INTRODUCTION

The reported incidence rate of esophageal varices in patients with liver cirrhosis is 30–60%, depending on the severity of portal hypertension (1, 2). Esophageal variceal hemorrhage is the second most common cause of death in patients with cirrhosis (3), with a 20–35% mortality rate (4-6); it develops at a 10–30% rate in patients with varices (7). Esophageal varices are mainly supplied by an enlarged left gastric vein (LGV) originating from the portal vein (PV).
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or splenic vein (SV), which runs to the esophagogastric junction along the lesser curvature of the stomach.

As PV pressure increases, LGV flow in the PV system transforms to a portal systemic shunt by changing the flow direction from normal (toward the liver: hepatopetal) to reversed (away from the liver: hepatofugal) before exceeding the normal LGV diameter (8), which causes gradual formation of esophageal varices. Therefore, early detection of the change in the LGV flow direction is clinically important to predict esophageal varix development. The relationship between LGV diameter and esophageal varices for ultrasonography (US) and magnetic resonance imaging (MRI) has been previously evaluated (9, 10); however, the LGV flow direction has not been evaluated.

The time-spatial labeling inversion pulse (Time-SLIP) method of MRI is a form of spin labeling, providing quantitative and selective inflow information by placing the inversion pulse before data acquisition and background suppression (11). A Time-SLIP can be used for the selective suppression of the blood signal when placed on a vessel of no interest. More than one Time-SLIP can be placed arbitrarily, independent of the imaging area; this Time-SLIP method enables selective visualization of the vessel of interest. Therefore, it can demonstrate the change in the LGV flow direction. The method of placement of Time-SLIP for visualizing selective PV flow supplied by the superior mesenteric vein (SMV) and the inferior mesenteric vein (IMV) or SV has been reported (12, 13). The source of hepatofugal LGV flow resembles the source of PV flow. The improved methods could facilitate relatively easy imaging of the hepatofugal LGV flow. However, the source of hepatopetal LGV flow is obscure, and how to place Time-SLIP for the selective visualization of hepatopetal LGV is unknown.

Herein, non-contrast-enhanced MR angiography (MRA) with respiratory-triggered three-dimensional (3D) balanced steady-state free-precession (bSSFP) sequence with Time-SLIPs for imaging was used (14). We aimed to selectively visualize hepatopetal LGV by non-contrast-enhanced MRA in healthy subjects and determine the optimal visualization protocol. In addition, we compared the accuracies of the methods of non-contrast-enhanced MRA for the LGV detection using Doppler US as the gold standard.

MATERIALS AND METHODS

Subjects

This study was approved by the Institutional Review Board and conducted in accordance with the ethical standards of the World Medical Association (Declaration of Helsinki). Thirty-one healthy adult subjects (20 males and 11 females; average age, 34.6 years; range, 25–52 years) were included in the study. None of the subjects had viral hepatitis and none were excessive drinkers. Written informed consent was obtained from all subjects before MRI. All subjects underwent US after MRI examinations. The MRI and US examinations were performed after the subjects fasted for at least 4 hours.

MRI

All examinations were conducted with the subject in the supine position using a 3T MRI system (Vantage, Toshiba Medical Systems, Otawara, Japan), equipped with a pair of phased array coils that had 16 channel outputs, placed at the front and back of the abdomen.

First, to locate the heart, stomach, spleen, liver, SMV, PV and SV, coronal 2D single-shot fast spin echo images were acquired (repetition time [TR]/echo time [TE] = 17910 ms/80 ms, flip angle [FA] = 90, field-of-view [FOV] = 400 × 350 mm², matrix size = 288 × 320, number of slices = 22–30, slice thickness = 4 mm, gap = 1 mm, acceleration factor = 2, acquisition time of 18 × 2 seconds during two breath-holds at the beginning of expiration) for Time-SLIP placement.

For non-contrast-enhanced MRA, 3D bSSFP with respiratory triggering was acquired in the coronal plane. Detailed tagged and untagged regions using Time-SLIP are shown in Figure 1. Respiratory triggering with an external belt sensing the respiratory cycle was conducted at the beginning of expiration using a bellows wrapped around the abdomen to reduce motion artifacts. All subjects were instructed to breathe calmly and regularly by listening to a recorded voice (“inhale … exhale”). The short tau inversion recovery method (TI = 220 ms) was used for fat suppression. A bSSFP scan was conducted with the following parameters: TR/TE/FA = 4.8 ms/2.4 ms/90° and receiver bandwidth = 781 Hz/pixel. TR and TE were the shortest, and an FA of 90° was the maximal value to clear the specific absorption rate limitation. The phase-encoding order was centric for both acquisition methods. Parameters related to spatial resolution were the same among sequences as follows: FOV = 330 × 200 mm², matrix size = 256 × 256, slice thickness = 2 mm, number of slices = 38–50 without a gap, number of acquisitions = 1. Parallel imaging was applied in the phase direction with a factor of two, resulting in the collection of 128 phase-encoding lines per respiratory-trigger in a centric
order in both acquisitions. The actual spatial resolution was $1.3 \times 0.8 \times 2 \text{ mm}^3$, and final images were reconstructed into an apparent spatial resolution of $0.6 \times 0.4 \times 1 \text{ mm}^3$ with zero filling.

**Application of the Time-SLIP**

The LGV usually runs from the esophagogastric junction to the portal-SV junction or its vicinity. Two methods for LGV visualization with hepatopetal flow were as follows (Fig. 1). In the first method A, one Time-SLIP was placed on the whole abdomen to suppress signals using $\text{TI} = 1513 \text{ ms}$. The other Time-SLIP was placed on the spatial area from the gastric fornix to the gastric upper body to recover signals of the area, which was considered the source of LGV.

**Fig. 1. Application of Time-SLIP.**

A. Method A: one Time-SLIP was placed on whole abdomen to suppress signals. Other Time-SLIP was placed on spatial area from gastric fornix to upper gastric body to recover signals of area, which was considered source of LGV. B. Method B: nonselective inversion recovery pulse inverts all magnetization in region. One Time-SLIP was placed on center of esophagogastric junction in axial plane to recover signals of area, which was considered source of LGV. C. Method C: nonselective inversion recovery pulse inverts all magnetization in region. One Time-SLIP was placed on lower abdomen to recover inflow signal from mesenteric veins to portal vein. D. Method D: nonselective inversion recovery pulse inverts all magnetization in region. One Time-SLIP was placed on spatial area containing lesser curvature of stomach, excluding gastric fornix to gastric upper body. LGV = left gastric vein, Time-SLIP = time-spatial labeling inversion pulse.
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the LGV. The Time-SLIP thickness was 80 mm, and TI was 1500 ms. In the second method B, nonselective inversion recovery pulse inverts all magnetization in the region using TI = 1513 ms. One Time-SLIP was placed on the center of the esophagogastric junction in the axial plane to recover signals of the area, which was considered as the LGV source. The Time-SLIP thickness was 30 mm, and TI was 1500 ms.

When the LGV flow is hepatofugal, it is supplied by the SMV and IMV flow and/or SV flow. First, in the method for visualizing the LGV with hepatofugal flow supplied by SMV and IMV (method C), nonselective inversion recovery pulse inverts all magnetization in the region using TI = 1513 ms. One Time-SLIP was placed on the lower abdomen to recover the inflow signal from the mesenteric veins to PV. The Time-SLIP thickness was 150 mm, and TI was 1500 ms. Second, it is necessary to place a Time-SLIP at the spatial area containing the spleen to visualize the LGV with hepatofugal flow supplied from SV (12). However, when the set tagged region contained the spleen, it inevitably included part of the spatial area from the gastric fornix to the gastric upper body, which is a hepatopetal LGV flow source. Therefore, we tried to visualize veins that were distributed around the gastric fornix, cardia, and esophagus derived from the LGV with hepatofugal flow by placing the tagged region on the SV and LGV away from the gastric fornix to the gastric upper body (method D). As the method for visualizing those veins, nonselective inversion recovery pulse inverts all magnetization in the region using TI = 1513 ms. One Time-SLIP was placed on the spatial area containing the lesser curvature of the stomach, excluding the gastric fornix to the gastric upper body. TI was 1500 ms.

US Examination

All measurements were performed using an ultrasound system (Aplio™ XG, Toshiba Medical Systems) by the same physician to avoid interobserver errors. A convex transducer with a center frequency of 3.75 MHz (PVT-375BT, Toshiba Medical Systems) extended operating frequency range

![Fig. 2. Definition of LGV with hepatopetal flow on non-contrast-enhanced MRA. A. Method A: LGV is well visualized (arrow). B. Method B: LGV is also well visualized (arrow). C. Method C: LGV is not visualized. D. Method D: veins distributed around fornix, cardia, and esophagus are not visualized. MRA = MR angiography](image-url)
was used. The LGV generally accompanies the left gastric artery (LGA). The vessel accompanying the LGA, which was connected to the PV or SV, was considered as the LGA. Color Doppler US was performed using the color map with a standardized Doppler protocol to detect blood flow direction. The Doppler flow velocity range was set in the maximum range of 8–15 cm/s to detect the LGV flow signal because the blood flow velocity is slow (15). The LGV flow direction was measured during short time breath-holding.

**Definition of LGV with Hepatopetal Flow on Non-Contrast-Enhanced MRA**

Images using maximum intensity projection (MIP) and source images were evaluated to define the LGV with hepatopetal flow (Fig. 2). We estimated whether the vessel visualized by non-contrast-enhanced MRA was the LGV using anatomical information of the bSSFP without the Time-SLIP. The “visualized LGV”, was defined as the LGV identified in the suppressed signal area on source images; and “clearly visualized LGV”, as the LGV identified in the suppressed signal area on both MIP and source images. Subsequently, we evaluated the vein with method C or method D. Method D was used for veins around the fornix and cardia. When the LGV was visualized with method A or B, and not visualized with method C or D, the visualized vessel was defined as the LGV with hepatopetal flow. Two radiologists evaluated the images, and reached a consensus. All assessments were performed using a free available DICOM viewer (YAKAMI Software, Kyoto, Japan).

**Correlation between Non-Contrast-Enhanced MRA and US**

To exclude a possibility that the vessel selectively visualized in method A or B was not even partly in accordance with true LGV, we estimated whether the correlation was in agreement with the running course, flow direction of the LGV between non-contrast-enhanced MRA and US, on the basis of US findings.

**RESULTS**

The image acquisition was successfully conducted in all subjects. The scan time for each method ranged from about 3 to 5 minutes. Including preparations, the total scan time was approximately 25–30 minutes.

**Detection of LGV on Non-Contrast-Enhanced MRA**

Left gastric veins on method A and B were visualized in 30 (97%) and 24 (77%) (Fig. 3), and clearly imaged in 21 (68%) and 17 (55%) of 31 subjects, respectively. For only one subject in whom the LGV was not visualized by method A, the LGV was also not visualized by method B.

**LGV Visualization with Hepatofugal Flow**

There were no visualized LGVs and veins distributed around the fornix, cardia, and esophagus in methods C and D.

**Detection of Blood Flow and Its Direction on US**

The LGV was detected in 23 (74%) of 31 subjects. The LGV flow direction was hepatopetal in 22 of 23 subjects. In one subject, the LGV could be detected, but the LGV flow direction could not be determined.

**Correlation between Non-Contrast-Enhanced MRA and US**

The LGV was visualized in all 22 subjects by method A, and the LGV with hepatopetal flow was detected by US. In all the subjects, the running course of LGV was similar.
in method A and US. In one subject, The LGV was not visualized by method A and US was unable to confirm the flow direction (Fig. 4).

**DISCUSSION**

The ability of non-contrast-enhanced MRA has been demonstrated with combined usage of a respiratory triggered 3D bSSFP sequence and Time-SLIP (16, 17). Using this technique, flow direction of a particular vessel could be visualized by placing the Time-SLIP appropriately. However, there have been few reports of vessel flow direction using the technique only regarding PV (12, 13).

Herein, two different methods were used to depict the LGV with hepatopetal flow using non-contrast-enhanced MRA. Consequently, the rate of visualization of the LGV with hepatopetal flow by method A was higher than those by method B. Visualization of the vessels differs with slight differences in the range of the tagged region (18). A bright signal flow for visualizing the LGV with hepatopetal flow by method B was not considered as sufficient, because the tagged region containing the source area of the LGV flow by method B was smaller than that by method A (18). The use of a wider tagged region on method B than the current setting could improve the visualized rate of the LGV with hepatopetal flow, because this increases bright signal flow into the LGV. Alternatively, The LGV with hepatopetal flow may not be visualized, because the wider tagged region includes the LGV. Therefore, this study could not expand the tagged region on method B. Further studies in which the tagged region is set only from the gastric fornix to the gastric upper body are required to resolve this problem.

In one subject, the LGV could not be detected by method A; and method C and D did not confirm hepatofugal flow. In that subject, the LGV was detected on US, but the LGV flow direction could not be determined using Doppler US. The results suggested that LGV did not show hepatofugal flow but stagnant flow. The subject was considered to have a healthy liver and clinically normal portal pressure. In general, it is considered that the LGV flow is hepatopetal and rarely stagnant in healthy persons with normal portal pressure. Our results suggested that the LGV flow direction can be determined by method A with C and D, even when the LGV is not visualized by method A because of stagnant or hepatofugal flow.

The LGV detection rate by US was lower than that by non-contrast-enhanced MRA using both methods (A and B), although the LGV detection rate by US in this study was almost similar to that of a previous report (15). Occasionally, the LGV may not be visualized with US because of limited acoustic windows compounded by overlying gas-filled bowels and habitus (19, 20). Herein, the LGV could not be visualized by US in 8 of 31 subjects due to these limitations. Advantages of non-contrast-enhanced MRA over US are largely related to its unrestricted field of view and insensitivity to these limitations. Successful the LGV detection with US depends on the operator’s skill. Alternatively, MRI is not influenced by the operator’s skill and patient’s condition. Thus, MRI is more effective than US in determining the LGV flow direction.

This study had several limitations. First, this study included only healthy subjects and the LGV flow direction in patients with portal hypertension has not been assessed. Second, since the sample size of this study was small, the data may have been influenced by selection and verification biases. Third, because US-based identification of the LGV was not possible in some subjects, we were unable to estimate agreement in the running courses between non-
contrast-enhanced MRA and US. Therefore, it is possible that the visualized vessel on non-contrast-enhanced MRA was not in agreement with true LGV in those subjects. However, the possibility of disagreement is very low, because we also evaluated the anatomical relationship between the LGV and surrounding structures using 3D bSSFP without Time-SLIP. Fourth, depicting the hepatofugal LGV flow but not the hepatopetal LGV flow in the blood flow of SV was probable, because a part of SV was inevitably visualized to partially enter the spleen within the tagged regions on method A or B. However, we believe that the SV blood flow did not mix in the LGV, because the distributed veins derived from the LGV with hepatofugal flow were not visualized by method D. Finally, we did not conduct the correction of LGVs flow velocity by US, because an angle between the Doppler beam and the LGV was so large that it was not often possible to measure correct flow velocity of the LGV (21).

In conclusion, The LGV with flow direction can be more selectively visualized by non-contrast-enhanced MRA using bSSFP sequence with Time-SLIPs compared with US. Method A was superior to method B regarding the visualization of the LGV with hepatopetal flow. It is necessary to place the tagged region on the spatial area from the gastric fornix to the gastric upper body for visualizing the LGV with hepatopetal flow (method A). Further studies are needed using non-contrast-enhanced MRA to recognize the LGV with flow direction in a larger population of chronic liver disease patients with various degrees of portal hypertension.

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