Foremost among the worrisome lesions are malignant neoplasms, both epithelial and mesenchymal in origin. Hepatoblastoma, angiosarcomas, epithelioid hemangioendothelioma, perivascular epithelioid cell tumors (PEComas), mucinous cystic neoplasms (MCNs), intraductal papillary mucinous neoplasm (IPMN), and primary hepatic lymphoma are some of the malignant lesions described in this article. Imaging appearances of other benign lesions such as biliary hamartomas and inflammatory pseudotumors are also discussed.

**Hepatoblastoma**

Hepatoblastoma, although rare, is the most common primary malignant hepatic neoplasm in children, comprising two-thirds of primary liver tumors in this population. However, the incidence of hepatoblastoma has been increasing worldwide. Most cases occur in children less than 5 years old. Adult cases are very rare with fewer than 70 adult cases reported in the literature.

Hepatoblastoma is more common in children born prematurely. It is almost never associated with chronic liver disease. Most cases are sporadic. However, there is an increased incidence in patients with certain genetic abnormalities such as Beckwith-Wiedemann syndrome, trisomy 18, trisomy 21, Li-Fraumeni syndrome, type 1a glycogen storage disease (von Gierke disease), and familial adenomatous polyposis. Screening for hepatoblastoma is considered in such individuals. A recommended strategy is abdominal ultrasound (US) and alpha fetoprotein (AFP) levels every 3 months until 6 to 8 years of age.
Most patients are asymptomatic. Many patients present with an abdominal mass and a diagnosis suspected by imaging and subsequently confirmed by biopsy. Occasionally, nonspecific symptoms such as pain, nausea, and vomiting are seen. Large tumors may present after rupture with acute abdominal pain and intra-abdominal hemorrhage. Serum AFP is usually elevated. Very rarely, patients may also present with sexual precocity due to synthesis of ectopic β-human chorionic gonadotropin. Rarely, ectopic production of adrenocortical hormones and/or ectopic production of parathyroid hormone may lead to Cushing syndrome and hypercalcemia, respectively.

Hepatoblastomas are embryonal tumors that develop from pluripotent cells with the potential to differentiate along several lines. Hence, histologically, it is a complex tumor that may include fetal or embryonal hepatocytes and heterogeneous tissues including cartilage, muscle fibers, squamous epithelium, and bone. Consequently, the imaging features are varied and depend on the histological makeup of the tumors. The most common site of metastasis is the lung, while lymph node involvement is rare.

On plain radiographs, hepatomegaly may be recognized in patients with large tumors. Occasionally, these tumors also show specks of calcification on radiographs. US is usually the first imaging study performed in children and hence the most common modality in which the diagnosis of hepatoblastoma is first suggested. These masses are usually large solitary masses of varying echogenicity (Fig. 1). Occasionally, they are multifocal, sometimes with smaller satellite lesions. Areas of necrosis and hemorrhage are common. Rarely, hepatoblastomas may infiltrate the entire liver and present as diffusely enlarged liver with heterogeneous and inhomogeneous echotexture. Further evaluation with cross-sectional imaging such as computed tomography (CT), magnetic resonance imaging (MRI), and histological confirmation with liver biopsy is necessary.

Calcifications may be seen on CT scans, particularly in tumors which have mixed mesenchymal elements. Hepatoblastomas show heterogeneous pattern of enhancement, generally hypo-enhancing compared with normal liver. Tumor thrombus in portal and hepatic veins are common.

MRI is usually the preferred modality by most pediatric radiologists due to lack of radiation. However, MRI does pose challenges, often requiring sedation and general anesthesia in the pediatric population. Depending on the histologic makeup of the tumor, the tumors are either homogeneous or heterogeneous with varying amounts of hemorrhage, fibrosis, and septae (Fig. 2). Septations appear on MRI as hypointense band both on T1- and T2-weighted images. Hepatoblastomas have imaging features similar to those of fibrolamellar carcinoma which should be considered in the differential. The latter condition usually occurs in young adults and teenagers.
Imaging plays a key role in diagnosis and staging of hepatoblastomas. The PRETEXT (PRE-Treatment EXTent of tumor) system is used for risk stratification. In the PRETEXT system, the extent of liver disease is based on the number of sectors of liver involved by the tumor. Annotation is a part of pretext staging to help guide surgical approach which includes assessment of involvement of hepatic veins, vena cava, portal veins, extrahepatic extension, caudate lobe involvement, multiplicity, tumor rupture, and distant metastases. MR angiography is commonly performed to help in planning surgical approach. For patients whose tumor is not resectable at diagnosis, repeat radiologic evaluation after initial chemotherapy is termed POSTTEXT and uses the same criteria as PRETEXT. Assessment of downstaging and timing of resection is assessed on posttreatment cases as well as suitability for liver transplantation or other form of targeted treatment.

The 5-year survival of hepatoblastoma has dramatically increased from 30% reported in the 1980s to approximately 75 to 90% reported in contemporary studies reflecting improvements in pretreatment staging, chemotherapy, surgical techniques, and introduction of liver transplantation for unresectable disease.

Angiosarcoma

Angiosarcomas account for less than 1% of all primary liver tumors. However, it is the most common primary hepatic sarcoma. It almost always portends a poor prognosis. The majority of the untreated patients die within 6 months and most patients die within a year after surgical resection. Angiosarcomas have received attention in recent decades because of its frequent association with several environmental carcinogens such as Thorotrast, vinyl chloride, arsenic compounds, and radiation. Both hemochromatosis and anabolic steroids have been associated with hepatic angiosarcoma. It usually presents in older patients more than 50 years old and is more common in men. Patients often present with abdominal pain, ascites, weight loss, fatigue, jaundice, and occasionally Budd–Chiari syndrome. Thrombocytopenia and hepatomegaly are common in these patients. Metastatic disease is also frequent and is seen in the spleen, lymph nodes, lungs, bones, and adrenal glands. Liver failure and intra-abdominal bleeding are common causes of death.

Hepatic angiosarcomas are primarily composed of vascular or lymphatic endothelial cells and are characterized by abundant blood vessels. Hemorrhage can occur spontaneously. Angiosarcomas form numerous poorly defined variable sized nodules. The entire liver is frequently involved. Better differentiated areas may show large, cavernous spaces lined by atypical endothelial cells. Tumor cells grow along and into the sinusoids and branches of the portal and hepatic veins. These histologic features determine imaging appearances seen on CT and MRI. Although biopsy is commonly utilized to diagnose angiosarcoma, a relatively frequent complication is postbiopsy hemorrhage.

The lesions are generally ill defined, multifocal, and show heterogeneous areas including areas of hemorrhage. On MRI, these lesions may be of mixed signal on precontrast T1- and T2-weighted images particularly due to hemorrhage. Larger dominant masses demonstrate heterogeneous internal architecture. Nonenhanced CT images can show areas of hemorrhage. On postcontrast studies, enhancing regions show progressive enhancement on portal and delayed phase imaging. Lack of washout helps in differentiating angiosarcoma from multifocal hepatocellular carcinoma (HCC) in patients with cirrhosis.

Epithelioid Hemangioendothelioma

Epithelioid hemangioendothelioma is a low-grade malignant vascular neoplasm affecting middle-aged patients with a slight female preponderance. Patients often present...
with nonspecific symptoms including abdominal pain, weight loss, or malaise. Despite being a low-grade tumor, there is a high rate of regional and distant metastases. Histologically, these lesions may be diagnosed as cholangiocarcinoma due to significant overlapping features. Accurate diagnosis may require staining for markers of endothelial cells such as ERG transcription factor, factor VIII-related antigen, and CD 31/34 antigens, which are usually negative in cholangiocarcinoma. Recognition of classic radiologic findings may also be useful in providing accurate diagnosis.

These lesions tend to be multifocal, predominantly peripheral in distribution sometimes coalescing into individual nodules. Occasionally these lesions calcify. Subcapsular lesions usually produce hepatic capsular retraction.

On US, the lesions are predominantly hypoechoic or with mixed echotexture.

On MRI, the lesions are usually well-demarcated, hyperintense on T2-weighted sequences. Occasionally, a two-layer halo is described with alternating bands of hypo- and hyperintense signals on T2-weighted images (target sign). The lesions are hypointense on noncontrast T1-weighted sequences. They are hypoenhancing on postcontrast studies sometimes with a ring peripheral enhancement or target pattern of enhancement. Coexistence of the “target sign” on both postcontrast and T2-weighted sequences are considered highly specific for epithelioid hemangioendothelioma. Progressive enhancement and lack of washout is common. Some lesions show minimal or no enhancement on postcontrast studies. Another sign which has been considered specific for epithelioid hemangioendothelioma is the lollipop sign, which consists of a contrast-enhanced vessel.
ending in a hypoattenuating or hypointense lesion resembling a lollipop.\textsuperscript{20} CT enhancement patterns are very similar to that seen on MRI.

**Hepatic Perivascular Epithelioid Cell Tumor**

PEComas represent a group of neoplasms that show both smooth muscle and melanocytic differentiation.\textsuperscript{21} PEComas can show a range of biological behavior ranging from benign tumors to aggressive sarcomas. They can arise in any anatomical site. For example, in the lungs, lymphangioleiomyoma/lymphangioleiomyomatosis and cell sugar tumor of the lung belong to this group of tumors. In the kidneys, angiomylipoma belong to this group of tumors.\textsuperscript{22} PEComas arising in the kidney and liver are commonly composed of epithelioid or spindle cells, mature fat, and thick-walled vessels. Hepatic PEComas are very rare.

Due to its rarity, the natural history of this condition has not been elucidated. However, it has been noted that the lesion is found mainly in females. Clinical presentation is usually with nonspecific abdominal pain. However, most are incidentally diagnosed on imaging for other indications.\textsuperscript{23–25} Tuberous sclerosis is associated with approximately 5 to 15% of hepatic PEComas.

Presence of macroscopic fat in a tumor should alert the radiologist to this condition (\textsuperscript{\textcircled{8}} Fig. 8). However, this is not very specific as some HCC may contain fat. On US, these lesions tend to be expansile masses which are hypoechoic. On CT and MRI, the lesions tend to be large masses and show arterial hyperenhancement. There may or may not be washout on the delayed phase images (\textsuperscript{\textcircled{9}} Figs. 9 and 10). Although most of these tumors are benign, a few of these lesions are malignant and can metastasize (\textsuperscript{\textcircled{10}} Fig. 10).\textsuperscript{23} The imaging features can overlap with those of other more common conditions such as focal nodular hyperplasia as shown in the example in \textsuperscript{\textcircled{9}}, a HCC as shown in the example in \textsuperscript{\textcircled{10}}, or a hemangioma. Biopsy is generally required for diagnosis. Radiologists should be alert to any atypical findings such as normal background liver, macroscopic fat, and enhancement and washout characteristics which do not conform to well-known classic features.

**Hepatic Mucinous Cystic Neoplasms**

Hepatic MCNs were previously referred to as biliary cystadenomas. These are rare cystic tumors that occur within the liver parenchyma, or less frequently, in the extrahepatic bile ducts. The tumors grow to a large size and require surgical intervention in most cases. MCNs occur in adults and more often in women. These lesions are characterized by presence of ovarian stroma. They are rare, accounting for less than 5% of cystic hepatic masses.\textsuperscript{26,27}

While most MCNs arise from the intrahepatic biliary system, up to 10% of these lesions are extrahepatic in location.\textsuperscript{27,28} On ultrasonography, an MCN is seen as a cystic lesion with thickened and irregular walls with wall nodularity. Internal debris due to hemorrhage or secretions is common.\textsuperscript{29} On a CT and MRI scans, these lesions are complex unilocular or multilocular cystic mass with septations. There may be variable amounts of nodular and solid enhancing intracystic components. These lesions may also cause biliary dilatation by causing extrinsic compression.\textsuperscript{27} MRI characterizes cyst fluid content by varying signal intensities on T1-weighted images depending on cyst fluid protein content.\textsuperscript{30} The cyst wall is usually thickened and nodular. Presence of low-grade metabolic activity on positron emission tomography (PET-CT) may be seen in MCN and does not necessarily suggest malignant degeneration (\textsuperscript{\textcircled{11}} Fig. 11).

The preferred treatment for noninvasive MCNs is resection since malignant transformation is seen in up to 15% of patients.\textsuperscript{29,31} Formal surgical resection with negative margins is recommended. Extrahepatic MCNs (\textsuperscript{\textcircled{12}} Fig. 12) are resected along with the associated bile duct followed by biliary diversion.\textsuperscript{28}

**Hepatic Mucinous Cystic Neoplasms with Associated Invasive Carcinoma (Cystadenocarcinoma)**

MCNs with associated invasive carcinoma were previously referred to as cystadenocarcinomas. These are usually found in older adults. Although these lesions can metastasize, their prognosis is better than that associated with cholangiocarcinoma.

MCNs with associated invasive carcinoma generally have a thick wall that may show large enhancing soft tissue masses protruding from the internal cyst lining. Advanced tumors may show a predominantly solid and enhancing component. PET-CT may show intense increase in metabolic activity (\textsuperscript{\textcircled{13}} Fig. 13). Needle biopsy is not typically performed due to its low diagnostic accuracy and risk of tumor
**Fig. 8** Magnetic resonance imaging (MRI) images in a 63-year-old female patient with multifocal hemangioendothelioma. (A) T2-weighted images show well-demarcated lesions with alternating bands of hypo- and hyperintense signals compatible with target sign (arrows). (B–E) Axial T1-weighted sequences show hypointense lesion in the precontrast phase (B) with progressive enhancement into the delayed phase. Please note the appearance of a target sign in the portal venous phase study (D).

**Fig. 9** A 53-year-old male with hepatic perivascular epithelioid cell tumor (PEComa). (A) Noncontrast study shows a lesion in the right liver with macroscopic fat (asterisk). (B) Postcontrast arterial phase image shows an adjacent component with hyperenhancement. (C) Postcontrast delayed phase image shows washout of the enhancing component (arrow).
When imaging findings are nondiagnostic, and invasive carcinoma is suspected, surgical resection is required to confirm the diagnosis.

### Intraductal Papillary Neoplasm of the Bile Ducts

Intraductal papillary neoplasm of the bile duct (IPNB) is a recently recognized entity characterized by intraductal mucin-producing tumor. This is considered a biliary equivalent of IPMN in the pancreas and is shown to demonstrate stepwise progression to cholangiocarcinoma. Many different terms have been used to describe IPNB including biliary papilloma, mucin-producing bile duct tumor, and biliary papilla.

![Fig. 10](image1.png) **Fig. 10** A 49-year-old female with hepatic perivascular epithelioid cell tumor (PEComa). Axial precontrast (A) and postcontrast (B–D) images of the liver show an expansile well-demarcated mass in the left liver. The lesion shows arterial hyperenhancement (B) and becomes iso-intense to the liver in the delayed phase (D).

![Fig. 11](image2.png) **Fig. 11** A 59-year-old female with malignant hepatic perivascular epithelioid cell tumor (PEComa). Axial precontrast (A) and postcontrast (B–D) images of the liver show a heterogeneous well-demarcated mass in the right liver. The lesion shows areas of necrosis, pseudocapsule, and washout in the delayed phase images (D).

![Fig. 12](image3.png) **Fig. 12** A 54-year-old female with mucinous cystic neoplasm. (A) Axial computed tomography (CT) scan without contrast and (B) contrast-enhanced image showing thin septations and a solid mass. (C) Fused fluorodeoxyglucose (FDG) positron emission tomography (PET)-CT shows low-grade FDG avidity. The mass was resected and was benign.
tract IPMN. These tumors are distinguished from MCN by lack of ovarian stroma and communication with bile ducts. Most patients with IPNB are asymptomatic. When they do present, they present with obstructive symptoms such as jaundice and recurrent cholangitis. IPNB is more common in East Asia and shows a slight male preponderance.35,36

On imaging, these patients demonstrate dilation of both upstream and downstream biliary ducts caused by intraductal obstructive growth and copious mucin production. This finding is thought to be specific for IPNB and is not seen in other conditions (Fig. 14). Some lesions are predominantly cystic and are thought to arise from peribiliary glands. These lesions may resemble MCN. Communication with biliary tree is a key finding in establishing the diagnosis in such cases. Although IPNB is rare, a radiologist familiar with its manifestation can help with early diagnosis when surgical resection can be curative. MRI with MR cholangiopancreatography is the most sensitive imaging modality for detection of the lesion, followed by CT and US.37 Complete surgical resection with negative margins (R0 resection) is the standard treatment goal given the high potential for malignancy and for recurrent cholangitis and obstructive jaundice in nonmalignant cases.38 Tumors involving the extrahepatic bile ducts are treated with Whipple’s surgery, whereas tumors confined to intrahepatic bile ducts undergo hepatic resection depending on the degree of biliary involvement highlighting the importance of accurate radiological staging.11

**Primary Hepatic Lymphoma**

Hepatic lymphoma can either be primary or secondary. To be classified as primary hepatic lymphoma, the disease should

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**Fig. 14** A 65-year-old female with mucinous cystic neoplasm with associated invasive carcinoma. Axial T2-weighted (A) image shows a high signal cystic mass with surrounding intermediate T2 hyperintensity mass which enhances after injection of intravenous (IV) contrast (C). (D) Fused axial positron emission tomography (PET)/computed tomography (CT) on same patient shows the tumor is intensely metabolically avid. A second focus in segment 7 (arrow) represented metastatic disease.
be limited to the liver and hepatic hilar lymph nodes, with no distant involvement. Primary hepatic lymphoma is rare and account for less than 1% of all cases of non-Hodgkin lymphoma. The most common subtype of primary hepatic lymphoma is diffuse large B cell lymphoma.

Imaging appearances of hepatic lymphoma is varied. They most commonly present as a large solitary hypoenhancing masses on CT and MRI (Fig. 15). Up to 33% are multifocal and can present as multiple small discrete nodules (Fig. 16). They can be isointense or hyperintense compared with the liver on T2 imaging. Other patterns include diffuse infiltration (with or without hepatomegaly) and a mass in the porta hepatis. They are usually intensely metabolically avid on fluorodeoxyglucose PET-CT.

**Inflammatory Pseudotumor**

Inflammatory pseudotumor refers to a heterogeneous group of mass-forming lesions that can involve many organs and that are characterized by a prominent inflammatory infiltrate. In the literature, these have been described as synoni-

mous with inflammatory myofibroblastic tumor and xanthogranulomatous tumors or plasma cell granulomas of the liver. These present most commonly in young adults with males affected more than females. The patients usually present with systemic symptoms such as vague upper abdominal discomfort, loss of weight, malaise, and intermittent fever. Liver function tests are usually not useful.

On CT and MRI, these lesions have been described as having a targetoid appearance in a significant proportion of patients studied and suggest this to be a characteristic finding. These tumors have a wide range of imaging appearances with no specific feature that can clinch the diagnosis on radiology alone. Generally, inflammatory pseudotumors have nonspecific imaging findings. These tend to have ill-defined edges, are hypoenhancing, and can resemble common neoplastic conditions such as metastases or cholangiocarcinoma (Fig. 17). In view of the lack of specific imaging findings, percutaneous biopsy is key to establish diagnosis. A prudent clinician would consider this diagnosis in patients presenting with significant systemic symptoms and hepatic tenderness.
Conflict of Interest
None declared.

References
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Fig. 18  A 59-year-old female with primary hepatic lymphoma. (A) Ultrasound scan image shows numerous ill-defined hyperechoic lesions scattered throughout the liver (white arrows). (B) Noncontrast computed tomography (CT) in narrow windows show numerous hypoattenuating lesions (white arrows). (C) Contrast-enhanced CT scan demonstrates the lesions enhance to a lesser degree than the liver (black arrows) leading to a heterogeneous appearance.

Fig. 19  (A) Coronal T2-weighted magnetic resonance imaging (MRI) and (B) axial T2-weighted MRI in a 62-year-old patient showing numerous cystic lesions measuring 2–15 mm scattered throughout the liver (white arrows). These represent Von Meyenburg complexes or multiple biliary hamartomas.

Fig. 20  Axial contrast-enhanced computed tomography (CT) scan of a 44-year-old male who presented with abdominal pain and general malaise. Hypoattenuating defined lesions (arrows) with ill-defined edges. Biopsy showed no evidence of malignancy but numerous inflammatory cells. A diagnosis of inflammatory pseudotumor was suggested. The patient improved on a course of corticosteroid therapy.


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