Contrast enhanced ultrasound of hepatocellular carcinoma

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Abstract
Sonazoid (Daiichi Sankyo, Tokyo, Japan), a second-generation of a lipid-stabilized suspension of a perfluorobutane gas microbubble contrast agent, has been used clinically in patients with liver tumors and for harmonic gray-scale ultrasonography (US) in Japan since January 2007. Sonazoid-enhanced US has two phases of contrast enhancement: vascular and late. In the late phase of Sonazoid-enhanced US, we scanned the whole liver using this modality at a low mechanical index (MI) without destroying the microbubbles, and this method allows detection of small viable hepatocellular carcinoma (HCC) lesions which cannot be detected by conventional US as perfusion defects in the late phase. Re-injection of Sonazoid into an HCC lesion which previously showed a perfusion defect in the late phase is useful for confirming blood flow into the defects. High MI intermittent imaging at 2 frames per second in the late phase is also helpful in differentiation between necrosis and viable hypervascular HCC lesions. Sonazoid-enhanced US by the coded harmonic angio mode at a high MI not only allows clear observation of tumor vessels and tumor enhancement, but also permits automatic scanning with Sonazoid-enhanced three dimensional (3D) US. Fusion images combining US with contrast-enhanced CT or contrast-enhanced MRI have made it easy to detect typical or atypical HCC lesions. By these methods, Sonazoid-enhanced US can characterize liver tumors, grade HCC lesions histologically, recognize HCC dedifferentiation, evaluate the efficacy of ablation therapy or transcatheter arterial embolization, and guide ablation therapy for unresectable HCC. This article reviews the current developments and applications of Sonazoid-enhanced US and Sonazoid-enhanced 3D US for diagnosing and treating hepatic lesions, especially HCC.

Key words: Sonazoid; Contrast-enhanced ultrasonography; Contrast-enhanced three-dimensional ultrasonography; Hepatic tumor; Hepatocellular carcinoma; Fusion image

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INTRODUCTION
Since the late 1990s the first-generation agent Levovist (SH U 508A; Schering AG, Berlin, Germany), which consists of air-microbubbles, has been introduced as an ultrasound (US) contrast agent to enhance liver lesions mainly in Japan but also in other countries. In addition to its vascular phases (arterial and portal), this contrast agent has a late phase (parenchyma-specific phase, delayed phase) and can accumulate for up to 20 min within the liver [1-3]. Using both phases, Levovist-enhanced US is useful for characterization of liver tumors [4-13], evaluation of the efficacy of ablation therapy or transcatheter arterial embolization [14-16], and for guiding ablation therapy [17-20], in unresectable hepatocellular carcinoma (HCC). However, the parenchyma-specific contrast of Levovist was effective only when imaging was performed at high acoustic power using a high mechanical index (MI), and the effect was transient. Therefore, it cannot be used in real-time examinations in the late phase, and visualization of the whole liver is limited to a single scan [21,22].

SonoVue (Bracco, Milan, Italy) is widely available in Europe and China, replacing Levovist for radiology applications. Unlike Levovist, SonoVue contrast visualization can be achieved with low MI imaging, a far less destructive technique [23-29]. Thus, low MI techniques with SonoVue enable continuous real-time imaging. SonoVue is thought to be a truly intravascular, blood-pool contrast agent, with no interstitial equilibrium phase. SonoVue is minimally phagocytosed by reticuloendothelial cells (Kupffer cells) [30,31]. Parenchyma-specific contrast images with SonoVue can be seen only 3 to 5 min after the injection. Therefore, SonoVue requires repeated injections during intraoperative contrast-enhanced US to perform a whole liver examination [26,32]. SonoVue is thus not approved in Japan.

A newly developed second-generation ultrasound contrast agent, Sonazoid (Daiichi Sankyo, Tokyo, Japan), a lipid-stabilized suspension of perfluorobutane gas microbubbles was exclusively approved for clinical use in patients with liver lesions and for phase inversion harmonic gray-scale US in Japan in January 2007. Sonazoid is associated with a low occurrence of side effects, i.e. diarrhea 1.6%, albuminuria 1.6% and neutropenia 1.0% [33]. There is no contraindication for patients with renal dysfunction or iodine-allergy while one contraindication of Sonazoid was reported for patients allergic to eggs [34].

Like Levovist, contrast-enhanced US with Sonazoid has two phases of contrast enhancement: vascular and late (parenchyma-specific phase, delayed phase). Both Levovist and Sonazoid microbubbles enhance the liver parenchyma during the late phase, i.e. more than 5 min after injection of the contrast agent, and most of the malignant tumor tissues are not enhanced. Based on the results obtained in rabbit liver tumor models, Sonazoid microbubbles are retained within hepatic reticuloendothelial cells for 10-30 min after injection and enhance the liver parenchyma in late phase contrast-enhanced US at a low (0.4) or moderate MI setting (0.6) [35]. In clinical use for human beings, enhancement of liver parenchyma and non-enhancement of malignant tumors in the late phase of Sonazoid-enhanced US is not completely understood but is thought to be due to phagocytosis by reticuloendothelial (Kupffer) cells or to adherence of microbubbles to the hepatic sinusoids and tumor vascular spaces [29,36,37], or recirculation of Sonazoid microbubbles [38]. Because malignant tumors contain few or no reticuloendothelial cells [39], they appear as perfusion defects in late phase Sonazoid-enhanced US [37,39,40].

It has been recently noted that more Sonazoid microbubbles than Levovist are taken up by reticuloendothelial cells in the rat liver [40]. The high stability of Sonazoid has overcome the deficiency of Levovist and microbubbles of Sonazoid have the advantage of long-lasting imaging and strong contrast effects that allows detailed observations [37,41]. Moreover, in the late phase of Sonazoid-enhanced US, we scanned the whole liver using this modality at a low MI without destroying the Sonazoid microbubbles, and this method allowed detection of small malignant lesions as perfusion defects. This method also allows the detection of small hypervascular HCC lesions that cannot be detected by conventional US [37]. Moreover, we can use Sonazoid in combination with low MI and high MI to obtain the advantages of both conditions. Because of these features, Sonazoid is widely used in Japan.

This article mainly reviews the current developments and applications of Sonazoid-enhanced US and Sonazoid-enhanced three dimensional (3D) US for diagnosing and treating hepatic lesions, especially HCC.

HEPATIC LESIONS
Injection dose of Sonazoid
The recommended dose of Sonazoid for liver tumor enhancement is 0.015 mL/kg. This dose was decided on the basis of clinical research conducted approximately 7 years ago [33]. With the development of US equipment, the image quality of Sonazoid-enhanced US is sufficiently good at doses lower than this recommended dose. Injection of the recommended dose of Sonazoid enhances the liver parenchyma excessively, which in turn sometimes induces the attenuation of the deep portion of the liver. This phenomenon prolongs the duration of washout of the malignant tumor in the late phase, resulting in a prolonged examination time. Therefore, most authors injected a decreased dose of Sonazoid to evaluate the vascularity of liver lesions, especially HCC [37,39,42-44] (Table 1).

Sonazoid-enhanced US at a low MI
Sonazoid-enhanced US at a low MI is easy to use, and allows visualization, even using non-high-end equipment, and therefore, dependence on the operator’s skills/equipment is decreased, which may facilitate the widespread use of contrast-enhanced US [45].

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Figure 1  A 67-year-old man with recurrent HCC (maximum diameter 25 mm) in segment IV. A: Fusion image combining arterial phase contrast-enhanced CT (right side) and early phase Sonazoid-enhanced US employing the coded phase inversion (CPI) mode at a low MI (left side). Arterial phase contrast-enhanced CT shows a high attenuation area (arrowheads) adjacent to the non-enhanced area treated by radiofrequency ablation (RFA) (arrows) in segment IV. Early phase Sonazoid-enhanced US by the CPI mode at a low MI shows homogeneous enhancement in the recurrent viable area (arrowheads) and no enhancement in the necrotic area (arrows). These enhanced and non-enhanced areas correspond well to the arterial phase contrast-enhanced CT; B: Late phase Sonazoid-enhanced US by CPI mode at a low MI shows a perfusion defect (wash out) in both the recurrent viable area and the necrotic area (arrowheads). It is difficult to differentiate between the necrotic and viable areas because both appear as a perfusion defect; C: Defect on re-perfusion image with Sonazoid-enhanced US by the CPI mode at a low MI shows tumor enhancement in the recurrent viable area which previously showed a perfusion defect in the late phase (see B) (arrowheads). The necrotic area appears as a perfusion defect (arrows).

Table 1  Sonazoid-enhanced US at a low mechanical index (MI)

| Authors          | Injection volume of Sonazoid | MI in late phase | Scan starting time in late phase |
|------------------|------------------------------|------------------|----------------------------------|
| Moriyasu et al[36] | 0.015 mL/kg                 | 0.3-0.5          | 10 min after injection           |
| Korenaga et al[40] | 0.01 mL/kg                  | 0.2-0.3          | 30 min after injection           |
| Hatanaka et al[44] | 0.01 mL/kg                  | 0.2              | At least 10 min after injection  |
| Maruyama et al[42] | 0.0075 mL/kg                | 0.24-0.3         | 5-10 min after injection         |
| Numata et al[39]  | 0.2 mL/body                  | 0.2              | At least 5 min after injection   |

All examinations were conducted using contrast-enhanced phase inversion harmonic US.

Sonazoid-enhanced US at a low MI made it easy to prevent destruction of the microbubbles around the tumor and adjacent to the liver parenchyma, which in turn made it easy to obtain perfusion images of hepatic lesions in the vascular phase (Figure 1A). Sonazoid-enhanced US at a low MI also made it possible to scan the whole liver in the late phase, facilitating detection of perfusion defect images of hepatic malignant lesions[37,39,46]. Employing both vascular and late phases, Sonazoid-enhanced US is useful for characterization of liver tumors[39], histological grading of HCC lesion[42], early recognition of HCC dedifferentiation[47], and guiding ablation therapy for unresectable HCC[44,46]. Xia et al[39] reported that Sonazoid-enhanced US was significantly more sensitive than dynamic CT in depicting the residual tumor blood supply to HCCs 1 wk after (transcatheter arterial chemoembolization (TACE) (P < 0.01, Chi-squared test). Sonazoid-enhanced US appeared to be a highly sensitive and accurate modality for evaluating responses of HCCs shortly after TACE.

Sonazoid-enhanced US at a low MI allows the detection of small viable HCC lesions that cannot be detected by conventional US. However, it was difficult to differentiate between necrotic and viable areas because both appeared as perfusion defects in the late phase[37] (Figure 1B). To solve this problem, Kudo et al[43] re-injected Sonazoid into HCCs which had previously shown a perfusion defect in the late phase. They devised a method called “defect re-perfusion imaging” which confirms blood flow into the defects (Figure 1C). Using this method, Hatanaka et al[44] reported that Sonazoid-enhanced US has a higher sensitivity and accuracy for the diagnosis of hepatic malignancies than contrast-enhanced CT. There were significant differences in sensitivity and accuracy between Sonazoid-enhanced US and contrast-enhanced CT (P < 0.05, respectively).

Korenaga et al[45] reported that Sonazoid-enhanced US was useful for estimating the histological grades of HCCs. Results of late phase images of Sonazoid-enhanced US and MRI with superparamagnetic iron oxide (SPIO) [Ferucarbotran (Resovist), Bayer, Osaka, Japan] matched perfectly (100%) in all of the moderately and poorly differentiated HCCs. They conclude that Sonazoid-enhanced US is a modality that can potentially replace SPIO-MRI. This result is similar to a report by Inoue et al[48]. In the late phase of Sonazoid-enhanced US, all of the moderately and poorly differentiated HCCs appeared hypoechoic and were detected as perfusion defects, whereas the majority (9 of 13 cases, 69.2%) of the well-differentiated HCCs were of an isoechoic pattern[42]. We must remain aware that we cannot detect the majority of well-differentiated HCCs as perfusion defects in the late phase of Sonazoid-enhanced US.
Advantages and disadvantages of Sonazoid-enhanced US at a low MI and a high MI

Vascular phase Sonazoid-enhanced US: Table 2 shows the setting conditions of Sonazoid-enhanced US at a low MI and a high MI. Table 3 compares the images of Sonazoid-enhanced US at a low MI and a high MI for evaluating the vascularity of liver lesions.

Sonazoid-enhanced US at a low MI permits visualization of tumor vessels and tumor enhancement of liver lesions. However, not only relatively large vessels, such as tumor vessels and portal veins, but also microvessels within the liver parenchyma are rapidly filled with Sonazoid microbubbles under low MI conditions, which in turn permit rapid enhancement of the tumor and liver parenchyma simultaneously. This phenomenon induces a short duration of detailed observation for tumor vessels and tumor enhancement in the vascular phase of Sonazoid-enhanced US. Microflow imaging (MFI) is one of the methods of solving this problem.

Late phase Sonazoid-enhanced US: A low MI setting enables repeated observation of the whole liver in real-time while maintaining detailed observation for tumor vessels and enhancing liver lesions.

Grades are classified as poor, good and very good. >> or >>: Low MI is superior to high MI; << or <<: High MI is superior to low MI.

| Parameters | Low MI | High MI |
|------------|--------|---------|
| Transducer | 3.5 or 4.0-MHz convex transducer | Coded harmonic angio (A) |
| Imaging mode | Coded phase inversion (A), Contrast pulse sequence (B), Pulse inversion harmonic (C), Amplitude modulation (D) | Coded harmonic angio (A) |
| MI | 0.12-0.3 (average 0.2) | 0.7-1.0 (average 0.8) |
| Frame rate | 7.13 frames | 2 - 13 frames |
| Focus position | Bottom of the liver (A, D), Bottom of the lesion (B, C) | Bottom of the lesion (A) |

A: LOGIQ 7 ultrasound system (GE Healthcare, Milwaukee, WI). Both imaging software programs consist of phase inversion harmonic and coded technology. These programs restrict signal components from tissues and emphasize signals from blood vessels; B: ACUSON Sequoia ultrasound system (Mochida Siemens Medical System, Tokyo, Japan); C: Aplio-XV ultrasound system (Toshiba, Tokyo, Japan); D: LOGIQ E9 ultrasound system (GE Healthcare, Milwaukee, WI).

Table 3 Comparison between Sonazoid-enhanced US images at low and high MI for evaluation of liver lesion vascularity

| Parameters | Low MI | High MI |
|------------|--------|---------|
| Vascular phase images | | |
| Tumor vessel image | Good | < Very good |
| Tumor enhancement image | Good | < Very good |
| Tumor enhancement image of the lesion located in deep portion or high echoic lesions | Poor | << Good |
| Real time image (frame rate) | Good (high frame rate) | > Good to poor (high to low frame rate) |
| Late phase images | | |
| Scan starting time | At least 10 min | < At least 5 min |
| Perfusion defect image of the lesion | Good | < Very good |
| Perfusion defect images of the lesion located in deep portion or high echoic lesion | Poor | << Good |
| Real-time image (frame rate) | Very good (high frame rate) | >> Poor (low frame rate) |
| Examination time | More than 10 min | < More than 5 min |
| Injection volume of Sonazoid | Half or the volume recommended | < Quarter or third of the volume recommended |

Grades are classified as poor, good and very good. >> or >>: Low MI is superior to high MI; << or <<: High MI is superior to low MI.
time. However, because of US attenuation, we cannot
evaluate whether the hepatic lesion located in the deep
portion of the liver exhibits a perfusion defect or not.
Moreover, due to enhancement of background B-mode,
we cannot adequately evaluate whether or not the high
echoic lesions exhibit perfusion defects in the late phase
of Sonazoid-enhanced US at a low MI (Figure 3A and B).

In the late phase of Sonazoid-enhanced US, we
scanned the whole liver at a high MI with CHA mode,
which enables us to observe a perfusion defect image
clearly even when a hyper-echoic nodule is located in a
deep portion of the liver (Figure 3C). However, the
opportunity to scan with a high MI is limited because of
microbubble destruction.

**Sonazoid-enhanced US at combined high MI and low MI**

**Procedure:** To obtain the advantages of both the low
and the high MI setting of Sonazoid-enhanced US, we
combined the high with the low MI setting for detection
and diagnosis of hypervascular HCC lesions.

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**Figure 2** A 76-year-old man with HCC (maximum
diameter 40 mm) in segment VIII. A: Early phase
Sonazoid-enhanced US by CHA mode at a high
MI shows intratumoral vessels (small arrows), right
portal vein (arrow), hepatic artery (curved arrow),
and hepatic vein (arrowhead); B: Accumulation
maximum intensity holding image in the early
phase. Sonazoid-enhanced US by CHA mode at
a high MI more clearly shows the serial images of
intratumoral vessels (small arrows), right portal vein
(arrow), hepatic artery (curved arrow), and hepatic
vein (arrowhead) than the images in A.

**Figure 3** A 75-year-old man with multiple HCC
lesions (maximum diameter 22 mm, 12 mm
and 10 mm, respectively) in segment VI. A:
Conventional US shows one hyper-echoic HCC
lesion alone; B: Late phase Sonazoid-enhanced
US by CPI mode at a low MI shows two perfusion
defects not detected by conventional US (arrows).
However, one hyper-echoic lesion located in the
deep portion far from the skin surface cannot be
visualized by Sonazoid-enhanced US by CPI mode
at a low MI; C: Late phase Sonazoid-enhanced 3D
US by CHA mode at a high MI shows three HCC
lesions as clear perfusion defects, as depicted on
tomographic ultrasound images in plane A, which
can be translated from front to back (arrows and
arrowheads). The hyper-echoic HCC lesion which
was not detected by Sonazoid-enhanced US at a
low MI is clearly seen (arrowheads).
Figure 4  A 70-year-old man with newly developed HCC (maximum diameter 15 mm) in segment VI and residual viable lesion in segment V (maximum diameter 8 mm). A-B: Arterial phase contrast-enhanced CT shows a high attenuation area in segment VI (arrowhead) (A) and a high attenuation area (arrowhead) adjacent to the non-enhanced area (arrow) in segment V (B); C: Conventional US shows a hypo-echoic tumor in segment VI (arrowheads); D: Early phase Sonazoid-enhanced US by CHA mode at a high MI shows intratumoral vessels and homogeneous tumor enhancement (arrowheads); E: Middle phase Sonazoid-enhanced US by CHA mode at a high MI shows slightly hypo-echoic (wash out) but homogeneous tumor enhancement (arrowheads); F: Late phase Sonazoid-enhanced US by CPI mode at a low MI shows a perfusion defect (wash out) (arrowheads); G-J: Late phase Sonazoid-enhanced US by CHA mode on a high MI intermittent image shows a perfusion defect (G, H) (arrowheads). Intratumoral vessels (I) and homogeneous enhancement (J) are then seen later (arrowheads); K: Conventional US shows a hypo-echoic tumor in segment V (arrowheads); L: Late phase Sonazoid-enhanced US by CPI mode at a low MI shows a perfusion defect (arrowheads). It is difficult to differentiate between the necrotic and viable areas because both appear as a perfusion defect. The normal liver parenchyma is enhanced; M-N: Late phase Sonazoid-enhanced US by CHA mode at a high MI shows intratumoral vessels in the right side of the lesion (viable area) (arrow) and no enhancement in the left side of the lesion (necrotic area). Arrowheads indicate the margin of the lesion. This enhanced area corresponds closely to the area of high attenuation seen in B.
We injected only 0.2 mL/body in order to easily destroy the Sonazoid microbubbles within and around the tumor (Table 1). However, no significant differences in the imaging quality of Sonazoid-enhanced US were observed when the same amount of contrast agent was injected to the patients with different body weight. We consider this amount to be sufficient to evaluate the vascularity of hepatic tumors using the CHA mode at a high MI in the early (20-60 s after injection) (Figure 4) and middle (80-120 s after injection) phases (Figure 4E), using the coded phase inversion (CPI) mode at low MI in the late phase (more than 5 min after injection) (Figure 4F), and finally, using the CHA mode with high MI intermittent imaging in the late phase[53] (Figure 4G-J).

High MI intermittent imaging: Recently, we used intermittent imaging at 2 frames per second with CHA mode at a high MI (0.7-1.2) to depict the features of tumor vascularity in the late phase. We called this method “high MI intermittent imaging”. As a result of using high acoustic power, high MI intermittent imaging destroyed the microbubbles in and around the lesions, and allowed replenishment of microbubbles in the capillary bed between image acquisitions, which resulted in visualization of both tumor enhancement and the distribution of tumor vessels[37]. When we scanned the tumor lesion with high MI intermittent imaging, the Sonazoid microbubbles within and around the tumor may have been destroyed immediately and the tumor vessels and tumor enhancement may have been seen because of back flow into the tumor vessels and vascular spaces. In contrast, if the tumor became necrotic as a result of transcatheter arterial embolization, ablation therapy, or spontaneous necrosis, the lesions would exhibit neither tumor vessels nor enhancement when scanned using high MI intermittent imaging (Figure 4M and N). We then scanned the lesions again by this method to differentiate necrotic areas from newly developed or residual viable tumor areas detected as perfusion defects by Sonazoid-enhanced US at a low MI (Figures 4-6). There were no vascular spaces in the necrotic area, whereas the viable areas contained vascular spaces. We considered these enhanced lesions to be viable HCC lesions and treated them by percutaneous radiofrequency ablation (RFA) therapy guided by late phase Sonazoid-enhanced US (Figure 5) or early phase Sonazoid-enhanced US (Figure 6).

Advantages of this method include that sufficient information can be obtained in patients with multiple lesions. If the multiple lesions are located close to each other, the enhancement patterns can be observed simultaneously during a single scanning process, and if they are located in different lobes or segments, repeated scanning is possible, because of the long duration of late phase Sonazoid-enhanced US imaging (Figure 4). This method is regarded as an alternative to additional injection of contrast agents when tumor vascularity requires further observation. This method can be used when there is a lack of medical personnel or to save time. We finally concluded that high MI intermittent imaging in the late phase is potentially useful for evaluating the enhancement patterns of focal liver tumors[38] and differentiation between necrosis and viable hypervascular HCC lesions after treatment[37].

Sonazoid-enhanced 3D US

Acquisition and reconstruction of Sonazoid-enhanced 3D US: Sonazoid-enhanced 3D US is being used as a clinical diagnostic method in Japan, especially for liver lesions[84]. With the assistance of CHA mode and high MI contrast conditions, which reduce microbubbles in microvessels but not in relatively large vessels, such as tumor vessels and portal veins, Sonazoid-enhanced three-dimensional (3D) US facilitated detailed observation of tumor vessels. Compared with Sonazoid-enhanced (two dimensional) US, Sonazoid-enhanced 3D US allows viewing the volume of interest (VOI) in three orthogonal planes, supplying more spatial information and facilitating easier anatomic assessment.

The process of Sonazoid-enhanced 3D US includes two steps: data acquisition and image reconstruction. With equipment development, a volume probe which scans automatically with internal sectorial mechanical tilt movement to obtain the data has become available. This type of probe supplies us a convenient and fast means for data acquisition in the contrast-enhanced phase. The scanning parameters, including volume angle and number of scanning frames, need to be adjusted before the examination.

After data acquisition, the raw data are stored in the hard disk and image reconstruction can be performed at any time. Two important functions in reconstruction of imaging are to obtain tomographic images and sonographic angiograms. In tomographic images, enhancement of the VOI can be demonstrated in three orthogonal dimensions, and in each dimension, the enhancement of VOI can be presented in multiple parallel planes with an adjustable inter-plane distance (Figure 6). Sonographic angiograms are reconstructed using a rendering mode, such as “gray surface”, “texture”, “maximum intensity”, or “average intensity” in the GE LOGIQ 7 ultrasound system. Different rendering modes can be employed to depict different enhancement characteristics[58] (Figure 7).

Characterization of liver lesions by Sonazoid-enhanced 3D US: Sonazoid-enhanced US has been proved useful for characterizing liver lesions, such as HCC, metastases, hemangioma and focal nodular hyperplasia[39]. In addition to the information from Sonazoid-enhanced US, Sonazoid-enhanced 3D US also supplies important spatial views of any part of the VOI. From the early to the late phase, Sonazoid-enhanced 3D US shows tumor vascularity and changes in tumor enhancement patterns in a multi-planar image[10], Tortuous vessels within the lesion can be clearly depicted (Figure 7).

In a retrospective study on 139 liver lesions, the accuracy of Sonazoid-enhanced 3D US and that of contrast-
enhanced 3D CT for differentiating liver lesions were assessed. The sensitivity was 83% or higher with both modalities, the specificity was 87% or higher with Sonazoid-enhanced 3D US and 92% or higher with contrast-enhanced 3D CT, the positive predictive value was at least 71% with both modalities, and the area under the receiver operating characteristics curve ($\text{Az}$) was at least 0.89 with US and at least 0.92 with CT. Inter-reader agreement was good to excellent ($\kappa \geq 0.76$) with both modalities.

Evaluation of the effect of ablation therapy by Sonazoid-enhanced 3D US: To evaluate the effect of ablation therapy and to detect the residual tumor exactly in the early phase are very important for further treatment. Immediately after the RFA procedure and during the early follow-up period on contrast-enhanced CT images, the presence of hyper-attenuation due to dehydration produced by coagulative necrosis, and periablational enhancement due to reactive hyperemia, arteriovenous shunts, or fibrosis/giant cell reaction, hinder accurate use of this modality for evaluating RFA.[57-59] Compared with CE-CT images, contrast-enhanced US images acquired soon after RFA have shown very few artifacts and have thus been proven to be useful for early evaluation of the therapeutic efficacy of RFA.[60-62]

Sonazoid-enhanced 3D US images allowed observers to compare HCC tumors before treatment with ablated areas after treatment employing a 3D visualization. When the lesion sites were evaluated on Sonazoid-enhanced 3D US images after ablation, the ablation was evaluated as adequate if a non-enhancing area in the early, middle, and late phases covered the hypervascular enhancement seen in the early and middle phases before ablation. Residual tumor on Sonazoid-enhanced 3D US images was diagnosed when an area within the tumor was detected with hypervascular enhancement in the early and middle phases, and was hypoechogenic or isoechoic in the late phase of Sonazoid-enhanced 3D US after ablation (Figure 6).

In a previous study, 63 cases of HCC were treated by RFA. Sonazoid-enhanced 3D US was performed 5-7 d before and 1 d after RFA. Contrast-enhanced 3D CT was performed 5-7 d before and 1 mo after the ablation, and during the follow-up period. When 1-mo contrast-enhanced 3D CT scans were used as the reference standard, the sensitivity, specificity and accuracy of 1 d Sonazoid-enhanced 3D US for detecting residual tumor were 100%, 97% and 97%, respectively, and the Kappa value for agreement between the findings of the two modalities was 0.65. By demonstrating the ablated areas and residual tumors in three dimensions, Sonazoid-enhanced 3D US was shown to be helpful in evaluating the therapeutic effect of RFA for HCC lesions at an earlier time than contrast-enhanced 3D CT.[63] Sonazoid-enhanced 3D US can be used also to evaluate the immediate therapeutic effect of high-intensity focused ultrasound (HIFU) on small HCC lesions. This modality has the potential to en-
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A

B

C

D

E

F

G

H

I

J
able us to detect residual portions of HCCs and perform additional HIFU ablation of the untreated portions\[64\].

**Fusion images combining US with dynamic CT or US with Gd-EOB-DTPA**

For detection of HCC lesions, a combination of modalities using different imaging techniques is recommended to increase the sensitivity and specificity of diagnosis. To achieve this combination of different imaging modalities, we used the fusion feature which can fuse and synchronize US images with multiplanar reconstruction CT or MRI images on the same screen in real time using LOGIQ E9 (GE Healthcare, Milwaukee, WI). A new ultrasound transducer together with a navigation system and dynamic positioning system (volume navigation system: “VNav”) make it possible to combine current ultrasonic images with uploaded CT or MRI data\[65\].

Fusion images combining conventional US and arterial phase contrast-enhanced CT can detect hypervascular HCC lesions easily (Figures 1A and 8). On the contrary, fusion images combining conventional US and hepatobiliary phase contrast-enhanced MRI with
gadolinium-ethoxybenzyl-diethylenetriamine (Gd-EOB-DTPA; Primovist®, Bayer Schering Pharma AG, Berlin, Germany) obtained at 20 min after injection can exhibit small hypervascular (typical) HCC lesions or non-hypervascular (atypical) HCC lesions as hypo-intense areas (Figure 9). Recently, Kim et al. reported that hepatobiliary phase contrast-enhanced MRI with Gd-EOB-DTPA obtained at 20 min after injection can detect smaller HCC lesions better than contrast-enhanced CT.

In our experience, hepatobiliary phase contrast-enhanced MRI with Gd-EOB-DTPA can detect small non-hypervascular (atypical) HCC lesions not detectable by contrast-enhanced CT (Figure 10). Hepatobiliary phase contrast-enhanced MRI with Gd-EOB-DTPA is sensitive for detecting atypical HCC lesions. These lesions were seen as hypo-intense areas in this phase of contrast-enhanced MRI with Gd-EOB-DTPA. In cases of advanced cirrhosis with repeated treatment using various forms of ablation therapy, conventional US detects many hypo- or hyper-echoic areas in the liver parenchyma, resulting in difficulties recognizing targeted small HCC lesions. However, fusion images combining hepatobiliary phase contrast-enhanced MRI with Gd-EOB-DTPA and conventional US can be used to recognize such atypical lesions.
HCC lesions not clearly detected by contrast-enhanced CT. Hepatobiliary phase contrast-enhanced MRI with Gd-EOB-DTPA as the reference standard can detect atypical HCC lesions which appear as hypo- or hyper-echoic lesions on conventional US (unpublished data).

CONCLUSION
Sonazoid, a lipid-stabilized suspension of perfluorobutane gas microbubbles, has been approved for clinical use in Japan only since January 2007. Like Levovist, contrast-enhanced US with Sonazoid has two phases of contrast enhancement: vascular and late (parenchyma-specific phase, delayed phase). The high stability of Sonazoid has overcome the deficiency of Levovist, and microbubbles of Sonazoid have the advantage of long-lasting imaging and strong contrast effects allowing detailed observations.

Contrast-enhanced US and contrast-enhanced 3D US...
using Sonazoid are useful in making the diagnosis and assisting with treatment of hepatic, especially HCC, lesions. Moreover, fusion images combining US with CT or MRI allows correct selection of small HCC lesions which appear as hypo- or hyper-echoic lesions on the US.

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