Gadoxetic Acid–enhanced Magnetic Resonance Imaging and Contrast-enhanced Ultrasonography in the Diagnosis of Hepatocellular Carcinoma

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ABSTRACT

Hepatocellular carcinoma (HCC) is the most common primary malignancy of the liver and one of the leading causes of cancer-related deaths. Early diagnosis of HCC is crucial to achieve good outcome. Advances in imaging technology enable detection of early disease, accurate tumour staging, treatment planning, and post-treatment monitoring, as well as an update of management guidelines. This review focuses on the development of gadoxetic acid–enhanced magnetic resonance imaging and contrast-enhanced ultrasonography in the diagnosis of HCC.

Key Words: Carcinoma, hepatocellular; Gadolinium ethoxybenzyl DTPA; Magnetic resonance imaging; Sonazoid; Ultrasonography

中文摘要

釓塞酸增強磁共振成像和超聲造影的肝癌診斷

曹子文
肝細胞癌是肝臟中最常見的原發性惡性腫瘤，是癌症相關死亡的主要原因之一。及時診斷肝細胞癌是達至良好治療結果的關鍵。成像技術的進展有利於早期疾病檢測、準確腫瘤分期、治療計劃和治療後的監測，以及對治療指引的更新。本文重點回顧釓塞酸增強磁共振成像和超聲造影在診斷肝癌的進展。

INTRODUCTION

Hepatocellular carcinoma (HCC) is the sixth most common cancer, accounting for >90% of all primary liver cancers.¹ Early diagnosis of HCC enables effective treatment and reasonable 5-year survival of 50% to 70%.² Risk factors for HCC include chronic viral hepatitis infection, alcoholic and non-alcoholic fatty liver disease, and other types of chronic inflammatory liver diseases that lead to cirrhosis.³⁴ The risks of developing chronic hepatitis B and hepatitis C infections are 2.5% and 2% to 8% per year, respectively.³⁵ Before 2000, the diagnosis of HCC was...
made primarily by histopathological analysis. Advances in imaging and the risk of percutaneous biopsy (such as tumour seeding and bleeding) have resulted in an increased use of non-invasive methods. In 2001, the European Association for the Study of the Liver and the European Organisation for Research and Treatment of Cancer accepted non-invasive criteria based on a combination of imaging and laboratory findings for the diagnosis of HCC. HCC is the only malignancy for which a radiological diagnosis is acceptable without histological confirmation.

Gadoxetic acid–enhanced magnetic resonance imaging (Gd-EOB-MRI) and contrast-enhanced ultrasonography (CEUS) have gained popularity as the investigative tool of choice for HCC. Gadoxetic acid is a hepatocyte-specific contrast agent, commonly used in the detection and characterisation of hepatic lesions, particularly in patients at risk of developing HCC. Its sensitivity is superior to other modalities such as contrast-enhanced (CE) computed tomography (CT) and magnetic resonance imaging (MRI) with extracellular agents (ECA).

For ultrasonography, second-generation contrast agents such as SonoVue (Bracco, Milan, Italy) and Sonazoid (GE Healthcare, Oslo, Norway) are most commonly used. They offer better diagnostic capability (than conventional grey-scale ultrasonography), safety profile (than iodinated or gadolinium chelate agents), and applicability in HCC management.

GADOXETIC ACID–ENHANCED MAGNETIC RESONANCE IMAGING

Gadoxetic acid possesses dual properties by providing information on both the vascular phase (during dynamic contrast enhancement) and hepatobiliary phase (HBP). It has a higher affinity for protein binding and therefore an increased signal intensity during enhancement. While normal liver parenchyma progressively enhances due to the hepatocyte uptake of gadoxetic acid, most HCCs appear hypointense on the HBP. This leads to a greater lesion-to-liver contrast ratio and thus greater sensitivity in detecting small HCCs and greater specificity in differentiation from other focal liver lesions, and provides additional information on the multistep hepatocarcinogenesis process. Early detection and treatment of HCCs can improve patient outcome. Gd-EOB-MRI has also replaced fine-needle biopsy in the diagnosis of atypical lesions.

Mechanism and Technique

Gadoxetic acid is an ionic contrast medium with a linear molecular structure. Its enhancement effect is mediated by gadoxetate, an ionic complex formed by gadolinium and the ethoxybenzyl diethylenetriamine pentaacetic acid ligand (EOB-DTPA), which has a lipophilic property. Thus, gadoxetic acid possesses dual properties of extracellular and hepatobiliary proponents. After intravenous injection, Gd-EOB-DTPA distributes within the vessels and in the interstitial spaces during the dynamic enhancement phases (arterial phase, portovenous phase [PVP], and transition phase). After the PVP, progressive uptake of the contrast by normal hepatocytes peaks at approximately 20 to 40 minutes after injection. The contrast is eliminated by the renal and hepatobiliary tracts in similar amounts (50% each).

Biliary excretion typically commences after 10 minutes of injection in healthy individuals. The uptake by hepatocytes occurs through transport proteins in the sinusoidal membrane (organic anion-transporting polypeptide [OATP] 8, 1B1, and B3), and biliary excretion occurs through proteins in the canalicular membrane (MRP2) later. Gd-EOB-DTPA acts similarly to extracellular gadolinium chelates during the early dynamic enhancement phases, but it provides additional information during the HBP, during which normal hepatocytes concentrate the contrast medium while HCCs do not, resulting in a greater lesion-to-liver contrast ratio.

Gd-EOB-DTPA has a high protein-binding capability and increases the T1 relaxivity. This produces superior enhancement and enables a reduction of dose as compared with other extracellular gadolinium-based contrast media.

The recommended imaging protocols include non–contrast-enhanced sequences, T1-weighted in-phase and opposed-phase sequences, fast T2-weighted fat-saturated sequences, diffusion-weighted imaging (DWI), and pre-contrast T1-weighted fat-saturated sequences. These are followed by intravenous bolus injection of Gd-EOB-DTPA in a dose of 0.1 ml/kg (0.025 mmol/kg) at a rate of 1 ml/s, followed by saline solution flush (20 ml) at the same infusion rate. This corresponds to one-half of the dose of ECA usually used in abdominal studies. After injection, a T1-weighted fat-saturated gradient-echo sequence is obtained in the arterial phase (after 15-20 seconds), PVP (after 50-60 seconds), transition phase (after 120 seconds), and HBP (after 20 minutes). The total scan time may be reduced by
performing the T2-weighted sequence and the diffusion sequence between the transition phase and HBP. Gd-EOB-DTPA is well tolerated and is eliminated through the renal and hepatobiliary tracts. In patients with terminal renal dysfunction, it can be eliminated by dialysis. Nonetheless, there is still a possibility of nephrogenic systemic fibrosis despite the low exposure to gadolinium. Careful evaluation of risks and benefits is needed in patients with severe renal impairment. The compound does not cross the intact blood-brain barrier and diffuses through the placental barrier only in a small concentration. Data about exposure to Gd-EOB-DTPA during pregnancy are not available. No effect to the infant during breastfeeding is expected. Dose adjustment is not required in elderly patients or patients with hepatic or renal dysfunction. An increased bilirubin level is associated with a reduction in the enhancement effect in the liver during the HBP.

Nonetheless, Gd-EOB-DTPA may be associated with hypersensitivity reactions or other idiosyncratic reactions characterised by cardiovascular, respiratory, or cutaneous manifestations, or shock. Its reported side-effects are similar to those of non-specific gadolinium chelates, including nausea (1%), headache (0.9%), lumbar pain (0.5%), vertigo (0.4%), vasodilation (0.6%), dysgeusia, and pain at the injection site.

Detection of Small Hepatocellular Carcinoma
Imaging diagnosis of small HCC (≤2 cm) is a challenge, as typical features of arterial hyperenhancement and washout are uncommon and overlap with other benign or premalignant lesions. Small HCC may be sub-classified as early and progressed HCC. Histologically, early HCC is difficult to distinguish from high-grade dysplastic nodules. The presence of stromal invasion is the key to diagnosis. Early HCC consists of well-differentiated tumour cells that grow by replacement and thus has a vaguely nodular appearance with an indistinct margin. Progressed HCC consists of moderately differentiated tumour cells with an expansile growth pattern and thus has a discrete nodular appearance with a tumour capsule.

In a meta-analysis of patients with chronic liver disease, the per-lesion sensitivity was significantly lower in subcentimetre HCC than larger HCC on CT (31% vs. 82%) and MRI (48% vs. 88%). Gd-EOB-MRI has a higher sensitivity than CECT in detecting HCC because of a higher lesion-to-liver contrast ratio (Figure 1). Gd-EOB-MRI can detect additional HCCs in patients initially diagnosed as having a single lesion on dynamic CT. Accurate detection can reduce the risk of recurrence and improve overall survival. Gd-EOB-MRI is superior to CT or ECA-MRI in detecting small HCCs. A combination of Gd-EOB-MRI and DWI has even higher sensitivity in detecting small HCCs than either method alone.

Hepatocellular Carcinoma Versus Dysplastic Nodule
In cirrhotic livers, atypical enhancement patterns in small HCCs are not uncommon. It is difficult to differentiate early HCC from a non-malignant or premalignant dysplastic nodule owing to overlapping pathological and radiological features. The HBP of Gd-EOB-MRI can be used to distinguish between the two (Figure 2). As OATP expression decreases during hepatocarcinogenesis, HCC is hypointense to background liver parenchyma on the HBP. Signal intensities on DWI and HBP are useful to differentiate between HCC and dysplastic nodule, and between high-grade and low-grade dysplastic nodules. HCC and a lesser proportion of high-grade dysplastic nodules are hyperintense on DWI, whereas low-grade dysplastic nodules are not.

Hyperintensity on the HBP is suggestive of benign hepatocellular lesions such as focal nodular hyperplasia. Most HCCs are hypointense on HBP, though 5% to 20% of HCCs are iso- to hyper-intense on HBP due to genetic mutations, which cause a paradoxical over-expression of OATP, most commonly in moderately differentiated HCC. Common features of hyperintense HCC on HBP include focal geographic defects of contrast uptake (indicating intratumoural necrosis or heterogeneous histological differentiation) and a hypointense rim (indicating a peritumoural capsule).

Hepatocellular Carcinoma Versus Vascular Pseudolesion
Vascular pseudolesions are non-tumourous perfusion alterations (such as arteriportal shunts) and can mimic small HCC. As cirrhosis progresses, sinusoidal capillarisation and obliteration of hepatic venules lead to arteriportal shunting. These shunts usually appear as subcapsular wedge-shaped transient parenchymal enhancement on the arterial phase. In cirrhotic livers, the shunts typically have a centrally located, round or oval appearance due to architectural distortion of
Figure 1. Gadoxetic acid–enhanced magnetic resonance imaging of the liver showing a 9-mm hepatocellular carcinoma in segment IVb (arrows), with very subtle hyperenhancement in the (a) early arterial phase, capsule enhancement and washout in the (b) portovenous phase, distinct hypoenhancement in both the (c) transitional and (d) hepatobiliary phases. Contrast-enhanced computed tomography showing near isoenhancement of the lesion in the (e) arterial, (f) portovenous, and (g) delayed phases.

Figure 2. Gadoxetic acid–enhanced magnetic resonance imaging of the liver showing a dysplastic nodule (arrows) with mild T1-hyperintensity in the (a) in-phase, signal loss in the (b) opposed phase, hypoenhancement in the (c) late arterial, (d) portovenous, and (e) transitional phases, and isointensity in the (f) hepatobiliary phase.
the background parenchyma. On Gd-EOB-MRI, vascular pseudolesion is typically isointense on the HBP due to intact hepatocyte function, whereas most small hypervascularised HCCs are hypointense on HBP (Figure 3). Nonetheless, >10% of vascular pseudolesions are relatively hypointense on HBP, it is suggested that low lesion-to-liver signal intensity ratio on the HBP and hyperintensity on DWI are more likely to indicate HCC than vascular pseudolesion.

**Hepatocellular Carcinoma Versus Non-hepatocellular Tumours**

Patients with chronic hepatitis and cirrhosis are at risk of developing intrahepatic cholangiocarcinoma (ICC), which is the second commonest primary hepatic malignancy after HCC. Differentiating ICC from HCC is crucial as their management and prognoses are different. On CECT or ECA-MRI, ICC typically shows peripheral or weak enhancement on the arterial phase and centripetal or persistent enhancement on the PVP and delayed phases. Nonetheless, small ICCs in patients with cirrhosis display arterial enhancement and/or venous washout more frequently than those in the normal liver. On Gd-EOB-MRI, both ICC and HCC are hypointense on HBP due to the lack of functioning hepatocytes. On Gd-EOB-MRI, features suggestive of ICC include the absence of fat or capsule appearance, central hypointensity on T2-weighted images, lesser degree of arterial enhancement, and target appearance on DWI and HBP images (Figure 4).

Imaging features of metastases vary according to the primary histology. Hepatic metastases from adenocarcinoma (such as colorectal cancer) are usually hypovascular and have arterial rim-like enhancement, whereas metastases from neuroendocrine tumour, renal cell carcinoma, melanoma, and breast cancer are hypervascular and have arterial enhancement.

**Figure 3.** Gadoxetic acid–enhanced magnetic resonance imaging of the liver showing a vascular pseudolesion (arrows) with vague hyperenhancement in the (a) arterial phase and subsequent isoenhancement in the (b) portovenous, (c) transitional, and (d) hepatobiliary phases.

**Figure 4.** Gadoxetic acid–enhanced magnetic resonance imaging of the liver showing an intrahepatic cholangiocarcinoma (arrows), with subtle peripheral enhancement and arteriportal shunting with large perfusion alteration in the left lobe in the (a) arterial phase, centripetal enhancement in the (b) portovenous and (c) delayed phases, and a target appearance with hypointense rim in the (d) hepatobiliary phase.
with delayed washout, like HCC. Regardless of the primary histology, metastases are commonly identified as hypointense lesions on HBP images, owing to the lack of functioning hepatocytes within the tumour.\textsuperscript{53} Thus, Gd-EOB-MRI may not be superior to CECT or ECA-MRI in differentiating metastasis from HCC. Nonetheless, metastasis is more likely in the presence of multiple focal lesions in a non-cirrhotic liver, especially in patients with known malignancy.\textsuperscript{54}

Haemangioma is the most common benign hepatic pathology. On CECT and ECA-MRI, a typical haemangioma shows early peripheral nodular enhancement with subsequent centripetal and prolonged enhancement. On Gd-EOB-MRI, haemangioma is typically hypointense on HBP because of the absence of hepatocytes,\textsuperscript{55} and of low signal intensity on the transition phase as the parenchymal enhancement gradually increases after the PVP, which is known as pseudowashout (Figure 5). According to the Liver Imaging-Reporting and Data System (LI-RADS) guidelines, washout should be determined in the PVP only and not in the transition phase.\textsuperscript{11} Haemangiomas are typically moderately to markedly hyperintense on T2-weighted images and hyperintense on DWI with high apparent diffusion coefficient.\textsuperscript{56,57}

Hepatic angiomyolipoma is a rare benign mesenchymal tumour and comprises variable proportions of thick-walled vessels, smooth muscle cells, and adipose tissue. It has diverse imaging features and mimics HCC to show arterial enhancement, intralesional fat, and washout.\textsuperscript{58,59} In addition, lipid-poor angiomyolipoma has no detectable fat component on imaging or chemical shift imaging.\textsuperscript{58,60} On Gd-EOB-MRI, both entities show similar dynamic enhancement patterns, but angiomyolipoma tends to be more homogeneously hypointense and of lower signal intensity in HBP, as it does not contain hepatocytes, whereas HCC contains hepatocytes with various degrees of malignant change.\textsuperscript{61}

**Gd-EOB-MRI Versus Other Modalities**

Gd-EOB-MRI provides higher per-lesion accuracy for HCC diagnosis than CECT, CT hepatic arteriography, CT arterioportography, and ECA-MRI.\textsuperscript{62-64} It provides maximum lesion conspicuity for HCC,\textsuperscript{35,65,66} and characterises atypical HCCs.\textsuperscript{53,67-69} It adds value in HCC diagnosis when combined with other imaging modalities.\textsuperscript{35,68,70-74} Gd-EOB-MRI is more cost-effective than ECA-MRI or CECT; its direct costs are lower and can generate more quality-adjusted life years.\textsuperscript{75}

**Gd-EOB-MRI as a Biomarker**

HCC that is hyperintense on HBP has a higher grade of tumour differentiation and vice versa.\textsuperscript{76-80} HCC with iso- / hyper-enhancement on HBP has a lower risk for vascular invasion and recurrence,\textsuperscript{81,82} and lower levels of expression of poor-prognostic immunohistochemical / progenitor cell markers including alpha-fetoprotein, protein induced by vitamin K absence or antagonist II, epithelial cell adhesion molecule, cytokeratin 10, and glypican-3.\textsuperscript{13,82,84} Signal intensity, morphology, and signal heterogeneity in HBP are poor prognosticators.\textsuperscript{59,86}

For hypovascular nodules on dynamic imaging, hypointensity on HBP indicates high-risk lesions that later transform into overt hypervascular HCC.\textsuperscript{87-92} The presence of hypovascular HBP hypointense nodules

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**Figure 5.** Gadofacetic acid–enhanced magnetic resonance imaging of the liver showing a hepatic haemangioma at segment VIII (arrows), with peripheral nodular hyperenhancement in the (a) arterial phase, contrast fill-in in the (b) portovenous phase, hypoenhancement or pseudowashout in the (c) transitional phase, and hypointensity in the (d) hepatobiliary phase.
predicts multi-centric recurrence after resection.93

**CONTRAST-ENHANCED ULTRASONOGRAPHY**

Colour and power Doppler ultrasonography have a low signal-to-noise ratio and thus can only depict blood flow in relatively large vessels and cannot detect intralesional vascularity.94 Conventional ultrasonography is limited in diagnosis of HCC, as it depends on a characteristic vascular enhancement pattern. Ultrasound contrast agents can overcome this limitation.

Ultrasound contrast agents consist of microbubbles containing air or various gases within a shell. When administered into the vasculature, the agents enhance the backscatter of the ultrasound waves by resonance within sonic windows.95 This results in a marked amplification of the signals from the blood flow and provides additional information about the microvasculature.94

CEUS has been applied to HCC management, including surveillance, diagnosis, CEUS-guided treatment, treatment response evaluation, and follow-up.96 CEUS is safer and more assessable than CECT or CEMRI for real-time dynamic assessment of vascular perfusion. CEUS with Sonazoid provides an additional Kupffer phase, which is similar to MRI with superparamagnetic iron oxide. HCCs in cirrhotic livers usually do not harbour reticuloendothelial (Kupffer) cells, which differ from normal and cirrhotic liver parenchyma. This leads to a visualised defect in Sonazoid uptake in the postvascular or Kupffer phase.97-100

The Japan Society of Hepatology and Asian Pacific Association for the Study of the Liver guidelines have incorporated CEUS with Sonazoid into HCC management. The guidelines allow confirmation of HCC when CEUS shows a hypervascular lesion and / or a defect in the Kupffer phase, even if a lesion does not display the typical arterial hyperenhancement and subsequent washout on either CECT or CEMRI. However, CEUS has been removed from the latest American Association for the Study of Liver Diseases guidelines because of potential false-positive HCC diagnosis in patients with ICC and because Sonazoid is not licensed for use in the liver in the United States.101 The role of CEUS in differentiation of HCC and ICC remains controversial. On CEUS, compared with HCCs, ICCs enhance to a lesser degree and at a later time after injection, as well as washing out more quickly. In experienced hands, CEUS and CECT are comparably accurate in diagnosing ICC.102-105

**Ultrasound Contrast Agents**

The first-generation ultrasound contrast agent, Levovist (Bayer Schering Pharma, Berlin, Germany), consists of air within a shell of galactose microparticles (99.9%) / palmic acid (0.1%). The lack of stability of the microparticles hampers its commercial use.95 Second-generation ultrasound contrast agents such as SonoVue, Sonazoid, Definity (Lantheus Medical Imaging, North Billerica [MA], USA), and Optison (GE Healthcare, Princeton, NJ, USA) successfully stabilised microbubbles by replacing air with a more inert and slowly diffusing gas such as sulphur hexafluoride or perfluorobutane. Definity consists of octafluoropropane gas within a lipid shell, and Optison consists of octafluoropropane within an albumin shell. Both are approved for cardiac application only.95,106 SonoVue consists of sulphur hexafluoride (SF₆) within a phospholipid shell. SF₆ is an inert molecule that does not interact with any other molecule in the body. After destruction of the microbubble, SF₆ gas is excreted through the lungs only. The shell consists of a monolayer of an amphiphilic phospholipid. The outer side of the shell is in contact with blood and has hydrophilic properties, whereas the inner side has hydrophobic properties and the shell can contain SF₆ gas with stability.94 As only 7.3% of SonoVue is phagocytosed by Kupffer cells, a parenchyma-specific Kupffer phase cannot be obtained. Repeated injections are required to evaluate the entire liver, and thus its use is not approved in Japan.107 Sonazoid consists of perfluorobutane within a hydrogenated egg phosphatidylserine shell. In contrast to SonoVue, 99% of Sonazoid is phagocytosed by Kupffer cells and thus can be used to obtain both vascular phase and Kupffer phase images. The entire liver can be evaluated during the Kupffer phase, with microbubbles trapped by the Kupffer cells leading to a homogeneous enhancement in normal functioning liver parenchyma.95 Lesions lacking functioning Kupffer cells, including small malignant tumours, appear as defects. Currently, Sonazoid has been approved only in Japan and Korea for the evaluation of focal liver lesions.13

**Mechanism and Technique**

The enhancement patterns of CEUS and CECT or CEMRI are not comparable, as the ultrasound contrast agent is retained only within the blood vessels, whereas CT and MRI contrast agents move into the extracellular space until the concentration is balanced between the
intravascular space and the extracellular space.\textsuperscript{108}

The recommended dose of Sonazoid for evaluation of the liver is 0.015 ml/kg. Nonetheless, image quality is maintained at doses lower than the recommendation.\textsuperscript{107} For SonoVue, after reconstitution as directed, 1 ml of the resulting dispersion contains 8 μl sulphur hexafluoride in the microbubbles, equivalent to 45 μg. The common single injection volume of dispersion is 2.4 ml, although half-dose and repeat injections may be administered as needed.\textsuperscript{13}

Vascularity of focal lesions can be evaluated during the arterial phase (10-20 seconds after injection and lasting 30-45 seconds). The PVP starts at 30-45 seconds and lasts for 2-3 minutes, whereas the late phase starts at 2-3 minutes and lasts for 4-6 minutes. On the PVP or late phase images, the degree of washout between the focal liver lesion and the adjacent liver parenchyma can be compared.\textsuperscript{108} Washout is defined as the transition of iso- or hyper-enhancement to hypoenhancement as compared to the adjacent normal liver parenchyma.\textsuperscript{109} When using Sonazoid, a Kupffer-phase image can be additionally obtained from 10-15 to 120 minutes after injection.\textsuperscript{110,111} The equilibrium phase of CECT or CEMRI does not exist on CEUS, as the ultrasound contrast agent is a pure intravascular contrast agent, and no concentration equilibrium can be achieved.\textsuperscript{106}

The mechanical index is defined as the peak rarefractional (or negative) pressure divided by the square root of the ultrasound frequency. At a very low mechanical index, microbubbles stay static and only play a role in the scattering of the ultrasound beam. As the mechanical index increases, microbubbles oscillate at their resonance frequency linearly (<0.2) or nonlinearly (0.2-0.5). When the mechanical index is >0.5, microbubbles oscillate strongly and expand, resulting in disruption of the bubbles. CEUS images can be created from either the signals of the nonlinear oscillation of microbubbles or from microbubble destruction.\textsuperscript{94} In first-generation ultrasound contrast agents, a high mechanical index of >0.7 is used, and CEUS images are created using signals from microbubble destruction. As a result, only intermittent scanning can be performed for a few seconds and images are recorded frame by frame. In second-generation ultrasound contrast agents, a low mechanical index of <0.3 is used, and thus continuous, real-time scanning is possible.\textsuperscript{112} To detect specific signals from a small amount of ultrasound contrast agent, the use of the contrast-specific ultrasound mode is essential.\textsuperscript{106}

**Safety Considerations**

Ultrasound contrast agents are excreted via the lungs only after the destruction of the microbubbles and therefore are not nephrotoxic. They are not iodinated and have no effect on thyroid function. Nonetheless, they can be regarded as foreign materials by the immune system and hypersensitivity reactions may occur.\textsuperscript{113} The incidence of severe hypersensitivity reaction is about 0.002% in large-scale abdominal application studies.\textsuperscript{114,115} The overall incidence of hypersensitivity reaction of ultrasound contrast agents is less than that of iodinated CT contrast agents and is similar to that of MRI gadolinium chelate contrast agents.\textsuperscript{113}

SonoVue is contraindicated in patients with acute coronary syndrome or clinically unstable ischaemic cardiac disease, right-to left shunts, severe pulmonary hypertension, uncontrolled systemic hypertension, and adult respiratory distress syndrome.\textsuperscript{116} Sonazoid is contraindicated in patients with right-to-left shunts, severe pulmonary hypertension, and adult respiratory distress syndrome. Sonazoid should be avoided or used with extreme caution in patients with egg allergies, as its shell is made of hydrogenated egg phosphatidylserine sodium. The safety of SonoVue and Sonazoid has not been evaluated in pregnant women; both should be avoided in women who are breast-feeding or in patients younger than 18 years.\textsuperscript{13}

Insonation of microbubbles may cause harmful effects to cells or tissue, such as microvascular rupture, haemolysis of red blood cells, increased heating around the ultrasound contrast agent, and killing of phagocytic cells that have engulfed the contrast agent.\textsuperscript{115} The European Federation of Societies for Ultrasound in Medicine and Biology recommends use with caution, as damage to the microvessels of the eye or brain can be clinically harmful.\textsuperscript{117} A mechanical index of >0.4 rapidly accelerates this harmful biological effect and hence it should be maintained as low as possible.\textsuperscript{115}

**Diagnosis of Hepatocellular Carcinoma**

On CEUS, 93.5% to 97% of HCCs in cirrhotic livers exhibit arterial hyperenhancement in comparison with the surrounding liver tissue. Hyperenhancement in the arterial phase is usually homogeneous and intense, but may be inhomogeneous in larger nodules (>5 cm),
because of regional necrosis. A thin, perilesional, rim-like hyperenhancement is seen in about 5% to 34.6% of HCCs, which may represent the tumour capsule or blood vessels around the lesion. Most HCCs show earlier enhancement than the surrounding liver tissue. The detection rates of hyperenhancement in lesions ≤1.0 cm, 1.1-2.0 cm, and 2.1-3.0 cm are 67%, 83%-88%, and 92%-100%, respectively. CEUS has a relatively low ability to determine the characteristics of smaller lesions.

Washout in the PVP to late phase is characteristic of HCC and is more common in larger lesions (up to 80.4% in PVP and 95.3% in late phase). In lesions measuring 1-2 cm, only 53.5% exhibit washout in the PVP and 69% to 90.7% in the late phase. Washout is observed more frequently and quickly in HCCs with poorer grades of differentiation, compared with well-differentiated HCCs, which tend to be iso-enhanced in the late phase. Compared with other liver malignancies such as ICC and metastatic liver cancer, HCC usually has less marked washout in the late phase. In HCC, washout tends to start later (60 seconds after injection), and in about 25% of cases, washout appears only after 180 seconds. Therefore, it is important to observe nodules in cirrhosis for >4 minutes to increase the sensitivity for the diagnosis of HCC (Figure 6).

Pathologically, large regenerative nodules and low-grade dysplastic nodules generally show arterial and capillary supply similar to that detected in the adjacent cirrhotic nodules, whereas high-grade dysplastic nodules and HCCs may show abnormally increased arterial supply. 33.3% to 60% of high-grade dysplastic nodules show arterial hyperenhancement, whereas 40% to 66.7% show hypo-enhancement. Washout is seldom seen in the late phase for high-grade dysplastic nodules, in contrast to typical HCCs.

The sensitivity, specificity, and positive predictive value of CEUS in diagnosing HCC are 88.8%, 89.2%, and 91.3%, respectively. Diagnostic ability is associated with nodule size; sensitivity for nodules of 1.0-2.0 cm, 2.1-3.0 cm, and 3.1-5.0 cm is 69%-80%, 97%, and 100%, respectively, and the specificity is 82%-87%, 97%, and 100%, respectively.

**Hepatocellular Carcinoma Surveillance**

Conventional ultrasonography is non-invasive, low cost, has no radiation exposure, and is easily accessible, but its diagnostic accuracy is insufficient for HCC, particularly for small lesions.

CEUS is not recommended as the sole imaging tool to screen for HCC, because its arterial phase is too short to examine the entire liver, and washout in the PVP or late phase may not be always detected in small or well-differentiated HCCs.

In CEUS with Sonazoid, HCCs appear hypoechoic on the Kupffer phase, and the entire liver can be assessed with a single injection (Figure 7). Its specificity in diagnosing HCC is higher than that of conventional B-mode ultrasonography (97.8%-98.2% vs. 89.2%-94.9%). The positive and negative predictive values have been reported as 99% and 97%, respectively. Histologically advanced HCC might appear as more hypoechoic than the adjacent liver parenchyma on the Kupffer phase, which is comparable to the signal intensity difference in the HBP of Gd-EOB-DTPA-enhanced MRI. In addition, the defect reperfusion image in both Kupffer-phase and arterial phase images can be evaluated simultaneously and with the same slice by reinjection of Sonazoid during the Kupffer phase. CEUS with Sonazoid has higher diagnostic accuracy (95%) than CECT (82%) in the depiction of malignant hepatic lesions using the defect reperfusion technique.

In patients with compensated hepatitis C virus–related liver cirrhosis, CEUS with Sonazoid is a cost-effective screening tool when the annual incidence of HCC is >2% and the sensitivity of CEUS in detecting HCC is >80%. Nonetheless, most centres prefer to use CEUS as a problem-solving tool, particularly for nodules measuring 1-2 cm.

**Hepatocellular Carcinoma Intervention**

In lesions with atypical patterns, ultrasound-guided percutaneous biopsy is recommended for definitive pathological diagnosis. CEUS prior to biopsy procedures can increase the diagnostic yield by 10% and decrease the false-negative rate, especially in large tumours with areas of necrosis. CEUS can localise the optimal site for biopsy by demonstrating regions of vascularised viable tumours and by avoiding regions of necrosis.

Common HCC treatments include surgical resection and liver transplantation, ethanol ablation, radiofrequency ablation, and microwave ablation, particularly for small or focal recurrent / residual HCC lesions. Survival after...
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ablation in Child-Pugh A patients is 50% to 70% at 5 years, comparable to that after resection.\(^{138-140}\) Prior to percutaneous therapy, CEUS can be used to assess HCC lesion size, margins, and its relationship with the surrounding structures, and to plan the treatment strategy.\(^{128}\) CEUS can also guide the real-time puncture during the arterial phase, PVP, late phase and in the instance of Sonazoid contrast, the Kupffer phase. For multiple lesions, ultrasound contrast agents can be

Figure 6. (a) Pre-contrast injection ultrasonography of the liver showing a well-defined hyperechoic lesion at segment VI (arrow). Contrast-enhanced ultrasonography with Sonazoid showing avid homogeneous arterial enhancement of the lesion (b) in the arterial phase, and contrast washout (c) in the portovenous phase indicating a hepatocellular carcinoma (arrows).

Figure 7. Conventional ultrasonography of the liver showing two isoechoic lesions (arrows) that are distinctly hypoechoic (arrows) in the Kupffer phase on contrast-enhanced ultrasonography with Sonazoid (Courtesy of Prof. MJ Kim, Department of Radiology, Yonsei University College of Medicine).

Figure 8. Immediately after radiofrequency ablation for a hepatocellular carcinoma, the ablative zone (arrows) is hypoechoic on (a) conventional ultrasonography and has a thin rim enhancement on (b) contrast-enhanced ultrasonography with SonoVue, indicating post-ablative hyperaemia.
administered repeatedly to guide percutaneous therapy at multiple sites.\textsuperscript{128,141}

Fusion imaging with conventional ultrasonography and CECT or CEMRI can be used for guiding biopsy or ablation of inconspicuous lesions. About 83\% of HCCs that are inconspicuous on fusion imaging with conventional ultrasonography are well visualised on CEUS.\textsuperscript{142}

**Treatment Response**

The use of CECT or CEMRI to detect residual viable tumour or recurrent HCC for treatment response evaluation is widely accepted.\textsuperscript{143,144} CEUS is regarded as a competent alternative for this purpose.\textsuperscript{143,145-151}

Intra-procedural CEUS is effective and comparable to early follow-up CECT in the assessment of percutaneous ablative therapies and adequacy of ablative margins.\textsuperscript{145,148,152,153} Intra-procedural CEUS reduces the numbers of incomplete treatments and re-treatments and the total cost of radiofrequency ablation for HCC.\textsuperscript{152} Nonetheless, post-procedural reactive hyperaemia commences soon after ablative therapy, lasting up to a few weeks, potentially obscuring small-volume residual viable tumour. Uniform rim enhancement can be seen up to 30 days after radiofrequency ablation and should not be misdiagnosed as marginal tumour recurrence (Figure 8).\textsuperscript{106}

The role of CEUS versus CECT or MRI for long-term surveillance remains controversial.\textsuperscript{155} According to the modified Response Evaluation Criteria in Solid Tumors, viable HCC is defined as uptake of contrast agent in the arterial phase of CEUS, whereas complete response is defined as disappearance of any intratumoural arterial enhancement in HCC.\textsuperscript{149} CEUS and CECT show conflicting results depending on the timing of follow-up; their results are comparable at 1 month or earlier, although the sensitivity of CEUS in detecting both local recurrence and new intrahepatic recurrence in long-term follow-up is lower than that of CECT.\textsuperscript{154} For response evaluation after transarterial chemoembolisation, the accuracy of CEUS ranges from 72.6\% to 100\% and that of CECT from 61\% to 94\%.\textsuperscript{143} The suboptimal performance of CEUS in long-term follow-up can be attributed to its difficulty in providing an overview of the entire liver to detect HCC progression and intrahepatic recurrence even after reinjection of contrast.\textsuperscript{128,153,154} Nonetheless, when CECT or CEMRI is contraindicated or inconclusive, CEUS may be an alternative to assess tumour progression and intrahepatic recurrence.\textsuperscript{106,128}

**Guidelines on Management of Hepatocellular Carcinoma**

Most guidelines recommend multiphase multidetector row CT and / or contrast-enhanced ECA-MRI as the standard imaging modalities for diagnosis of HCC, based on the typical arterial phase hyperenhancement and washout on the PVP or delayed phase. Gd-EOB-MRI is recommended as the primary diagnostic imaging modality for the diagnosis of HCC by the Japan Society of Hepatology, the Korean Liver Cancer Study Group and National Cancer Center, and the LI-RADS, which is acknowledged by the American Association for the Study of Liver Diseases.\textsuperscript{18} The guidelines set by the European Association for the Study of the Liver-European Organisation for Research and Treatment of Cancer and the Asian Pacific Association for the Study of the Liver are being updated.

According to the 2014 Japan Society of Hepatology guidelines, a non-invasive diagnosis of HCC can be made when a mass shows: (1) arterial hypervascularity and venous washout, (2) arterial hypervascularity without venous washout but with hypointensity on the HBP, or (3) arterial hypovascularity on Gd-EOB-MRI but with hypervascularity on CEUS with Sonazoid and / or defect in the Kupffer phase. Haemangioma must be excluded; it can exhibit pseudowashout in the transition phase (the late dynamic phase between PVP and HBP usually in a 3-minute delayed scan), and hypointensity on the HBP as well as defect in the Kupffer phase on CEUS with Sonazoid.\textsuperscript{18}

According to the 2014 Korean Liver Cancer Study Group and National Cancer Center guidelines, hypointensity on the HBP cannot be regarded as an alternative to washout, but hypoenhancement on the 3-minute delayed scan can be considered as washout.\textsuperscript{18}

In contrast, the LI-RADS versions 2014 and 2017 stipulate that washout appearance should only be described on the PVP, as hypointensity on the transition phase alone is partially attributed to the progressive contrast uptake of the background liver parenchyma. It incorporates hypointensity on the HBP as one of the ancillary features of malignancy, thus allowing the observation to be upgraded to LR-4, or probably HCC.\textsuperscript{12}

The difference in adoption of Gd-EOB-MRI into
various guidelines reflects the preference of either high sensitivity or high specificity. For example, when hypointensity in the transition phase and / or HBP is considered as an alternative to washout, the sensitivity is increased at the cost of increasing the likelihood of false positive, such as haemangioma and ICC. In countries where liver transplantation is not a prevalent treatment for HCC, the potential increase in false positive and treatment thereof may be considered acceptable.

Compared with Gd-EOB-MRI, CEUS is not well-accepted in management guidelines worldwide. This can be attributed to the limited availability of Sonazoid and hence its user experience.

CEUS and Gd-EOB-MRI are complementary in HCC management. In Hong Kong, both modalities tend to be reserved as problem-solving tools. CEUS with Sonazoid can be a cost-effective strategy in HCC surveillance, and Gd-EOB-MRI is more cost-effective than ECA-MRI and CECT in HCC diagnosis. In addition, CEUS is complementary to conventional ultrasonography during interventional procedures. The use of Gd-EOB-MRI and CEUS in HCC management should be prospectively validated, based on differences in disease incidence, healthcare system, and cost worldwide.

CONCLUSION

Gd-EOB-MRI improves the ability to detect dysplastic nodules, diagnose early or atypical HCCs, differentiate HCC from vascular pseudolesions and non-hepatocellular tumours, and predict outcome. It is superior to CECT and ECA-MRI and has been included in various HCC management guidelines. CEUS can play a role in HCC surveillance, problem solving in HCC diagnosis, guide interventional procedures, and assess treatment response. Sonazoid has an additional benefit of providing the Kupffer phase, during which the entire liver can be assessed from a single injection.

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