Congenital Absence of Gall Bladder

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Summary:
Congenital absence of gall bladder is a very rare entity found in clinical practices. Due to misinterpretation of sonographic findings as they are not familiar with the condition it possess a great difficulty in management the patient.

Though the patient present with the feature of acute cholecystitis, conservative treatment is enough to cure the disease. Modern radiological intervention make the diagnosis confirm, thus preventing unnecessary surgical procedure.

Here we report two cases of Congenital absence of gall bladder presenting as acute cholecystitis ultimately diagnosed and treated.


Introduction
Biliary system variants are relatively common but is isolated gallbladder agenesis is a rare entity with an estimated incidence of 10–65 per 100,000. Females are more commonly affected (ratio 3:1), typically presenting in the 2nd or 3rd decade of life. Despite an absent gallbladder, half of patients present with symptoms similar to biliary colic, which is poorly understood. Symptomatic congenital absence of gall bladder puts the diagnostic dilemma and treatment difficulties.

Case Report:
1. A 35 years old woman, mother of two children, of Noakhali District, Bangladesh, presented with severe upper abdominal pain, bouts of vomiting, slight abdominal distention. She was non-icteric, mildly anaemic.

Physical examination revealed good health, remarkable tenderness over the Right hypochondriac region. Murphy's sign was positive. She was mildly dehydrated. Patient was normotensive and non diabetic. Examination of other systems revealed normal findings.

Routine laboratory examinations (Complete Blood Count, Erythrocyte sedimentation rate, blood biochemistry Panel) revealed nothing abnormal except Neutrophilic leucocytosis.

Liver function tests, serum Creatinine, Random blood sugar were within normal limit.

Ultrasoundographic scan done but the comment was “gall bladder could not visualized may be due to bowel gas” (Fig.-1)

Contrast enhanced computed tomography (Fig.-2) revealed absence of gall bladder without any remarkable change in the extra and intra hepatic biliary apparatus.

Common bile duct was normal in diameter with no calculus present. Magnetic Resonance Cholangiopancreatography done later (Fig.-3) which revealed absent gall bladder, normal intra and extra hepatic biliary channels. Common bile duct normal in diameter.

1st Patient (Printed with Permission)
Diagnostic laparoscopy done and gall bladder could not
found. Common bile duct and Right and Left hepatic
duct were found normal.

No other congenital abnormalities like annular pancreas
or atresia found. The patient was treated conservatively
with nothing per oral, intravenous fluid, Broad spectrum
antibiotics, Nasogastric Suctions, Analgesics and
antispasmodics.

2. The second patient 32 years old Bangladeshi woman,
mother of two children, of Comilla District Bangladesh,
admitted in a local clinic with severe upper abdominal
pain, fever, nausea, and anorexia. Physical examination
revealed good health, remarkable tenderness over the
Right hypochondriac region. Murphy’s sign was positive. She was mildly dehydrated. Patient was
hypertensive, non insulin dependent diabetic. Examination of other systems revealed normal.

Routine laboratory examinations revealed nothing
abnormal except Neutrophilic lucocytosis.

Liver function tests, serum Creatinine were within
normal limit.

Ultrasoundographic scan done and the comment was
“contracted gallbladder “ (Fig.-4)
The patient underwent laparotomy but during the
procedure gall bladder was absent. The extra hepatic
biliary apparatus were normal.

Fig.-1: Ultrasonographic Scan

Fig.-2: CT Scan

Fig.-3: MRCP Scan

Fig.-4: Ultrasonographic Scan

Fig.-5: CT Scan
Discussion:
Gallbladder agenesis is a rare entity with an estimated incidence of 10–65 per 100,000\(^1,2\). The incidence is noted to be higher (up to 90 per 100,000) in studies based on autopsy reports\(^3\). The first reports of cases of gallbladder agenesis date back to 1701 and 1702 by Lemery and Bergman\(^1,2,4\). The pathogenesis is related to embryonic development due to failure of the gallbladder and cystic duct to bud off from the common bile duct during the fifth week of gestation\(^1\).

Prior authors have classified patients into three groups. The first group consists of asymptomatic anatomical abnormalities seen incidentally on autopsy. The second group presents with symptoms of biliary colic (54%), dyspepsia (34%) and/or jaundice (27%), and the third presents in childhood with other associated severe fetal anomalies\(^2,4\).

The exact prevalence of each of the three groups is variable based on published reports. It is thought that approximately 70% of cases are usually isolated anomalies, although some cases appear to be familial and are associated with more severe anomalies\(^5,8,9\). In an interesting series of 34 cases (29 children and 5 adults) of congenital gallbladder agenesis, the most common anomalies associated were involving the genitourinary tract followed by gastrointestinal and cardiovascular malformations. Family history was negative in all, suggesting a sporadic occurrence\(^10\).

Historically, all cases were identified intraoperatively. In a review of 9 cases by Cho et al.\(^11\), all patients underwent a laparotomy, which failed to identify the gallbladder. However, now with the increased frequency of advanced imaging, cases are being diagnosed more often and, more importantly, before any surgical intervention.

However, given that patients with gallbladder agenesis tend to present symptoms suggestive of biliary colic, a number of them are still diagnosed intraoperatively. Due to a lack of awareness of the diagnosis, this entity remains a diagnostic challenge\(^2,12\). In those cases which are diagnosed intraoperatively, patients often are exposed to complications from prolonged exploration\(^13\), and it is suggested to abort the procedure rather than complete further exploration if a gallbladder is not found on laparoscopy since open exploration for possible ectopic gallbladder increases the risk of complications\(^14\). Intraoperative ultrasound can demonstrate an ectopic gallbladder but is not always available\(^13\). A follow-up with more advanced imaging techniques should be the next option to truly identify gallbladder agenesis as the sole abnormality to guide management further.

It is therefore important to consider the presence of this unusual entity when the nonvisualization of the gallbladder is suggested on ultrasound\(^15\). However, as is known, ultrasound is highly dependent not only on the operator but also on other factors such as body habitus or presence of bowel gas obscuring visualization. Cases of gallbladder agenesis have been reported as ‘contracted/fibrotic gallbladder’ on ultrasound\(^14\).

HIDA scans, which are also usually performed in patients with cholecystitis, in this case are unhelpful since nonvisualization of the gallbladder remains typical of cystic duct obstruction, as well as of agenesis\(^13\). MRCP is considered the test of choice if there is suspicion. It is also helpful in demonstrating an ectopic gallbladder along with other possible anomalies of the biliary tract system\(^8\). In terms of treatment, there are no specific guidelines on how to manage these cases. Interestingly, one author notes that 98% of patients had resolution of symptoms after exploratory, nontherapeutic surgery\(^5\). It is unclear how these patients would have had symptom resolution in the absence of exploration.

Conclusions:
Gallbladder agenesis presents as a significant diagnostic challenge. With the advances in imaging, more cases of gallbladder agenesis are being diagnosed incidentally and outside of the operating room. Clinicians should have a strong index of suspicion if nonvisualization is suggested by an ultrasound. A positive HIDA scan can be seen in the presence of gallbladder agenesis in the absence of cholecystitis. MRCP is considered the investigation of choice if there is suspicion. It is also helpful in demonstrating an ectopic gallbladder along with other possible anomalies of the biliary tract system. Management is usually conservative with smooth muscle relaxants.
References:


