Arterial Shear Stress Reduces Eph-B4 Expression in Adult Human Veins

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Vein graft adaptation to the arterial environment is characterized by loss of venous identity, with reduced Ephrin type-B receptor 4 (Eph-B4†) expression but without increased Ephrin-B2 expression. We examined changes of vessel identity of human saphenous veins in a flow circuit in which shear stress could be precisely controlled. Medium circulated at arterial or venous magnitudes of laminar shear stress for 24 hours; histologic, protein, and RNA analyses of vein segments were performed. Vein endothelium remained viable and functional, with platelet endothelial cell adhesion molecule (PECAM)-expressing cells on the luminal surface. Venous Eph-B4 expression diminished (p = .002), Ephrin-B2 expression was not induced (p = .268), and expression of osteopontin (p = .002) was increased with exposure to arterial magnitudes of shear stress. Similar changes were not found in veins placed under venous flow or static conditions. These data show that human saphenous veins remain viable during ex vivo application of shear stress in a bioreactor, without loss of the venous endothelium. Arterial magnitudes of shear stress cause loss of venous identity without gain of arterial identity in human veins perfused ex vivo. Shear stress alone, without immunologic or hormonal influence, is capable of inducing changes in vessel identity and, specifically, loss of venous identity.

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†Abbreviations: RNA, ribonucleic acid; PECAM, platelet endothelial cell adhesion molecule; HIC, human investigation committee; TUNEL, terminal deoxynucleotidyl transferase dUTP nick end labeling; H&E, hematoxylin & eosin; SDS-PAGE, sodium dodecyl sulfate-polyacrylamide gel electrophoresis; Cleaved Caspase-3; Cleaved PARP (poly ADP ribose polymerase); ECL, enhanced chemiluminescence; DAPI, 4',6-diamidino-2-phenylindole; RT-PCR, real time polymerase chain reaction; GAPDH, glyceraldehyde 3-phosphate dehydrogenase; EPCs, endothelial progenitor cells.

Keywords: vein graft adaptation, EphB4, Ephrin-B2, osteopontin, shear stress, bioreactor, Saphenous vein

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INTRODUCTION

While the arterial autograft is the gold standard for both cardiac and peripheral vascular bypass, it is not always possible to use arterial conduits, for example in lower extremity vascular reconstruction. As such, the autogenous saphenous vein remains the best conduit for arterial reconstruction in the peripheral system [1]. Vein graft adaptation to the arterial environment, commonly known as “arterialization,” is characterized by vessel wall thickening with deposition of smooth muscle cells and extracellular matrix in all layers of the vessel [2,3]. This remodeling process is thought to be an essential response to the different arterial environment, including increased wall shear stress, stretch force, transmural pressure, and oxygen tension [4-6]. Despite this physiologic adaptation, 30 to 50 percent of vein grafts eventually fail with significant clinical morbidity and mortality for patients [7,8]. The failure of the PREVENT III and IV trials shows that our understanding of the biology of venous remodeling is incomplete and needs additional approaches for this important clinical need [9,10].

The Eph receptors — and their ligands the Ephrins — are differentially expressed on arteries and veins [11]. Ephrin-B2 is a determinant of arterial fate during embryologic development and persists as an arterial marker on adult vessels; alternatively, Eph-B4 is a determinant of venous fate in development and persists as a marker of venous identity on adult veins [11,12]. We have previously reported that Eph-B4 expression is reduced, but Ephrin-B2 expression is not induced, during human and rat vein graft adaptation [13], i.e., venous identity is lost without development of arterial identity.

Previous studies have examined the effects of shear stress on isolated cells, including endothelial cells, progenitor cells, stem cells, or cells in tissue-engineered grafts [14-18]. We have previously shown that bioreactors can control shear stress to alter the thickness of endothelial-lined prosthetic grafts, as well as enhance the culture of tissue-engineered vessels with pulsatile stimulation in vitro [19-21]. However, it is not clear whether these studies are adequate models of saphenous vein segments used by vascular surgeons for bypass in adult humans. We hypothesized that, using a bioreactor that can precisely control the magnitude of shear stress, we can test whether changes in magnitudes of shear stress can induce changes in vessel identity, i.e., changes in expression of Eph-B4 or Ephrin-B2 in adult human saphenous vein segments ex vivo. Therefore, we examined whether a bioreactor, with a flow circuit that reproduces arterial or venous magnitudes of shear stress while minimizing the effects of variations and high magnitudes of pressure, can support short-term viability in an explanted adult human saphenous vein segment and whether changes in vessel identity occur during this period [20].

MATERIALS AND METHODS

Flow Model

Segments of adult human saphenous veins that were surgically harvested but not used for surgical bypass were removed from the operating room (HIC Protocol #9908011041, Yale University Human Investigation Committee). Vein remnants were then transported in Dulbecco’s Modified Eagle Medium (DMEM)-based media (11995-065; Invitrogen, Grand Island, NY) and brought immediately to the laboratory. Veins were inspected for feasibility of incorporation into the bioreactor, and structural parameters were measured. The vein segment was interposed in a reversed fashion to avoid native valve obstruction and secured on each end with silk suture between custom glass cannulae of various internal and external diameters depending on the vein sample and placed within the glass bioreactor (Figure 1).

The flow system consists of a 300+ mL capacity glass chamber with inflow and outflow ports with two equally sized compliance vessels on either side of the bioreactor to equalize pressure in the system and an intake 0.22µm air filter on the distal chamber.
These were all connected with platinum-cured silicone tubing (MasterFlex, Cole-Palmer, Vernon Hills, IL). A digital programmable, peristaltic roller-pump (MasterFlex) was used to push media along the system, which was placed in an incubator (37°C; 21% O₂, 5% CO₂).

Flow rates, pressure, and pressure-drop across the system were verified in the bioreactor with interposition of a flow meter and pressure gauges, with data acquired via PowerLab 26T LTS (ML4856) system and LabChart (MLU60/8) software (AD Instruments, Colorado Springs, CO). The flow meter was interposed directly after the tubing exit from roller pump. Pressure meters were placed before and after the venous segment, allowing measurement of both the upstream pressure as well as the pressure change across the vein (Figure 1). A thin-walled, distensible silicone tube with a 4mm internal diameter was used as a sample for calibration of the Powerlab system. Flow rates for each specimen were estimated to achieve desired wall shear stress (WSS) via the Hagen-Poiseuille formula

\[ \tau_{\text{mean}} = \frac{4\mu Q}{\pi R^3} \] [1]

where \( \tau \) = wall shear stress, \( Q \) = volume flow rate, \( \mu \) = viscosity of fluid, and \( R \) = inner radius of cylindrical tube. The arterial environment was set at a wall shear stress of 20 dynes/cm², and the venous environment was set at a wall shear stress of 3 dynes/cm², independently of pressure [22]. For determination of the flow rate for each environment, a derivation of this formula was used to approximate the shear stress:

\[ \Delta Q = \frac{\pi P d^4}{128 \mu l} \] [2]

where \( P \) = pressure difference, \( d \) = diameter of vein, \( \mu \) = viscosity of fluid, and \( l \) = length of vein. Using a given vein’s radius, a standard length of 10 cm, constant viscosity of 3.8cP, and the measured pressure-drop across the system, the flow rate for the de-
sired WSS was calculated. Vein tautness was adjusted to prevent noticeable vein distention or bowing during various flow rates. Finally, venous segments were applied to the bioreactor in “static” conditions for additional control comparison; to prevent vessel death, media was circulated at a minimal rate (1 mL/min) to maintain intraluminal exposure to the flow media and oxygenation as well as prevention of desiccation that occurs with 0 mL/min.

Endothelial cell basal media (CC-3156, Lonza, Ltd.) with 150mL Fetal Bovine Serum (FBS) (HyClone, SV3001403, Thermo Scientific, Wilmington, DE) per 1000mL media and Penicillin & Streptomycin (P&S) (10,000 U/mL, Gibco, 15140-122) was used for internal circulation. Xanthum Gum (XG; Sigma-Aldrich Co., LLC) was sterilized and added as a thickening agent for the internal media to the desired viscosity of 3.8cP to approximate human blood and was tested with a glass capillary viscometer. Medium was circulated for 24 hours at calculated flow rates. The external media bathing the vein within the bioreactor chamber consisted of a DMEM solution with FBS and P&S. Specimens were removed from the bioreactor, excluding approximately 1 cm of vein from each attachment site, in order to avoid effects from the connections.

Histology
Vein segments were fixed in 30 percent sucrose solution overnight for dehydration and then 4 percent formaldehyde solution. Specimens were subsequently analyzed with H&E, Van Gieson, and TUNEL staining per standard protocols of the Yale Histology Service at the Yale School of Medicine.

Western Blot Analysis
Equal amounts of protein initially isolated from each specimen and controls with a lysis buffer were loaded and run in SDS-PAGE, then probed with antibodies (antibodies for PECAM-1, ClvCasp3, and ClvPARP; Cell Signaling Technology, Inc., Danvers, MA, Catalog # - 3528, 9661, and 9541, respectively). Membrane signals were detected using ECL detection reagent (GE Healthcare, Denville scientific). As a positive control for apoptosis, staurosporine (1µm) was applied to human umbilical endothelial cells (HUVEC) for 5 minutes.

Immunofluorescence Analysis
Specimens were fixed in formaldehyde as above. Unstained and sectioned samples were de-paraffinized and dehydrated with three xylene washes of 5 minutes each, two 100 percent ethanol washes of 10 minutes each, and two 70 percent ethanol washes of 10 minutes each. Antigen unmasking was achieved with sodium citrate buffer. Primary antibody treatment was performed according to the manufacturer’s instructions, and concentrations optimized when needed (antibody for EphB4, ABCAM, Cambridge, MA, Catalog # - ab64820). Secondary detection was performed using Donkey Anti-Rabbit IgG (H&L) Alexa Fluor 568 secondary antibody (A-21206, Invitrogen) and counterstained with DAPI. Images were acquired with an AxioImager A1 (Carl Zeiss, Inc., Thornwood, NY).

Relative quantification of immunofluorescence images was performed (MetaMorph, Molecular Devices, LLC, Sunnyvale, CA). Each sample was compared to static conditions.

RT-PCR Analysis
RNA was isolated from cells or tissue using TRIzol Reagent (Invitrogen), and RNA was cleaned using the RNeasy Mini kit (Qiagen, Germantown, MD). Total RNA quantification quality was measured with a spectrophotometer (Nanodrop, Thermo Scientific). RT was performed using the SuperScript III First-Strand Synthesis Supermix (Invitrogen) according to the manufacturer’s instructions. Real-time quantitative PCR was performed using SYBR Green Supermix (Bio-Rad Laboratories, Hercules, CA) and amplified for 40 cycles using the iQ5 Real-Time PCR Detection system (Bio-Rad Laboratories). Correct target amplification and exclusion of nonspecific amplification was confirmed by 2 percent agarose gel electrophoresis, and primer efficiencies were
determined by melting curve analysis. All samples were normalized by GAPDH amplification. Primers are listed in Table 1.

**Statistical Analysis**

Statistical analysis with paired t-tests was performed with Sigma Plot 11.0 software (Systat Software Inc., San Jose, CA). P < 0.05 was considered significant.

**RESULTS**

**Flow Model and Vein Samples Configured to Analyze Shear Stress Independent of Pressure**

All vein segments were trimmed to as close to 10 cm as possible, and the diameter at each end measured. The flow rate was calculated based on the measured diameter, vein length, constant viscosity, and measured pressure drop across the system according to Equation 2. Calculated examples of flow rates for venous (3 dynes/cm²) and arterial (20 dynes/cm²) shear are listed in Table 2. The mean flow rate was 24.8 mL/min for venous shear conditions, and the mean flow rate was 200.6 mL/min for arterial shear conditions. Although the bioreactor can be adjusted to near physiologic arterial pressures (~100/85mmHg) with post-vein clamping, pressures were held near constant and minimal to prevent confounding on the effects of shear stress.

**Retention of Endothelial Cells without Apoptosis after Venous and Arterial Shear Stress**

The effect of arterial magnitudes of shear stress (20 dynes/cm²) on venous struc-
ture and endothelium was examined (n = 6). General architecture was preserved in veins at baseline and after 24 hours of arterial flow in the bioreactor, with thin elastic laminae present at both time points (Figure 2A). Multiple endothelial cells were easily identifiable on the luminal surface of the veins, at both 0 and 24 hours after exposure to arterial flow (Figure 2B). To confirm that the cells were endothelial cells, immunofluorescence staining for PECAM was performed; PECAM-reactive cells were seen at both 0 and 24 hours after exposure to arterial flow, consistent with endothelial identity (Figure 2C). Similar to the maintained PECAM immunoreactivity between 0 and 24 hours of arterial shear stress, PECAM protein was also detected via Western blot after 24 hours of arterial shear stress (n = 4, Figure 2D). Den- sitometry showed similar PECAM protein amounts after 24 hours of arterial magnitudes of shear stress. These results are consistent with retention of the endothelial cells within the saphenous vein segment under arterial and venous magnitudes of shear stress in the bioreactor.

Vein Viability and Function after Venous and Arterial Shear Stress

The effect of arterial shear stress on vein viability was also examined. Western blots for cleaved caspase-3 and cleaved PARP were performed on paired (0 and 24 hours) individual vein samples to assess apoptosis; no apop-

Table 1. Primers used in PCR analysis.

| Primer   | Sequence                                      |
|----------|-----------------------------------------------|
| Eph-B4   | 5'-GTCTGACTTTGGCCCTTTCCC-3'/5'-TGACATCACCTCCACATCA-3' |
| Ephrin-B2| 5'-CTGCTGGATCAACCAGGAAT-3'/5'-GGGCCCTCTCAGACCTTG-3' |
| GAPDH    | 5'-CCAGGGCCGCACTACGA-3'/5'-GCCAGCGAGCCACATC-3' |
| 28S      | 5'-GGTAGGATGCGAGTGCTCTAGT-3'/5'-AGTTGATTCGAGGTTGAGTT-3' |
| Osteopontin | 5'-TTGCAGTGATTTTGCTTTTG-3'/5'-GCCACAGCATCTGGATT-3' |

Figure 3. Vein viability after venous or arterial shear stress. A. Western blot for cleaved caspase-3 or GAPDH in representative vein samples at 0 or 24 hours of venous or arterial shear stress. Positive control; staurosporine-killed HUVEC. N = 6. B. Western blot for cleaved PARP or GAPDH in representative vein samples at 0 or 24 hours of venous and arterial shear stress. Positive control; staurosporine-killed HUVEC. N = 6. C. Photomicrographs showing representative TUNEL staining of matched vein samples at 0 or 24 hours of arterial (upper panels) or venous (lower panels) shear stress. Arrows indicate TUNEL positive cells. Scale bar represents 200 µm. N = 8.
Apoptosis was detectable at either baseline or after 24 hours of arterial shear stress (n = 6, Figures 3A,B).

To determine whether venous magnitudes of shear stress were associated with apoptosis in the vein, veins were examined with TUNEL at baseline and after 24 hours of shear stress. Very few TUNEL positive cells were seen, with only a minimal increase in adventitial staining at both 0 and 24 hours (n = 8, Figure 3C). These results are consistent with lack of apoptosis with exposure to either venous or arterial magnitudes of shear stress in the bioreactor.

Metabolic health of the saphenous vein segment was assessed using immunoreactivity of mTOR, a synthetic protein reflecting global cellular metabolism [23], at baseline and after 24 hours of arterial shear within the same vessel. Although mTOR immunoreactivity was variable between the samples, there was a trend (p = 0.08) toward increased mTOR immunoreactivity after 24 hours of arterial shear (n = 8, Figure 4A).

Since mTOR may reflect metabolic activity of both the endothelial cells and the smooth muscle cells in the vein, we assessed endothelial cell metabolic health using eNOS immunoreactivity [24], at baseline and after 24 hours of arterial shear within the same vessel. Western blots for 8 paired samples at 0 and 24 hours of arterial shear showed no difference in eNOS immunoreactivity at 0 and 24 hours of arterial shear stress (n = 8, Figure 4C).

Reduced EphB4 Expression with Arterial but not Venous Shear Stress

To determine the effect of shear stress on markers of cellular identity, segments of the same saphenous vein were exposed to either arterial or venous magnitudes of shear stress and then PCR was performed to assess cell identity; we also assessed the effect on osteopontin expression, a marker of vein graft adaptation [25]. Exposure to an arterial magnitude of shear stress for 24 hours was associated with decreased number of Eph-B4 mRNA transcripts (n = 10, P = .002, Figure 5A), no change in Ephrin-B2 mRNA transcripts (n = 9, P = .268, Figure 5B), and increased number of osteopontin mRNA transcripts (n = 7, P = 002, Figure 5C). Exposure to a venous magnitude of shear stress did not result in decreased number of Eph-B4 mRNA transcripts (n = 4, P = .665), and there was a trend toward increased number of both Ephrin-B2 (n = 6, P = .06) and osteopontin (n = 4, P = .398) transcripts (Figure 5D-F). Segments exposed to static conditions demonstrated no significant change in Eph-B4 (n = 3, P = .995), ephrin-B2 (n = 3, P = .973), or osteopontin (n = 3, P = .984) expression at 24 hours compared to time 0 (Figure 5G-I); the lack of change in osteopontin expression confirms the static condition as a control.

Immunofluorescence examining Eph-B4 also showed diminished Eph-B4 im-

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Table 2. Parameters of vein and flow conditions.

| Parameter | Mean     | Min | Max |
|-----------|----------|-----|-----|
| Radius (mm) | 1.8 ± 0.5 | 1   | 2.5 |
| Pressure, mean (mmHg) | Upstream | Post-vein | Drop |
| Venous shear (3 dynes/cm²) | 5.4 | 3.1 | 2.3 |
| Arterial shear (20 dynes/cm²) | 37.6 | 7.8 | 29.8 |
| Flow (mL/min) | Mean     | Min | Max |
| Venous shear (3 dynes/cm²) | 24.8 ± 20.9 | 3.7 | 58.1 |
| Arterial shear (20 dynes/cm²) | 200.6 ± 133.2 | 83.7 | 387.5 |

Flow rate calculated based on measured radius, measured pressure change, and approximate length of 10cm using Equation 2.
munoreactivity after 24 hours of arterial compared with venous magnitudes of shear stress (n = 3, Figure 6). Relative quantification of immunofluorescence images demonstrated a trend toward loss of Eph-B4 protein in the arterial samples compared to static, although not statistically significant.

**DISCUSSION**

We show that human saphenous veins treated with arterial shear stress for 24 hours retain their endothelial monolayer without apoptosis. In addition, saphenous veins exposed to arterial, but not venous, magnitudes of shear stress have reduced Eph-B4 expression, no change in Ephrin-B2 expression, and increased osteopontin expression. These changes recapitulate the previously described changes in vessel identity that occur during venous adaptation to the arterial environment [3,13] and confirm that shear stress is capable of inducing these changes.

The endothelium plays a critical role in the prevention of vein graft neointimal hy-
perplasia and thrombosis, with functions that include sensing of shear stress, regulation of inflammation, and prevention of platelet adherence, thrombosis, and plaque formation [26-29]. Surgical manipulation of the vein is thought to be an important component of endothelial injury that leads to neointimal hyperplasia and ultimately vein graft failure [30,31]. There are several potential mechanisms of venous injury during the surgical harvest and implantation, including devascularization of the adventitia leading to ischemia; storage in non-physiologic solutions; high-pressure distention resulting in endothelial damage, dissection, and cell loss; and possibly tissue crushing with poor instrumentation or suturing techniques [32-36]. As such, the viability of the vein — and especially the endothelium — has been a persistent question in vascular surgery. We show that endothelial cells are present, without apoptosis, after routine surgical vein harvest. We also confirm the presence of endothelial cells with molecular markers (Figure 2). It is known that some endothelial cells are lost during surgical harvest. Repopulation in vivo may be from the remaining endothelial cells, or potentially from circulating endothelial progenitor cells after reimplantation, or even pannus ingrowth in young patients [15,37-39]. Nonetheless, while it is possible that the endothelium is lost upon implantation of the vein into the arterial system in vivo, our results show that at least some endothelial cells can survive the perioperative harvest period and persist after exposure to arterial magnitudes of shear stress.

Our finding that venous Eph-B4 is reduced without induction of Ephrin-B2 under...
conditions of arterial shear stress (Figure 5) is consistent with previous reports showing loss of venous identity without gain of arterial identity in both human, rat, and mouse vein grafts exposed to the arterial environment [13,40]. Since we examined the response in whole veins, and Eph-B4 is found in both endothelial cells as well as smooth muscle cells [11], we have not identified the putative cell source of these changes. Previous work with endothelial and endothelial progenitor cells (EPC) have shown that shear stress can alter cell identity [19,41]. However, our study used whole saphenous veins taken from adult patients, with cardiovascular disease; as such, differences between our findings and these reports using cultured, presumably healthy cells of either venous or arterial origin, may be expected. Moreover, our finding that Ephrin-B2 expression does not increase with arterial shear stress may reflect our examination of adult vessels [13], as adult vessels may be deficient in delta-notch signaling upstream of Ephrin-B2, as reported by Kondo et al. in a vein graft model using aged rats [42]. Nevertheless, our findings mirror those described by Kudo et al. in adult human and aged rat vein grafts [13,40], suggesting the utility of our bioreactor model to recapitulate the adaptive response seen in vivo during vein graft adaptation to arterial shear stress. Interestingly, we show that osteopontin expression is induced with arterial magnitudes of shear stress (Figure 5C), showing the viability of these specimens in the bioreactor. However, the lack of increased osteopontin expression under venous shear stress conditions (Figure 5F) may reflect the variability and small number of human specimens, or the possibility that venous magnitudes of shear stress do not induce vein graft adaptation.

Our finding that a venous magnitude of shear stress was not associated with significantly diminished Eph-B4 expression (Figure 5B) is consistent with the presence of venous identity in adult veins. Laminar shear stress is considered atheroprotective [42,43], whereas disturbed or turbulent shear stress is associated with atherosclerosis as well as vein graft neointimal hyperplasia [44]. Veins are exposed to low magnitudes of laminar shear stress in vivo, suggesting that the laminar character of shear stress is important for normal venous endothelial homeostasis. However, the failing vein graft in vivo is also associated with low magnitudes of arterial shear stress [45]; these findings suggest that the magnitude of the shear stress changes may be more important than the frequency in the regulation of vein graft identity. Diminished venous endothelial Eph-B4 expression is associated with an angiogenic and mitogenic phenotype characterized by increased secretion of smooth muscle cell mitogens and reduced nitric oxide production [46], suggesting that vein
graft adaptation to the arterial environment may be mediated directly by the effects of changes in shear stress magnitude on Eph-B4.

There are several limitations to our in vitro experiments. Firstly, the artificial in vitro environment cannot completely model all aspects of the in vivo system, including lack of an immune system, as well as the circulating elements such as platelets or EPCs. Secondly, although we have examined shear stress as a variable, the roles of pressure and oxygen tension have not been examined. Finally, the extended viability of the veins in the flow system has not yet been examined beyond 24 hours. Nonetheless, acute shear stress has been shown to elicit numerous changes in cell structure and function, including cytoskeletal remodeling and activation of signaling cascades that are different with exposure to chronic shear stress [47]. Whether the cells of intact vessels experience the same acute changes remains unclear.

Many other investigators have examined veins in ex vivo perfusion circuits. For example, Hoenicka et al. [48] have reported a similar bioreactor system to characterize the metabolic changes that occur during vessel perfusion. Importantly, this study examined the larger bovine saphenous vein and did not assess effects on vessel identity; nevertheless, their demonstration of increased endothelial survival in the presence of shear stress is consistent with our data. Similarly, Gusic et al. [49] showed that shear stress regulates intimal hyperplasia, whereas transmural pressure regulates medial hypertrophy in perfused veins, but they also did not report effects on vessel identity. Recently, Be rrard et al. [50] treated human saphenous veins in a perfusion system similar to ours for 7 days and showed that both Eph-B4 and Ephrin-B2 expression decrease in response to arterial magnitudes of shear stress but not pressure. Our data is complementary to this study, showing that arterial magnitudes of shear stress increases osteopontin expression and that these expression changes do not occur with static controls (Figure 5). In addition, we show lack of apoptosis in the perfused veins (Figure 3). Importantly, we also show preservation of eNOS and mTOR (Figure 4), downstream effectors of the Eph-B4 pathway. Lastly, multiple adjustable aspects of our system allow for the close approximation of laminar flow without high pressure gradients, compression effects, pinch shear or distention, preventing confounding effects on vessel structure or function.

Our finding that Eph-B4 expression was diminished with treatment using an arterial but not a venous magnitude of shear stress shows the importance of shear stress in the upstream regulation of Eph-B4 expression as well as in the regulation of vein graft adaptation. It is not currently clear whether shear stress directly induces changes in venous endothelial Eph-B4 signaling, such that Eph-B4 is a direct mechanosensing molecule, or whether Eph-B4 is part of the mechanotransduction cascade.

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