Guidelines and Good Clinical Practice Recommendations for Contrast Enhanced Ultrasound (CEUS) in the Liver – Update 2020

WFUMB in Cooperation with EFSUMB, AFUMB, AIUM, and FLAUS

Aktualisierte Leitlinien und Empfehlungen für die gute klinische Praxis für CEUS der Leber

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Introduction

The present, updated document describes the fourth iteration of recommendations for the hepatic use of contrast enhanced ultrasound (CEUS), first initiated in 2004 by the European Federation of Societies for Ultrasound in Medicine and Biology (EFSUMB) [1]. The previous updated editions of the guidelines reflected changes in the available contrast agents and updated the guidelines not only for hepatic but also for non-hepatic applications.

The 2012 guideline requires updating as previously the differences of the contrast agents were not precisely described and the differences in contrast phases as well as handling were not clearly indicated. In addition, more evidence has been published for all contrast agents. This update also reflects the most recent developments in contrast agents, including the United States Food and Drug Administration (FDA) approval as well as the extensive Asian experience, to produce a truly international perspective.

These guidelines and recommendations provide general advice on the use of ultrasound contrast agents (UCA) and are intended to create standard protocols for the use and administration of UCA in liver applications on an international basis to improve the management of patients.

Bibliography

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ABSTRACT

The present, updated document describes the fourth iteration of recommendations for the hepatic use of contrast enhanced ultrasound (CEUS), first initiated in 2004 by the European Federation of Societies for Ultrasound in Medicine and Biology (EFSUMB). The previous updated editions of the guidelines reflected changes in the available contrast agents and updated the guidelines not only for hepatic but also for non-hepatic applications. The 2012 guideline requires updating as previously the differences of the contrast agents were not precisely described and the differences in contrast phases as well as handling were not clearly indicated. In addition, more evidence has been published for all contrast agents. The update also reflects the most recent developments in contrast agents, including the United States Food and Drug Administration (FDA) approval as well as the extensive Asian experience, to produce a truly international perspective. These guidelines and recommendations provide general advice on the use of ultrasound contrast agents (UCA) and are intended to create standard protocols for the use and administration of UCA in liver applications on an international basis to improve the management of patients.
World-wide commercial availability of Ultrasound Contrast agents

Availability of UCA for clinical use is based on the approval by regulatory agencies specific to the territory of intended use. Currently there are four agents that are available internationally for the use in the liver, listed here with their manufacturers.

- Definity/Luminity – Lantheus Medical Imaging, Inc., North Billerica, MA, US
- SonoVue/Lumason – Bracco Suisse SA, Geneva, Switzerland
- Optison – GE Healthcare AS, Oslo, Norway
- Sonazoid – GE Healthcare AS, Oslo, Norway

The approval of these agents varies throughout the world along with the approved indications. ICUS in collaboration with WFUMB has developed an interactive map (Fig. 1).

Indications, contraindications, safety considerations

The indications and contraindications are different among different UCAs; detailed information can be found in the official package insert of the drug.

Safety considerations

UCAs can be administered safely in various applications with minimal risk to patients [4, 7–11]. They are not excreted through the kidneys and can be safely administered to patients with renal insufficiency with no risk of contrast-related nephropathy or nephrogenic systemic fibrosis. There is no additional need for biochemical assessment or fasting prior to injection, and there is no evidence of any effect on thyroid function, as UCAs do not contain iodine [2, 3]. UCAs have a very low rate of anaphylactoid-type reactions (1/7000 patients, corresponding to 14/100,000, or 0.014 %) [7, 11–13] significantly lower than the rate with current iodinated computed tomography (CT) agents (35–95/100,000 patients, 0.035–0.095 %) [14] and gadolinium-based contrast agents at 4/64 (6.3 5) [15]. Serious anaphylactoid-type reactions to UCAs are observed in approximately 1/10,000 exposures, 0.01 % [5, 11].

SonoVue data pooled from 75 completed studies (of 6307 patients) in Europe, North America and Asia showed that the most frequent adverse events were headache (2.1 %), nausea (0.9 %), chest pain (0.8 %), and chest discomfort (0.5 %). All other adverse events occurred at a frequency of less than 0.5 % [16]. Most adverse events were mild and resolved spontaneously within a short time without sequelae. Most cases of allergy-like events and hypotension occurred within a few minutes following injection of the agent. The overall reported rate of fatalities attributed to SonoVue is low (14/2447 083 exposed patients; 0.0006 %) and compares favorably with the risk for fatal events reported for iodinated contrast agents (approximately 0.001 %). In all reported fatalities after use of an UCA, in both cardiac and non-cardiac cases, an underlying patient medical circumstance played a major role in the fatal outcome.

The intravascular administration of UCAs has been evaluated in a total of 7082 children described in 15 studies and in a European survey of 4131 children with 0.8 % reported adverse events, mostly related to bladder catheterization [17, 18]. Intravenous CEUS is also used in the pediatric population [19] and in numerous other documented areas [5]. The Food and Drug Administration (FDA) in the United States of America (USA) recently approved the use of Lumason for pediatric liver imaging [20], which is an important development. This application is, however, still off label in pediatric imaging in many countries. A significant reduction of ionizing radiation exposure is likely to be achieved in many areas by using CEUS in pediatric patients [19, 21].
Most recently it was shown that the use of SonoVue appears to be safe in pregnant women [22].

**RECOMMENDATION 1**
Intravenous use of UCAs in adult populations is safe (LoE 2) (Pro 28, Abs 0, Against 0).

**RECOMMENDATION 2**
Intravenous use of UCAs in pediatric populations is safe (LoE 3) (Pro 28, Abs 0, Against 0).

**RECOMMENDATION 3**
Intracavitary use of UCAs is safe (LoE 2) (Pro 27, Abs 1, Against 0).

Liver CEUS: Scanning technique and basic image interpretation

The study procedure is well documented in previous CEUS Guidelines [2, 3] and has been described in detail in a recent WFUMB position paper [23]. Prior to performing a liver CEUS study, it is necessary to review the patient’s clinical history, laboratory data and any prior imaging findings [2, 3].
Study procedure

Before CEUS, cysts and calcifications must be identified by conventional US, since these structures do not exhibit contrast enhancement and could therefore be erroneously interpreted as a malignant infiltration if only scanned in the late phase (LP). When cysts are missed by the baseline examination, it is necessary to carefully review both the contrast and the reference image, and to analyze the B-mode pattern of the liver tissue after the disappearance of the microbubbles.

Image interpretation

CEUS of the liver has three overlapping vascular phases after the injection of UCA, because of the dual blood supply of the liver, i.e. hepatic artery and portal vein (respectively 25%–30% and 70%–75% of liver blood flow in non-cirrhotic conditions) (Table 1).

- The arterial phase (AP) provides information on the degree and pattern of the arterial vascular supply of a focal liver lesion (FLL). Early arterial enhancement pattern and vascular architecture are best seen in slow replay of a stored cine loop.
- The portal venous phase (PVP) represents the arrival of UCA through the portal system, resulting in diffuse and maximal enhancement of normal liver parenchyma.
- The late phase (LP) lasts until the clearance of the UCA from the circulation and depends on the type and dose of UCA, total scanning time, acoustic power output and on the sensitivity of the US system.
- The post-vascular phase is only observed with Sonazoid and represents uptake of the UCA by phagocytic cells, e.g. Kupffer cells.

Slight/moderate variations of timing may occur, particularly in the case of cardiac dysfunction and in patients with vascular liver disease. Vascular architecture and phase-specific contrast enhancement of the lesion compared to the adjacent liver parenchyma are the most important diagnostic features for the characterization of FLLs [2, 3].

Differences between CEUS and other contrast-enhanced imaging modalities (CECT, CEMRI)

UCAs comprise gas-filled particles (microbubbles) and differ in fundamental respects from the agents used in contrast enhanced CT (CECT) and contrast enhanced magnetic resonance imaging (CEMRI), and for this reason play a complementary problem-solving role for indeterminate FLLs. Unlike CT and MR agents, microbubbles are not excreted by the kidneys. With the exception of Sonazoid, UCAs are purely intravascular agents. Therefore, CEUS should be considered as the first contrast imaging modality in patients with renal insufficiency. UCAs can be safely administered more than once during the same examination. While the dynamic phases of liver enhancement with UCA resemble those of CECT with iodinated agents and CEMRI with gadolinium chelates, imaging is real-time with US. Other important differences exist and are well described in the literature [24–27]. For FLL characterization, an overall improvement in sensitivity and specificity is found for CEUS over CECT [28–33]. CEUS, in addition, is reported to be invaluable in providing characterization of indeterminate FLL on CT, MR imaging and positron emission tomography (PET) [34–36]. It is also reported that CEUS should be the subsequent imaging modality for all CT- and MR-indeterminate nodules before biopsy is undertaken [37].

Detection of malignant FLL: Transabdominal approach

Conventional US is the most frequently used modality for the primary imaging of abdominal organs, including the liver, but is less sensitive in the detection of FLL than CECT, CEMRI or intraoperative US. A number of studies [38–47] have reported that CEUS has a considerably higher sensitivity of up to 80%–90% in detecting liver metastases, comparable to that of CECT [48] and CEMRI [40]. Furthermore, some reports have shown that CEUS is of particular usefulness with metastases ≤10 mm [49, 50]. CEUS has dramatically increased the capability of US for detection of FLL and has the potential to be incorporated into the diagnostic algorithm for malignant FLL.

Study procedures

The study procedure is described above. A second contrast administration (reinjection technique) can be used to confirm the metastatic nature of focal areas of contrast washout by demonstrating AP enhancement within the areas of contrast washout.
Detection of metastatic lesions
The typical and almost invariable appearance of metastases is focal contrast washout. The enhancement patterns observed during the AP has limited clinical utility in lesion detection [2, 3, 23].

With vascular phase agents (Sonovue/Lumason, Definity/Lumity, Optison), several studies have shown that the accuracy in the detection of liver metastases is comparable to that of CECT and CEMRI, when scanning conditions allow a complete imaging of all liver segments [44]. However, it should be noted that most of the studies have used initial and/or follow-up imaging (mostly CT examinations and sometimes MR imaging, and intra-operative US) as a reference standard, and very few reports include histologic or pathologic confirmation. Nonetheless, as CT and MR imaging are currently the modalities of choice for metastatic FLL detection, comparison of CEUS with these techniques seems reasonable in evaluating the diagnostic efficacy of CEUS. In addition, histologic confirmation of every malignant FLL in patients with clear imaging diagnosis might not be ethically appropriate. According to a meta-analysis including 828 metastases from 18 studies, overall sensitivity of CEUS for diagnosis of metastases was 91% (95% CI: 87−95%) [30].

RECOMMENDATION 6
CEUS can be used for liver metastases detection as part of multimodality imaging approach (LoE 2, weak recommendation) (Pro 31, Abs 0, Against 0).

Detection of hepatocellular carcinoma (HCC) and intrahepatic cholangiocellular carcinoma (ICC)
With all UCAs, most HCC show AP hyperenhancement (APHE), but the short duration of APHE makes adequate assessment of the whole liver impracticable. The LP lasts long enough for a detailed examination, but the appearances of HCC are variable. Important-ly, not all HCCs demonstrate contrast washout in the LP, limiting the sensitivity of CEUS for HCC detection. CEUS for staging of HCC is not recommended except for patients with portal vein tumor thrombus [51, 52].

With the post-vascular phase UCA (Sonazoid), scanning the entire liver at 10 min or later after injection helps to detect malignant nodules since typical HCC shows as an enhancement defect [53–58]. However, approximately half of well differentiated HCCs do not show enhancement defects in the post-vascular phase [124].

ICCs behave in virtually the same manner as metastases, washing out rapidly and appearing as defects in the LP, regardless of the appearance in the AP [59]. This pattern may facilitate detection of satellite nodules adjacent to a larger lesion that were not visualized on conventional US.

RECOMMENDATION 7
Routine use of CEUS for the surveillance of patients at risk for HCC is not recommended (LoE 4, strong recommendation) (Pro 29, Against 2, Abs 0).

Characterization of FLL in the non-cirrhotic liver
The probability of a FLL being benign (including inflammatory) or malignant depends on the symptoms and past medical history. An incidentally detected FLL in otherwise healthy and asymptomatic persons is likely benign [60, 61], whereas with pre-existing malignant disease, the probability of malignancy is significantly higher [1]. In patients with supportive symptomatology, FLLs may raise suspicion for phlegmonous inflammation or abscess formation.

The primary aim of CEUS in patients with a non-cirrhotic liver is to differentiate benign from malignant FLLs [29, 33, 45, 56–58, 62–78]. Thus, CEUS is useful to facilitate the clinical decision as to whether a sonographically detected liver lesion needs further investigation or surgery [79].

RECOMMENDATION 10
CEUS is recommended as the first-line imaging technique for the characterization of incidentally detected, indeterminate FLL at US in patients with non-cirrhotic liver and without a history or clinical suspicion of malignancy (LoE 1, strong recommendation) (Pro 30, Abs 2, Against 0).

CEUS for characterization of focal liver lesions
Before starting liver CEUS, it is necessary to review the patient’s clinical history, laboratory data and any prior imaging. The entire liver and the FLL should be interrogated using conventional B-Mode and color Doppler US in order to obtain reproducible information regarding segmental localization, size and relation to vessels and other anatomical landmarks as well as to guarantee optimal examination quality and ascertain whether underlying cirrhosis is present. The range of tumor types differs between cirrhotic and non-cirrhotic livers, with description of the characterization of FLL discussed separately for each.

RECOMMENDATION 8
Routine use of CEUS for staging of HCC is not recommended (LoE 4, strong recommendation), (Pro 31, Against 0, Abs 0).

RECOMMENDATION 9
Before performing CEUS to characterize FLLs, it is recommended to perform a systematic liver examination using B-Mode and Doppler US (LoE 5, strong recommendation) (Pro 32, Abs 0, Against 0).

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Focal nodular hyperplasia (FNH)

On CEUS, FNH typically appears as a hyperenhancing homogeneous lesion in all phases. The hyperenhancement might be only mild during the PVP and LP [67, 77, 97–100]. Hyperenhancement is usually marked in the AP [100], with a rapid fill-in from the center outwards (a spoke-wheel pattern) (70 %) or sometimes with an eccentric vascular or multilocular arterial supply (30 %) [77, 84]. A centrally hypo- or non-enhancing located scar may be seen in the LP. This, together with the direction of filling of the lesion in the AP if recognizable (centrifugal vs. centripetal), is an important feature to distinguish FNH from shunt (high-flow) hemangiomas. In distinction to an FNH, HCA and a hypervascular malignant FLL show washout as the most important CEUS feature [90].

In the vast majority of cases (93.5 %) iso- or only slight hyper-enhancement of FNH is observed in post-vascular phase compared with the surrounding liver parenchyma, whereas in the remainder (6.5 %) hypoenhancement is observed [101, 102]. Overall sensitivity of CEUS for diagnosis of FNH is 88 % (95 % CI: 81–94 %) according to a large meta-analysis of 365 FNH from 18 studies [30]. Several studies have suggested that diagnostic accuracy of CEUS for diagnosis of FNH is a “matter of size”, with accuracy decreasing in patients with lesion size > 30 mm [2, 3].

Hepatocellular adenoma (HCA)

HCA is a rare benign and sometimes estrogen-dependent hepatic neoplasm. Typical imaging characteristics of HCA are displayed in smaller lesions < 50 mm [77, 90]. At CEUS, HCA shows homogeneous arterial hyperenhancement, typically with rapid, complete, peripherally dominated filling without a spoke-wheel pattern and without a peripheral globular enhancement pattern, which often enables the correct differential diagnosis, except in telangiectatic and inflammatory HCA [87]. However, HCCs and hyperenhancing metastases may exhibit a similar arterial enhancement pattern, making the differentiation impossible during the AP. In the early PVP, HCA usually become isoenhancing or, more rarely, remain slightly hyperenhancing [77, 97]. Previous bleeding episodes or

Hemangioma

After focal fatty sparing, hemangioma is the second most common benign solid lesion of the liver [61, 81]. In asymptomatic patients with a normal appearing liver on US and without findings or history of malignant or chronic liver disease, a well-circumscribed, round-shaped hyperechoic and homogeneous FLL < 30 mm without intratransitional vessels at color Doppler and without halo sign is diagnostic of hemangioma. CEUS or other contrast-enhanced imaging modalities are not recommended for further characterization [82, 83]. CEUS is indicated when a definitive diagnosis of a hemangioma cannot be achieved using conventional US, as the addition of CEUS markedly improves the diagnostic accuracy in 90–95 % of cases [29, 66, 76].

The typical CEUS feature of a hemangioma is peripheral, discontinuous nodular (syn.: globular) enhancement in the AP with progressive centripetal partial or complete fill-in [84–87]. Complete fill-in occurs only in 40–50 % cases during the LP. This filling-in is often more rapid in smaller lesions and the entire lesion may be hyperenhancing in the AP. Persistent iso- or hyperenhancement is sustained through the LP [76, 88–92]. On post-vascular imaging using Sonazoid, hemangiomas appear iso- to hypoenhancing relative to the surrounding liver parenchyma, and may resemble metastatic tumors and HCCs [93, 94]. Overall sensitivity of CEUS for diagnosis of haemangioma is 86 % (95 % CI: 81–92 %) according to a meta-analysis including 612 cases from 20 studies [30].

Atypical appearances, in particular LP hypoenhancement (UCA washout) or lack of centripetal fill-in, have been described and may be explained by the destruction of microbubbles that are not adequately replenished due to very long bubble transit times within the lesion [95]. Hemangiomas with arteriovenous shunts (also called high flow or shunt hemangiomas) show rapid homogeneous hyperenhancement in the AP and therefore can be confused with focal nodular hyperplasia (FNH), or even with hepatocellular adenomas (HCA) or HCCs [76]. They are almost always hyperenhancing in the PVP and LP. Thrombosed hemangiomas show lack of enhancement and can be confused with malignancy if only identified during the later CEUS phases [76, 96].

For differential diagnosis of FLL, CEUS is superior to CT and equivalent to MR imaging [30, 33, 45, 64, 68]. CEUS has been shown to be the most cost effective imaging modality in some countries in Europe [80].

Benign solid FLLs

In addition to contrast enhancement of the FLL compared to the adjacent tissue, vascular architecture during AP can further characterize FLL. The enhancement patterns are summarized in Table 2.

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necrotic portions exhibit intratumoral nonenhancing areas in larger HCA. In most cases, washout occurs in the LP requiring biopsy to exclude malignancy [90]. Due to the different sub-types of HCA, characterization and differentiation (for example from FNH and HCC such as inflammatory subtype) may be difficult using CEUS as well as MRI and biopsy (HCA < 50 mm) or surgery (≥ 50) are indicated for final diagnosis [91]. Liver-specific contrast-enhanced MRI may be helpful when HCA is suspected at CEUS to exclude multilocularity. No studies are available for the diagnosis of HCA using Sonazoid.

### Table 2  Enhancement patterns of benign focal liver lesions in the non-cirrhotic liver.

<table>
<thead>
<tr>
<th>lesion</th>
<th>Arterial phase</th>
<th>portal venous phase</th>
<th>late phase</th>
<th>post-vascular phase</th>
</tr>
</thead>
<tbody>
<tr>
<td>hemangioma</td>
<td>peripheral nodular enhancement</td>
<td>partial/complete centripetal fill in</td>
<td>Incomplete or complete enhancement</td>
<td>iso/slightly hypo-enhancing</td>
</tr>
<tr>
<td></td>
<td>small lesion: complete, rapid centripetal enhancement</td>
<td>nonenhancing regions</td>
<td>nonenhancing regions</td>
<td></td>
</tr>
<tr>
<td>FNH</td>
<td>hyperenhancing from the center, complete, early</td>
<td>hyperenhancing</td>
<td>iso/hyperenhancing</td>
<td>iso/slightly hyper- or hypoenhancing</td>
</tr>
<tr>
<td></td>
<td>spoke-wheel arteries</td>
<td>unenhanced central scar</td>
<td>unenhanced central scar</td>
<td></td>
</tr>
<tr>
<td>hepatocellular adenoma</td>
<td>hyperenhancing, complete</td>
<td>isoenhancing</td>
<td>isoenhancing</td>
<td></td>
</tr>
<tr>
<td></td>
<td>nonenhancing regions</td>
<td>hyperenhancing</td>
<td>slightly hypoenhancing</td>
<td></td>
</tr>
<tr>
<td>focal fatty infiltration</td>
<td>isoenhancing</td>
<td>isoenhancing</td>
<td>isoenhancing</td>
<td>isoenhancing</td>
</tr>
<tr>
<td>focal fatty sparing</td>
<td>isoenhancing</td>
<td>isoenhancing</td>
<td>isoenhancing</td>
<td>isoenhancing</td>
</tr>
<tr>
<td>abscess</td>
<td>peripheral enhancement, no central enhancement</td>
<td>hyper-/isoenhancing rim, no central enhancement</td>
<td>hypoenhancing rim, no central enhancement</td>
<td>hypoenhancing rim</td>
</tr>
<tr>
<td></td>
<td>enhanced septa</td>
<td>hypoenhancing rim</td>
<td>enhanced septa</td>
<td></td>
</tr>
<tr>
<td></td>
<td>hyperenhanced liver segment</td>
<td>hyperenhanced liver segment</td>
<td></td>
<td></td>
</tr>
<tr>
<td>simple cyst</td>
<td>nonenhancing</td>
<td>nonenhancing</td>
<td>nonenhancing</td>
<td>nonenhancing</td>
</tr>
</tbody>
</table>

Focal fatty change

Focal fatty changes, either by fat infiltration or fatty sparing are usually shown on conventional B-Mode US as oval or polygonal areas located along the portal bifurcation or close to the hepatic hilum and gallbladder. On visualization of possible focal fat infiltration, atypical location or history of malignancy should prompt further characterization to exclude malignant lesions. Focal fatty change shows the same degree of enhancement (isoenhancing) as the surrounding liver parenchyma during all phases [92, 103]. Typically, a centrally located artery can be identified [82, 92, 104].

Infection

The CEUS findings in phlegmonous inflammation are variable. During the early stage of infection, lesions often appear hyperenhancing, while mature lesions develop non-enhancing foci as liquefaction progresses. Mature liver abscesses on CEUS show enhancement of the margins and frequently of the septae in the AP, which sometimes can be followed by PVP hypoenhancement. The most prominent feature on CEUS is the nonenhancement of the liquefied portions combined with arterial rim enhancement [105–109]. Diffuse hyperenhancement of the affected liver sub-

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segment(s) in the AP and LP washout of liver parenchyma surrounding the nonenhancing necrotic area have been described in the majority of cases [109].

The appearances of granulomas and focal tuberculosis on CEUS are variable, which make it hard and sometimes impossible to differentiate these from malignancy [107, 110–112].

Other solid benign liver lesions

A range of other, very rare, solid benign liver lesions can be seen including the following entities:

- Active hemorrhage (including spontaneous, traumatic and iatrogenic liver bleedings) demonstrates contrast extravasation whereas hematomas appear as non-enhancing areas.
- Inflammatory pseudotumor is a rare disease whose definite diagnosis is usually only made at surgery. It may show arterial enhancement and LP hypoenhancement, falsely suggesting malignancy.
- Hepatic angiomylipoma is a rare benign mesenchymal tumor. It appears homogeneous in most cases and strongly hyperechoic at baseline US. CEUS shows arterial hyperenhancement [76, 113].
- Cholangiocellular adenomas (CCA or bile duct adenoma) are rare lesions that are usually small (90 % < 1 cm). CEUS may show strong arterial hyperenhancement and early washout in the PVP and LP (they lack portal veins), falsely suggesting malignancy [114, 115].
- Hepatic epithelioid hemangioendothelioma (HEHE) often manifests as multinodular FLL. On CEUS, HEHE shows rim-like hyperenhancement in the AP and hypoenhancement in the PVP and LP, a sign of malignancy [96, 116, 117]. Some patients show centrally located unenhanced areas. In contrast, all hemangioma and FNH show hyper- or isoenhancement in the PVP and LP, which is their most distinguishing feature.

For liver trauma we refer to the recently published EFSUMB Guidelines and Recommendations for the Clinical Practice of Contrast-Enhanced Ultrasound (CEUS) in Non-Hepatic Applications: Update 2017 [4, 5].

Malignant solid FLLs

In patients with a non-cirrhotic liver, metastases are more common than primary liver malignant tumors, though conventional US is occasionally helpful to show the malignant nature of an FLL, by demonstrating a hypoechoic halo and infiltration of intrahepatic vessels. Contrast enhanced imaging is necessary to determine the malignant nature under many circumstances, which is true for US, CT and MRI [45, 118]. Contrast washout in the PVP and LP is the most important feature to determine malignancy [2, 3]. Almost all metastases show this feature, regardless of the enhancement pattern in the AP. Very few exceptions to this rule have been reported, mainly in liver metastases of neuroendocrine tumors and atypical HCC (Table 3).

Hepatocellular carcinoma in the non-cirrhotic liver

Hepatocellular carcinoma (HCC) is the most common primary liver malignancy and most of the patients are at risk with known or unknown liver cirrhosis [119]. There is little literature on the value of CEUS in the diagnosis of HCC in the non-cirrhotic liver. Generally, the enhancement patterns of HCC in the non-cirrhotic liver on CEUS are similar to HCC in the cirrhotic liver, but size at time of diagnosis tends to be larger [60]. HCA and FNH are the main dif-

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<th>Enhancement patterns of malignant focal liver lesions in the non-cirrhotic liver.</th>
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HCC in the non-cirrhotic liver are usually hyperenhancing in the AP, typically with a chaotic vascular pattern [121] and variably iso- or hypoenhancing in the PVP and LP [88]. Hyperenhancement in the AP is often homogenous but starts predominantly along the periphery [122]. The fibrolamellar variant of HCC has nonspecific appearances at CEUS. According to expert opinions and case reports, they show rapid wash-in with a heterogeneous pattern in the AP and early PVP and early and marked washout thereafter [123, 124].

**Cholangiocarcinoma**

**(intrahepatic cholangiocellular carcinoma, ICC)**

ICC is the second most common primary malignant liver tumor and usually arises in healthy liver parenchyma. The different treatment approaches and prognosis, necessitate that ICC is distinguished from HCC [125]. Although rarely observed in Europe and America, more frequently seen in Asia, where combined HCC-ICC also exists [126].

In distinction to the late enhancement on CECT or CEMRI, ICC has a variety of patterns in the AP at CEUS but all show washout in the LP [59, 113, 127]. The typical pattern of malignancy is better displayed by CEUS than by CECT or CEMRI [128, 129].

There is controversy on the differential diagnosis of HCC and ICC with CEUS [35, 52, 93, 130–137]. Compared to HCC, ICC shows a less intense enhancement in the AP and shows early (< 60 seconds) and marked washout compared to a typically late and mild washout in HCC [93]. ICC can be subcategorized into three types: mass-forming, periductal infiltrating, and intraductal growing. Mass-forming ICC can exhibit four enhancement patterns in the AP: peripheral irregular rim-like enhancement, heterogeneous hyperenhancement, homogeneous hyperenhancement, and heterogeneous hypoenhancement [113, 127]. Mass-forming ICC usually shows washout in PVP and invariably shows marked hypoenhancement in the LP followed by complete hypoenhancement in the postvascular phase [53, 138].

During the AP, periductal infiltrating ICCs appear heterogeneously enhancing, intraductal growing ICCs exhibit homogeneous hyper-enhancement in most cases. Both lesions show marked washout during PVP and LP [138].

**Metastases**

Liver metastases are the most common malignant lesions of liver, arising mainly from cancers of the gastrointestinal tract, breast, pancreas or lung. CEUS markedly improves the detection of liver metastases compared to conventional US. Liver metastases can be detected and characterized reliably as hypoenhancing lesions during the PVP and LP, with few exceptions. Washout is of marked degree and with early onset, usually before 60 sec after UCA injection. In the LP, very small metastases may be conspicuous and lesions that were occult on B-Mode US can be detected [45]. Due to lack of Kupffer cells, metastatic lesions on post-vascular phase imaging with Sonazoid are clearly demarcated and completely hypoenhancing [53, 136, 137, 139–141].

Metastases usually show at least some contrast enhancement in the AP, exhibiting sometimes a marked and chaotic enhancement. Rim or halo enhancement is often seen [2, 3].

**Lymphoma**

Lymphoma shows variable arterial enhancement but characteristic fast and marked washout in the PVP and LP, predictive of malignancy [87, 142, 143].

**Focal cystic liver lesions (benign and malignant)**

Focal cystic liver lesions (FCLL) represent a wide spectrum of benign and malignant disease [114]. Benign FCLL include simple cysts, hematoma and hemorrhagic hepatic cysts [144, 145], abscess, bilomas, hydatid cysts [146], cystic cavernous hemangiomas [147] and cystic HCA and other rare entities [86, 148]. Malignant FCLL include cystic HCC [149], cystic lymphoma, cystic metastases as typically seen in neuroendocrine tumors [150] and other rare entities. Simple cysts are completely non-enhancing on CEUS, and CEUS is not indicated for assessment of simple cysts but useful to evaluate complicated or atypical cysts [145].

With complex cystic masses, CEUS characterizes the vascular flow within septa in cystadenoma and cystadenocarcinoma [1, 151]. Such septa are often visualized by US but not by CT and MRI. Some atypical cystic lesions may have a solid appearance at conventional US, thus mimicking a malignant lesion, particularly a cystic metastasis or biliary cystadenocarcinoma [1, 32].

The CEUS distinguishing feature in the differential diagnosis of hepatic cystadenoma (HBCA) from hepatic cystadenocarcinoma (HBCAC) is the honeycomb septal hyperenhancement during the AP for HBCA, and hypoenhancement during the PVP and LP for HBCAC [1].

**RECOMMENDATION 14**

If CEUS has definitively characterized a benign FLL, further investigations are not recommended to confirm the diagnosis (LoE 1, strong recommendation) (Pro 26, Abs 5, Against 0).

**RECOMMENDATION 15**

CEUS can be used to characterize hepatic abscess in the appropriate clinical setting (LoE 2, weak recommendation) (Pro 24, Abs 2, Against 1).

**CEUS for characterization of FLLs in liver cirrhosis**

**Study procedure**

In addition to the general recommendations for the study of FLL, important aspects should be followed if the liver is cirrhotic. Since the AP is crucial to observe for characterization of FLL in cirrhosis, good visualization of the nodule is important. Despite the use of a low MI, microbubbles can be disrupted and acoustic output power should then be reduced, while maintaining sufficient signal intensi-
ty to allow contrast persistence until the very LP (beyond 3–4 minutes); crucial for detecting washout and establishing a diagnosis of HCC. Furthermore, when the arterial/early PVP is complete (after 60 seconds), it is recommended to image the lesion intermittently (usually brief scan every 30 to 60 seconds), rather than continuously, to minimize microbubble destruction that may cause problems in the identification of subtle or late washout.

Image interpretation and evaluation

The key feature for the diagnosis of HCC in liver cirrhosis is AP hyperenhancement (APHE), followed by late onset mild washout (>60 sec after injection) [152–156]. This pattern of washout in a HCC is seen in more than 97% of cases according to a large retrospective series [157]. Arterial hyperenhancement is usually homogeneous and intense in HCC, but may be inhomogeneous in larger nodules (>5 cm) that are necrotic. Rim enhancement is atypical for HCC. Washout is observed overall in about half the cases of HCC, but rarely in small nodules (20–30% in those 1–2 cm, 40–60% in those 2–3 cm) [49, 132, 158]. Washout is observed more frequently in HCC with poorer grades of differentiation than in well-differentiated HCC, which tend to be isoenhancing in the LP [159–162]. ICC risk is increased in patients with liver cirrhosis, but only 1–2% of newly detected FLL in a cirrhotic liver are ICCs [126, 163].

Hypoenhancement in the LP is usually less marked in HCC than in other primary tumors or in liver metastases [159, 164]. Furthermore, the washout tends to start later in HCC, usually not before 60 seconds after injection [159, 164] and appearing only after 180 seconds in up to 25% of cases [159, 164]; consequently, it is important to observe nodules in cirrhosis until late (>4 minutes). Early washout (<60 seconds) has been reported to occur in poorly differentiated HCC or to suggest a non-hepatocellular malignancy [159, 160, 162, 164], most often a peripheral ICC. For details regarding the CEUS Liver Imaging Reporting and Data System (LI-RADS) classification we refer to the published literature [52, 128, 129, 133–135, 152, 156, 165, 166]. Sensitivity of CEUS for diagnosis of HCC is 88% (95% CI: 84–92%) according to a meta-analysis including 1333 HCCs from 19 studies [30].

RECOMMENDATION 16
CEUS can be utilized in first line to characterize FLL found in patients with liver cirrhosis to establish a diagnosis of malignancy (CEUS LR-M) or specifically of HCC (CEUS LR-S), but CT or MR imaging remain required for accurate staging unless contraindicated (LoE2, weak recommendation) (Pro 29, Abs 0, Against 0).

RECOMMENDATION 17
CEUS can be utilized when CT or MR imaging is inconclusive, especially in FLL in cirrhotic liver not suitable for biopsy, to assess the probability of a lesion to be an HCC (LoE3, weak recommendation) (Pro 29, Abs 0, Against 0).

Characterization of portal vein thrombosis

CEUS is superior to color Doppler US for the diagnosis of portal vein thrombosis [167]. Acute bland thrombus is typically “avascular” and shows as a void within the enhancing liver in all phases of CEUS but best visualized during the PVP. A “tumor in vein” has the same enhancement characteristics as the tumor from which it originated, including rapid AP hyperenhancement and washout [26, 167–173]. Differential diagnosis between partially occlusive/recanalized bland thrombus and “tumor-in-vein” is more challenging. For reliable differentiation, careful assessment of the arrival time of the UCA to the vein is needed. Early arrival of UCA into the lesion in the portal vein at about the same time as opacification of hepatic arteries suggests tumor but this behavior is not specific to HCC. Tumor in peripheral portal veins may be mistaken for tumor nodules, erroneously downstaging the patient. Avoidance is facilitated by real-time imaging while sweeping through the liver, especially in the PVP, to depict the tubular configuration of the tumor and its continuity with more central portal or hepatic veins.

The tumor source of a malignant portal vein thrombus may be obvious, or it may be identified with the assistance of CEUS. A suspicious thrombus within the portal vein may be amenable to US guided biopsy, targeting, if possible, any enhancing regions within the thrombus [167, 169, 173, 174].

RECOMMENDATION 18
CEUS can be utilized for the selection of FLL(s) in a cirrhotic liver to be biopsied when they are multiple or have different contrast patterns (LoE4, weak recommendation) (Pro 28, Abs 0, Ag 0).

RECOMMENDATION 19
CEUS can be used to monitor changes in enhancement patterns in FLL in cirrhotic liver requiring follow-up (LoE4, Weak Recommendation) (Pro 29, Abs 0, Ag 0).

Contrast enhanced intraoperative ultrasound (CE-IOUS)

Several studies using different UCAs have shown that contrast enhanced intraoperative US (CE-IOUS) enhances tumor detection and allows to assess the region for resection, where previously a pre-treatment colorectal liver metastasis was present but has re-
gressed [175–182]. In particular it has proven valuable for the differential diagnosis between HCC and dysplastic nodule, using both SonoVue [176] and Sonazoid [180]. In addition, CE-IOUS may have an important impact on surgical strategy depending on the attitude of the surgeon [183]. The study procedure and image interpretation of CE-IOUS examination are the same as for the transabdominal approach described in the above sections [175]. The most important difference is that CE-IOUS is performed during the surgical procedure and uses an intraoperative transducer, which, because of its higher frequency, may require a higher UCA dosage.

**RECOMMENDATION 21**
CE-IOUS can be used to detect and characterize FLLs not detected at preoperative imaging (LoE 3, strong recommendation) (Pro 27, Abs 0, Against 0).

**RECOMMENDATION 22**
CE-IOUS is recommended to assess the region for resection, where previously a pre-treatment colorectal liver metastasis was present but has regressed (LoE 2, strong recommendation) (Pro 27, Abs 0, Against 0).

**CEUS for guiding biopsy**
Ultrasound is an established technique to guide biopsy of FLLs, with excellent safety profile and good overall accuracy [184, 185]. Ultrasound is inferior to MR and CT in detecting liver lesions, but with CEUS the sensitivity is comparable to CECT and CEMRI, but importantly, CEUS enables real-time guidance of the biopsy procedure. The addition of CEUS could potentially increase the diagnostic outcome of percutaneous biopsies for four different reasons [186–198]:
- Biopsy can be made from perfused areas to avoid necrosis or avascular tissue [186–192].
- Biopsy can be made of poorly visualized or "invisible" lesions on B-mode US [188–190, 193–198].
- Biopsy can be avoided completely if a CEUS study unequivocally shows typical features of benign FLL or HCC in an appropriate patient population [2, 3].
- The combination of image fusion techniques and CEUS may have a synergistic effect on both modalities. CEUS fusion has been proven to visualize lesions invisible at conventional US fusion in a substantial number of cases [197, 198].

**Study procedure**
Depending on the contrast agent, a two-step procedure is recommended. Typically, the first UCA dose is injected to characterize the target lesion and select a zone for biopsy; and the second dose used for the CEUS-guided biopsy itself. Dual-screen contrast imaging is recommended with simultaneous contrast imaging on one side to visualize the lesion and conventional B-mode imaging on the other to track the needle. Biopsy should be performed during the contrast phase in which the lesion is best visualized. CEUS prior to a suggested US-guided biopsy of a FLL can help by avoiding biopsy in the case of a diagnostically unequivocal CEUS result. Conventional unenhanced US guidance is adequate for biopsy of most tumors detected on real-time US and there is no rationale for substituting with CEUS-guidance for routine use.

**RECOMMENDATION 23**
CEUS-guidance should be attempted to biopsy FLLs that are invisible or inconspicuous at B-mode imaging (LoE 1, strong recommendation) (Pro 27, Abs 2, Against 0).

**RECOMMENDATION 24**
CEUS-guidance should be considered in FLLs with potential necrotic areas or if previous biopsy resulted in necrotic material (LoE 4, weak recommendation) (Pro 29, Abs 0, Against 0).

**Intracavitary uses**
Intracavitary CEUS (ICCEUS, intracavitary administration of UCAs) is increasingly used as an adjunct to US-guided interventional techniques [199–202]. Concepts and techniques have been published [203–205]. Liver abscess drainage and biliary drainage procedures using ICCEUS have been described in detail. A systematic review covering the role of CEUS in relation to percutaneous intervention has been published [190].

**Study procedure**
The standard dosage for intracavitary CEUS is approximately 1 drop UCA per 10 mL normal saline but may vary with the anticipated distribution volume (e.g. ascites). Higher concentrations are possible for problem solving decisions, but accurate imaging is dependent on correct dilution dosage; high concentrations will result in acoustic shadowing. Higher frequency transducers (linear, endoscopic US) demand higher concentrations.

**CEUS guided biliary interventions**
CEUS-guided percutaneous cholangiography can delineate the biliary tree via drainage catheters, T-tubes placed intraoperatively, or during endoscopic access [206–216]. Intraoperatively, 3D-intracavitary CEUS can aid the surgeon to plan resection lines [206–216].
Intracavitary CEUS of the biliary tree is used during the intervention procedure:
- to demonstrate puncture or cannulation success
- to evaluate for communication with other structures, e.g., intestine [214], pleural cavity [206, 207], gall bladder, vessels [210], abscesses or others
- to evaluate the level of obstruction [216]
- after the interventional procedure
- to evaluate for dislodgement or occlusion.

Importantly, intracavitary CEUS reduces or obviates radiation exposure during the intervention. In addition, it has a positive impact on patient logistics during both procedure and follow-up by making transportation of the patient to an X-ray fluoroscopy room unnecessary when catheter dislodgement is suspected, allowing bedside investigation with a portable US system.

**CEUS for the abscess drainage**

Image-guided liver abscess management, normally with US or CT guidance, is a standard procedure, with advantages of both efficiency and effectiveness, allowing percutaneous abscess drainage with either a needle or catheter, with concurrent lavage [184]. During image guided intervention, correct placement of the needle or the drainage catheter can be confirmed using intracavitary CEUS [205]. Communication with other abscess cavities or other structures can be demonstrated or excluded (e.g., bile duct, pancreatic pseudocyst, peritoneal cavity, pleural space), resulting in additional interventions in a number of instances (biliary drainage, pleural drainage, additional abscess interventions in complex cases, pseudocyst intervention etc.). During follow up, the cavity size can be evaluated, and dislodgement of a drainage catheter can be identified or excluded [217, 218].

**RECOMMENDATION 25**

ICCEUS can be used for delineation of the liver abscess cavity, identification of correct drain position and of communication with other structures (LoE 3, weak recommendation) (Pro 28, Abs 1, Against 0).

**RECOMMENDATION 26**

ICCEUS can be used to guide transhepatic biliary interventions (LoE 3, weak recommendation) (Pro 27, Abs 1, Against 1).

**CEUS for interventional tumor ablation**

Ultrasound is the most commonly used imaging modality for guiding ablation therapies in patients with liver tumors [219, 220]. US allows real-time precise placement of the ablation needle in any visualized target lesion. The procedure is safe, rapid and cost-effective in comparison to other imaging modalities, allowing positioning within the target in a short time [184]. The adjunctive use of CEUS is recommended for pre-treatment evaluation of the ablation target, and also for peri-procedural assessment of treatment results [184, 190]. CEUS-guided ablation of liver tumors may be dispensed with when the lesion target is well recognizable at conventional B-mode US.

**Pre-treatment CEUS**

Pre-treatment evaluation includes assessment of ablation target size, vascularization and tumor margins. For ablative treatment of undetected or inconspicuous target lesions at unenhanced US, availability of CEUS-guided technology and possibly real-time CEUS fusion imaging is of pivotal importance [196, 221–224]. Often, two intravenous injections are required: the first to identify the target lesion and plan treatment, the second for correct positioning of the ablation needle. For CEUS, dual screen is recommended to allow the simultaneous real-time visualization of the probe insertion with both conventional B-mode US and CEUS.

**RECOMMENDATION 27**

CEUS prior to US guided ablation procedure is recommended as a complement to US, CT and MRI for treatment planning (LoE 2, strong recommendation) (Pro 27, Abs 2, Against 0).

**RECOMMENDATION 28**

CEUS-guidance is recommended for the US guided ablation of tumors that are invisible or inconspicuous on US (LoE 2, strong recommendation) (Pro 27, Abs 2, Against 0).

CEUS performed 10–15 minutes after ablation treatment should be considered for immediate evaluation of therapeutic efficacy and for early detection of residual viable tumor, which allows for instantaneous CEUS-guided re-ablation under the same anesthesia. This technique has been proven to decrease both the number of second ablation sessions and the tumor recurrence rate during follow-up [225–227]. Similarly, CEUS fusion imaging with CT/MR imaging for periprocedural assessment of ablation has been reported to enable immediate repeat ablations under CEUS or CEUS-CT/MR fusion imaging-guidance, and decrease long term local tumor progression [228–232].

**Study procedure**

After cessation of ablation, a 5–10 minutes period is necessary before performing CEUS to allow the hyperechoic “cloud” of gas produced during the ablation to diffuse into tissue. In cases of residual tumor, a second CEUS injection must be used to allow correct insertion of the ablation needle.
RECOMMENDATION 29
CEUS is recommended for the evaluation of the treatment effect after ablation and guidance for immediate US guided retreatment of residual tumor (LoE 2, strong recommendation) (Pro 26, Abs 0, Against 2).

Post-treatment CEUS
CEUS is a reliable method for evaluation of the ablation margin and detection of tumor recurrence, potentially reducing the number of CT examinations needed during follow-up [225, 229, 232–239].

Follow-up CEUS
The purpose of first post-ablation CEUS is to evaluate immediate treatment response of the target lesion (by size, perfusion, safety margin and residual viable tumor) and to look for complications (such as hemorrhage, hepatic infarction, bile duct dilatation, abscess, biliary tumor, etc.) [226, 240]. Regular CEUS follow-up weeks to months after ablation can detect local recurrence and new lesions [190, 221, 222, 224, 239]. Frequently, more than one injection is required to evaluate multi-ablated lesions, any suspicious areas or new lesions.

RECOMMENDATION 30
CEUS is recommended as the priority imaging method in the follow-up after ablation treatment to identify residual or recurrent tumor at appropriate time intervals (LoE 2, strong recommendation) (Pro 24, Abs 3, Ag 0).

In the early post-ablation evaluation (within the first 30 days), a thin, uniform enhancing hyperemic rim is visible along the periphery of the necrotic region, similar to the findings on CECT. Due attention must be taken not to confuse this with recurrence.

Monitoring Medical Tumor treatment response
Neoangiogenesis is an important target for novel anticancer treatments and many new antiangiogenesis or antivascular treatments aim at destroying or limiting the growth of tumor vessels [241, 242]. Dynamic contrast enhanced US (DCEUS) has emerged for monitoring the response to these drugs [243]. Initially, such monitoring relied on qualitative analyses only. More recently, robust and quantitative features have been developed. To achieve successful results, standardization and strict control of scanner settings are needed [243].

Methodology and equipment for quantification
Measurements of contrast kinetics are performed using a time-intensity curve (TIC) analysis of dynamic contrast enhancement. Background subtraction is necessary to compensate for attenuation effects [244] and extract reliable time-based features, such as time to peak, mean transit time, etc. However, the nonlinear compression applied to the original signals (required to display them on video monitors) distorts amplitude-based TIC features (e.g., peak intensity and area under the curve) [245]. The majority of reports have used uncompressed, post beamformed data (radio-frequency data are not required since the phase information is not essential). TIC based on such raw data sets allow for accurate assessment of both time-based and amplitude-dependent features. All manufacturers that supply built-in analysis packages on their scanners use this type of data but off-line software packages are also available [246]. For details of administration of UCA and quantitative analysis we refer to the EFSUMB position paper [243].

Assessment of antiangiogenic treatment
Since antiangiogenic treatments frequently induce necrosis without causing tumor shrinkage, functional imaging techniques are particularly suitable for the early assessment of response, a task for which both the RECIST and World Health Organization (WHO) size criteria [247, 248] are unsatisfactory. Studies of various types of tumors such as HCC, gastrointestinal stromal tumor (GIST) or renal cell carcinoma (RCC) treated with antiangiogenic therapies have confirmed that DCEUS may allow early prediction of response to treatment [91, 96, 249–262]. The first multicenter study including more than 500 patients in 19 centers [263] correlated DCEUS with progression free survival (PFS) and overall survival (OS). A decrease of 40 % of AUC at one month was correlated to PFS and OS in patients treated with thyrosine kinase inhibitors [264].

RECOMMENDATION 31
DCE-US can be used in the quantitative assessment of response to targeted therapies in patients with malignant tumors of the liver (LoE 2, weak recommendation) (Pro 22, Abs 5, Against 0).

Paediatric liver lesions
Ultrasound imaging is the ideal imaging technique for many areas in pediatrics, and should always be the first line imaging modality whenever practicable [19]. The advantages of US are established: it is child-friendly, easy to use in the difficult child, repeatable, and has limited safety issues. Moreover, in liver imaging, the relatively fat free body habitus of the child renders US an ideal technique for assessment of FLLs. CEUS in the assessment of pediatric liver lesions has been investigated by a number of groups, predominantly in Europe and almost exclusively using the agent SonoVue, with some reports from North America using Definity, using these agents off-label in children. There has been approval from the FDA to use Lumason in the assessment of FLLs in children. This is likely to increase the use of CEUS as a first line imaging method for...
the incidental FLL, as well as for the assessment of malignancy, recurrence and treatment response in the pediatric population [265].

Experience with the assessment of pediatric FLL has been based on the extensive investigations of adult FLL, with initial experience using SonoVue in pediatric practice mainly centered around the investigation of indeterminate FLLs seen on an US examination [265]. A single study applying adult criteria for the CEUS diagnosis of FNH and HCA found good concordance with MR and CT imaging [266]. More extensive experience has been documented with blunt abdominal trauma of the liver, and in the follow-up of focal areas of injury in the liver [267–269]. Experience using CEUS in the assessment of the liver transplant recipients is limited, with studies reporting success mainly with areas of infarction and abscesses formation, with vascular "Doppler rescue" useful [270–272]. For details we refer to the EFSUMB position paper “Role of Contrast-Enhanced Ultrasound (CEUS) in Paediatric Practice: An EFSUMB Position Statement” [19].

Safety and Dose

Most extensive assessment of safety in children has been with SonoVue, with reports of severe anaphylaxis in 1/137 (0.6 %) patients studied [273] and two minor delayed adverse reactions in 2/305 (0.7 %) patients in a review of local clinical practice [274]. Current evidence suggests that the safety profile of using SonoVue in children is similar to that in adults.

The recommended dose from the FDA for Lumason in assessing FLL is based on body weight, 0.03 mL per kg, not exceeding 2.4 mL per injection. On a more practical basis, other authors have indicated a dose regime for the liver that is age based, with adult doses for children over 12 years of age (SonoVue 2.4 mL), half the adult dose between 6 and 12 years (1.2 mL), and one quarter the adult dose under the age of six years (0.6 mL), well within the safety margins of dose finding studies in adults [274]. These relatively small doses may continue to decrease as imaging technology improves [19].

Indications for pediatric liver CEUS

- The child with any incidentally discovered FLLs on an US examination should also have the opportunity to be assessed with a CEUS examination. The CEUS examination requires an intravenous line, as would a CECT or CEMRI examination. Limited evidence suggests the CEUS assessment of a FLL is as accurate in the child as in the adult, without the morbidity of ionizing radiation, iodinated contrast, gadolinium-based contrast or the need for sedation or general anesthesia.
- Any indeterminate FLL in a child with underlying chronic liver disease from any cause on a follow up program should have a CEUS examination prior to any other imaging, with the possibility of avoiding further imaging if the lesion is categorically benign.
- The assessment of vascular complications following liver transplantation benefits from the ‘Doppler rescue’ of a CEUS examination before resorting to a CT examination.
- Initial investigation of blunt abdominal trauma in children should involve a CT examination except for the most trivial trauma. Follow-up of the identified areas of localized, low-energy liver trauma may be readily assessed with a CEUS examination, where a complicating traumatic pseudoaneurysm is readily identified, and progressive healing of lacerations or hematomas can be recorded on serial investigations.

RECOMMENDATION 32

CEUS assessment of FLLs in children is consistent with findings in the adult patients and it should be used to characterize these lesions (LoE 2b, strong recommendation) (Pro 25, Abs 0, Against 0).

RECOMMENDATION 33

CEUS follow-up of traumatic liver injuries in children should be utilized for the assessment of complications, reducing ionizing radiation exposure (LoE 2, strong recommendation) (Pro 26, Abs 1, Against 0).

Documentation

All US examinations should principally be documented both by wording and by image storage to ensure high-quality patient care [275]. However, there exists little scientific evidence to substantiate how this should be done. This is a matter of expert opinion and practice differs substantially worldwide. Each image lesion should be described in terms of size, localization (liver segment), and contrast enhancement in all phases [276]. The operator should record the temporal behavior and degree of enhancement relative to surrounding tissue (non-enhanced, hypo-enhanced, iso-enhanced or hyper-enhanced), as well as the UCA distribution (homogeneous or heterogeneous) [23].

The written report must include type and dose of UCA applied. Furthermore, the enhancement pattern in all phases should be described with a conclusion regarding diagnosis and follow-up. It is important to report washout timing in actual seconds or minutes. Real-time video clips should be recorded, preferably digitally, in a format which enables later retrieval and comparison [277]. The clips should ideally show the whole examination, but at least the AP and other clinically relevant parts of the scan should be recorded and stored. Cine loops can be supplied with still images of relevant findings. Finally, the clips and images should be archived permanently.

RECOMMENDATION 34

The user of a CEUS examination should report the type and dose(s) of contrast agent, the enhancement pattern and clinically relevant findings in a written format (LoE 5, strong recommendation) (Pro 27, Abs 1, Against 0).
Clinical training and education

Adequate knowledge and hands-on training are prerequisites for building competence in the use of CEUS. Therefore, it is of great importance to maintain high quality education and high professional standards in the practice of CEUS. In 2006, three levels of training requirements were defined by EFSUMB [261], with Appendix 14 specifically addressing the use of CEUS [262]. It is recommended that CEUS should be performed by operators who have obtained adequate expertise with both conventional US and CEUS. They should be familiar with these techniques and the distribution of pathologies within their local medical environment; they should be recognized as competent by local standards and the relevant medicolegal framework. Some federations offer dedicated CEUS courses on a regular basis, often in collaboration with the UCA- or US-equipment industry and it is beneficial for any user of CEUS to attend such educational activities [278]. It is advised that investigators intending to start using CEUS attend relevant courses and spend time under the supervision of an expert. Ideally, their own department should have sufficient volume of examinations to maintain adequate numbers of cases with various pathologies. Furthermore, it is advised that the manufacturers are consulted to maintain up-to-date CEUS scanner software. The practice of CEUS also requires knowledge of UCA administration, of contra-indications and necessary skills to handle possible side effects within the medicolegal framework of the country of practice.

Errors and artifacts in CEUS of the liver

Errors may occur in the CEUS of the liver due to the limitations of CEUS and other factors, such as contrast dose, MI, image artifacts, background noise, pseudoenhancement, unintended microbubble destruction, attenuation, shadowing, prolonged heterogeneous liver enhancement. These errors may result in lesion mischaracterization. It is recommended that CEUS imaging interpretation should be performed in conjunction with analysis of patient’s clinical history, symptoms and laboratory values. In select cases when CEUS imaging results are discordant, correlation with other imaging modalities or tissue sampling might be advised.

Conflicts of Interest

Christoph F Dietrich: Speaker honoraria, Bracco, Hitachi, GE, Mindray, Supersonic, Pentax, Olympus, Fuji, Ge, Boston Scientific, AbbVie, Falk Foundation, Novartis, Roche; Advisory Board Member; Hitachi, Mindray, Siemens; Research Grant, GE, Mindray, Supersonic
Richard G Barr: Speaker honoraria, Bracco, Hitachi, GE, Mindray, Supersonic, Philips, Siemens, Lantheus, Canon; Advisory Board – Lanteus and Bracco Diagnostics; Board Member – ICUS; Research Grants – Siemens Ultrasound, Philips Ultrasound, GE Ultrasound, GE Medical, Mindray, Supersonic Imagine, B and K Ultrasound
Annalisa Berzigotti: Research grant, Bracco
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References


Strobel D, Bernatik T, Blank W et al. Diagnostic accuracy of CEUS in the differential diagnosis of small (<20 mm) and subcentimetric (<10 mm) focal liver lesions in comparison with histology. Results of the DEGUM multicenter trial. Ultraschall in Med 2011; 32: 593–597

Bernatik T, Seitz K, Blank W et al. Unclear focal liver lesions in contrast-enhanced ultrasound—lessons to be learned from the DEGUM multicenter study for the characterization of liver tumors. Ultraschall in Med 2010; 31: 577–581


Dietrich CF, Ignea A, Trojan J et al. Improved characterization of histologically proven liver tumors by contrast enhanced ultrasound during the portal venous and specific late phase of SHU 508A. Curr Opin Radiol 2004; 53: 401–405


Jones EC, Chezmar JL, Nelson RC et al. The frequency and significance of small (less than or equal to 15 mm) hepatic lesions detected by CT. Am J Roentgenol 1992; 158: 535–539


Kim TK, Noh SY, Wilson SR et al. Contrast-enhanced ultrasound (CEUS) liver imaging reporting and data system (U-I-RADS) 2017 – a review of important differences compared to the CT/MRI system. Clin Mol Hepatol 2017; 23: 280–289


Heller E, Gorg C. Focal liver lesions in patients with malignant haematological disease: value of B-mode ultrasound in comparison to contrast-enhanced ultrasound – a retrospective study with N = 61 patients. Z Gastroenterol 2013; 51: 558–567


Kessler T, Bayer M, Schwopp C et al. Compounds in clinical Phase III and beyond. Recent Results Cancer Res 2010; 180: 137–163


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