatoduodenal ligament and inter-aortocaval region.

Patient had uneventful postoperative recovery and discharged on postoperative day 7. Final histopathology report showed dense fibrosis (Figure 2) arranged in a storiform pattern along with oblitative phlebitis and dense transmural lymphoplasmacytic infiltrate along with eosinophil and neutrophil (Figure 3) [7]. Lymph-nodes were harvested and all showed sinus histiocytic changes only. All these features were suggestive of IgG4 sclerosing cholecystitis. For confirmation, we performed Immunohistochemistry (IHC) for IgG4 and CD138 which showed IgG4 positive 80 plasma cell and CD138 was positive in plasma cells. IgG was inconclusive. Serum IgG4 was 0.972 (reference range 0.03-2.01)
Discussion

IgG4-cholecystitis is a total masquerader having a similar constellation of signs and symptoms as of gall bladder cancer. Etiopathogenesis of this entity is still not clear, but mostly it is stated that this is manifestation of autoimmune process driven by lymphoplasmacytic interaction in the organ involved. Depending upon the organ or system involved, it really poses difficulty in making final diagnosis based on clinical and radiological features. To circumvent this dilemma, we need a complete and in-depth understanding of the clinical presentations of IgG4 related diseases. This will eventually lead to decrease unnecessary invasive forms of surgical interventions and ultimately benefits the patient and helps in reducing the futile exercise. It is prudent to put this rare entity in the list of differential diagnosis along with gall bladder cancer whenever any abnormal or suspicious features are there on clinically/ radiologically [7].

As per Rui Zhang, et al. the incidence of misdiagnosis of IgG4-C is 9.63% [8]. This entity is more common in older people having more predilection towards male [9]. Patients with IgG4 cholecystitis on sonography shows hypoechoic, diffuse, circumferential thickening of the gallbladder Wall. As in our case, sonography showed gallstones along with, asymmetrical thickening of 6 mm in gall bladder.

On CECT scan, it is mainly the delayed enhancement in IgG4 related cholecystitis, which helps in differentiating it from gall bladder cancer. In our case, there was asymmetrical thickening of 6 mm along with porcelain bladder.

On MRI, it showed low-signal smooth diffuse gallbladder wall thickening on T2 weighted MR images, and delayed enhancement post-contrast [4,10,11].

This thickening or enlargement of organ on imaging is generally due to infiltration of organs by lymphocytes and plasma cells along with coexisting fibrosis.

As per Deshpande, et al. [12], the diagnostic criteria for diagnosing IgG4 related diseases are as mentioned in Table 1.
surgical complications which can be completely avoided if we overtreat IgG4 cholecystitis at the cost of other post operative embarking upon any modality of treatment. Any misdiagnosis will cancer. Therefore, a firm and confident diagnosis is required before and regional lymphadenectomy is mainly done for gall bladder after steroid treatment, the extent of surgery is definitely less as has been tried in Mayo clinic [15]. Even if patient require surgery trial has not been yet performed. Role of immune-modulators also relapse rate are yet not documented, as no randomised controlled cancer. As both these entities have completely different options of treatment. We should not rely solely upon clinical and radiological picture. Histopathological pictures completes the diagnosis confirmation. Role of serum IgG4 is still controversial, and needs more data to establish its role. Our case report will surely add some useful information in literature as an isolated case of IgG4 sclerosing cholecystitis without pancreas involvement.

Conclusion

IgG4 cholecystitis is an immune mediated disease whose pathophysiology is still not completely understood. Every clinician should keep possibility of IgG4 cholecystitis in mind whenever any patient with abnormal gall bladder thickening or gall bladder mass is encountered in their clinical practice, As both these entities have completely different options of treatment. We should not rely solely upon clinical and radiological picture. Histopathological pictures completes the diagnosis confirmation. Role of serum IgG4 is still controversial, and needs more data to establish its role. Our case report will surely add some useful information in literature as an isolated case of IgG4 sclerosing cholecystitis without pancreas involvement.

References


