Evaluation of hepatic cystic lesions

Marten A Lantinga, Tom JG Gevers, Joost PH Drenth

Abstract

Hepatic cysts are increasingly found as a mere coincidence on abdominal imaging techniques, such as ultrasonography (USG), computed tomography (CT) and magnetic resonance imaging (MRI). These cysts often present a diagnostic challenge. Therefore, we performed a review of the recent literature and developed an evidence-based diagnostic algorithm to guide clinicians in characterising these lesions. Simple cysts are the most common cystic liver disease, and diagnosis is based on typical USG characteristics. Serodiagnostic tests and microbubble contrast-enhanced ultrasound (CEUS) are invaluable in differentiating complicated cysts, echinococcosis and cystadenoma/cystadenocarcinoma when ultrasonography (USG), computed tomography and magnetic resonance imaging show ambiguous findings. As a result, serodiagnostic tests and CEUS reduce the need for invasive procedures. USG screening of the liver and both kidneys combined with extensive family history taking remains the cornerstone of diagnostic decision making in PLD. In conclusion, an amalgamation of these recent advances results in a diagnostic algorithm that facilitates evidence-based clinical decision making.

Key words: Coincidental hepatic cystic lesions; Cystic liver disease; Complicated cyst; Polycystic liver disease; Diagnostic algorithm

INTRODUCTION

Hepatic cystic lesions represent a comprehensive heterogeneous cluster with regard to pathogenesis, clinical presentation, diagnostic findings and therapeutic management (Table 1). Hepatic cystic lesions predominantly remain asymptomatic and are found as a mere coinci-
dence on abdominal imaging techniques, such as ultrasoundography (USG), computed tomography (CT) and magnetic resonance imaging (MRI)[1,2]. The use of these techniques has greatly increased over the last years, and as a corollary, there has been an increase in incidental findings of asymptomatic hepatic cystic lesions[3]. In most cases, hepatic cystic lesions will follow a benign course[4]. However, it is essential to differentiate benign cysts from potentially harmful cysts, such as echinococcosis, cystadenoma and cystadenocarcinoma, which require specific treatment[5,6,7]. Currently, clinicians must also be aware of changes in the epidemiology of certain hepatic cystic lesions. Echinococcosis has spread to previously non-endemic Western European countries[8]. For this reason, the early and accurate diagnosis of cysts is crucial. To facilitate the diagnostic process, we provide an overview of the wide spectrum of mono- and polycystic liver diseases based on literature published over the last five years.

**LITERATURE SEARCH**

We searched the electronic database PubMed using the following search terms: “liver” and “cyst” and “diagnosis”. We limited our search to articles that were written in English, published between November 2007 and November 2012 and available in full text. A total of 992 articles were identified. For the purpose of this review, we included articles with a main focus on the evaluation of hepatic cystic lesions in humans. Screening the titles and abstracts identified 252 articles meeting these inclusion criteria (Figure 1). Additionally, we searched the reference lists from all eligible reviews for additional leads.

**SIMPLE CYSTS**

**Pathogenesis**

Simple cysts arise congenitally from aberrant bile duct cells and contain a clear, bile-like fluid[9]. Because bile duct epithelium covers the simple cyst inner lining, it is hypothesised that simple cysts arise during embryogenesis when intrahepatic ductules fail to connect with extrahepatic ducts[4,10,11].

**Clinical features**

The prevalence of simple cysts ranges from 2.5% to 18% and increases with age[11,12]. More than half of individuals older than 60 years are likely to have one or more simple cysts. Cysts are small in most patients but can grow to over 30 cm in selected cases. In a small fraction of patients, symptoms, such as abdominal pain, early satiety, nausea and vomiting, arise as a result of a mass effect[12]. Physical examination may reveal a palpable abdominal mass or hepatomegaly[11]. Complications such as haemorrhage, rupture and biliary obstruction are uncommon but are more likely in larger cysts[12,13]. Intracystic haemorrhage is a rare complication of simple cysts and usually presents with severe abdominal pain[14,15], although asymptomatic presentations are also observed[15,16].

**Laboratory findings**

Laboratory findings are predominantly normal, but a minority of patients have raised serum γ-glutamyl-transferase (γGT)[17]. Several studies have shown that serum and cyst fluid levels of carcinoembryonic antigen (CEA) and cancer antigen 19-9 (CA 19-9) may be elevated[18]. CA 19-9 is expressed in the simple cyst inner epithelial lining and leads to elevated cyst fluid and serum CA 19-9 levels[17]. CA 19-9 is not helpful in the differential diagnosis of intracystic haemorrhage[19].

**Diagnostic features**

Most simple cysts are diagnosed incidentally on USG (Figure 2A), CT (Figure 2B) or MRI. The diagnosis of a simple cyst is based on the following USG criteria: anechoic (i.e., fluid filled cavity), no septations, sharp smooth borders, strong posterior wall echoes (indicating a well-defined fluid/tissue interface), spherical or oval shaped and a relative accentuation of echoes beyond the cyst compared to echoes at a similar depth transmitted through normal adjacent hepatic tissue (Table 2)[20,21]. CT shows a sharply defined homogeneous hypodense lesion (Figure 2B)[21]. MRI T1-weighted sequence shows low signal intensity, whereas the T2-weighted sequence shows extremely high signal intensity, which does not enhance after contrast injection[22]. USG has a reported sensitivity and specificity of approximately 90% for diagnosing a simple cyst[23], and recent advances in CT and MRI technology might result in even higher sensitivity rates[22,24]. However, because CT is accompanied with a radiation load and both CT and MRI come at a significantly higher cost, USG remains the most accurate, non-invasive and cost-effective imaging modality for diagnosing simple cysts.

In case of an intracystic haemorrhage (i.e., complicated cyst), USG typically shows a hyperechoic echo pattern combined with internal echoes that mimic septations or solid portions (Figure 3)[25]. In contrast, CT visualises intracystic haemorrhage as a high-density area[26], whereas MRI depicts it as a high signal intensity on T1- and T2-weighted sequences[23,27]. Neither CT nor MRI has additional diagnostic value compared to USG in the diagnosis of cystic bleeding[28]. The recent development of microbubble contrast-enhanced ultrasound (CEUS) enables us to visualise vascular flow within septa or solid components of cysts, which is absent in simple cysts with intracystic

<table>
<thead>
<tr>
<th>Table 1 Differential diagnosis of cystic lesions in the liver</th>
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<tbody>
<tr>
<td>Monocytic disease</td>
</tr>
<tr>
<td>Simple cyst</td>
</tr>
<tr>
<td>Echinococcosis</td>
</tr>
<tr>
<td>Cystic echinococcosis</td>
</tr>
<tr>
<td>Alveolar echinococcosis</td>
</tr>
<tr>
<td>Cystadenoma</td>
</tr>
<tr>
<td>Cystadenocarcinoma</td>
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<tr>
<td>Polycystic disease</td>
</tr>
<tr>
<td>Autosomal dominant polycystic kidney disease</td>
</tr>
<tr>
<td>Autosomal dominant polycystic liver disease</td>
</tr>
</tbody>
</table>

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haemorrhage. Therefore, CEUS can accurately characterise these cysts when USG, CT and MRI show ambiguous findings.

**Therapy**

The management of most simple cysts relies on a “wait-and-see” policy, and no further treatment is required in these cases. If there are symptoms, aspiration-sclerotherapy is the preferred treatment. Laparoscopic or open surgical fenestration techniques are similarly or even more effective in reducing symptoms but have a significantly higher morbidity and mortality rate.

**ECHINOCOCCOSIS**

Echinococcosis is a zoonosis caused by larval stages of taeniid cestodes (tapeworms) belonging to the *Echinococcus* species. Two of the six known species cause solitary cystic lesions in humans: (1) *Echinococcus granulosus* (*E. granulosus*), responsible for cystic echinococcosis (CE); and (2) *Echinococcus multilocularis* (*E. multilocularis*), responsible for alveolar echinococcosis (AE).

Echinococcosis-related deaths are uncommon in developed countries. For example, there were 41 echinococcosis-associated deaths in the United States over an 18-year study period. However, echinococcosis is considered to be an emerging disease in Europe. Thus, CE and AE are diseases with a considerable global disease impact, as indicated by a substantial loss in disability-adjusted life years.

**Cystic echinococcosis**

**Pathogenesis:** Humans become infected by acting as intermediate hosts of *E. granulosus* after ingestion of *Echinococcus* eggs, which are excreted by infected carnivores (dogs and other canids). Infection is typically observed in areas containing large numbers of the intermediate hosts of the parasite (sheep and goats) that are in close contact with the final host (herding dogs).

**Clinical features:** Although CE has a worldwide geographic distribution, the highest prevalence of CE is observed in the Western United States.
found in the temperate zones, including the Mediterranean, Central Asia, Australia and some parts of America[44].

Because cyst growth in the liver is slow (ranging from 1-5 millimetres in diameter per year), CE can remain asymptomatic for a long time. In approximately 90% of cases, the primary presentation is a spherical, fluid-filled vesicle with an inner cellular layer and an outer laminated layer located in the liver, lungs or both[40]. Symptoms occur when cysts exert mass effects within the organ or surrounding tissues or rupture, often presenting as a sudden onset of abdominal pain. Secondary cholangitis (rupture into the biliary tree), biliary obstruction and intraperitoneal rupture followed by anaphylaxis are common complications of CE and require hospitalisation[41]. Worldwide mortality rate estimates vary between 2.2%-5.0%[43,46], although the exact mortality rate of CE in developed countries remains unknown.

**Diagnostic features:** The diagnosis of CE is based on the following criteria: endemic region history, clinical findings (e.g., abdominal pain, fever, chest pain, and dyspnea), pathognomonic USG features and positive immuno diagnostic tests[47]. USG shows a round or oval-shaped, anechoic or atypical (i.e., snowflake-like inclusions or floating laminated membranes) echo pattern with multiple septa confined by a laminated border (Table 2)[43]. USG has a reported specificity of 90% and is used in combination with CT when surgical treatment is considered. MRI has not been proven to be cost-effective and has no added value[48]. The currently used serodiagnostic tests to reveal *E. granulosus* antibodies have a sensitivity of 93.5% and specificity of 89.7%[49].

**Therapy:** The treatment of CE, including surgery (open or laparoscopic), percutaneous treatments [e.g., puncture aspiration injection re-aspiration (PAIR) method] and chemotherapy[50], is indicated to reduce symptoms and prevent complications[51]. PAIR is the treatment of choice for CE, as a recent review showed that PAIR resulted in parasitological clearance (i.e., negative serodiagnostic tests) in 95.8% of cases[52].

### Table 2 Ultrasonography features for the diagnosis of monocytic diseases of the liver

<table>
<thead>
<tr>
<th>Feature</th>
<th>Simple cyst</th>
<th>Cystic echinococcosis</th>
<th>Alveolar echinococcosis</th>
<th>Cystadenoma and cystadenocarcinoma</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Shape</strong></td>
<td>Spherical or oval</td>
<td>Anechoic</td>
<td>Round or oval</td>
<td>Irregular</td>
</tr>
<tr>
<td><strong>Appearance</strong></td>
<td>No septa</td>
<td>Multiseptated</td>
<td>Hyperechogenic outer ring and hyperechogenic centre</td>
<td>Irregular</td>
</tr>
<tr>
<td><strong>Wall</strong></td>
<td>Strong posterior wall echoes</td>
<td>Relative acentration of echoes</td>
<td>Dorsal shadowing (calculated areas)</td>
<td>Round or oval</td>
</tr>
<tr>
<td><strong>Posterior acoustic feature</strong></td>
<td></td>
<td></td>
<td></td>
<td>Hypoechogenic with hyperechogenic septations</td>
</tr>
</tbody>
</table>

1Fluid-filled cavity; 2Snowflake-like inclusions or floating laminated membranes; 3Compared to echoes at a similar depth transmitted through normal adjacent hepatic tissue.
due to an infiltrative neoplastic growth with potential metastasis to adjacent and distant (e.g., lungs, spleen, bone, and brain) organs[60,64].

**Diagnostic features:** Typical USG aspects are observed in 70% of cases and include irregular shape and border, hyperechoicogenic outer ring and hypoechoicogenic centre, multivesicular appearance and dorsal shadowing due to calcified areas (Table 2)[47]. Atypical USG aspects include small hyperechoicogenic nodules (amorphous AE), large lesions with massive necrosis (pseudocyst) and small calcified lesions (inert AE)[65]. In contrast to CE, MRI is superior to CT in detecting AE lesion margins[68,69]. Similar to CE, high diagnostic sensitivity (90%-100%) and specificity (95%-100%) are attained with serodiagnostic tests, and in 80%-95% of cases, AE can be differentiated from CE with the help of serologically obtained purified Echinococcus antigens[64].

**Therapy:** The approach to the management of AE resembles that of a hepatic malignancy. The cornerstone of treatment for AE includes radical surgery followed by a 2-year period of chemotherapy[66]. A recent study concluded that AE can be cured in 42% of cases by complete surgical removal of the parasitic mass. Early diagnosis could even improve this rate further[61].

**CYSTADENOMA AND CYSTADENOCARCINOMA**

**Pathogenesis**

Cystadenoma and cystadenocarcinoma are biliary cyst tumours that originate from the biliary epithelium[64]. Analogous to simple cysts, cystadenoma is considered to be a congenital disorder[61]. The exact mechanism of carcinogenesis in cystadenoma remains unknown. Several studies have suggested that cystadenocarcinoma develops from the ectopic remnants of primitive foregut sequestered within the liver[64]. In contrast, the malignant transformation of cystadenoma into cystadenocarcinoma is considered to be an alternative mechanism of carcinogenesis, as some cystadenocarcinomas may co-exist with cystadenoma[64]. This hypothesis is supported by the observation that the presence of cystadenoma increases the chance of developing cystadenocarcinoma[68].

**Clinical features**

Less than 5% of all cystic lesions of the liver are cystic neoplasms[6]. The clinical presentation of cystadenoma and cystadenocarcinoma is asymptomatic or tends to mimic symptoms of simple cysts or echinococcosis[60,67]. Studies have reported a predominance in women, with a mean age of onset varying from 40-60 years[64,65]. Cystadenomas appear to be slow growing, but exact growth rates are unknown. One case series evaluated 75 patients and recorded a variability in cyst size from 1.5-35 cm[64]. One study involving 63 cases diagnosed with cystadenocarcinoma demonstrated infiltrative growth in neighbour-
not performed. In contrast, CEUS can be helpful in differentiating cystadenoma and cystadenocarcinoma from complicated cysts when USG, CT or MRI is inconclusive. CEUS characterises the vascular flow within septa in cystadenoma and cystadenocarcinoma, which is absent in complicated cysts\(^{[29-31]}\). Nonetheless, surgical resection remains the golden standard for diagnosing cystadenoma and cystadenocarcinoma when CEUS is not available.

**Therapy**
The primary treatment of cystadenoma and cystadenocarcinoma is hepatic resection. A study in which 66 cases of cystadenocarcinoma were subjected to hepatic resection described a 3-year survival rate of 74%\(^{[9]}\).

### PCLD AND ADPKD

#### Polycystic liver disease
Polycystic liver disease (PLD) is arbitrarily defined as the presence of >20 liver cysts\(^{[7]}\). Autosomal dominant polycystic liver disease (PCLD) and autosomal dominant polycystic kidney disease (ADPKD) are two distinct genetic disorders associated with the development of polycystic livers\(^{[78]}\). Liver function, as judged by parameters of liver synthesis, is not affected in PLD, as functional hepatic tissue remains unaffected\(^{[77,79]}\).

#### Pathogenesis
During embryogenesis, the intrahepatic bile ducts are formed from a cylindrical layer of cells (i.e., ductal plate) surrounding each portal vein. Incorrect involution of the ductal plate results in ductal plate malformation (DPM)\(^{[80,81]}\). DPM consists of excess embryonic bile duct structures in a ductal plate configuration that does not communicate with the normally developed intrahepatic bile ducts. The progressive dilatation of these excess intrahepatic structures during life results in multiple liver cysts\(^{[82]}\). Similar to simple cysts, these cysts contain a clear, bile-like fluid and an inner lining of cholangiocytes\(^{[83]}\).

#### Genetics
PCLD was historically considered a phenotypic variant of ADPKD\(^{[84]}\). However, the presence of PLD in the absence of renal cysts led to the belief that PCLD should be regarded as a separate entity\(^{[85]}\). The discovery of a familial form of PLD\(^{[85]}\), genetically distinct from the heterozygous mutation in genes PKD1 and PKD2 identified in ADPKD\(^{[87]}\), ultimately led to the identification of heterozygous mutations in genes encoding SEC63 and PRKCSH\(^{[88-90]}\). Mutation analysis identified a heterozygous mutation in PRKCSH (15%) and SEC63 (5%) in approximately 20% of studied PCLD cases\(^{[91]}\). In contrast, a PKD1 mutation was found in 85% of cases of ADPKD, and a PKD2 mutation was found in the remaining cases\(^{[92]}\).

PRKCSH and SEC63 encode hepatocystin and SEC63 proteins, respectively. Hepatocystin acts in the folding process of proteins, while SEC63 acts as part of the endoplasmic reticulum translocon\(^{[93]}\). Unfortunately, the exact mechanism of cystogenesis in PCLD remains unclear. Polycystin 1 and 2, encoded by PKD1 and PKD2, respectively, are important for adequate functioning of the primary cilium\(^{[94]}\). Its therefore suggested that primary cilia play a central pathogenic role in the mechanism of hepatic cystogenesis in ADPKD\(^{[95]}\).

#### Clinical features
The extra polarisation of 137 identified PCLD cases in a specific adherence region (the Netherlands) led to an estimated PCLD prevalence of 1 per 158000\(^{[97]}\). This number is most likely an underestimation of the true prevalence because only symptomatic patients referred to tertiary centres were included in this study, and PCLD often remains asymptomatic\(^{[98]}\). ADPKD is the most common monogenetic disorder, with a worldwide estimated prevalence of 0.10%-0.25%\(^{[99]}\), and it is responsible for approximately 8%-10% of cases with end-stage renal disease\(^{[100]}\). Although ADPKD is primarily characterised by the presence of renal cysts\(^{[101]}\), liver cysts are considered the most prevalent extra-renal manifestation of ADPKD\(^{[99,100]}\). Indeed, one study involving 230 ADPKD cases found an overall prevalence of 83%\(^{[102]}\). However, the exact prevalence of PLD in ADPKD is still unknown. PCLD is predominantly confined to the liver, but a few renal cysts can also be present, which leads to difficulties in the accurate differentiation between PCLD and ADPKD\(^{[103]}\). Although renal cysts in ADPKD ultimately lead to renal failure, renal function remains unaffected in the presence of PCLD-associated renal cysts\(^{[104]}\).

PLD is predominantly discovered during the fourth or fifth decade of life and is more severe in females\(^{[77,96,103,104]}\). PCLD tends to lead to a higher number and greater volume of liver cysts\(^{[79]}\). The number of pregnancies, increased age and severity of renal disease are considered additional risk factors for liver cyst growth in ADPKD\(^{[105]}\). PLD is mainly asymptomatic, but mechanical complaints can arise in a subset of patients\(^{[79,106]}\). Complications such as intracystic haemorrhage and infection are rare and typically occur in large cysts\(^{[106]}\).

#### Laboratory findings
PLD causes increased yGT and AP levels in both PCLD and ADPKD patients\(^{[77]}. Occasionally, increased serum aspartate aminotransferase (AST) is also found in ADPKD\(^{[107]}\). Renal function remains intact in PCLD, whereas ADPKD patients show a rise in serum creatinine due to impaired renal function\(^{[108]}\).

#### Diagnostic features
PLD is detected with the use of USG, CT or MRI. USG, which is accurate, non-invasive and low cost, is the preferred imaging modality for both PCLD and ADPKD\(^{[108,109]}\). Currently, there are no generally accepted USG criteria for PCLD. One study suggested that the diagnosis can be made in case of a positive family history of PCLD and the presence of >4 liver cysts\(^{[78]}\). However,
diagnosing ADPKD is usually relatively straightforward when enlarged bilateral cystic kidney lesions are present in combination with a positive family history for ADPKD[109]. In case of a negative family history, screening direct family members with USG can be helpful to reveal asymptomatic ADPKD. Because mutation analysis for ADPKD has no clinical implications, its use is limited to family members of ADPKD patients involved in kidney donation programs. In 2009, the Pei USG criteria were developed because the original Ravine USG criteria for diagnosing ADPKD appeared to be insufficient[10,11]. Table 3 gives an overview of the USG criteria for diagnosing ADPKD when the causative gene is unknown. For example, in case of a positive ADPKD family history, diagnosis can be made when ≥ 3 renal cysts are unilaterally present in individuals aged 15 to 39 years[116]. ADPKD should be considered when there are ≥ 10 bilateral renal cysts present in the absence of other renal or extra-renal disease that can cause renal cysts[106]. When PCLD or ADPKD criteria are not met, multiple simple cysts are most likely responsible for the hepatic cystic lesions.

ADPKD is characterised by an increased risk of developing vascular manifestations. Hypertension occurs in approximately 50%-70% of patients, and almost half of these hypertensive patients are reported to have left ventricular hypertrophy (LVH)[112]. Mitral valve prolapse is observed in 25% of patients and intracranial aneurysms in 4%-12% of patients[112]. As a result, magnetic resonance angiography (MRA) must be performed when ADPKD patients have a positive family history of intracranial aneurysms because the rupture of aneurysms is reported to be responsible for 4%-7% of deaths in affected ADPKD families[113]. In contrast to ADPKD, several studies have shown that PCLD patients do not appear to have an increased risk of vascular malformations. One study involving 19 PCLD cases reported hypertension in 10.5% of cases, mitral valve prolapse in 0% and aneurysms in 5.3%[79]. Another study involving 58 PCLD cases found mitral valve prolapse in 1 case (2.6%)[114]. Subsequently, targeted screening is not advised for PCLD.

Therapy

The main objective of therapy is to reduce liver cyst volume to diminish mass effect-related symptoms[115]. Hence, the only indication for reducing cyst volume is when a PLD patient reports symptoms that can be linked to the polycystic liver[118].

Surgical procedures, such as aspiration-sclerotherapy and fenestration, are indicated when PLD consists of large cysts confined to a limited part of the liver. In more extensive disease, segmental hepatic resection or even liver transplantation is imperative to relieve symptoms[117]. Future medical therapies include somatostatin analogues, as several clinical trials with lanreotide and octreotide achieved polycystic liver volume reduction in PCLD and ADPKD[118-123].

CONCLUSION

Cystic lesions of the liver encompass a wide spectrum of disorders. As a result of the frequent use of abdominal imaging techniques in recent years, the incidence of so-called coincidental cysts has increased. Simple cysts are the most prevalent and have a tendency to follow a benign course. However, complicated cysts, echinococcosis and cystic neoplasms (e.g., cystadenoma and cystadenocarcinoma), which cause a diagnostic enigma, demand accurate diagnosis in the early stage because specific treatment could be required. Furthermore, the presence of multiple hepatic cystic lesions must raise the suspicion of PCLD or ADPKD and requires further screening.

USG remains the most accurate, non-invasive and cost-effective imaging modality for diagnosing simple cysts. Despite recent advances (e.g., contrast-enhanced CT and DWI), distinguishing complicated cysts from echinococcosis and cystic neoplasms remains impossible with USG, CT or MRI alone. Because of an ever-increasing spread of Echinococcus to previously non-endemic regions and its initial quiescent phase after primary infection, it is necessary to exclude echinococcosis. Serodiagnostic tests have high sensitivity and specificity to reveal Echinococcus antibodies. Subsequently, CEUS can be used to accurately and reliably exclude cystic neoplasms by demonstrating the absence of any enhancement within the hepatic cystic lesion. Therefore, when CEUS is available, it reduces the need for surgical resection.

The detection of multiple liver cysts requires USG screening of both kidneys and extensive family history taking regarding the occurrence of ADPKD or PCLD. When PCLD or ADPKD criteria are not met, multiple simple cysts are most likely responsible for the hepatic cystic lesions. PCLD or ADPKD could eventually be diagnosed through USG follow-up.

To summarise, we developed a diagnostic algorithm by integrating recent advances with conventional diagnostic tools (Figure 4). Our diagnostic algorithm facilitates evidence-based clinical decision making when clinicians are confronted with coincidental hepatic cystic lesions on USG. Further development of USG- and MRI-based techniques, such as CEUS and DWI, will probably lead to further improvement of hepatic cystic lesion characterisation.

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**Table 3  Ultrasonography criteria for the diagnosis of autosomal dominant polycystic kidney disease**

<table>
<thead>
<tr>
<th>Family history positive¹</th>
<th>Unknown genotype</th>
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<tbody>
<tr>
<td>Age (yr)</td>
<td></td>
</tr>
<tr>
<td>≥ 15 and ≤ 39</td>
<td>≥ 3 unilateral renal cysts</td>
</tr>
<tr>
<td>≥ 40 and ≤ 59</td>
<td>≥ 2 bilateral renal cysts</td>
</tr>
<tr>
<td>≥ 60</td>
<td>≥ 4 bilateral renal cysts</td>
</tr>
</tbody>
</table>

¹Exclude autosomal dominant polycystic kidney disease when < 2 unilateral renal cysts and ≥ 40 years of age.

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Figure 4 Diagnostic algorithm. Diagnosis of hepatic cystic lesions after detection on ultrasonography. E. granulosus: Echinococcus granulosus; E. multilocularis: Echinococcus multilocularis; CEUS: Contrast-enhanced ultrasound; PCLD: Polycystic liver disease; ADPKD: Autosomal dominant polycystic kidney disease.

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