Wall Shear Stress Effects on Endothelial-Endothelial and Endothelial-Smooth Muscle Cell Interactions in Tissue Engineered Models of the Vascular Wall

Dalit Shav*, Ruth Gotlieb, Uri Zaretsky, David Elad, Shmuel Einav

Department of Biomedical Engineering, Tel Aviv University, Tel Aviv, Israel

Abstract

Vascular functions are affected by wall shear stresses (WSS) applied on the endothelial cells (EC), as well as by the interactions of the EC with the adjacent smooth muscle cells (SMC). The present study was designed to investigate the effects of WSS on the endothelial interactions with its surroundings. For this purpose we developed and constructed two co-culture models of EC and SMC, and compared their response to that of a single monolayer of cultured EC. In one co-culture model the EC were cultured on the SMC, whereas in the other model the EC and SMC were cultured on the opposite sides of a membrane. We studied EC-matrix interactions through focal adhesion kinase morphology, EC-EC interactions through VE-Cadherin expression and morphology, and EC-SMC interactions through the expression of Cx43 and Cx37. In the absence of WSS the SMC presence reduced EC-EC connectivity but produced EC-SMC connections using both connexins. The exposure to WSS produced discontinuity in the EC-EC connections, with a weaker effect in the co-culture models. In the EC monolayer, WSS exposure (12 and 4 dyne/cm² for 30 min) increased the EC-EC interaction using both connexins. WSS exposure of 12 dyne/cm² did not affect the EC-SMC interactions, whereas WSS of 4 dyne/cm² elevated the amount of Cx43 and reduced the amount of Cx37, with a different magnitude between the models. The reduced endothelium connectivity suggests that the presence of SMC reduces the sealing properties of the endothelium, showing a more inflammatory phenotype while the distance between the two cell types reduced their interactions. These results demonstrate that EC-SMC interactions affect EC phenotype and change the EC response to WSS. Furthermore, the interactions formed between the EC and SMC demonstrate that the 1-side model can simulate better the arterioles, while the 2-side model provides better simulation of larger arteries.

Introduction

The close proximity between the endothelial cells (EC) that compose the intima of blood vessels and the smooth muscle cells (SMC) composing the tunica media enables both these cells to interact with each other. Extensive research on the effects of physiological hemodynamic forces has found that WSS causes functional switching of the endothelial phenotype from a quiescent atheroprotective phenotype under physiological and elevated levels of WSS (10–20 dyne/cm²) to an atherogenic phenotype at low WSS (0–4 dyne/cm²) [1–3]. However, the impact of these SMC-EC interactions on the response of EC to WSS is still unclear and requires more research.

Most research on the effects of WSS on EC biology has been done with EC alone, that only forms EC-EC interactions and without the interactions of EC with its neighboring SMC (EC-SMC interactions) [4]. To investigate the interactions between EC and SMC, several co-culture techniques and models have been introduced. The main four co-culture models are: (1) direct culture of EC on SMC (1-side model) as illustrated in Figure 1B; (2) a culture of SMC and EC on opposite sides of different membranes (2-side models) as illustrated in Figure 1C; (3) a culture of EC on collagen gels or other polymers containing SMC (3 dimensional (3D) model) as illustrated in Figure 1E; and (4) the use of conditioned media as illustrated in Figure 1D, which is a useful method for assessing EC-SMC interactions through secretory mechanisms [5–7].

The effect of WSS on various EC-EC interactions has mainly been studied on EC monolayers (Table 1). A short exposure to WSS was shown to induce a discontinuity in vascular endothelial (VE)-Cadherin staining at the border of the EC. This change was not accompanied by a change in VE-Cadherin expression [8–11]. Similarly, in vivo, VE-Cadherin expression was weak and discontinuous in areas of disturbed flow, but significantly higher and continuous where flow was laminar [8,12]. In addition, connexin 43 (Cx43), which forms gap junctions between the EC (EEGJ), was found to be minimally expressed or absent when exposed to physiological WSS, whereas the gap junctions Cx37 and Cx40 were found to be up-regulated when exposed to physiological WSS. This altered pattern of Cx expression in EC was supported by in vivo findings [13–15]. Moreover, acute WSS increased EC permeability, which was mediated by nitric oxide (NO) [16], and occludin phosphorylation [17] on the other hand, chronic WSS...
decreased EC permeability, which was mediated by NO as well [18]. The Platelet Endothelial Cell Adhesion Molecule-1 (PECAM-1) was found to participate in a mechano-sensory complex that responded to WSS activation [19]; on the other hand, staining of PECAM-1 was reduced in the presence of SMC in a 1-side model [20].

The effect of WSS on the interactions between EC and SMC has been shown to reduce the inflammatory response induced by the proliferating SMC on the EC. This was evidenced by the reduced expression levels of adhesion molecules such as P-selectin and the vascular adhesion molecule (VCAM) [21,22], a reduced fibrinolytic and coagulatory response [23], reduced cell migration and proliferation rate [4,24], as well as the EC and SMC gene profiles [25,26] following exposure to WSS. Table 2 summarizes these main EC-SMC interactions. In addition, Wallace et al. found that EC attached to SMC using integrin α5β1 and lacked focal adhesions in the 1-side model [27], but it was still attached firmly to SMC with very limited cell loss following exposure to high WSS [20,28]. By contrast, in a 2-side model, the presence of

---

**Table 1.** A summary of published endothelial-endothelial interactions examined in different vascular wall models.

| Model Type                  | EC-EC Interactions                                                                 | EC source   | SMC source   | References               |
|-----------------------------|-----------------------------------------------------------------------------------|-------------|--------------|--------------------------|
| **EC monolayer model**      | Adherens junctions: VE-Cadherin and catenin biochemistry and effect of WSS         | PTAEC, BAEC, HUVEC |             | [8,11,49]               |
|                             | Gap junctions: connexins 37, 40, 43 endothelium derived hyperpolarizing factor, permeability between the cells (dye transfer) and effect of WSS | HAEC, bEnd.3, BAEC |             | [13,42,50]               |
|                             | Endothelium permeability (tight Junctions: Zonula occludens proteins) and the effect of WSS | Reviewed by [17, 18] |             |                         |
|                             | PECAM-1 role as a WSS sensor and in inflammatory response.                        | BAEC        |             | [19], reviewed by [51]  |
| **1-Side**                  | LDL uptake/infiltration                                                            | BAEC        | BASMC        | [52,53]                  |
| **PECAM-1**                 |                                                                                   | PCASMC      |              | [20]                     |
| **2-Side**                  | LDL uptake and water infiltration                                                  | HUVEC       | HUSMC        | [52]                     |
| **3D-model**               | Effect of WSS on myoendothelial Junctions (Cx37, 40 and 43) expression            | HAEC        | HASMC        | [35]                     |

Abbreviations: PTAEC-pig thoracic aorta EC, BAEC-bovine aortic EC, HUVEC-Human umbilical vein EC, HAEC-Human aortic EC, bEnd.3-PymT-transformed mouse endothelial cells clone bEnd.3, PCAEC-pig carotid artery EC, HUSMC-Human umbilical SMC, HASMC-Human aortic SMC, BASMC-Bovine aortic SMC, PCASMC-Pig carotid artery SMC.

doi:10.1371/journal.pone.0088304.t001
SMC increased the number of adherent ECs, as well as the total focal adhesion area in EC [29].

The direct communication of EC-SMC through myoendothelial gap junctions (MEGJ) has mainly been studied without exposure to WSS (Table 2). Isakson et al. reported the presence of Cx37, Cx40 and Cx43 between the 2 cell types using the 2-side model. In this model, the EC and SMC form direct connections via cellular projections through the membrane pores [30]. Further studies have explored different biochemical activations and regulations of the MEGJ in this model without the effect of WSS [31–34]. Johnson and Nerem used a 3D model and found an up-regulation in Cx37, Cx40 and Cx43 expression in the presence of SMC, and that Cx37 expression was even higher following exposure to WSS, whereas Cx43 was down-regulated by the flow. The increased expression of connexins of EC in the presence of SMC was probably due to the role of the connexins in the MEGJ [35]. However, the presence of the MEGJ has not been studied in the 1-side model and it is not clear whether the interactions between the EC and SMC in the different models are similar; moreover, the effect of WSS exposure on the EC-SMC connections in the 1-side and 2-side model awaits investigation.

Most studies on WSS effects on the EC-SMC interaction have focused on the inflammatory response of both EC and SMC. However, those studies have not considered that in vivo, the EC-SMC interactions may vary between different types of blood vessels, affecting the response of EC to WSS. Based on the understanding of these interactions in vivo, we hypothesized that for the studies of the effects of mechanical forces on the endothelium the 1-side model can simulate better the arterioles, while the 2-side model provides better simulation of larger arteries. Thus, the main goal of this work was to study the ways in which the interactions between EC and their neighboring cells (i.e. EC-EC interactions) affect the response of EC-SMC interactions) to WSS stimulation. To support our hypothesis we compared the response of EC to WSS using 3 different models: (1) EC monolayer, (2) 2-side model, and (3) 1-side model. The models were exposed to a low WSS of 4 dyne/cm², a physiological WSS of 12 dyne/cm² and a control level (0 dyne/cm²). We analyzed the morphology of the models and the expression levels of proteins responsible for the cell interactions with their neighboring cells.

### Methods

#### Custom-designed Wells for the in-vitro Cell Culture Models

The cell cultures for the in vitro models of the vascular wall in this work were grown in modified wells that were previously developed in our laboratory [36]. Briefly, these wells are constructed from a well bottom and a cylinder. The well bottom holds a stretched polytetrafluoroethylene (PTFE) membrane that can be used as a substrate for the cell culture and the cylinder is

| Model Type | EC-SMC Interactions | EC source | SMC source | References |
|------------|---------------------|-----------|------------|------------|
| 1-Side - (Fig. 1B) | EC adhesion to SMC | PCAEC, HAEC, HUVEC | PCASM, HASMC | [20,27,28,54] |
| | WSS effect on LDL uptake | BAEC | BASMC | [55] |
| | EC elastic modulus | HCAEC | HASMC | [56] |
| | WSS effect on coagulation and fibrinolytic factors | HSVEC | HSVMSC | [23] |
| | WSS effect on SMC and EC gene profile | HSVEC | HSVMSC | [25,26] |
| 2-Side - (Fig. 1C) | Myoendothelial Junctions (Cx37, 40 and 43) organization and regulation in Static conditions | MAEC, RPAEC, HUVEC | MASMC, RPASMC, HUSMC | [30–34] |
| | WSS effect on EC and SMC migration and proliferation | HUVEC | HUSMC | [4,24,57,58] |
| | SMC differentiation | BAEC, HAEC | BASMC, HASMC | [6,59,60] |
| | Effect of WSS on SMC phenotype | HUVEC | HUSMC | [61] |
| | SMC and EC adhesion to matrix | HUVEC | HUSMC | [29,62] |
| | Effect of WSS on EC gene and proteins expression of adhesion molecules | HUVEC | HUSMC | [22,63,64] |
| | Direct contacts through Jagged 1 protein importance in SMC differentiation | HCAEC | HCASM | [65] |
| | Effect of WSS on EC gene expression | HUVEC | HUSMC | [26] |
| Conditioned Medium – (Fig. 1D) | WSS on EC change SMC proliferation and migration | HUVEC | HUSMC | [57] |
| | SMC differentiation | BAEC, HUVEC | BASMC, HUSMC | [6,66,67] |
| 3D-model - (Fig. 1E) | Effect of WSS on myoendothelial Junctions (Cx37, 40 and 43) expression | HAAEC | HASMC | [35] |
| | Effect of WSS on EC expression of adhesion molecules | HUVEC | HUSMC | [21] |
| | Effect of WSS on EC proliferation and migration | HCAEC | HASMC | [68] |
| | WSS effect on SMC phenotype | BPAEC | BPASM | [69,70] |

Abbreviations: PTAEC-pig thoracic aorta EC, BAEC-bovine aortic EC, HUVEC-Human umbilical vein EC, HAEC-Human aortic EC, PCAEC-Pig carotid artery EC, HCAEC-Human carotid artery EC, BPAEC-Bovine pulmonary artery EC, HASMC-Human aortic SMC, HASMC-Human muscular SMC, HASMC-Human aortic SMC, BASMC-Bovine aortic SMC, PCASM-Pig carotid artery SMC, HSVMSC-Human saphenous vein EC, MASMC-Mouse aorta SMC, RPASMC-Rat pulmonary artery SMC, BPAEC- Bovine pulmonary artery SMC.

doi:10.1371/journal.pone.0088304.t002
assembled in such a way as to hold the medium above the membrane. The cylinder is constructed so that the well bottoms can be assembled on both sides (Fig. 2). This makes it possible to culture cells on both sides of the membrane. Once the co-cultured layers become confluent and ready for flow experiments the cylindrical medium holder can be disassembled from the well bottom (Fig. 2) and installed in the flow chamber for flow experiments. Upon completion of the experiments, the medium holder can be re-assembled on the well bottom for either biological treatment of the cells or for further culture. The PTFE membrane (Millipore) was coated using 10% fibronectin (Biological Industries, Beit Haemek, Israel) solution prior to cell culturing.

Human Umbilical Vein Endothelial Cells and Human Umbilical Artery Smooth Muscle Cells

Human umbilical vein endothelial cells (HUVEC) (PromoCell) were cultured in endothelial cell media (PromoCell) supplemented with 100 U/ml Penicillin, 0.1 mg/ml Streptomycin and 12.5 U/ml Nystatin (Biological Industries, Beit Haemek, Israel). Human umbilical aortic smooth muscle cells (SMC) (Lonza) were cultured in MCDB-131 Medium supplemented with 5% Fetal Bovine Serum, 2 mM L-Glutamine, 100 U/ml Penicillin, 0.1 mg/ml Streptomycin and 12.5 U/ml Nystatin (Biological Industries, Beit Haemek, Israel), 2 ng/ml Insulin (Sigma Aldrich), 0.5 ng/ml Epithelial Growth Factor (Sigma Aldrich), 3 ng/ml Basic-Fibroblasts Growth Factor (bFGF) (PeproTech Inc., USA) and 1 μg/ml Hydrocortisone (Sigma Aldrich). Both cell types as well as the co-culture were cultured at 37°C and 5% CO₂ in a humidifier incubator. In both cell types, only passages 4–8 were used for this study.

The in-vitro Vascular Wall Models

The 2-side co-culture model. The PTFE membrane was installed in the custom-designed well facing downwards and was placed into a plastic tissue culture dish containing the SMC growth medium (Fig. 2B bottom and Fig. 2D left). About $7.5 \times 10^4$ SMCs were seeded on the PTFE membrane on the lower side of the well. After 24 hours, the membrane was washed to remove excess non-adherent SMC and the well was re-assembled so that the SMC

**Figure 2. The custom made co-culture wells for the *in vitro* vascular wall models.** A – Diagram of the parts of the well. B – Assembled well, top – upper side up, bottom – face down. C – Picture of the disassembled well bottom. D – Pictures of assembled well, right – upper side up, left – face down. doi:10.1371/journal.pone.0088304.g002
cells would be facing downwards and completely submerged in the HUVEC growth medium (Fig. 2B top and Fig. 2D right). Then, 10^5 HUVEC were seeded on the membrane side facing upward. After 24 hours, both layers of EC and SMC became confluent and ready for the WSS experiments.

**The 1-side co-culture model.** The PTFE membrane was installed in the custom-designed well with the upper face upwards and was placed into a plastic tissue culture dish containing the SMC growth medium. About 7.5 × 10^4 SMC were seeded on the PTFE membrane on the upper side (Fig. 2B top and Fig. 2D right). After 24 hours, the membrane was washed to remove excess non-adherent SMC and the medium was replaced by HUVEC growth medium. Then, 10^5 HUVEC were seeded on top of the SMC layer (upper side). After 24 hours both layers of EC and SMC became confluent and ready for WSS experiments.

The endothelial monolayer model. The PTFE membrane was installed in the custom-designed well with the upper face upwards and was placed into a plastic tissue culture dish containing the HUVEC growth medium (Fig. 2B top and Fig. 2D right). About 10^5 HUVEC were seeded on the PTFE membrane (upper side). At this step we ensured that the medium completely covered the underside of the membrane with the cell culture and without any gas bubbles. After 24 hours the HUVEC layer became confluent and ready for WSS experiments.

Once confluent layers were obtained (approximately 16 hours after the EC were seeded), the custom-designed wells were disassembled and the well bottom with the in vitro vascular wall model was assembled in the flow chamber, so that the HUVEC side would be exposed to physiological flows while the other side (with no cells or with SMC) would stay in static conditions. Prior to conducting the experiments, all models were inspected under phase contrast microscopy to verify that the cells had reached at least 90% confluency.

**Flow System for Wall Shear Stress Experiments**

A custom designed parallel plate flow chamber with a rectangular cross section was built to apply uniform laminar WSS on 3 samples of the tissue-engineered vascular wall models. Since the height (h = 1 mm) of the flow chamber is much smaller than either its length (l = 100 mm) or its width (w = 35 mm) the fluid mechanics of the channel can be described as a parallel plate flow chamber. Thus, the WSS ($\tau_w$) is given by:

$$\tau_w = \frac{6\mu Q}{bh}$$  

where $Q$ is the flow rate and $\mu = 0.007$ dyne s/cm$^2$ is the fluid viscosity for the medium at 37°C.

In addition, a computational fluid dynamics (CFD) model was developed to validate the uniform distribution of the WSS in the flow chamber, using the computational fluid dynamics package of FLUENT (Fluent Inc., Lebanon, NH). The flow profiles at different locations were computed and compared for a uniform inlet velocity of $U_i = 0.3125$ m/s, which corresponds to a flow rate of $Q_i = 1.2$ L/min, for which $Re = 1628$ and the analytically calculated WSS equaled 24 dyne/cm$^2$. The WSS distributions on the bottom of the flow chamber were computed for three axial locations that corresponded to the center of each of the three well bottoms and showed a uniform WSS field of 24 dyne/cm$^2$ over the entire surface of all three well bottoms.

For the WSS experiments, the well bottoms were inserted into the bottom part of the flow chamber so that the PTFE membrane was in the same plane as the chamber bottom, allowing for laminar flow on the HUVEC side of the vascular wall models. The flow chamber was then connected with tubes to a peristaltic pump and a medium reservoir. The flow chamber and the medium reservoir were maintained at 37°C and 5% CO$_2$ in a humidifier incubator during exposure to WSS of the endothelial side of the models. The flow chamber allows for up to 3 wells in each flow experiment. All experiments were performed using a flow rate of 600 ml/min resulting in a WSS of 12 dyne/cm$^2$ for physiological WSS, a flow rate of 200 ml/min resulting in a WSS of 4 dyne/cm$^2$ for low WSS and, static conditions (i.e. 0 dyne/cm$^2$) as the control. The WSS experiments were conducted for 30 min in the case of the connexins protein quantification, and for 30 and 60 min for visualization of cell morphology, VE-Cadherin and FAK. All experiments were performed using all three models of the vascular wall as described above and all the experiments were performed at least three independent times with three wells in each test.

**Cell Morphology (Immunofluorescence Stains)**

The vascular wall models were exposed for 30 min and 60 min to 0, 4 and 12 dyne/cm$^2$, and then fixed in a 4% paraformaldehyde solution for 10 min. This was followed by blocking, using a phosphate buffered solution (PBS) containing 10% normal goat serum and 0.1% Triton X-100 solution for 30 min followed by incubation overnight at 4°C with the following primary antibodies: a mouse anti VE-Cadherin (Santa Cruz Biotechnology) and a rabbit anti FAK (Abcam). This was followed by an incubation of 30 min with a secondary antibody: alexa fluor 647 conjugated goat anti rabbit, alexa fluor 488 conjugated goat anti mouse (Jackson ImmunoResearch) and TRITC conjugated phalloidine (Sigma Aldrich). Between each step the samples were rinsed 3 times with PBS. The samples were then mounted between 2 large microscope glass slides using a DAPI mounting gel (Santa Cruz Biotechnology) in order to visualize the cell nucleus. The slides were stored in a covered box at 4°C until microscopy examination. The cells were examined under a Zeiss LSM510 confocal microscope. Analysis and quantification of the images was done using the LSM image browser software (Zeiss) and the provided measuring tools.

**Protein Quantification (Flow Cytometry)**

The vascular wall models were exposed for 30 min to 0, 4 and 12 dyne/cm$^2$ and then the cells were trypsinized and centrifuged for 5 min at 7 g to separate the cells from the trypsin solution. The cell suspensions of both HUVEC and SMC were then fixed and permeabilized using ice cold methanol for 10 min at −20°C. The samples were incubated first with the appropriate primary antibody, followed by incubation with the appropriate secondary antibodies for 30 min each at room temperature. Following each step, 1 ml of PBS was added to the cell suspension and the resulting solution was then centrifuged for 5 min at 7 g to separate the cells from the solution. The solution was gently poured out of the test tube, leaving the cell pellet at the bottom of the test tube. The cells were mixed to form a homogenous cell suspension using a vortex after the next solution was added. In order to differentiate the different cells, each sample was labeled for VE-Cadherin (which is positive for HUVEC and negative for SMC) using a mouse anti VE-Cadherin antibody (Santa Cruz) and for one of the following primary antibodies: rabbit anti Cx37 alexa fluor 647 conjugated (BIOS), rabbit anti Cx43 alexa fluor 488 conjugated (BIOS) followed by the secondary antibodies: alexa fluor 647 conjugated goat anti rabbit and alexa fluor 488 or alexa fluor 647 conjugated goat anti mouse. Positive control samples were labeled with primary and secondary antibodies and negative control samples were labeled with secondary antibody alone. Samples were analyzed using a BD LSR digital flow cytometer. The
Flowing software (Turko University, Finland) was used for data analysis.

**Statistical Analysis**

The results were evaluated using a 2-way analysis of variance (SPSS, IBM v19). A P value of less than 0.05 was considered statistically significant. Post-hoc Tukey-Kramer tests supplied by the SPSS software were used to perform pair wise multiple comparisons between the data obtained from the different WSS applied on the different models.

**Results**

We investigated the attachment of EC to the matrix as well as the interactions of EC-EC and EC-SMC in response to WSS stimulation. For this purpose we compared the morphology and amount of proteins immediately following short term exposure to different WSS in three different models of the vascular wall made up of EC monolayer and two co-cultures of EC and SMC in which EC were cultured directly over a confluent layer of SMC or on the opposite side of a membrane with a culture of SMC.

**Endothelial-matrix Interactions**

The focal adhesion kinase (FAK), which is an important protein in the focal adhesion complex that attaches cells to the matrix, was used to evaluate the EC-matrix interaction. Stains of HUVEC actin stress fibers and FAK, following the exposure to a WSS of 12 dyne/cm² for 0 and 60 min are shown in Figure 3. The FAK stains shows that the focal adhesions are located at the end of the stress fibers, forming a wide band at the cell borders as can be seen in Figure 3-A to C, with fewer and smaller apparent plaques in the 1-side model in the control compared to the amount of plaques in the EC monolayer and the 2-side models (Fig. 3i - B). Following exposure to a WSS of 12 dyne/cm², the HUVEC became a little more elongated and the stress fibers aligned with the flow direction in all three models (Fig. 3i). The stress fibers aligned with the cell direction and formed longer and less peripheral fibers (Fig. 3ii A to C). Following exposure to WSS, in the 1-side model, the FAK stain started to resemble the other models (Fig. 3ii - B).

The length and amount of FAK plaques were measured from the images of the vascular wall models. For each field the average plaque length and amount of plaques per cell were calculated and used to estimate the equivalent average total amount of FAK per cell (Fig. 4). The results showed that in all three models, the exposure to a WSS of 12 dyne/cm² led to an increase in the amount of FAK in a time dependent manner. In static conditions and following exposure to 30 min of 4 dyne/cm² (LSS) the amount of FAK in the 1-side model was very low (significantly different from the other models). Following an exposure of 30 min to 12 dyne/cm² (PSS30) the amount of FAK in this model increased (significantly); however it was still lower than in the other two models. Following an exposure of 60 min to 12 dyne/cm² (PSS60), however, the amount of FAK was similar to that of the EC monolayer model. The amount of FAK in the 2-side model was a little higher in all modes of WSS exposure (Fig. 4).

**Endothelial-endothelial Interactions**

The VE-Cadherin, which is the main adherens junction of the endothelium, was used to evaluate the EC-EC interaction, since these junctions are the major contributors to the cell–cell adhesion force. In addition, the gap junction proteins Cx37 and Cx43 were used to evaluate the EC-EC communication.

Figure 5A shows the endothelial plane of the models when stained for VE-Cadherin. The VE-Cadherin marks the line where HUVEC are connected to one another, demonstrating the cell borders, and showing a confluent, mature endothelium in all three models. The width of the VE-Cadherin mark in the 1-side and 2-side models was much thicker than in the EC monolayer model. The VE-Cadherin in these two models seemed to be either double or had protrusions between two adjacent cells. Measurement of the stain thickness showed that the distribution of the thickness values of the stain in the EC monolayer model was around 1 μm whereas that of the 1-side and 2-side models was around 2 μm, hence showing a significance increase in the thickness of the stain that marks the connection between the cells (Fig. 5B). This thickness remained visible following exposure to WSS, however a slight (non-significant) decrease in this value emerged in the 1-side model.

Following exposure to a WSS of 12 dyne/cm² for 60 min, the VE-Cadherin stain seemed less connected (Fig. 5A-D, E and F), and took on a fenestrated appearance. This discontinuity in the 1-side model was not as large as in the other models (Fig. 5A-E). Quantification of the intensity of the stain was done using a scale where 100% symbolizes a fully continuous line and a 0% symbolizes no line. The results showed that under static conditions, in all three models the intensity of the stain was around 80%; following exposure to a WSS of 12 dyne/cm² the EC monolayer and 2-side monolayer had a 50% score for both periods of 30 min and 60 min of WSS exposure. The 1-side model integrity varied significantly only following 60 min of exposure to a WSS of 12 dyne/cm² to a value of 65%. Following exposure to a WSS of 4 dyne/cm² for 30 min, only the EC monolayer showed a significance difference in the intensity of the stain to about 65% (Fig. 5C).

The quantification of VE-Cadherin expression using FACS indicated a significantly higher amount of protein in the EC monolayer than in the two co-culture models. This difference remained significant following exposure to WSS in the 2-side model, where the exposure to WSS did not change the amount of VE-Cadherin expression significantly. A slight increase in the amount of VE-Cadherin expression in the 1-side model (not significant) emerged following exposure to a WSS of 12 dyne/cm² with another increase following a WSS of 4 dyne/cm² (not significant). Following exposure to a WSS of 12 dyne/cm² a small reduction in the expression of VE-Cadherin was found in the EC monolayer model with a non-significant reduction in the case of 4 dyne/cm² (Fig. 5D).

EC-EC communications increased following exposure to flow, as can be seen by the increase in the amount of Cx43 and Cx37, which formed EEJ in the EC monolayer model following exposure to a WSS of 12 dyne/cm² and 4 dyne/cm² (Fig. 6 and 7), with a greater increase following exposure to 4 dyne/cm² in the case of Cx43 (Fig. 6).

**Endothelial-smooth Muscle Cell Interactions**

The myoendothelial gap junction proteins Cx37 and Cx43 form direct channels between EC and SMC. Thus, the amount of Cx37 and Cx43 was used to evaluate the EC-SMC direct interactions in the co-culture models. In order to calculate a statistic response of the cells, each sample was normalized to the EC-monolayer in static conditions (control) sample. The normalized results of the three repeats of the same test (non dependant repeats) were then averaged and presented (Fig. 6 and 7).

In static conditions, the amount of Cx43 showed a slight increase in the 2-side model compared to the EC monolayer and a significant increase in the 1-side model (Fig. 6), and there was a significant increase in the amount of Cx37 in both the 2-side and
1-side models, although there was no significant difference between the 2-side and 1-side models.

Following exposure to a WSS of 12 dyne/cm² the amount of both Cx43 and Cx37 did not change significantly in either the 1-side and 2-side models (Fig. 6 and 7). However, following exposure to 4 dyne/cm² the amount of Cx43 increased (significantly only in the 2-side model) (Fig. 6) and decreased (significantly only in the 1-side model) (Fig. 7).

Discussion

In this study we have studied how WSS affect the interactions of the endothelium with its surrounding cells, which constructs the vascular wall. This was done using two different co-culture models of the vascular wall as well as a control model, which lacks the interaction with SMC. The results show that EC attaches to the SMC using a different FA complex, and not using FAK. Following exposure to a WSS of 12 dyne/cm² the FAK became longer and increased in its amount in a time dependent manner. In the 1-side model, this exposure caused the appearance of the FAK until it resembled the EC monolayer model. This effect was more pronounced when there was no distance between the two cell types (caused by the membrane between the cells in the 2-side model). In the case of Cx43, a WSS of 4 dyne/cm² increased Cx43 expression compared to the control in the EC monolayer (demonstrating higher EEGJ interactions); however, in the co-culture models (where it formed EC-SMC interactions), the expression of Cx37 was reduced, with a greater difference in the 1-side model, showing that again, a distance between the 2 type of cells due to the presence of the membrane, limits the effect of WSS on the EC-SMC interactions, as demonstrated by the amount of VE-Cadherin and stain. However, when exposed to WSS, the interactions with the SMC reduced the discontinuity in EC-EC interactions seen in the VE-Cadherin stain. In the presence of SMC the EC formed direct EC-SMC interactions using both Cx37 and Cx43 to form MEGJ. When exposed to a WSS of 12 dyne/cm² both Cx43 and Cx37 expression were elevated in the EC monolayer, suggesting higher connectivity of EEGJ, however without a significant effect in the co-culture models where it formed EC-SMC interactions. When exposed to a WSS of 4 dyne/cm² the amount of Cx43 was elevated in all 3 models, with the highest in the EC monolayer and lowest in the 1-side model, suggesting that the interactions with SMC reduce the effect of WSS. This effect was more pronounced when there was no distance between the two cell types (caused by the membrane between the cells in the 2-side model). In the case of Cx37, a WSS of 4 dyne/cm² increased Cx37 expression compared to the control in the EC monolayer (demonstrating higher EEGJ interactions); however, in the co-culture models (where it formed EC-SMC interactions), the expression of Cx37 was reduced, with a greater difference in the 1-side model, showing that again, a distance between the 2 type of cells due to the presence of the membrane, limits the effect of WSS on the EC-SMC interactions.
FAK expression between EC cultured on fibronectin coated plastic proteins. In their model, however, there was no difference in expression levels and less mRNA in several focal adhesion complexes used to attach to the different matrices. Moreover, our method may have been application of WSS [39]. This discrepancy can be attributed to the findings reported by Wang et al where there was an increase of the total focal adhesion area (as measured using paxillin) in EC [29].

Following exposure to physiological WSS, the FAK stain became stronger and is easier to detect, showing an increase in the EC-matrix interaction. It is important to notice that this increase is probably not due to new formation of FAK but rather the recruitment and growth of existing FAK to the growing focal adhesion complexes. The antibody used for the FAK staining was a phosphorylated Y397 FAK (Abcam), suggesting that the increased stain was induced by increased phosphorylation of the FAK. Such phosphorylation was found in previous studies shortly after exposure to physiological WSS, and was accompanied by an increased amount of FAK at the focal adhesion sites [37,38].

Our finding of an increase in the amount of FAK runs counter findings reported by Davies et al, where no differences in the total adhesion area of EC to the matrix were found following the application of WSS [39]. This discrepancy can be attributed to the quantification methods. Davies et al quantified the area of the adhesion complex as the distance between the two cell types through pores present in the thin membranes [6,26,30]. In the 1-side model, the EC form a separate layer on top of the SMC and the cells do not migrate into the underlying SMC layer; hence the two cell types remain as two separate monolayers as well, though without a detectable distance.

In order to analyze the EC-Matrix interactions, we examined the FAK morphology of EC. In the 1-side model, the EC attached to the SMC directly and to the ECM produced by the SMC. This ECM differed from the fibronectin coated membrane that served as the attachment matrix to the EC in the EC monolayer and 2-side models. Thus the limited FAK stain (Fig. 3i-B) and reduced amount of FAK in the 1-side model (Fig. 4) might originate in the different FA complexes used to attach to the different matrices. Our finding is congruent with the work of Wallace et al, where EC cultured directly over SMC lacked focal adhesions, had reduced expression levels and less mRNA in several focal adhesion proteins. In their model, however, there was no difference in FAK expression between EC cultured on fibronectin coated plastic and EC cultured on SMC. They found that EC was mainly attached to SMC via α5β1 integrin whereas attachment to plastic involved both α5β1 and αvβ3. This difference in focal adhesion composition resulted in a different signaling cascade that drives different functions of the cell when exposed to WSS [27].

On the other hand, in a 2-side model as compared to the EC monolayer, an increase in the total amount of FAK was found in all WSS exposure modes (Fig. 4). This is similar to the finding reported by Wang et al where there was an increase of the number of adherent ECs, and an increase of the total focal adhesion area (as measured using paxillin) in EC [29].

and the effect is more similar to the that of the EC-EC interactions. The limited EC-SMC interactions found in the 2-side model resemble the limited interactions found in vivo in larger arteries, compared to the higher amount of MEGJ found in the 1-side model, which resembles the in vivo interactions in smaller arteries.

Thorough Inspection of the Z slices of the confocal images confirmed that the SMC and the endothelial layers remained as two separate monolayers on the opposite sides of the membrane in the 2-side model without migration of cells into the membrane or as 2 layers one on top of the other in the 1-side model. The distance between the two cell types in the 2-side model was measured to be roughly 20 μm. Although this is a large distance between EG and SMC, gap junctions have been reported to occur between the two cell types through pores present in the thin membranes [6,26,30]. In the 1-side model, the EC form a separate layer on top of the SMC and the cells do not migrate into the underlying SMC layer; hence the two cell types remain as two separate monolayers as well, though without a detectable distance.

In order to analyze the EC-Matrix interactions, we examined the FAK morphology of EC. In the 1-side model, the EC attached to the SMC directly and to the ECM produced by the SMC. This ECM differed from the fibronectin coated membrane that served as the attachment matrix to the EC in the EC monolayer and 2-side models. Thus the limited FAK stain (Fig. 3i-B) and reduced amount of FAK in the 1-side model (Fig. 4) might originate in the different FA complexes used to attach to the different matrices. Our finding is congruent with the work of Wallace et al, where EC cultured directly over SMC lacked focal adhesions, had reduced expression levels and less mRNA in several focal adhesion proteins. In their model, however, there was no difference in FAK expression between EC cultured on fibronectin coated plastic and EC cultured on SMC. They found that EC was mainly attached to SMC via α5β1 integrin whereas attachment to plastic involved both α5β1 and αvβ3. This difference in focal adhesion composition resulted in a different signaling cascade that drives different functions of the cell when exposed to WSS [27].

On the other hand, in a 2-side model as compared to the EC monolayer, an increase in the total amount of FAK was found in all WSS exposure modes (Fig. 4). This is similar to the finding reported by Wang et al where there was an increase of the number of adherent ECs, and an increase of the total focal adhesion area (as measured using paxillin) in EC [29].

Following exposure to physiological WSS, the FAK stain became stronger and is easier to detect, showing an increase in the EC-matrix interaction. It is important to notice that this increase is probably not due to new formation of FAK but rather the recruitment and growth of existing FAK to the growing focal adhesion complexes. The antibody used for the FAK staining was a phosphorylated Y397 FAK (Abcam), suggesting that the increased stain was induced by increased phosphorylation of the FAK. Such phosphorylation was found in previous studies shortly after exposure to physiological WSS, and was accompanied by an increased amount of FAK at the focal adhesion sites [37,38].

Our finding of an increase in the amount of FAK runs counter findings reported by Davies et al, where no differences in the total adhesion area of EC to the matrix were found following the application of WSS [39]. This discrepancy can be attributed to the quantification methods. Davies et al quantified the area of the adhesion complex using digitized analysis of cells images, whereas we examined the images of a stained single protein in the adhesion complex. The increase we observed was due to a change in the protein complex. By contrast, in Davies’ work, no change in this complex was described. Moreover, our method may have been
more sensitive in assessing the adhesion plaques. However, Davies et al noticed the same increase in the size of the focal adhesion complex as we did [39].

In the 1-side model (Fig. 3ii) following exposure to WSS there was a dissociation of the VE-Cadherin stain (D–F). The stain thickness distribution, as measured from the images of the VE-Cadherin, in the different models with and without exposure to WSS. The thickness distribution in the EC monolayer was around 1 μm, whereas in the co-culture models it was roughly 2 to 3 μm. The difference across models was significance. B - The continuity in the vascular wall models, as estimated from the VE-Cadherin images, following exposure to a WSS of 12 dyne/cm² for 30 and 60 min and a WSS of 4 dyne/cm² for 30 min and without WSS exposure. VE-Cadherin continuity was high in static conditions and declined when the cells were exposed to a WSS of 12 dyne/cm², with a significant effect in the 1-side model alone following 60 min of exposure. Following exposure to a WSS of 4 dyne/cm² this continuity only declines significantly in the EC monolayer model. D - VE-Cadherin expression in the endothelial cells of the different models following exposure to 30 min of a WSS of 4 and 12 dyne/cm² and without exposure to WSS. The change in VE-Cadherin expression was significant when comparing the EC monolayer model to the two co-culture models; however the change in the amount of VE-Cadherin did not vary significantly in each model following exposure to WSS. The presented data is normalized to the EC-monolayer result in static conditions (control) sample. LSS - WSS of 4 dyne/cm² for 30 min; PSS30 - WSS of 12 dyne/cm² for 30 and PSS60 - of 12 dyne/cm² for 60 min. (± - P<0.05 compared to the same model without WSS. * - P<0.05 compared to the EC monolayer model following the same exposure to WSS).

doi:10.1371/journal.pone.0088304.g005

**Figure 5. VE-Cadherin analysis.** A - The VE-Cadherin marks the border of the cells, with a wider mark of the cell borders in the 1 and 2-side models (B and C). Following exposure to WSS there was a dissociation of the VE-Cadherin stain (D–F). B - The thickness distribution, as measured from the images of the VE-Cadherin, in the different models with and without exposure to WSS. The thickness distribution in the EC monolayer was around 1 μm, whereas in the co-culture models it was roughly 2 to 3 μm. The difference across models was significance. C - The continuity in the vascular wall models, as estimated from the VE-Cadherin images, following exposure to a WSS of 12 dyne/cm² for 30 and 60 min and a WSS of 4 dyne/cm² for 30 min and without WSS exposure. VE-Cadherin continuity was high in static conditions and declined when the cells were exposed to a WSS of 12 dyne/cm², with a significant effect in the 1-side model alone following 60 min of exposure. Following exposure to a WSS of 4 dyne/cm² this continuity only declines significantly in the EC monolayer model. D - VE-Cadherin expression in the endothelial cells of the different models following exposure to 30 min of a WSS of 4 and 12 dyne/cm² and without exposure to WSS. The change in VE-Cadherin expression was significant when comparing the EC monolayer model to the two co-culture models; however the change in the amount of VE-Cadherin did not vary significantly in each model following exposure to WSS. The presented data is normalized to the EC-monolayer result in static conditions (control) sample. LSS - WSS of 4 dyne/cm² for 30 min; PSS30 - WSS of 12 dyne/cm² for 30 and PSS60 - of 12 dyne/cm² for 60 min. (± - P<0.05 compared to the same model without WSS. * - P<0.05 compared to the EC monolayer model following the same exposure to WSS).
gap (2–3 μm) between the endothelial cells. This could indicate a higher permeability level of the monolayer, suggesting breaks in the tight junctions or even a leaky endothelium [17]. This is in line with the quantification of the lower amount of VE-Cadherin expression in the co-culture models, indicating lower connectivity between the endothelial cells. This suggests that the presence of SMC can lead to a more inflammatory phenotype of the HUVEC. A similar phenotype can be found in vivo in areas of disturbed flow and inflamed endothelium, where VE-Cadherin expression is weak and discontinuous; however, it is significantly higher and continuous where the flow is laminar. Moreover, the areas of disturbed flow in vivo have been shown to be correlated with high permeability to plasma protein and cholesterol, which highlight the contribution of VE-Cadherin to intercellular widening, and the high permeability to macromolecules in these areas [8,12]. In addition, Lavender et al. found a decrease in PECAM-1 staining in a 1-side model as compared to EC monolayer model which was related directly to the EC-SMC interactions. Since PECAM-1 and VE-Cadherin are both located at the contacts of EC-EC, this a reduction is consistent with our findings [20].

Following the application of physiological WSS the VE-Cadherin stain ceased to be continuously distributed at the periphery of the cells and appeared in clusters at the cell borders. This was consistent in all 3 models; however, in the 1-side model the effect seemed weaker (Fig. 5C) and the discontinuity was only significant after longer exposure to WSS (i.e. 60 min). A discontinuity in VE-Cadherin staining of this type was found in vitro previously, following short exposure to disturbed pulsatile flow or following long exposure to reciprocating flow and physiological WSS [8,11]. This change was not accompanied by a change in VE-Cadherin expression [8–10]; however there was a change in the cellular distribution of VE-Cadherin within the cell, allowing for cell alignment while still attached to each other [10]. Crucially, VE-Cadherin expression did not change within the models following exposure to the different WSS; hence the difference between the co-culture models and the EC monolayer model was still present following exposure to WSS.

Following exposure to 4 dyne/cm² the discontinuity was not as strong as after exposure to 12 dyne/cm² in all 3 models; however, in both co-culture models this change was not significant compared to the same model in static conditions. This suggests that the EC-SMC interactions might change the EC response to WSS and the alignment sequence of the cells. In addition, the change in the focal adhesion composition could possibly be attributed to the reduced amount of fenestration in the stains of the VE-Cadherin following exposure to 12 dyne/cm² in the 1-side model (Fig. 5A–E) compared to the other 2 models. Moreover, WSS was found to induce FAK phosphorylation and to increase the trans-electrical resistance of EC monolayer, suggesting that FAK may play a role in EC permeability [40]. This is supported by a recent finding where inhibition of FAK activity resulted in a leaky vessel, abnormal VE-Cadherin distribution and reduced VE-Cadherin phosphorylation. On the other hand, activation of FAK was related to increased endothelium barrier properties [41]. These data suggest that FAK is important in the control of cell-cell junctions and the barrier properties of the EC monolayer. It also provides a possible mechanism for the reduced expression of VE-Cadherin in the 1-side model and the increased thickness of the VE-Cadherin stain in this model.
Previous studies have shown that Cx37 is up-regulated when exposed to physiological WSS whereas Cx43 is up-regulated following exposure to pathological WSS, and hence exhibits different response to WSS stimuli of the different connexins, with independent regulation mechanisms [35]. Johnson and Nerem used a model of SMC in collagen gel and found an up-regulation in Cx37 expression in the presence of SMC which was even higher following exposure to WSS. In the case of Cx43, there was no protein expression in the EC monoculture either with or without flow. By contrast, in the presence of SMC there was high expression of the protein, which was down-regulated by the flow. These results demonstrate that the presence of SMC increases the expression of connexins in EC, which is probably due to the role of the connexins in MEGJ [35]. In our system, the amount of both Cx43 and Cx37 increased in the presence of SMC prior to exposure to WSS, suggesting that the excess amount of connexins was used to form MEGJ with the SMC. The amount of Cx43 was significantly higher in the 1-side model, which proves that the introduction of the membrane, which increases the distance between the cells, creates a barrier between the two cell types; however, such a difference was not found with Cx37.

The effect of exposure to WSS on the EC monolayer model was a 1.5 fold elevation in the amount of connexins in all WSS modes (partly for Cx37 in the case of a WSS of 12 dyne/cm²). Since the exposure time to WSS was only 30 min, this change is likely to correspond to the transient change found following the onset of the flow [42]. On the other hand, in the presence of SMC, with MEGJ formed, the exposure to a WSS of 12 dyne/cm² did not induce any change in the amount of connexins. A possible explanation is that the higher amount of connexins from the MEGJ needs a longer exposure to WSS prior for a change to occur or that following exposure of the WSS the change in MEGJ requires longer exposure to WSS compared to EEGJ. A possible reason is that the connexins may move from MEGJ to EEGJ as an immediate response, before any change in the total amount of the protein takes place. Moreover, the amount of Cx37 in the EC monolayer following exposure to WSS of 12 dyne/cm² appeared to have been equivalent to that found in the co-culture models without the WSS stimulation. It is important to note that in the EC monolayer model, Cx37 and Cx43 form gap junctions between the EC; thus differences in amount are indicative of the connectivity and communication of EC-EC (i.e. EEGJ).

Subsequent to a WSS of 4 dyne/cm² the response of both co-culture models was consistent with what has been reported in previous works. The elevation of Cx43 expression was higher in the 2-side model than in the 1-side model; however it was the highest in the EC monolayer model. On the other hand, although Cx37 expression was down-regulated in the co-culture models, and was the lowest in the 1-side models, it was elevated compared to the control in the EC monolayer model. These changes in the magnitudes of the response show that the response of the 2-side model had more similarities with the EC monolayer response.

In our system we found that the presence of SMC changes the response of the cells to flow. However, the presence of a membrane between the cells apparently reduces the effect of the presence of SMC in both the connexins and the VE-Cadherin.
response. This change could not be ascribed to the changes in the amount of contact between the cells, since the change in the amount of Cx43 between the models was not notably high. On the other hand, the membrane found between the two cell types in the 2-side model increases the distance between the cells, so that the diffusion time of the secondary messengers’ increases, hence limiting or delaying the response of the cells. In addition, this change could be due to the different local adhesion complex formed by the EC on the SMC layer as compared to the FA complex on the membrane.

In vitro, the presence of MEGJ has been demonstrated in different animal models and different vascular sites; however, the frequency of MEGJ increases with decreasing vessel size [43–45]. Furthermore, their presence in some large arteries is not considered known [46,47]. In addition, the MEGJ function is more important in resistance arteries rather than in conduit vessels because of their participation in mediated endothelium dependent hyperpolarization [48]. Taken together, these observations suggest that the effect of WSS on the EC-SMC interactions through the MEGJ is an important factor in vascular functionality; however, the differences in the cellular interactions in the different models simulate the in vivo interaction found in different types of blood vessels; i.e. small arterioles would be better demonstrated using the 1-side model, whereas larger arteries would be better demonstrated using the 2-side model.

Conclusion

The data presented in this study demonstrate the effect of WSS on the EC-matrix, EC-EC and EC-SMC interactions. This was studied by measuring the changes in FAK, endothelial MEGJ and adherens junction proteins in response to WSS in two co-culture models and a control model of the EC monolayer. This is the first time that a reduction in the EC-EC interactions has been found in vitro in the presence of SMC, in both static conditions and following exposure to WSS, as demonstrated by the amount of VE-Cadherin. This was accompanied by increased distance between the endothelial cells, which suggest a less connected and more permeable EC monolayer that resembles the inflammatory endothelium in vivo. In addition, the amount of Cx37 and Cx43 increased in the presence of SMC, indicating the formation of direct EC-EC interactions through the formation of MEGJ. Following exposure to both physiological and low WSS, EC-EC communication through EEGJ increased in both the Cx37 and the Cx43 in the EC monolayer model. The EC-SMC interactions, however, did not vary following exposure to physiological WSS. Low WSS, on the other hand, increased EC-SMC interactions through Cx43 expression, whereas it reduced EC-SMC interactions through Cx37. The magnitude of this response varied between the different co-culture models, suggesting that the distance introduced by the membrane in the 2-side model reduces the EC-SMC interactions through Cx43. The response in the 2-side model was more similar to the EC-EC connexin interactions. These results show that EC-SMC interactions affect EC phenotype as well as the response of the cells to WSS and that the interactions formed between the cells depends on the distance between them. Altogether, this highlights the importance of cellular interactions to vascular wall functionality, and the importance of the interactions formed when simulating different types of blood vessels.

Author Contributions

Conceived and designed the experiments: DS DE SE. Performed the experiments: DS UZ RG. Analyzed the data: DS. Contributed reagents/materials/analysis tools: DS RG UZ. Wrote the paper: DS.

References

1. Ando J, Yamamoto K (2009) Vascular mechanobiology: endothelial cell responses to fluid shear stress. Circ J 73: 1983–1992.
2. Hahn C, Schwartz MA (2009) Mechanotransduction in vascular physiology and atherosclerosis. Nat Rev Mol Cell Biol 10: 53–62.
3. Chia JJ, Chien S (2011) Effects of disturbed flow on endothelial cells. Cell Commun Adhes 12: 261–270.
4. Wang YH, Yan ZQ, Qi YX, Cheng BB, Wang XD, et al. (2010) Normal shear stress and vascular smooth muscle cells modulate migration of endothelial cells through histone deacetylase 6 activation and tubulin acetylation. Ann Biomed Eng 38: 729–737.
5. Antounelli-Orridge A, Saunders KB, Smith SR, J’Amore PA (1989) An activated form of transforming growth factor beta is produced by cocultures of endothelial cells and pericytes. Proc Natl Acad Sci U S A 86: 4344–4348.
6. Fillinger MF, Sampson LN, Cronenwett JL, Powell RJ, Wagner RJ (1997) Coculture of endothelial cells and smooth muscle cells in bilayer and conditioned media models. J Surg Res 67: 169–178.
7. Truskey GA (2010) Endothelial Cell Vascular Smooth Muscle Cell Co-Culture Assay For High Throughput Screening Assays For Discovery of Anti-Angiogenesis Agents and Other Therapeutic Molecules. Int J High Throughput Screen 2010: 171–181.
8. Miao H, Hu XL, Shao YT, Yuan S, Zhao Y, et al. (2005) Effects of flow patterns on the localization and expression of VE-cadherin at vascular endothelial cell junctions: in vivo and in vitro investigations. J Vasc Res 42: 77–89.
9. Uropeo JA, Hollinger MK, Woolfalis MJ (2002) Regulation of VE-cadherin linkage to the cytoskeleton in endothelial cells exposed to fluid shear stress. Exp Cell Res 275: 240–247.
10. Shay-Salit A, Shushy M, Wolfowitz E, Yahav H, Breviario F, et al. (2002) VEGER receptor 2 and the adherens junction as a mechanical transducer in vascular endothelial cells. Proc Natl Acad Sci U S A 99: 9462–9467.
11. Noria S, Cowan DR, Gotlib AI, Langille BL (1999) Transient and steady-state effects of shear stress on endothelial cell adherens junctions. Circ Res 85: 504–514.
12. Chien S (2008) Effects of disturbed flow on endothelial cells. Ann Biomed Eng 36: 534–562.
13. Kwak BR, Silacci P, Stergiopoulos N, Hayoz D, Meda P (2005) Shear stress and cyclic circumferential stretch, but not pressure, alter connexin43 expression in endothelial cells. Cell Commun Adhes 12: 261–270.
14. García JE, Paul DL (1998) Connexin43 is highly localized to sites of disturbed flow in rat aortic endothelium but connexin37 and connexin40 are more uniformly distributed. Circ Res 83: 636–643.
15. Figureau SX, Duling BR (2009) Gap junctions in the control of vascular function. Annu. Rev Neurosci Signal 11: 231–266.
16. Tarbell JM (2003) Mass transport in arteries and the localization of astrocytosis. Annu Rev Biomed Eng 5: 79–118.
17. Tarbell JM (2010) shear stress and the endothelial transport barrier. Cardiovasc Res 87: 320–330.
18. Warboys CM, Eric Berson R, Mann GE, Pearson JD, Weinberg PD (2010) Acute and chronic exposure to shear stress have opposite effects on endothelial permeability to macromolecules. Am J Physiol Heart Circ Physiol 298: H1850–H1856.
19. Tsima E, Irani-Tehrani M, Kiosses WB, Dejana E, Schultz DA, et al. (2005) A mechanosensory complex that mediates the endothelial cell response to fluid shear stress. Nature 437: 426–431.
20. Lavender MD, Pang Z, Wallace GS, Niklasson LE, Truskey GA (2005) A system for the direct co-culture of endothelium on smooth muscle cells. Biomaterials 26: 4642–4653.
21. Chen CN, Chang SF, Lee PL, Chang K, Chen LJ, et al. (2006) Neutrophils, lymphocytes, and monocytes exhibit diverse behaviors in transendothelial and subendothelial migrations under coculture with smooth muscle cells in disturbed flow. Blood 107: 1933–1942.
22. Chia JJ, Chen LJ, Lee PL, Lee CI, Lo LW, et al. (2003) shear stress inhibits adhesion molecule expression in vascular endothelial cells induced by coculture with smooth muscle cells. Blood 101: 2667–2674.
23. Helenius G, Hagwall SH, Euguerra M, Fink H, Soderberg R, et al. (2008) Effect of shear stress on the expression of coagulation and fibrinolytic factors in both smooth muscle and endothelial cells in a co-culture model. Eur Surg Res 40: 322–337.
24. Wang HQ, Huang LX, Qu MJ, Yan ZQ, Liu B, et al. (2006) Shear stress protects against endothelial regulation of vascular smooth muscle cell migration in a coculture system. Endothelium 13: 171–180.
25. Heyderman-Hagwall S, Chiu JJ, Grander S, Welander S, Li YC, Yuan S, et al. (2006) DNA microarray study on gene expression profiles in co-cultured endothelial and smooth muscle cells in response to 4- and 24-h shear stress. Mol Cell Biochem 201c: 1–15.
46. Tolsa JF, Marino M, Peyter AC, Beny JL (2006) Role of membrane potential in
45. Heberlein KR, Straub AC, Isakson BE (2009) The myoendothelial junction:
44. Sandow SL, Hill CE (2000) Incidence of myoendothelial gap junctions in the
41. Zebda N, Dubrovskyi O, Birukov KG (2012) Focal adhesion kinase regulation of
38. Li S, Kim M, Hu YL, Jalali S, Schlaepfer DD, et al. (1997) Fluid shear stress
37. Li S, Butler P, Wang Y, Hu Y, Han DC, et al. (2002) The role of the dynamics of
35. Johnson TL, Nerem RM (2007) Endothelial connexin 37, connexin 40, and
34. Heberlein KR, Straub AC, Best AK, Greyson MA, Looft-Wilson RC, et al. (2010) Site-
specific connexin phosphorylation is associated with reduced heterocellular communication between smooth muscle and endothelium. J Vasc Res 47: 277–286.
33. Gairhe S, Bauer NN, Gebb SA, McMurtry IF (2011) Myoendothelial gap
junction influences gap junction organization. Circ Res 97: 44-51.
31. Straub AC, Johnstone SR, Heberlein KR, Rizzo MJ, Best AK, et al. (2010) Site- selective regulation of connexins 43 and 45. Am J Physiol Lung Cell Mol Physiol 301: L527–535.
30. Isaakson BE, Duling BR (2005) Heterocellular contact at the myoendothelial junction influences gap junction organization. Circ Res 97: 44-51.
29. Wang YH, Yan ZQ, Shen BR, Zhang L, Zhang P, et al. (2009) Vascular smooth muscle cells promote endothelial cell adhesion via microtubule dynamics and activation of paxillin and the extracellular signal-regulated kinase (ERK) pathway in a co-culture system. Eur J Cell Biol 88: 701–709.
28. Wallace CS, Champion JC, Truskey GA (2007) Adhesion and function of human endothelial cells co-cultured on smooth muscle cells. Ann Biomed Eng 35: 375–386.
26. Hastings NE, Simmers MB, McDonald OG, Wamhoff BR, Blackman BR (2007) Atherosclerosis-prone hemodynamics differentially regulates endothelial and smooth muscle cell phenotypes and promotes pro-inflammatory priming. Am J Physiol Cell Physiol 293: C1824–1833.
24. Tolsa JF, Marino M, Peyter AC, Beny JL (2006) Role of membrane potential in
23. Rizzo MJ, Best AK, Straub AC, Heberlein KR, Johnstone SR, et al. (2010) Site- selective regulation of connexins 43 and 45. Am J Physiol Lung Cell Mol Physiol 301: L527–535.
22. Straub AC, Hilliard M, Johnstone SR, Best AK, Yemen S, et al. (2006) Custom-designed wells and flow chamber for exposing air-liquid interface cultures to wall shear stress. Ann Biomed Eng 34: 1890–1895.
20. Li S, Butler P, Wang Y, Hu Y, Han DC, et al. (2002) The role of the dynamics of focal adhesion kinase in the mechanotransduction of endothelial cells. Proc Natl Acad Sci U S A 99: 3546–3551.
19. Li S, Kim M, Hu YL, Jalali S, Schlaepfer DD, et al. (1997) Fluid shear stress activation of focal adhesion kinase. Linking to mitogen-activated protein kinases. J Biol Chem 272: 30455–30462.
18. Davies PF, Robotowsky A, Grön ML (1994) Quantitative studies of endothelial cell adhesion. Directional remodeling of focal adhesion sites in response to flow forces. J Clin Invest 93: 2031–2038.
15. Shikata Y, Kass A, Kawkhuwarong K, DePaola N, Garcia JG, et al. (2005) Differential effects of shear stress and cyclic stretch on focal adhesion remodeling, site-specific FAK phosphorylation, and small GTPases in human lung endothelial cells. Exp Cell Res 304: 40–49.
14. Zebda N, Dubrovskyi O, Birukov KG (2012) Focal adhesion kinase regulation of mechanotransduction and its impact on endothelial cell functions. Microvasc Res 83: 71–81.
13. DePaola N, Davies PF, Prichard WF, Jr, Flores L, Harbeck N, et al. (1999) Spatial and temporal regulation of gap junction connexin43 in vascular endothelial cells exposed to controlled disturbed flows in vitro. Proc Natl Acad Sci U S A 96: 1315–1319.
12. Sandow SL, Tare M, Coleman HA, Hill CE, Parkington HC (2002) Involvement of myoendothelial gap junctions in the actions of endothelem-derived hyperpolarizing factor. Circ Res 90: 1108–1113.
11. Sandow SL, Hill CE (2000) Incidence of myoendothelial gap junctions in the proximal and distal mesenteric arteries of the rat is suggestive of a role in endothelium-derived hyperpolarizing factor-mediated responses. Circ Res 86: 341–346.
10. Heberlein KR, Straub AC, Isaakson BE (2009) The myoendothelial junction: breaking through the matrix? Microcirculation 16: 307–322.
9. Tolsa JF, Marino M, Peyter AC, Bengt JL (2006) Role of membrane potential in endothelium-dependent relaxation of isolated mouse main pulmonary artery. J Cardiovasc Pharmacol 47: 501–507.
8. de Wit C, Boettcher M, Schmidt VJ (2008) Signaling across myoendothelial gap junctions–fact or fiction? Cell Commun Adhes 15: 231–245.
7. Triggle CR, Samuel SM, Ravishanker S, Maeri S, Arunachalam G, et al. (2012) The endothelium: influencing vascular smooth muscle in many ways. Can J Physiol Pharmacol 90: 713–738.
6. Saez-Bach J, Donnert G, Kromer B, Werth S, Wojcik-Stothard B, et al. (2007) Regulation of endothelial barrier function during fluid-induced conversion to an arterial phenotype. Cardiovasc Res 75: 596–607.
5. Ebong EE, Kim S, DePaola N (2006) Flow regulates intercellular communication in HAE/HCC by activating functional Cx40 and Cx37 gap junctional channels. Am J Physiol Heart Circ Physiol 290: H2013–2025.
4. Fujitani K (2006) Platelet endothelial cell adhesion molecule-1 and mecha
otransduction in vascular endothelial cells. J Intern Med 259: 373–380.
3. Ding Z, Fan Y, Deng X (2009) Water filtration rate and infiltration/ accumulation of low density lipoproteins in 3 different modes of endothelial/ smooth muscle cell co-cultures. Sci China Life Sci 52: 1023–1029.
2. Sakai J, Karino T, Niwa K (2008) Flow-dependent accumulation of LDL in co-cultures of endothelial and smooth muscle cells in the presence of filtration flow through the cell layer. Clin Hemorheol Microcirc 36: 245–256.
1. Wallace CS, Truskey GA (2010) Direct-contact coculture between smooth muscle and endothelial cells inhibits TNF-alpha-mediated endothelial cell activation. Am J Physiol Heart Circ Physiol 299: H338–346.

58. Cao XF, Qian GL, Hu BS, Liu FQ (2010) [Prediction of potential suitable distribution area of Flaveria bidens in China based on niche models]. Ying Yong Sheng Tai Xue Bao 21: 3063–3069.
57. Feng JF, Cao J, Li X, Li M, Shandas R, et al. (2012) A new flow indicator for the analysis of endothelial cell-bound Jagged1 in Notch3-induced human coronary artery smooth muscle cell adhesion formation in coculture. Ann Biomed Eng 40: 410–416.