Dear Editor,

Liver sinusoidal endothelial cells (LSECs) are the most abundant non-parenchymal and highly specialized fenestrated cells in the liver. LSECs differ from endothelial cells of other capillaries due to the presence of open fenestrations and the absence of a basal lamina (Wisse et al., 1996; Braet and Wisse, 2002; Aird, 2007; Cogger et al., 2008). During the differentiation process, LSECs become fenestrated to facilitate the transfer of circulating nutrients, lipids, and lipoproteins between blood and the space of Disse for normal liver metabolism. Only particles smaller than the fenestrae can reach the hepatocytes or leave the space of Disse (Wisse et al., 1985). The size of fenestration changes upon different conditions to regulate the substrate exchange.

Previous studies reported that the diameter of the sinusoidal fenestrations varies between different species from 107 ± 1.5 nm (mean ± SEM) in the human liver (Wisse et al., 2008) to 141 ± 5.4 nm (mean ± SEM) in the mouse liver (Wisse et al., 2008), which was determined based on transmission electron micrographs of sections (Jacobs et al., 2010). However, considering the small size of fenestration, 100–150 nm in diameter, which is beyond the limit of resolution in light microscopy (Cogger and Couteur, 2009), in early stage, the morphology of fenestrations could only be studied primarily using conventional electron microscopy (Owen et al., 2010; Svistounov et al., 2012), however, with lack of 3D information.

Arsenic is a toxic metalloid and a common contaminant of drinking water. Drinking arsenic-contaminated water increases the risk of cardiovascular disease, lung disease, hepatic disease, and cancer in millions of people worldwide. Even at low levels, arsenic promotes angiogenesis and vascular remodeling in mice (Guha Mazumder, 2003; Mazumder, 2005; Navas-Acien et al., 2005). Pathological vascular remodeling, such as neovascularization, angiogenesis, and morphologic changes in vascular architecture, is a critical process in the development of vascular diseases including atherosclerosis, cardiovascular ischemic diseases, tumor vasculogenesis, and liver fibrosis. Different from angiogenesis of systemic vessels endothelial cells, LSECs angiogenesis, also called capillarization, is a dedifferentiation and maturation process with diagnostic hallmarks of LSEC defenestration and renewed surface expression of PECAM-1 and laminin-1 in response to environmental stress and aging (Straub et al., 2007). Liver capillarization precedes vascular remodeling of other liver vessels, such as hepatic arterioles and PBVP (peribiliary vascular plexus), causing blood flow shunting, vascular channel formation, and eventually liver fibrosis (Straub et al., 2007). Liver capillarization also affects the systemic vasculature and promotes atherogenesis by decreasing liver metabolism of lipids, lipoproteins, and glucose (Straub et al., 2007).

In this study, we investigated how the fenestration morphology of LSECs in mice changes in response to an arsenic stimulation. We utilized focused ion beam scanning electron microscopy (FIB-SEM) technique to reconstruct the 3D morphology (~20 × 20 × 1.5 µm³) of mouse liver sinusoidal with and without the arsenic stimulation. The three-dimensional data generated from FIB-SEM approach enabled us to perform morphologic comparison in 3D space and quantitative measurements and comparisons for the size and number of LSEC fenestrations.

FIB-SEM is a technique to generate high resolution three-dimensional images of biological samples in micrometer scale (Kizilyaprak et al., 2014). Samples are prepared by a similar method to transmission electron microscopy, typically by fixing the sample with aldehyde, staining with heavy metals such as osmium and uranium then embedding in an epoxy resin. The surface of the block of resin-embedded sample is imaged by detection of back-scattered electrons. Following imaging, the focused ion beam is used to trim a thin section (typically less than 30 nm) from the face of the block. After the section is trimmed, the sample block is raised back to the focal plane and imaged again. This sequence of sample imaging, section trimming and block raising can acquire many thousands of images with perfect alignment in an automated fashion and yield a 3D volume data of specimen.

The livers from both normal and As(III)-exposed mice (see Supplemental Materials and Methods) were firstly fixed, embedded in resin and sectioned for conventional transmission
electron microscopy (TEM) examination. The region of the sinusoidal endothelium was selected and imaged. In consistency with the previous observation (Straub et al., 2007), the LSECs in normal mice contain numerous sieve plates with open fenestræ (Fig. 1A). While, the liver sinusoids in the As(III)-exposed mice become defenestrated with continuous endothelium membrane (Fig. 1B). Besides, the density of hepatocyte microvilli in the space of Disse, which protrude through the LSEC fenestræ, becomes significantly increased in the As(III)-exposed liver sinusoids (Fig. 1B).

Since the size of fenestrations (>100 nm) is larger than the thickness of normal TEM sections (~70 nm), the overall morphology of fenestrations and their distribution along the sinusoidal vessel could not be resolved from a single TEM image. To get a more accurate observation of mouse liver sinusoidal fenestrations, we utilized focused ion beam scanning electron microscopy (FIB-SEM) technique to reconstruct the 3D morphology of mouse liver sinusoid in micrometer scale.

The SEM images of both normal and As(III)-exposed mouse liver sinusoid show perfect contrast and similar morphology observation (Fig. 1C and 1D) to that of TEM images (Fig. 1A and 1B). We collected 116 sequential FIB-SEM images (4096 × 4096) with the pixel size of 5 nm and the milling thickness of 15 nm to cover ~20 × 20 × 1.5 μm³ size of specimen for both normal and As(III)-exposed mouse liver sinusoid (Movies S1 and S2), respectively. After image alignment, denoising and reconstruction (see Supplemental Materials and Methods), the 3D morphology of mouse liver sinusoid was obtained (Fig. 1E and 1F), showing the morphology of LSECs and their fenestrations clearly in 3D space. Besides, the hepatocyte microvilli in the space of Disse are also well resolved. The small deformation factors for the rotation, magnification, and stretch corrections (Figs. 2G, 2H, and S1), which are less than 6%, 1%, and 1%, respectively, demonstrate the data collection stability of the FIB-SEM method and confirm the high quality of the 3D reconstructions.

At the current experiments, the pixel size of each FIB-SEM image is about 5 nm in XY direction and the FIB cutting step size is 15 nm in Z direction, and the size of live sinusoid fenestrations is approximately 50–150 nm in diameter, thus the fenestrations can be well resolved after the reconstructions by FIB-SEM method (Fig. 2A and 2B). In normal mice, most fenestrations are irregular holes with a relative large size (Fig. 2A) while, in As(III)-exposed mice, a majority of fenestrations appear as approximately round holes and most of them are relatively smaller than those in normal ones (Fig. 2B).

Based on the 3D morphologies of the liver sinusoids (Fig. 1E and 1F), we were able to count the number of fenestrations and measure their sizes statistically. Since the fenestration is irregular, we utilized a minimal rectangle to cover each fenestration and used the area of that rectangle as the approximate determinant of the fenestration size. From the statistic histogram of the numbers of fenestrations vs. their sizes, we could observe the different distributions of the fenestrations for the normal and As(III)-exposed specimen. For the normal specimen, the size distribution of the fenestrations is relatively uniform, while, for the As(III)-exposed specimen, more fenestrations tend to have a smaller size (Fig. 2C). We further counted the percentages of fenestrations in different sizes. For the normal specimen, the percentage of fenestrations with the size smaller than 15,000 nm² is 50% and 62% for the size smaller than 20,000 nm² (Fig. 2D). However, for the As(III)-exposed specimen, the percentage of fenestrations with the size smaller than 15,000 nm² is increased to 66% and 77% for the size smaller than 20,000 nm² (Fig. 2D). In total, the average area of fenestrations is about 18,000 nm² for the normal specimen and about 10,000 nm² for the As(III)-exposed one. As the result, the above quantitative analysis of the 3D morphologies of the liver sinusoids clearly reveals that the liver sinusoids become defenestrated upon As(III)-exposure.

The thicknesses of the epithelia layers are visually different between normal and As(III)-exposed specimen according to both TEM (Fig. 1A and 1B) and FIB-SEM (Fig. 2E and 2F) analyzes. We further measured the averaged thicknesses of the epithelia layers quantitatively and got the measurements of 200 nm and 270 nm for the normal and As(III)-exposed specimen, respectively (Fig. 2G). As a result, these comparisons suggest that after As(III)-exposure the thickness of the epithelia layers of live sinusoids is increased.

FIB-SEM has been recently emerging as an efficient technique to study the 3D high-resolution ultrastructure of cells and tissues by acquiring thousands of sequential surface images automatically in a perfect alignment. Here, we utilized FIB-SEM technique to achieve 3D reconstruction of...
**A**

**B**

**C**

**D**

**E**

**F**

**G**

**Figure Captions:**

**A** and **B**: Representative images of the epithelial layer in normal and arsenic-treated conditions, showing the changes in the appearance of the layer.

**C**: Bar graph showing the percentage of area of fenestration in different size categories for normal and arsenic-treated conditions.

**D**: Pie charts indicating the percentage distribution of area of fenestration into different size categories for normal and arsenic-treated conditions.

**E**: 3D reconstructions of the epithelial layer in normal and arsenic-treated conditions, highlighting the variations in the layer's structure.

**F**: 3D visualization of the position of the epithelial layer in the X, Y, and Z axes for normal and arsenic-treated conditions, showing the depth and width differences.

**G**: Graph showing the thickness of the epithelial layer in normal and arsenic-treated conditions, with error bars indicating the variability.
mouse liver SEC ultrastructure within 1.5 μm scale and 15 nm step size in Z direction and 5 nm pixel size in XY plane, and observed the ultra-structural changes upon the arsenic stimulation. By comparing the normal specimen with the As (III)-exposed one in three-dimensional space, we found that after As(III)-exposure, the sizes of live sinusoid fenestrations are reduced and the thicknesses of the epithelia layers are increased. Our present results provided a further support to understand the molecular mechanism of LSECs regulation and established a state-of-the-art approach to investigate LSECs in a large scale.

FOOTNOTES
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WL and FS designed research; WL and LW performed sample preparation; GJ and JZ collected EM data; WD performed image processing and 3D reconstruction; FS and WD analyzed data; FS, WL, and WD wrote the paper with the input from other authors.

All the authors declare that they have no conflict of interest. All institutional and national guidelines for the care and use of laboratory animals were followed.

Wenbo Li and Wei Ding have equal contributions to this work.

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