Arsenic-stimulated liver sinusoidal capillarization in mice requires NADPH oxidase–generated superoxide

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Environmental arsenic exposure, through drinking contaminated water, is a significant risk factor for developing vascular diseases and is associated with liver portal hypertension, vascular shunting, and portal fibrosis through unknown mechanisms. We found that the addition of low doses of arsenite to the drinking water of mice resulted in marked pathologic remodeling in liver sinusoidal endothelial cells (SECs), including SEC defenestration, capillarization, increased junctional PECAM-1 expression, protein nitration, and decreased liver clearance of modified albumin. Furthermore, the pathologic changes observed after in vivo exposure were recapitulated in isolated mouse SECs exposed to arsenic in culture. To investigate the role of NADPH oxidase–generated ROS in this remodeling, we examined the effect of arsenite in the drinking water of mice deficient for the p47 subunit of the NADPH oxidase and found that knockout mice were protected from arsenite-induced capillarization and protein nitration. Furthermore, ex vivo arsenic exposure increased SEC superoxide generation, and this effect was inhibited by addition of a Nox2 inhibitor and quenched by the cell-permeant superoxide scavenger. In addition, inhibiting either oxidant generation or Rac1-GTPase blocked ex vivo arsenic-stimulated SEC differentiation and dysfunction. Our data indicate that a Nox2-based oxidase is required for SEC capillarization and that it may play a central role in vessel remodeling following environmentally relevant arsenic exposures.

Introduction

Arsenic is a toxic metalloid that constitutes 0.0001% of the Earth’s crust and is a common contaminant of drinking water. Drinking arsenic-contaminated water increases risk of cardiovascular disease, lung disease, hepatic disease, and cancer in millions of people worldwide. The World Health Organization and the United States Environmental Protection Agency have set the drinking water standard for arsenic at 10 parts per billion (ppb) based on cancer risk. However, even at this low level, arsenic may increase the occurrence of cardiovascular disease in humans (1–3) and promotes angiogenesis and vascular remodeling in mice (4, 5). Epidemiological studies demonstrate that arsenic increases risk for many vascular pathologies, including peripheral vascular disease, ischemic heart disease, acute myocardial infarction, atherosclerosis, and hypertension (2, 6–8). In addition, vascular remodeling and shunting are commonly observed in arsenic-related liver diseases, such as portal hypertension and noncirrhotic liver fibrosis (9). The pathologic mechanisms that contribute to angiogenesis and vascular remodeling in response to arsenic exposures remain unresolved.

Liver sinusoidal endothelial cells (SECs) are unique endothelial cells in both their architecture and function. The sinusoids are the exchange vessels of the liver, and the SECs are distinguished by extensive fenestrations organized into sieve plates, a lack of a basement membrane, and low junctional expression of PECAM-1 (also known as CD31). The SEC architecture, including open fenestrations and weak junctional association between cells, provides a dynamic filtration system with low perfusion pressure that enables nutrients and macromolecular waste to pass freely to hepatocytes for efficient metabolism. The highly active, clathrin-mediated scavenging system in the SEC contributes significantly to endocytic clearance of waste from blood, including effective removal of pathogenic acylated or glycosylated proteins (10, 11). In addition, the SECs are antigen-presenting cells, and interactions with SECs and underlying hepatocyte microvilli may be important for naive T cell activation (12). Loss of these important SEC functions as a result of age or environmental stresses has been proposed to contribute to risk for systemic vascular diseases as well as hepatic pathogenesis (11, 13, 14).

The maintenance of the SEC phenotype is a critical, but poorly understood, process that requires both autocrine and paracrine cell signaling. Results of recent studies indicate that SEC fenestrations are maintained by constitutive VEGF-stimulated NO generation in the SECs and surrounding cells (15). SEC NO generation, in turn, promotes quiescence of surrounding stellate cells that are profibrotic when activated (16). In response to ethanol (17), oxidizing chemicals (18), surfactants (19), and aging (13), SECs dedifferentiate into a more regular endothelium in a process termed capillarization or pseudocapillarization. The hallmarks of capillarization are SEC defenestration, development of a laminin-rich basement membrane, and junctional expression of PECAM. Sinusoidal stellate cells are also induced to overexpress a laminin and collagen matrix that contributes to fibrosis (16). Capillarization precedes the development of many liver diseases, including both portal hypertension and liver fibrosis. In addition, loss of

Nonstandard abbreviations used: ppb, part(s) per billion; SEC, sinusoidal endothelial cell; tempol, 4-hydroxy-2,2,6,6-tetramethylpiperidine-1-oxyl.

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lipid metabolism or removal of acylated or glycolated proteins following capillarization may enhance atherosclerosis, insulin resistance, and possibly metabolic disease (13, 20, 21). Recently, environmental arsenic exposures (250 ppb) were shown to promote progressive SEC capillarization in intact mice through an unresolved mechanism (22, 23).

Superoxide and subsequent ROS generation by Nox-containing NADPH oxidase enzyme complexes in endothelial and smooth muscle cells are fundamental to neovascularization, angiogenesis, and vessel remodeling caused by a variety of endogenous and exogenous factors (24, 25). Arsenic stimulates Nox-based oxidase activity in cultured large vessel vascular cells (26, 27) through mobilization and activation of Rac1-GTPase (26) and increases Rac1 association with SEC membranes during capillarization (22). However, there are no reports of the functional role of this mobilization in capillarization, nor are there reports of a functional role for Rac1-stimulated SEC NADPH oxidase in capillarization in general. In the present study, we investigated the hypothesis that arsenic stimulates SEC Rac1 and oxidase activity to close fenestrations, promote capillarization, and disrupt the physiological function of scavenging modified proteins. The results from the in vivo experiments are the first to our knowledge to demonstrate that low-level arsenic exposure stimulates dose-dependent SEC capillarization through activation of p47phox-containing NADPH oxidase enzyme complexes. Ex vivo studies in primary SECs indicated that capillarization was a direct effect of arsenic on the SECs that required Rac1. In addition to identifying a mechanism for the vascular effects of arsenic, our results revealed an essential role for Nox2-based oxidase in the pathogenesis of capillarization.

**Results**

*Arsenic stimulated dose-dependent capillarization and PECAM-1 expression in vivo and ex vivo.* Previous studies demonstrated that 250 ppb arsenite in drinking water resulted in progressive SEC defenestration and capillarization that was maximal within 2 wk (23). The threshold for these pathologic changes was below the current maximal containment level of 10 ppb, and loss of porosity was dose dependent between 10 and 100 ppb (Figure 1A). The open dilated fenestrations and numerous sieve plates in control mice were lost, and the arsenite-exposed SECs formed tighter junctions. The loss in porosity was caused by a decrease in both number and size of fenestrations, and, as we have previously reported (22), there was no difference in the degree of arsenic effects between zones 1 and 3 of the sinusoidal beds. Capillarization was confirmed in confocal images of sections stained with anti–PECAM-1, demonstrating a reciprocal dose-response relationship for arsenic-stimulated junctional PECAM-1 expression relative to porosity (Figure 2A). As in previous studies (22, 23), there were no signs of SEC injury or cell death, nor were there any signs of injury in surrounding stellate cells or hepatocytes.

Ex vivo exposure of primary cultured SECs to 1–5 μM arsenite for 8 h recapitulated the in vivo observations, with arsenic causing defenestration and disorganized, consolidated sieve plates (Figure 1B). As seen in vivo, arsenite caused a concentration-dependent increase in junctional PECAM-1 expression as the sieve plates were lost (Figure 2B). These results demonstrate that direct SEC effects may account for arsenite-stimulated capillarization.

*Arsenic decreased SEC scavenger receptor function in vivo.* A major physiological role of the SECs is scavenging of small macromolecules, such as chylomicron and derivatized plasma proteins (11).
Comparison of liver clearance of biotinylated or FITC-labeled BSA with succinylated BSA was used to examine whether exposure to arsenic functionally impaired SECs. Only the labeled succinylated BSA was taken up in the liver sinusoids relative to labeled native BSA (Figure 3A). Arsenic decreased clearance of biotinylated succinylated BSA by 60% relative to untreated control mice (Figure 3B). A similar decrease was observed in isolated SECs exposed to arsenic ex vivo (Figure 3C). Thus, arsenic-stimulated capillarization was associated with functional loss of physiological scavenging of modified proteins by the SEC.

*p47phox knockout SECs are protected from arsenic-stimulated capillarization.* Oxidants and compounds that generate oxidants promote capillarization (13, 18). Because arsenic stimulates NADPH oxidase–derived ROS formation in large vessel endothelial cells (26), we tested whether a similar mechanism mediates arsenic effects in SECs. Mice deficient in the canonical Nox2-based oxidase subunit p47phox were exposed for 2 wk to 100 ppb arsenic in their drinking water to test the relevance of this oxidase to capillarization in vivo. Controls for these exposures included wild-type and arsenite-exposed mice that were maintained on the same antibiotic and housed on the same bedding as the immunocompromised p47phox knockouts. We found that sinusoids in the knockout mice did not capillarize in response to arsenic (Figure 4, A–C). In addition, ex vivo addition of arsenic failed to decrease porosity in SECs isolated from these knockout mice (Figure 4D). In contrast, addition of H$_2$O$_2$ bypassed the lack of oxidase activity in these cells and reduced porosity by 4-fold. The knockout mice were protected from other endpoints of capillarization, such as forming a basement membrane and increased hepatocyte microvilli in response to arsenic (Figure 4C). Finally, arsenic-increased nitration of sinusoidal proteins, an indicator of peroxynitrite formation, was absent in the exposed p47phox knockout mice (Figure 5). These results demonstrate that superoxide and H$_2$O$_2$ generated by p47phox-containing NADPH oxidase contributed significantly to capillarization.

SEC Nox2 oxidase activity is necessary and sufficient to mediate arsenic-induced capillarization. Primary SECs in culture were exposed to arsenite ex vivo to address whether SEC NADPH oxidase activity was sufficient to mediate the effects of arsenic on SEC function and capillarization. First, SECs loaded with superoxide-sensitive dihydroethidium increased their fluorescence, as measured by hydroethidium formation, within 30 min of arsenic exposure (Figure 6). This increase followed the same concentration dependence as decreased porosity (Figure 1) and increased junctional PECAM-1 expression (Figure 2). Preincubation of the cells with the superoxide scavenger 4-hydroxy-2,2,6,6-tetramethylpiperidine 1-oxyl (tempol) prevented arsenite-increased dihydroethidium fluorescence (Figure 6), prevented loss of scavenging function (Figure 3C), and both decreased porosity and increased junctional PECAM-1 expression (Figure 7). Likewise, preincubation...
of the cells with gp91ds-tat peptide (an inhibitor designed to disrupt Nox2-based oxidase protein complexes; ref. 28), but not the scrambled-tat control peptide, prevented the arsenite-stimulated superoxide generation and morphologic change (Figures 6 and 7). These results from isolated SECs, combined with the results from the p47phox knockout mice in Figures 4 and 5, suggest that the canonical p47phox containing Nox2-based oxidase, and its attendant superoxide generation, is necessary and sufficient to mediate arsenite-stimulated SEC capillarization and dysfunction.

Rac1-GTPase activity in arsenite-stimulated capillarization. Rac1 is an essential component of the Nox2-based oxidase required for arsenite-stimulated large-vessel NADPH oxidase activity (26). Arsenic mobilized Rac1 to the SEC plasma membrane during capillarization in vivo (22), but the function of this mobilization was unknown. Therefore, the role of Rac1 in arsenic-induced capillarization was tested by examining the effects of a chemical inhibitor of Rac1, NSC23766 (29), on arsenic-stimulated SEC capillarization and dysfunction. Overnight preincubation of the SECs with NSC23766 prevented arsenic-reduced porosity and arsenite-induced functional process in SEC capillarization. Because NSC23766 inhibits Rac1-GTPase activity but not membrane mobilization (29), these data suggest that the initial step in arsenic-stimulated oxidant signaling in SEC is upstream of Rac1 activation.

Discussion
Pathological vascular remodeling, such as neovascularization, angiogenesis, and morphologic changes in vascular architecture, in response to environmental insults is a critical process for the development of many vascular diseases, including atherosclerosis, cardiovascular ischemic diseases, tumor vasculogenesis, and liver fibrosis. The current studies focused on a unique form of vascular remodeling, sinusoidal capillarization, that is known to pathologically compromise the SECs and the effective exchange of nutrients and waste in the liver. Capillarization precedes remodeling and shunting in the progression of liver fibrosis (30, 31) and allows induction of profibrotic collagen expression in sinusoidal stellate cells (16). Decreased liver metabolism of lipids, glucose, and other nutrients promotes atherogenesis after capillarization in response to several environmental stressors and aging (13, 18). We previously demonstrated that arsenic stimulates SEC capillarization and remodeling of the liver vasculature (22, 23). Our present results provide support for an arsenic-stimulated mechanism that requires Nox2-based oxidase–generated oxidants to defenestrate and capillarize SECs as well as impair their physiological functions. In addition, this study is the first to our knowledge to demonstrate a functional consequence of this oxidase activity in SEC capillarization, a process that has been proposed to be oxidant mediated (18, 21). The potential public health implications of these findings are that they suggest that stimulation of Nox2-based oxidase in the SECs, and perhaps other vascular beds, is a fundamental mechanism for the etiology of arsenic-promoted hepatic and systemic vascular diseases.

There have been few reports of a direct physiologic or pathophysiologic role of SEC oxidase enzymes. Several studies demonstrated that oxidants, surfactants, and oxidizing conditions in aging promote capillarization to alter circulating lipid and lipoprotein profiles (13, 18, 19). However, the mechanisms for SEC oxidant generation other than direct chemical action were not investigated in these studies. In rats, high-level ethanol perfusions stimulate SEC and Kupffer cell superoxide production that is inhibited by the nonselective flavoprotein inhibitor diphenylenediiodonium chloride (17), and chronic coexposure to drinking water ethanol and high levels of injected arsenic (i.e., 100 ppm) promote oxidant-dependent vascular channeling and fibrosis (32). A proposed mechanism for the observed liver pathology was depletion of antioxidant glutathione (32), which was unlikely to occur during the shorter and lower-level arsenic exposure in the current study.

Dorman et al. proposed that either SEC or neutrophil Nox2 participates in sinusoidal injury secondary to hind-limb ischemia, since gp91ds-tat protected the hepatic parenchymal and endothelial cells from oxidative injury in this in vivo model (33). This model produced significant inflammation, and much of the parenchymal injury was mediated by activated neutrophils (33). In contrast, we...
have demonstrated that there was no increase in inflammatory cells in the livers of mice exposed for 2 wk to the higher arsenic dose of 250 ppb (23). Since arsenic-induced defenestration is fully manifested within 2 wk, these data suggest that arsenic directly affects the SECs or surrounding cells to induce capillarization rather than through a mechanism requiring leukocyte oxidase activity. In the current study, we used 2 independent strategies to test whether superoxide in response to arsenic exposure and that this superoxide was the central mediator of the oxidant response to arsenic. In contrast to the effects of the scrambled-tat peptide, in which the active targeting sequence in gp91ds-tat is rearranged, gp91ds-tat was completely effective in protecting the SEC line resulting in a 6-fold induction of Nox1, which increased NADPH oxidase activity and decreased arsenic-induced superoxide levels in wild-type cells. This data provide little support for a role of the other prevalent endothelial-expressed isozone, Nox4-based oxidase, in the response to arsenic, because this isoform does not require p47phox and thus should not be affected by either the p47phox knockout or gp91ds-tat (28, 34). There is a single in vitro report indicating that overexpression of constitutively active type 1 VEGF receptor in an immortalized SEC line resulted in a 6-fold induction of Nox1, which increased apoptosis (35). However, the cell line was not well characterized as retaining SEC phenotype, and Nox1 is generally expressed at low levels in endothelial cells (25, 28, 34). The data are consistent with SEC Nox2-based oxidase directly contributing to endothelial dysfunction and suggest that this oxidase is the dominant isozone for arsenic-stimulated superoxide generation and capillarization.

The data provide little support for a role of the other prevalent endothelial-expressed isozone, Nox4-based oxidase, in the response to arsenic, because this isoform does not require p47phox and thus should not be affected by either the p47phox knockout or gp91ds-tat (28, 34). There is a single in vitro report indicating that overexpression of constitutively active type 1 VEGF receptor in an immortalized SEC line resulted in a 6-fold induction of Nox1, which increased apoptosis (35). However, the cell line was not well characterized as retaining SEC phenotype, and Nox1 is generally expressed at low levels in endothelial cells (25, 28, 34). The in vivo data in the current study do not rule out contributions of the Nox2 oxidase in other liver cells, because stellate and Kupffer cells, as well as hepatocytes, express the Nox2 anchor (36, 37). Nonetheless, the ex vivo data confirm the
necessary and sufficient role for SEC Nox2-based oxidase in both capillarization and functional loss, and additional oxidants from other liver cells would only add to this pathologic change in vivo.

Arsenic-stimulated superoxide generation may promote defenestration and capillarization by ROS-mediated signaling or quenching NO that is required for maintaining the fenestrated SEC phenotype. The data in Figure 4D suggest that H$_2$O$_2$ is effective in promoting SEC defenestration and thus may be the second messenger for these arsenic actions in the SECs. However, fenestrated endothelium is found in tissues constitutively expressing high levels of VEGF and VEGF-stimulated NO (38). DeLeve et al. demonstrated that the SEC phenotype was maintained by paracrine stimulation through hepatocyte- and stellate cell–derived VEGF-stimulated NO production in either the stellate cells or the SECs (15). In reciprocal regulation, the SEC NO suppresses stellate cell activation and collagen expression that facilitates fibrosis (16). Loss of NO contributes to pathogenic angiogenesis, fibrosis, and portal hypertension in the liver (39). Peroxynitrite formed from the reaction of superoxide and NO has been proposed as a pathogenic mediator of liver perfusion defects caused by obesity and insulin resistance (21). Nitrosylation of protein tyrosine residues is a marker of peroxynitrite formation, and the observation that arsenite failed to increase protein nitrosylation in p47$^{phox}$ knockout mice (Figure 5) suggests that stimulation of superoxide generation and peroxynitrite formation is a primary mechanism for arsenic-induced remodeling of the liver vasculature and impaired function in the sinusoids. The data do not discriminate, however, between the significance of superoxide depletion of regulatory NO or peroxynitrite-mediated signaling to arsenic-stimulated SEC capillarization. Nonetheless, the fact that H$_2$O$_2$ is sufficient to reduce porosity suggests that both loss of NO and peroxynitrite formation are secondary mechanisms in arsenic-stimulated capillarization.

While the functional consequences of arsenic-stimulated, oxidant-dependent signaling in SECs are unique to this cell type, the initiation of this signaling may be common to endothelial cells in most vascular beds. We observed that environmental exposure to arsenic stimulates angiogenesis in Matrigel plugs and tumor xenographs in vivo (4, 40) as well as tube formation by human microvascular cells in cultured Matrigel (41). While the isoforms were not specifically identified, arsenic was found to stimulate large-vessel endothelial and smooth muscle cell NADPH oxidases (26, 27) and quench bradykinin-stimulated NO generation in aortic endothelial cells (42) or NO-dependent vasodilation in aortic rings (43). In addition, endothelial peroxynitrite formation appears to contribute to arsenic-promoted atherosclerosis in genetic mouse models (44). Thus, it is likely that the arsenic exposures in the current studies had systemic effects in most vascular beds. We observed that environmental exposure to arsenic stimulates angiogenesis in Matrigel plugs and tumor xenographs in vivo (4, 40) as well as tube formation by human microvascular cells in cultured Matrigel (41). While the isoforms were not specifically identified, arsenic was found to stimulate large-vessel endothelial and smooth muscle cell NADPH oxidases (26, 27) and quench bradykinin-stimulated NO generation in aortic endothelial cells (42) or NO-dependent vasodilation in aortic rings (43). In addition, endothelial peroxynitrite formation appears to contribute to arsenic-promoted atherosclerosis in genetic mouse models (44). Thus, it is likely that the arsenic exposures in the current studies had systemic effects in most vascular beds. However, other pathological endpoints, such as atherosclerotic plaque formation, vessel wall thickening, or peripheral vascular disease, would not be expected to occur in a mouse within the 2-wk exposure period examined in the current studies. Nonetheless, the importance of the SEC- and liver-specific observations is that...
capillarization is a significant and relatively early pathogenic event. Sustained capillarization and loss of SEC scavenging may precede arsenic-promoted systemic disease or enhance deleterious systemic effects of arsenic by impairing clearance of acetylated lipids or modified proteins that are known risk factors for vascular and metabolic disease. Further studies are needed to determine whether the arsenic-induced sinusoidal morphological changes and loss of SEC scavenger function are initial steps in arsenic-induced vascular disease or are limited to pathogenic changes in the liver.

The finding of morphological and functional changes in livers of mice exposed to concentrations of arsenic that promote human diseases other than cancer is notable, because mice are often several orders of magnitude less sensitive to humans to the carcinogenic and lethal effects of arsenic. However, cardiovascular effects may be more sensitive disease endpoints in humans drinking the same levels of arsenic used in the current studies (1, 2, 7), and our present findings suggest that the mouse is an appropriate model for studying the etiology of arsenic-promoted disease. In addition, these studies provide valuable information regarding gene environment interactions. Hsueh et al. suggested that genetic polymorphisms in the NADPH oxidase p22 subunit may contribute to arsenic-related hypertension in Taiwan (8).

In summary, the current data support an essential role for an NADPH oxidase in a form of pathogenic vascular remodeling that can contribute to liver disease as well as to systemic hypertension and atherogenesis. The Nox2-based oxidase in SECs appears to be directly activated by environmental exposures to arsenic, and this activation is central to arsenic-induced capillarization and loss of SEC scavenging function. The apparent requirement for Rac1-GTPase activity (Figure 8) suggests that Rac1 and oxidase activation are downstream of the target of arsenic in the endothelial cells and that peroxynitrite formation may be a pathogenic endpoint of arsenic signaling in SECs. However, future experiments are needed to identify the molecular targets on endothelial cells that initiates arsenic signaling for oxidase activation. Identifying this target may reveal important pathogenic mechanisms promoting SECs and possibly systemic endothelial dysfunction that contribute to the etiology of environmentally derived vascular diseases.

Methods

Animal exposure. Animal experiments were performed in agreement with institutional guidelines for animal safety and welfare at the University of Pittsburgh. Wild-type C57BL/6 Tac mouse exposures were performed as described previously (22, 23). Serial dilutions were used to prepare drinking water with sodium...
arsenate concentrations of 10, 50, 100, or 250 ppb. C57BL/6Ai p47phox knock-
out mice (Taconic Farms) were housed under pathogen-free conditions in auto-
claved microisolator cages with Alpha-dri bedding (Shepherd Specialty Papers)
to reduce footpad irritation and edema. The mice were treated with 380 mg
broad-spectrum sulfamethoxazole-trimethoprim antibiotic per 500 ml
drinking water according to the vendor’s instructions (Taconic Farms).
C57BL/6 mice housed in the same manner and treated with antibiotic served
as controls for the C57BL/6Ai p47phox knockout mice. There were no observ-
able infections during the arsenic exposures or at the time of necropsy. All
water was changed 3 times per week to maintain effective concentrations of
arsenite and antibiotic. Liver perfusions with collagenase, fixative, or modified
lipids and proteins were performed following sodium pentobarbital anesthe-
sia. All other mice were euthanized with CO₂.

SEC isolation and ex vivo culture. Nonparenchymal cells from C57BL/6 mouse livers were obtained from the laboratory of T. Billiar (University of
Pittsburgh, Pittsburgh, Pennsylvania, USA). The cells were isolated by stan-
dard collagenase digestion and liver cell separation, as described previously
(45, 46). Briefly, livers were perfused with buffered collagenase to obtain a
single-cell suspension. Parenchymal cells (hepatocytes) were removed
using low-speed centrifugation at 50 g. The remaining non-parenchymal
cells (SECs, Kupffer cells, and Stellate cells) were placed onto a 2-step 25%
and 50% Percoll gradient and centrifuged at 900 × g to give a single value per
mouse. Ultra-thin 70-nm sections were imaged on a JEM 1210 transmission
electron microscope (JEOL) at 80 kV to evaluate our 2 previous studies of the effects of arsenic on the SEC (22, 23). The percentage of positive pixels in the 3 images was aver-
egaged to give a single value per coverslip. Values were normalized by dividing
the percent thresholded area of PECAM expression by percent thresholded
area of nuclei. Porosity, the percent of open space in sinusoidal vessels or
cultured cells, was measured in representative x7,500 or x10,000 scanning EM
images of vessels and cells, respectively, as described previously (22, 23,
48). Briefly, images from 3-μm-thick liver sections were captured with a JSM-
6330F scanning electron microscope, and images of 10 millibular sinusoids
per mouse were measured for porosity using MetaMorph software (version
7.0, Molecular Devices). Within a defined millibular region of in represen-
tative sinusoids, total sinusoidal area and the open area of individual fenes-
trae (20–500 nm diameter) and gaps (>500 nm diameter) were quantified in
square microns. These open areas were summed, divided by the total areas
of the vessels examined, and multiplied by 100 to give the porosity as per-
cent of open area. The porosities of the 10 separate vessels were averaged
to give a single value per mouse. Ultrathin 70-nm sections were imaged on
a JEM 1210 transmission electron microscope (JEOL) at 80 kV to evaluate
changes in dihydroethidium oxidation normalized to percentage of
nuclear pixels, as described above. To scavenge superoxide, 1 mM tempol
(Sigma-Aldrich) was added for 10 min prior to adding arsenite. To inhibit

Microscopy and quantitative imaging. At necropsy, livers were either excised
and snap frozen in liquid nitrogen or perfusion fixed with glutaraldehyde in
preparation for immunofluorescent analysis of proteins or scanning EM
of tissue ultrastructure. Tissues for immunofluorescence and scanning EM
were prepared and imaged as previously described (22, 23, 47). For quantita-
tive immunofluorescence, 3 representative ×400 images from individual cov-
 erslips were captured by a microscopist blinded to treatments, as described
previously (22, 23). The percentage of positive pixels in the 3 images was aver-
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![Figure 7](https://example.com/figure7.png)

**Figure 7** Arsenite-stimulated defenestrated and junctional PECAM-1 expression is inhibited by tempol and gp91ds-tat peptide. Cultured SECs were preincubated with tempol (A and C) or with gp91ds-tat or scrambled-tat peptide (B and D) and then exposed to 2.5 μM for 8 h. At the end of exposure, groups of cells were analyzed for porosity (A and B), as in Figure 1B, or PECAM-1 expression changes (C and D), as in Figure 2B. Significant differences between groups were determined by ANOVA followed by Newman-Keuls post test (n = 5 cultures from 3 livers.). ***P < 0.001 versus control; ###P < 0.001 versus arsenite.
Modified albumin synthesis. FITC- or biotin-labeled native or succinylated BSA was prepared as follows. Briefly, a 10-M excess of FITC (Sigma-Aldrich) or biotin-NHS (Pierce), freshly prepared in DMSO, was added into 1 ml of BSA solution (100 mg/ml in 1 M NaHCO3). The reaction was allowed to proceed at 4°C for 1 h. The reaction mixture was then either thoroughly dialyzed against PBS, to obtain labeled native BSA, or reacted with 4 25-mg aliquots of succinyl anhydride at 15-min intervals. The succinylation continued overnight at 4°C, and then the reaction mixtures were purified by dialysis. The degree of succinylation of free amino groups was estimated to be 90% by reacting the remaining NH2 groups with 2,4,6-trinitrobenzene sulfonic acid (Pierce).

Modified albumin uptake. To demonstrate SEC-specific uptake, 150 mg/kg of FITC-labeled BSA or succinylated BSA was injected into the mouse tail vein in 200 μl saline. After 10 min, livers were excised, fixed in 4% paraformaldehyde, and sectioned for confocal microscopic analysis. Quantitative uptake of succinylated BSA was measured in control and arsenic-exposed mice by mixing a mixture of FITC-labeled acetylated LDL (BTI Technologies) and biotin-labeled succinylated BSA (150 mg/ml saline) into the vena cava over 3 min. Livers were then excised, snap frozen in liquid N2, and sectioned. To control for differential perfusion of liver lobes, microscopists blinded to treatment selected liver sections with equivalent levels of FITC-labeled acetylated LDL fluorescence. Total protein was extracted from these sections, and equal amounts were assayed for biotinylated albumin by immunoblotting. Likewise, cultured control SECs or SECs exposed to arsenic for 24 h were incubated with 20 μg/ml biotin-labeled succinylated BSA for 10 min, rinsed 3 times with PBS, and then extracted for total protein to determine modified albumin uptake. Biotin in the 75-kDa BSA protein band was detected on transfer membranes using HRP-conjugated anti-biotin antibody (Cell Signaling) and enhanced chemiluminescence. Densities of detected protein bands were quantified using NIH ImageJ software (version 1.28c; http://rsweb.nih.gov/ij/) and normalized to band densities of immunodetected β-actin in the same sample.

Materials. NSC23766 (EMD) was added to cells overnight to inhibit Rac1 GTPase activity prior to arsenite exposure. Primary antibodies used for immunodetection were rat monoclonal anti-mouse CD31 (BD); monoclonal anti-β-actin (Sigma-Aldrich); and polyclonal anti-nitrotyrosine (Invitrogen). Secondary antibodies included Alexa Fluor 488-conjugated goat anti-rat (Invitrogen), Alexa Fluor 594-conjugated goat anti-mouse (Invitrogen), Alexa Fluor 488-conjugated goat anti-rabbit (Invitrogen), and HRP-conjugated sheep anti-mouse (GE Healthcare). DRAQ5 was used to stain nuclei.

Statistics. Mice and SECs exposed to various dose of arsenic were analyzed using 1-way ANOVA followed by Dunnet’s or Newman-Keuls post test for difference between treatments. All statistical analyses were performed using GraphPad statistical software version 4.0. A P value less than 0.05 was considered significant.

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Figure 8
Rac1 inhibition prevents arsenite-stimulated defenestration and PECAM-1 expression. Cultured SECs were incubated overnight with or without 50 μM of Rac1-GTPase activity inhibitor NSC23766 (iRac1). The cells were then left untreated or exposed to 2.5 μM arsenite for 8 h. Cells were then fixed for scanning EM imaging of porosity (A and B), as in Figure 1B, or immunofluorescence imaging of PECAM-1 and DRAQ5 expression (C), as in Figure 2B. Significant differences between groups were determined by ANOVA followed by Newman-Keuls post test (n = 5 cultures from 3 livers). ***P < 0.001 versus control; ****P < 0.001 versus arsenite. Scale bar: 1 μm.