Platelet-derived microvesicles induce calcium oscillations and promote VSMC migration via TRPV4

Shan-Shan Li1,2#, Shuang Gao1#, Yi Chen1, Han Bao1, Zi-Tong Li1, Qing-Ping Yao1, Ji-Ting Liu1, Yingxiao Wang352, Ying-Xin Qi1,4,5#

1. Institute of Mechanobiology & Medical Engineering, School of Life Sciences & Biotechnology, Shanghai Jiao Tong University, Shanghai, China
2. School of Perfume and Aroma Technology, Shanghai Institute of Technology, Shanghai, China
3. Department of Bioengineering, Institute of Engineering in Medicine, University of California, San Diego, San Diego, United States
4. Key Laboratory for Biomechanics and Mechanobiology of Ministry of Education, School of Biological Science and Medical Engineering, Beihang University, Beijing, 100083, China
5. Beijing Advanced Innovation Center for Biomedical Engineering, Beihang University, Beijing, 100083, China

# These authors contributed equally to this manuscript.

© The author(s). This is an open access article distributed under the terms of the Creative Commons Attribution License (https://creativecommons.org/licenses/by/4.0/).
See http://ivyspring.com/terms for full terms and conditions.

Received: 2020.04.19; Accepted: 2020.11.18; Published: 2021.01.01

Abstract

**Rationale:** Abnormal migration of vascular smooth muscle cells (VSMCs) from the media to the interior is a critical process during the intimal restenosis caused by vascular injury. Here, we determined the role of platelet-derived microvesicles (PMVs) released by activated platelets in VSMC migration.

**Methods:** A percutaneous transluminal angioplasty balloon dilatation catheter was used to establish vascular intimal injury. Collagen I was used to activate PMVs, mimicking collagen exposure during intimal injury. To determine the effects of PMVs on VSMC migration *in vitro*, scratch wound healing assays were performed. Fluorescence resonance energy transfer was used to detect variations of calcium dynamics in VSMCs.

**Results:** Morphological results showed that neointimal hyperplasia was markedly increased after balloon injury of the carotid artery in rats, and the main component was VSMCs. PMVs significantly promoted single cell migration and wound closure *in vitro*. Fluorescence resonance energy transfer revealed that PMVs induced temporal and dynamic calcium oscillations in the cytoplasm of VSMCs. The influx of extracellular calcium, but not calcium from intracellular stores, was involved in the process described above. The channel antagonist GSK219 and specific siRNA revealed that a membrane calcium channel, transient receptor potential vanilloid 4 (TRPV4), participated in the calcium oscillations and VSMC migration induced by PMVs.

**Conclusions:** TRPV4 participated in the calcium oscillations and VSMC migration induced by PMVs. PMVs and the related molecules might be novel therapeutic targets for vascular remodeling during vascular injury.

Introduction

Vascular intimal injury occurs following cardiovascular disease treatments, such as bypass surgery, coronary vein graft, angioplasty and stent treatment [1]. As a consequence, intimal hyperplasia, even postangioplasty restenosis, may contribute to treatment failure [2,3]. Vascular smooth muscle cells (VSMCs), the dominant cellular component of arteries, are primarily present within the media layer of vessels under physiological conditions. However, it has been well established that abnormal VSMC media-to-intima migration is an important cellular process during intimal hyperplasia [4-6]. Therefore, studying the mechanism of abnormal VSMC migration during intimal hyperplasia is of great significance for improving the cure rate of cardiovascular disease.
Adult VSMCs retain the potential to alter their migrative properties, and this process can be regulated by extracellular matrix components, peptide growth factors, cytokines, RNA molecules, mechanical factors, ion signaling and other environmental cues [7-9]. For example, metalloproteinase-2 (MMP2) and metalloproteinase-9 (MMP9) participate in VSMC migration from the media to the intima following arterial injury and alter postinjury vascular remodeling [10,11]. Wu et al. revealed that kindlin-2 plays a critical role in VSMC proliferation, migration and intimal hyperplasia via Wnt signaling, and blocking the activity of kindlin-2 is an attractive therapeutic approach for vascular injury [12]. Blood flow and shear stress have also been found to abrogate the proliferative and migratory response of VSMCs in the early stages after injury [13,14]. In addition to these regulatory responses, numerous studies have revealed that the regulation of cell migration is critically dependent on calcium. Chemoattractants stimulate neutrophil migration by inducing repeated transient increases in intracellular calcium level [15]. Excitatory neurotransmitters initiate cell contraction, which is the key process in cell migration, by interacting with cell surface receptors to generate inositol 1,4,5-trisphosphate (IP3) [16]. IP3 binds to inositol 1,4,5-trisphosphate receptors (IP3Rs) on the sarcoplasmic reticulum to trigger calcium release, which sustains elevated levels of cytoplasmic calcium to regulate colonic smooth muscle contraction.

Although the studies described above have revealed many important factors that affect VSMC migration, the underlying mechanisms during intimal hyperplasia are still not fully understood. During vascular intimal injury, the adherence and accumulation of circulating platelets to the injured intima is an important pathological process [17,18]. Circulating platelets can be activated by exposure to collagen (mainly collagen I), which is caused by the injury of monolayer endothelial cells (ECs). Activated platelets can release a variety of heterogeneous platelet-derived microvesicles (PMVs) [17], which are mainly 100-1000 nm in diameter [19-21]. PMVs are capable of selectively carrying different types of biomolecules, such as membrane and cytoplasm proteins, lipids, RNAs, and other bioactive molecules, and transferring these biomolecules to recipient cells, thus participating in the regulation of recipient cell functions [17,18,22]. It has been reported that thrombin/collagen-induced PMVs can enhance the potential of early outgrowth cells (EOCs) to restore endothelial integrity by transferring chemokine receptor-4 (CXCR4) to EOCs after vascular injury [23]. PMVs have been reported to play critical roles in various cardiovascular diseases, including hypertension, atherosclerosis and thrombin formation [18,24]. Under conditions of intimal injury, the internal elastic lamina can be broken, and PMVs can thus directly contact the middle layer [25]. Even under conditions of mild intimal injury, the EC layer can be broken, and the fenestrae on the internal elastic lamina (1~3 µm in width) [26] can allow PMVs (10~1000 nm in diameter) to diffuse into the middle layer. However, the relationship between PMVs and VSMC migration during intimal hyperplasia remains poorly characterized. In this study, collagen I was used to activate PMVs, mimicking collagen exposure during intimal injury, and the roles of PMVs in VSMC migration were demonstrated. Our bioinformatic analysis, which was based on previously published proteomic data of collagen-induced PMVs [27], suggested that calcium may be the key node in the regulatory network of PMV-induced cell migration. We investigated the calcium dynamics induced by PMVs in live VMSCs at the single-cell and subcellular levels by using fluorescence resonance energy transfer (FRET) combined with a calcium biosensor. We then further identified the potential calcium channels that participate in PMV-induced calcium oscillations in VSMCs.

Materials and Methods

Animal model

The animal care and experimental protocols were conducted in accordance with the Animal Management Rules of China (55, 2001, Ministry of Health, China), and the study was approved by the Animal Research Committee of Shanghai Jiao Tong University.

Male Sprague Dawley rats were housed in a temperature-controlled room with a 12-h light/dark cycle and were given access to standard chow and tap water ad libitum. Vascular injury was established in 8-week-old rats, as previously described [3]. Briefly, a percutaneous transluminal angioplasty balloon dilatation catheter (Boston Scientific Corporation, Galway, Ireland) was used to establish vascular intimal injury. All the animals were anesthetized by isoflurane inhalation and treated under sterile conditions. The balloon was placed into the left carotid artery and repeatedly stretched to damage the blood vessels. The undamaged right carotid artery served as the self-control [3].

HE and immunofluorescence staining

Four weeks after balloon injury, the arteries were removed, fixed in 4% paraformaldehyde solution, embedded, and finally cut into 8-µm sections. HE staining [28] or immunofluorescence staining [29] was
performed as previously described. For immunofluorescence staining, the paraffin-embedded sections were permeabilized with 0.1% Triton X-100 for 10 min. After treatment with phosphate-buffered saline (PBS) containing 10% goat serum for 1 h, the sections were incubated with primary rabbit anti-von Willebrand Factor antibody (1:200, Dako, Copenhagen, Denmark A/S) and mouse monoclonal anti-α-SMA antibody (1:200, Dako, Copenhagen, Denmark A/S) for 24 h at 4°C. Secondary anti-mouse/rabbit IgG antibodies (1:1000, Cell Signaling Technology, Boston, MA) were used. The nuclei were stained with DAPI after immunofluorescence staining. The fluorescence images were acquired by using a fluorescence microscope (Olympus, Tokyo, Japan).

PMV isolation, collection and size analysis
PMVs were isolated according to the previous literatures [30,31]. As shown in Supplementary Figure 1A, whole blood was collected from the abdominal aorta of anesthetized rats into a syringe containing 100 μL/mL anticoagulant in 0.9% sodium chloride solution. The blood was centrifuged at 1500 rpm for 10 min to obtain platelet-rich plasma and then centrifuged at 2800 rpm for 15 min to obtain platelets. The platelets were resuspended using Tyrode solution. The blood was centrifuged at 2800 rpm for 15 min to remove the platelets, and the PMVs were collected at 20500 g/min for 90 min. The PMVs were analyzed with a flow cytometry column (FACSCalibur TM; BD Biosciences) combined with a platelet-specific marker (anti-rat CD41) to demonstrate the platelet origin[30,31], and with a NanoSight3000 high-sensitivity detection system (Malvern Panalytical, Malvern, England) to analyze the particle size.

PMV treatment
After the carotid artery balloon injury model was established, PMVs (6×10⁹ per ml) or normal saline (as a control group) were injected into the tail veins of the rats on alternate days for 2 weeks after the surgery. On day 14 after the surgery, the injured carotid artery was collected.

Cell culture and transfection
VSMCs were isolated from the thoracic aortas of male Sprague Dawley rats via an explant method, as previously described [32]. The VSMCs were cultured in Dulbecco’s modified Eagle’s medium (DMEM) (GIBCO, NY, USA) with 10% calf serum (FCS, GIBCO, NY, USA), 2 mM glutamine (GIBCO, NY, USA), 100 U/mL penicillin and 100 μg/mL streptomycin (BBI Company, Shanghai, China) at 37°C in a 5% CO₂ atmosphere. The VSMCs were characterized by antibody specific for α-smooth muscle actin (1:1000, Dako, Copenhagen, Denmark A/S). In all the experiments, the purity of the cell populations was universally higher than 95%, and only VSMCs between passages 4 and 7 were used.

For the transfection process, a FRET-based calcium biosensor, which includes a pair of fluorescent proteins (ECFP and YPet) and calmodulin-linked light chain protein kinase M13, was utilized. The structure of the biosensor is altered after binding to calcium ions [33]. Adenovirus vectors (Genepharma, Shanghai, China) were used to introduce the reconstructed plasmid containing the FRET-based calcium biosensor into VSMCs.

Microscopy, image acquisition and analysis
Cells were starved in DMEM without FCS for 6 h before PMV stimulation. Then, PMVs were gently added to the culture dish. During the imaging experiments, the cells were maintained in streptomycin-free medium to prevent possible effects on calcium ion channels. All the images were obtained by using a Leica inverted microscope (DMi8, Wetzlar, Germany) equipped with a charge-coupled device (CCD) camera (Andor iXon 897, Belfast, UK) and two emission filters controlled by a filter changer (480DF20 for ECFP and 535DF15 for YPet). During the image capturing process, a temperature-control system with CO₂ supplement was used to maintain cellular viability. Time-lapsed fluorescence images were acquired at 30-s intervals by Leica LASX software (Leica Biosystems GmbH). The emission ratio of FRET/ECFP was directly computed and analyzed. The max ratio and calcium oscillation frequency were further analyzed.

Scratch wound healing assay
To assess VSMC migration in vitro, scratch wound healing assays were performed. A wound healing cell migration assay was performed using 95% confluent cells, as described in previous studies [34,35]. A line was scratched across a monolayer of cells using a sterile 10-μL pipette tip. Images of the scratched line were immediately captured, and the cells were imaged again after 12 h and 24 h (4X objective, IX-71, Olympus, Japan). The wound area was measured using ImageJ software (NIH, USA). The cell migration abilities were calculated as (S₀ - Sₜ)/S₀, where S₀ is the wound area at the initial time point, and Sₜ is the wound area at the observation time point t (12 h or 24 h).
Solutions and chemicals

For the experiments that required calcium-free conditions, calcium-free DMEM (GIBCO, NY, USA) was used. The chemical reagents 2-aminoethoxydiphenyl borate (2-APB) (0.1 mM) [36] and nifedipine (10 μM) [37] were purchased from Sigma-Aldrich (St. Louis, MO). GSK219 (0.1 mM) [38] was obtained from Merck (NY, USA). All the drugs were preincubated with VSMCs for 20 min before the addition of the PMVs. The amount of drug administered was based on previous publications.

RNA interference

For RNA interference, VSMCs were transfected with 50 nM small interfering RNA (siRNA) fragments (si-1/2/3) or control nonsilencing siRNA (si-NC) (Genepharma, Shanghai, China) for 24 h using Lipofectamine 2000 (Invitrogen, Carlsbad, CA) according to the manufacturer’s instructions [39]. The sequences of the small interfering RNA fragments targeting TRPV4 (sequence accession number NM_023970.1) and si-NC are listed in Table S1.

Western blot

Lysates were separated by 10% SDS-PAGE. The proteins were detected using primary antibodies against TRPV4 (1:500, Alomone Labs, Jerusalem, Israel) and GAPDH (1:1000, Protein tech, Beijing, China). An HRP-labeled IgG was used as the secondary antibody at a 1:1000 dilution, and the bands were visualized using an ECL kit (Beyotime, Shanghai, China) and quantified with Quantity One software (Bio-Rad, Hercules, CA, USA).

Statistical analysis

All the values are expressed as the mean ± standard error (S.E.M.) of the mean, and all the data are consistent with variance after R studio software. The statistical analysis of the data was performed by unpaired Student’s t-test to identify significant differences between two mean values, and a value of P < 0.05 was considered statistically significant. The statistical analysis was performed using Excel (Microsoft Corporation, Washington, USA) and GraphPad 9.0 (GraphPad Software, San Diego, CA).

Results

Vascular intimal injury promotes VSMC migration and neointimal hyperplasia

Four weeks after intimal injury, HE staining revealed that compared with that of the self-contralateral noninjured artery, the neointimal layer of the intimal injured artery was markedly thickened, and the area of the vascular cavity was markedly reduced (Figure 1A). In addition, immunofluorescence staining revealed the expression of α-smooth muscle actin (α-SMA) in neointimal hyperplasia, which suggested that the main component of the vascular neointima was VSMCs (Figure 1B). During vascular intimal injury, collagen exposure activates platelets, which accumulate and release a variety of heterogeneous vesicles, including PMVs. *In vivo* immunofluorescence staining for CD41 (a marker of PMVs [40]) and α-SMA (a marker of VSMCs [41]) showed that CD41-positive particles were closely adjacent to α-SMA-positive cells, indicating that PMVs accumulate at the injured artery and contact VSMCs *in vivo* (Figure 1C). We then demonstrated the role of PMVs in VSMC migration and the potential mechanisms.

After using collagen I to simulate the pathological exposure of extracellular matrix components during intimal injury, PMVs were extracted, and the diameter of the PMVs was further examined by NanoSight3000. The results illustrated that the sizes of the PMVs were distributed within a range of 100~600 nm, and there were 3 peaks at 121.6 nm, 183.0 nm and 291.5 nm (Figure S1).

PMVs can adhere to VSMCs *in vitro* (Figure S2A). Our scratch wound data showed that PMVs induced a marked increase in the wound repair capabilities of VSMCs at 12 h and 24 h (Figure 1D). The averaged wound closure areas at 24 h are 78.42% and 29.64% smaller than the initial area at 0 h for the PMVs group and the control, respectively (Figure 1E). Moreover, the track results of the free migration of single cells revealed that compared with the control (Figure S2B, Video S1A), PMV treatment increased the migration distance of single VSMCs in a time-dependent manner, and a significant increase was detected at approximately 10.5 h (Figure S2B, Video S1B).

These results suggested that the PMVs released by collagen I-activated platelets promote the migration of VSMCs, which may participate in the intimal hyperplasia caused by intimal injury *in vivo*. We further conducted *in vivo* experiments to investigate PMV-induced neointimal hyperplasia in arteries. First, a carotid artery balloon injury model was established. Then, PMVs (6×10⁹ per 1 mL) or normal saline (as a control group) were injected through the tail veins of the rats on alternate days for 2 weeks after the surgery. Fourteen days after the surgery, the injured carotid artery was collected. Neointimal hyperplasia was examined with HE staining. As shown in Figure 1F, the thickness of the neointimal layer of the rats was considerably increased by PMVs.
**PMVs induce calcium oscillations in VSMCs**

Based on previously published proteomic data [27], IPA software was used to analyze the potential mechanism by which PMVs participate in VSMC migration. The results revealed that 31 proteins expressed in PMVs were correlated with calcium signaling (Figure 2A, Table S2), and cellular movement was one of the critical identified functions that are regulated by PMVs (Figure 2B). Then, the spatiotemporal characteristics of the calcium dynamics in VSMCs treated with PMVs were characterized with the aid of FRET real-time microscopy.

The representative heat map of the FRET/ECFP ratio showed that compared with the DMEM control,
PMVs triggered marked increases in the calcium levels in live VSMCs (Figure 2C, Video S2A-B). The average FRET/ECFP ratio increased by ~1.5-fold within 60 s and fluctuated at high levels for a period of 3000 s in VSMCs treated with PMVs (Figure 2D). Furthermore, the maximum ratio of FRET/ECFP in response to PMV treatment also significantly increased (1.3646 ± 0.0934 vs. 0.9869 ± 0.0149) (Figure 2E), and the frequency of calcium peaks upon PMV stimulation was significantly increased with those induced by the DMEM control (0.2733 ± 0.0681 vs. 0.0105 ± 0.0307) (Figure 2F). Emission spectral analysis showed that after PMV treatment, the ECFP emission peak at 475 nm decreased, and the YPet emission peak at 525 nm increased. The results revealed that PMVs enhanced YPet emission at the expense of ECFP emission, indicating a calcium-induced gain of FRET (Figure S3).

Calcium peaks that increase in amplitude modulation and frequency modulation are defined as calcium oscillations [42], and these results suggested that PMVs induced calcium oscillations in target VSMCs. There are two main sources of cytosolic calcium, the influx of calcium from the extracellular matrix and the release of calcium from intracellular stores; thus, the role of these two sources in PMV-induced calcium oscillation were explored in further studies.
Extracellular calcium participates in calcium oscillations and VSMC migration

To identify the calcium source, calcium-free medium was used to remove the extracellular calcium. The results showed that the calcium oscillations induced by PMV treatment were immediately abrogated in the calcium-free medium (Figure 3A, Video S3A), and the time-lapsed FRET/ECFP ratio line was horizontal and steady throughout the time course (Figure 3B). Compared with the DMSO control, PMV treatment led to significantly decreased frequencies of calcium peaks (0.0733 ± 0.0281 vs. 0.2733 ± 0.0681) and max FRET/ECFPs ratios (1.265 ± 0.0690 vs 1.8810 ± 0.1580) in the calcium-free medium (Figure 3C-D). Moreover, the wound healing assay revealed that the calcium-free medium also markedly reversed the migration of VSMCs induced by PMVs at 12 h and 24 h (Figure 3E-F).

Binding of IP$_3$ to IP$_3$Rs on the sarcoplasmic reticulum triggers the release of intracellular calcium [16]. Hence, 2-APB, an antagonist of IP$_3$Rs [36], was used to block the release of calcium from intracellular stores. The representative FRET/ECFP ratio maps are shown in Figure 3A (Video S3B-C). There was no significant difference in the average FRET/ECFP ratio (Figure 3B), the frequency of calcium peaks (0.2000 ± 0.014) (Figure 3C) or the max ratio (1.691 ± 0.1347) (Figure 3D) of the calcium dynamics between the
2-APB and DMSO control groups (Figure 3C-D).

These results suggested that oscillations were diminished by calcium-free medium, indicating that the influx of extracellular calcium was an important source of the calcium oscillations in the VSMCs.

TRPV4 channel, but not L-type voltage-gated calcium channel, mediates PMV-induced calcium oscillations

Since transient receptor potential vanilloid 4 (TRPV4) and L-type voltage-dependent calcium channel (L-VDCC) are widely reported to be abundantly expressed in VSMCs and related to important functions [42-44], the possible roles of these two channels that regulate the influx of extracellular calcium were further demonstrated.

GSK219 (GSK2193874, 0.1 mM), the specific antagonist of TRPV4, significantly reduced the calcium oscillations in response to PMV treatment (Figure 4A-D, Video S4B). Compared with the DMSO solvent control, GSK219 significantly decreased the max ratio of FRET/ECFP and the frequency of VSMC calcium peaks, and these values were 1.1770 ± 0.0635 vs. 1.8420 ± 0.1642 and 0.02 ± 0.0013 vs. 0.36 ± 0.098, respectively (Figure 4C-D, Video S4A-B). VSMC wound closure was suppressed at 12 h and 24 h by preincubation with GSK219 (Figure 4F). Nifedipine (Nife), the specific chemical inhibitor of L-VDCC, was used. Compared with the DMSO solvent control (Video S4A), Nife (10 μM) had no significant effect on the calcium oscillations (Figure 4A-B, Video S4C), the max ratio of FRET/ECFP (1.6341 ± 0.0633 vs. 1.8420 ± 0.1642) or the frequency (0.2901 ± 0.1056 vs. 0.36 ± 0.098) of the VSMC calcium peaks induced by PMVs. The wound closure rates induced by Nife and the DMSO solvent control, GSK219 significantly decreased the calcium oscillations and cell migration.

To further elucidate whether the effects of PMVs on VSMC migration and calcium oscillation were dependent on TRPV4, TRPV4 small interference RNA (siRNA) was transfected into VSMCs to knock down TRPV4. Then, FRET and wound healing were assessed. Three pairs of specific siRNAs were designed, and the most efficient siRNA, si-1, was identified (Figure 5A).

For further study, si-1 was labeled with the red fluorescent probe Cy3 (si-1-Cy3) and used to verify the effective silencing of TRPV4. Heat maps of the FRET/ECFP ratio revealed that in the si-1-Cy3 positive cells, the calcium oscillations were remarkably abolished and exhibited a steady state (Figure 5B, Video S5B); these results were not observed in the si-NC-positive cells (Figure 5B, Video S5A). After PMV treatment, the max ratio of FRET/ECFP was 1.2341 ± 0.0544 and the frequency of calcium peaks was 0.0125 ± 0.0307 in the si-1-Cy3-positive cells, indicating that these effects were all significantly abolished (Figure 5D-E); these results were not observed in the si-NC-positive cells (the max ratio was 2.0141 ± 0.1464 and the frequency was 0.3733 ± 0.06806). VSMC migration was also significantly suppressed by TRPV4 si-1 transfection (Figure 5F-G), and similar results were observed after incubation with the specific inhibitor GSK219.

Here, we validated that TRPV4 in VSMCs is essential for eliciting the calcium oscillations and VSMC migration triggered by PMVs.

Discussion

PMVs are submicroscopic (∼<1000 nm) membrane vesicles released by platelets during activation and selectively carry different molecules, including GP IIb/IIIa, GP Ib, P-selectin, and CXCR4 [45]. Low concentrations of PMVs are observed in normal circulation. Highly increased concentrations of PMVs may be an important indicator of thrombosis formation [18,24], atherosclerosis, hypertension and cardiopulmonary bypass [24]. PMVs have been studied in the pathogenesis of many diseases. Kim et al. reported that in vitro, PMVs promote human umbilical vein endothelial cell (HUVEC) survival and stimulate migration and elicit tube-like structure formation in angiogenesis [46]. PMVs stimulate
p42/p44 MAP kinase phosphorylation, cellular oncogene Fos (c-Fos) induction and DNA synthesis in a concentration-dependent manner to promote the proliferation and migration of coronary artery smooth muscle cells under conditions of thrombosis formation [47]. However, the molecular mechanisms underlying these functions of PMVs are still largely unknown.

**Figure 4.** The PMV-induced calcium oscillations in VSMCs and wound closure are abolished by the TRPV4 antagonist GSK219 but not by Nife. (A) The color images represent the time-lapse FRET images of the changes in cytoplasmic calcium upon PMV treatment of VSMCs pretreated with GSK219 (middle), Nife (bottom) and the DMSO control (top). The hot and cold colors represent high and low FRET ratios, indicating high and low levels of cytoplasmic calcium change, respectively. Scale bar: 30 μm. (B) The time courses represent the normalized FRET/ECFP ratio averaged over the cell body in VSMCs pretreated with GSK219 (n = 19, blue), Nife (n = 8, red) and DMSO (n = 12, orange) after PMV treatment, and all shadowed areas indicate the S.E.M. Comparison of the max normalized FRET/ECFP ratio (C) and frequency of cytoplasmic calcium oscillations (D) among VSMCs pretreated with GSK219 (n = 19, blue), Nife (n = 8, red) and DMSO control (n = 12, orange) after PMV treatment. (E) The migration of VSMCs pretreated with GSK219 (middle), Nife (bottom) and DMSO control (top) after PMV treatment. Scale bar: 200 μm. (F) The histogram shows the fold change in the level of GSK219- and Nife-pretreated VSMC migration relative to the that of DMSO control-pretreated VSMC migration. The values are shown as the mean ± S.E.M. for each condition (n = 6). * P < 0.05, ** P < 0.01. (G) The protein level of TRPV4 was increased after PMV treatment for 24 h. *** P < 0.001 vs. DMEM control (n = 5). (H) Immunofluorescence staining of TRPV4 (red) and nuclei (blue) of VSMCs after treatment with PMVs (top panel) or the DMEM control (bottom panel) for 5 min. Arrows indicate typical regions of dense and punctate TRPV4 staining. Scale bar: 40 μm. (I) TRPV4 inhibitor GSK219 attenuated neointimal hyperplasia in injured carotid arteries on Day 28 after the surgery. Scale bar: 50 μm.
Figure 5. TRPV4 siRNA abolished calcium oscillations and VSMC wound closure. (A) Western blot results indicate the interference efficiencies of 3 TRPV4 siRNA sequences. (B) Time-lapse FRET images of the changes in cytoplasmic calcium in VSMCs transfected with TRPV4 si-1-Cy3 or si-NC during the PMV treatment process. The hot and cold colors represent high and low FRET ratios, indicating high and low levels of cytoplasmic calcium change, respectively. Scale bar: 30 \( \mu m \). (C) The time courses represent the normalized FRET/ECFP ratio averaged over the cell body in VSMCs transfected with TRPV4 si-1-Cy3 (n = 16) or si-NC (n = 13) after PMV treatment, and all the shadowed areas indicate the S.E.M. Comparison of the max normalized FRET/ECFP ratio (D) and frequency of the cytoplasmic calcium oscillations (E) between the VSMCs transfected with TRPV4 si-1-Cy3 (n = 16) or si-NC control (n = 13) after PMV treatment. (F) The migration of VSMCs transfected with TRPV4 si-1 or si-NC control after 24 h of PMV treatment. Scale bar: 200 \( \mu m \). (G) The histogram shows the fold change in the level of TRPV4 si-1-transfected VSMC migration relative to that of the si-NC control-transfected VSMC migration. The values are shown as the mean ± S.E.M. for each condition (n = 6). ** \( P < 0.01 \), **** \( P < 0.0001 \).

It has been reported that platelets, which immediately accumulate at the sites of injury once the intact endothelium is damaged during vascular injury, become activated and release PMVs [48]. Our present research revealed that PMVs promote the migration of VSMCs, which may participate in neointimal hyperplasia during vascular injury, and calcium is the crucial molecule underlying this process.

Calcium is the simplest and most versatile second messenger involved in regulating various cellular functions, both under physiological and pathological conditions [49-51]. The calcium levels in the cytoplasm are usually low and become significantly increased when cells respond to stimulation, such as stimulation by growth factors and mechanical factors. In the present study, the increased calcium levels have been demonstrated as calcium influx from the extracellular environment. Increased calcium levels lead to further calcium binding to calmodulin to form a complex that activates downstream pathways, such as
phosphatidylinositol 3-kinase (PI3K), calcium-dependent protein kinases II (CaMKII), and myosin light chain kinase (MLCK) [38]. As a downstream molecule of calcium, CaMKII promotes VSMC migration during vascular injury through the posttranscriptional regulation of MMP9 [11,52]. Increased calcium activates PI3K/threonine-specific protein kinase (Akt) signaling via calmodulin in different cell lines [53,54]. In addition, MLCK causes changes in the fluorescence emission and the polarization excitation spectra of the enzyme by binding to calmodulin [55]. Due to its multifaceted roles, calcium signaling is the crucial coordinator of cell migration, this regulation occurs partly through local calcium pulses activating MLCK and modulating nascent focal adhesions [56]. In regulating the persistence of cell movement, calcium levels are highest at the rear and lowest at the leading edge of cells [57]. Previous literatures have demonstrated that extracellular calcium influx can activate small GTPases Cdc42 [58], Rac1 [59] and RhoA [60], and thus promote cell migration. Here we found that the protein levels of all these three molecules were significantly increased by the PMVs (Figure S5), which suggested that extracellular calcium influx induced by PMVs may regulate VSMC migration via small GTPases.

The concentration and real-time distribution of calcium can be visualized at the molecular level in single live cells. In our present study, a calcium biosensor based on FRET was used to detect the calcium dynamics induced by PMVs. The genetically encoded calcium biosensor based on FRET consists of a pair of fluorescent indicators [enhanced cyan fluorescent protein (ECFP) and a yield YFP for energy transfer (YPet)] and calmodulin-linked light chain protein kinase M13, and this biosensor has been shown to be a useful tool to characterize intracellular cytoplasmic calcium in live cells [63]. Upon increased binding of free calcium to calmodulin-linked light chain protein kinase M13, this biosensor has been used to a useful tool to characterize intracellular calcium dynamics in live cells.

VSMCs express a variety of ion channels in their cell membranes that mediate calcium influx in response to many environmental stimuli [64]. However, the identity of plasma membrane-associated calcium permeable pathways has not been reported in PMV-stimulated VSMCs. Among the calcium ion channels on VSMCs, the TRPV4 channels play crucial roles in regulating cellular functions [65,66]. TRPV4 channels are calcium-permeable nonselective cation channels on the cell membrane that are widely expressed in the cardiovascular system, including on endothelial cells, cardiac fibroblasts, and VSMCs [67]. In the present study, TRPV4 were indicated abundantly expressing in the neointimal hyperplasia (Figure S6). Hu et al. revealed that TRPV4 channels mediate the FSS-induced calcium influx and osteogenic differentiation of MSCs, and these effects were inhibited by the selective TRPV4 inhibitor HC-067047 and TRPV4-specific siRNA [32]. TRPV4 is required for the transforming growth factor-β (TGF-β)-induced differentiation of cardiac fibroblasts into myofibroblasts, which is critically involved in cardiac remodeling [68]. Activation of the TRPV4 channel at the plasma membrane appears to reflect the activation of existing channel structures with conformational changes.

Figure 6. Schematic drawing of the role of the calcium oscillations induced by PMVs in VSMC migration. Collagen-induced PMVs targeted TRPV4 and induced VSMC migration.
within the homotetrameric structure that lead to channel opening [69]. Cao et al. found that protein kinase A-mediated Ser-824 phosphorylation is required for TRPV4 activation in endothelial cells and other systems [29]. Note that the 2-APB investigated in the present study (Figure 3) is a typical inhibitor of IP3R that blocks intracellular calcium release from the endoplasmic reticulum [36,63] and thus suppresses cell migration. The FRET results indicate that 2-APB does not significantly affect the PMV-induced calcium increase, which further demonstrates that the calcium oscillation in the present study mainly occurs due to the influx of extracellular calcium. To reveal the mechanism of the effect of PMVs on TRPV4, we first conducted bioinformatics analysis with IPA software (Supplement Figure S7), and the results suggested that AGTR1, P2RY1 and Pka are the most important regulatory proteins upstream of TRPV4. Based on the 457 proteins previously reported to be expressed in PMVs by Dean et al. [27], the bioinformatics analysis suggested that 31 potential molecules in PMVs may participate in calcium regulation (Supplement Table 1). We further analyzed the connections between the three regulatory proteins upstream of TRPV4 (AGTR1, P2RY1 and Pka) and molecules from PMVs. As shown in Supplement Figure S7, the results identified the following possible pathways: 1) TGFβ1 in PMVs affected TRPV4 by regulating P2RY1 [70], and 2) CCL5 and CD36 in PMVs affected TRPV4 by regulating Pka [71,72]. The effects of microvesicles from endothelial cells and VSMCs on calcium oscillations and TRPV4 channel activation in VSMCs were examined with FRET, and the results are presented in Figure S8-S10. These results suggested that the regulatory mechanism of PMVs described above could not be the same in endothelial cell microvesicles or VSMC microvesicles. However, it still requires further studies to determine the specific stimulating composition in PMVs in order to fully understand the regulating mechanisms.

In conclusion, our findings identified the role of PMVs, specifically those induced by collagen exposure during vascular injury, in increasing VSMC migration (Figure 6). PMVs trigger the influx of extracellular calcium via the TRPV4 channel, which subsequently induces calcium oscillations and promotes VSMC migration. The study may provide new insights into the mechanism of abnormal VSMC migration after intimal injury and may have potential clinical applications for attenuating neointimal hyperplasia after intimal injury during vascular stent surgery. Further studies are encouraged to reveal the influences and mechanisms of antiplatelet drugs, such as aspirin [73-75] and clopidogrel [76], in PMV-induced calcium oscillations and in vivo regulatory functions.

**Abbreviation**

2-APB, 2-aminoethoxydiphenyl borate; α-SMA, α-smooth muscle actin; CXC4, chemokine receptor-4; DMEM, Dulbecco’s modified Eagle’s medium; ECs, endothelial cells; EOCs, early outgrowth cells; FRET, fluorescence resonance energy transfer; IP₃, inositol 1,4,5-trisphosphate; IP₃Rs, inositol 1,4,5-trisphosphate receptors; L-VDCC, L-type voltage-dependent calcium channel; MMP, metalloproteinase; siRNA, small interfering RNAs; PMVs, platelet-derived microvesicles; TRPV4, transient receptor potential vanilloid 4; VSMCs, vascular smooth muscle cells.

**Supplementary Material**

Supplementary figures and tables. http://www.thno.org/v11p2410s1.pdf
Supplementary video 1A. http://www.thno.org/v11p2410s2.wmv
Supplementary video 1B. http://www.thno.org/v11p2410s3.wmv
Supplementary video 2A. http://www.thno.org/v11p2410s4.avi
Supplementary video 2B. http://www.thno.org/v11p2410s5.avi
Supplementary video 3A. http://www.thno.org/v11p2410s6.avi
Supplementary video 3B. http://www.thno.org/v11p2410s7.avi
Supplementary video 3C. http://www.thno.org/v11p2410s8.avi
Supplementary video 4A. http://www.thno.org/v11p2410s9.avi
Supplementary video 4B. http://www.thno.org/v11p2410s10.avi
Supplementary video 4C. http://www.thno.org/v11p2410s11.avi
Supplementary video 5A. http://www.thno.org/v11p2410s12.avi
Supplementary video 5B. http://www.thno.org/v11p2410s13.avi

**Acknowledgments**

This research was supported by grants from the National Natural Science Foundation of China, nos. 11625209, 31700816 and 11222223.

**Competing Interests**

The authors have declared that no competing interest exists.
References

1. Wu B, Mottola G, Schaller M, Upchurch GR, Conte MS. Resolution of vascular injury: specialized lipid mediators and their evolving therapeutic implications. Circ Res. 2017; 121(15): 1873-1887.

2. Cai X. Regulation of smooth muscle cells in development and vascular disease.current therapeutic strategies. Expert Rev Cardiovasc Ther. 2006; 4(6): 799-809.

3. Tsaousi A, Williams H, Lyon CA, Taylor V, Swan A, Johnsen JL, et al. Wall shear stress-induced up-regulation of VSMC proliferation and is associated with intimal thickening. Circ Res. 2011; 108(4): 427-436.

4. Yao Y, Hu Z, Ye J, Hu C, Song Q, Da X, et al. Targeting AGGFI (angiogenic factor with G patc h and FHA domains 1) for blocking neointimal formation after vascular injury. J Am Heart Assoc. 2016; 6(6): e005889.

5. Hara T, Fukuda D, Tanaka K, Higashikuni Y, Hirata Y, Sagi S. Inhibition of activated factor X by rivaroxaban attenuates neointima formation after wire-mediated vascular injury. Eur J Clin Pharmacol. 2018; 82(6): 222-228.

6. Wu XL, Liu WW, Jiang H, Chen J, Wang JC, Zhu R, et al. Kindlin-2 siRNA miltiorrhiza-derived miRNAs suppress vascular remodeling through inhibition of VSMC proliferation and matrix metalloproteinase-2 expression. Mol Aspects Med. 2017; 58: 72-82.

7. Heath DE, Kang GC, Yeo C, Yin FP, Chan-Park MB. Biomaterials patterned with discontinuous microwalls for vascular smooth muscle cell culture: bioresorbable wall-less dialysis filters and stable cell culture culture. J Biomater Sci Polym Ed. 2016; 27(15): 139-159.

8. Li P, Li Y, Bi W, Wang G, You X, Zhao X, et al. MicroRNA-638 is highly expressed in human umbilical cord blood smooth muscle cells and inhibits PDK4-BH3-induced cell proliferation and migration through targeting orphan nuclear receptor N OR1. Cardiovasc Diabetol. 2013; 99(1): 185-193.

9. Bootman MD, Collins TJ, Mackenzie L, Roderick HL, Berridge MJ, Peppiatt CM. 2-aminoethoxydiphenyl borate (2-APB) is a reliable blocker of store-operated Ca2+ entry in a non-inhibitory isoform of InsP3-induced Ca2+ release. FASEB J. 2002; 7(6): 429-439.

10. Reid K, Guo TZ, Davies MF, Maze M, Nefedipine, an L-type calcium channel blocker, restores the hypertonic response in rats made tolerant to the alpha2-adrenergic agonist dexmedetomidine. J Pharmacol Exp Ther. 1997; 83(3): 903-999.

11. Cheng M, Bao W, Behm DJ, Brooks CA, Bury MJ, Dowdell SE, et al. Advanced transfection with Lipofectamine 2000 reagent: primary neurons, siRNA, and high-throughput applications. Methods. 2004; 32(3): 103-108.

12. Badimon L, Suades R, Fuentes E, Palomo I, Padro T. Role of platelet-derived microparticles as crosstalk mediators in atherothrombosis and future pharmacological targets: a review. Thromb Res. 2017; 157: 293-303.

13. Allahverdian S, Chaabane C, Boukais K, Francis GA. Smooth muscle cell fate: plasticity and plasticity in atherosclerosis. Cardiovasc Res. 2018; 114(4): 540-550.

14. Dalby B, Cates S, Harris A, Ohki EC, Tilkins ML, Price PJ, et al. Advanced transfection with Lipofectamine 2000 reagent: primary neurons, siRNA, and high-throughput applications. Methods. 2004; 32(3): 103-108.

15. Horstman LL, Ahn YS. Platelet microparticles: a wide-angle perspective. Crit Rev Oncol Hematol. 1999; 30(2): 111-142.

16. Kim HK, Song KS, Chung JL, Lee KR, Lee SN. Platelet microparticles induce microvascular angiogenesis in vitro. Br J Pharmacol. 2004; 142(3): 376-384.

17. Weber AA, Koppen HO, Schror K. Platelet-derived microparticles stimulate coronary artery smooth muscle cell mitogenesis by a PDGF-independent mechanism. Thromb Res. 2000; 98(5): 461-466.

18. Wu B, Mottola G, Schaller M, Upchurch GR, Conte MS. Resolution of vascular injury: Specialized lipid mediators and their evolving therapeutic implications. Mol Aspects Med. 2017; 58: 72-82.

19. Zhu M, Chen L, Zhao P, et al. Store-operated Ca2+ entry regulates glomma cell migration and invasion via modulation of Pyk2 phosphorylation. J Exp Clin Cancer Res. 2014; 33(1): 98.

20. Iamsukhonova O, Pla AF, Prevaskaya N. Molecular mechanisms of tumour invasion: regulation by calcium signals. J Physiol. 2017; 595(10): 3063-3075.

21. Liou J, Kim ML, Heo WD, Jones JT, Myers JW, James EF, et al. STIM is a Ca2+/calmodulin-dependent kinase II regulates vascular smooth muscle cell migration through a calcium store-depletion-induced Ca2+ influx. Curr Biol. 2005; 15(13): 1235-1241.

22. Scott JA, Xie LT, Li H, Li WW, He JB, Sanders PN, et al. The multifunctional Ca2+/calmodulin-dependent kinase II regulates vascular smooth muscle migration through matrix but an independent role of Ca2+ signaling. Am J Physiol Heart Circ Physiol. 2012; 302(1): H1253-H1264.

23. Pene F, Calessens Y-E, Muller O, Vigyne F, Mayeux D, Frydels F, et al. The role of the phosphodiesterase 5A gene in the development and apoptosis in multiple myeloma. Oncogene. 2002; 21(43): 6587-6597.

24. Xu J, Zhang QG, Li C, Zhang GY. Mitochondrial-mediated PI(3,5)K/Akt pathway activation. Hippocampus. 2007; 17: 525-537.
Malencik DA, Anderson SR, Bohnert JL, Shalitin Y. Functional interactions between smooth muscle myosin light chain kinase and calmodulin. Biochemistry. 1982; 21(17): 4031-4039.

Wei C, Wang X, Chen M, Ouyang K, Song LS, Cheng H. Calcium flickers steer cell migration. Nature. 2009; 457(7231): 901-905.

Kholmanskih SS, Koeller HB, Wynshaw-Boris A, Gomez T, Letourneau PC, Ross ME. Calcium-dependent interaction of Lis1 with IQGAP1 and Cdc42 promotes neuronal motility. Nat Neurosci. 2006; 9 (1):50-57.

Lam JGT, Vadia S, Pathak-Sharma S, McLaughlin E, Zhang X, Swanson J, et al. Host cell perforation by listeriolysin O (LLO) activates a Ca2+-dependent cPKC/Rac1/Arp2/3 signaling pathway that promotes Listeria monocytogenes internalization independently of membrane resealing. MBio. 2018; 29:270-284.

Masiero L, Lapidos KA, Ambukar I, Kohn EC. Regulation of the RhoA pathway in human endothelial cell spreading on type IV collagen: role of calcium influx. J Cell Sci. 1999; 112:3205-3213.

Giannone G, Dubin-Thaler BJ, Rossier O, Cai Y, Chaga O, Jiang G, et al. Lamellipodial actin mechanically links myosin activity with adhesion-site formation. Cell. 2007; 128(3): 561-575.

Zhu LJ, Kuchta PJ, Scott JA, Xie L, Luczak ED, Dibbern ME, et al. Oxidative activation of the Ca2+/calmodulin-dependent protein kinase II (CaMKII) regulates vascular smooth muscle migration and apoptosis. Vase Pharmacol. 2014; 60(2): 75-83.

Kim TJ, Joo C, Seong J, Vafabakhsh R, Berthon EL, Beron MW, et al. Distinct mechanisms regulating mechanical force-induced Ca2+ signals at the plasma membrane and the ER in human MSCs. Elife. 2015; 4:e04876.

Wu S, Jian MY, Xu YC, Zhou C, Almeidi AB, Liedtke W, et al. Ca2+ entry via alpha1A and TRPV4 channels differentially regulates surface expression of P-selectin and barrier integrity in pulmonary capillary endothelium. Am J Physiol. Lung C. 2009; 297(4): L650-L657.

Montell C. The TRP superfamily of cation channels. Sci. STKE. 2005; (272): re3-re3.

Pedersen SF, Owsianik G, Nilius B. TRP channels: an overview. Cell Calcium. 2005; 38(3-4): 233-252.

Randhawa PK, Jaggi AS. TRPV4 channels: physiological and pathological role in cardiovascular system. Basic Res Cardiol. 2015; 110(6): 54.

Adolph JK, Thoppil RJ, Luther DJ, Panachur SJ, Meszaros KE, Chilian WM, et al. TRPV4 channels mediate cardiac fibroblast differentiation by integrating mechanical and soluble signals. J Mol Cell Cardiol. 2013; 54: 45-52.

Jin M, Wu Z, Chen L, Jaimes J, Collins D, Walters ET, et al. Determinants of TRPV4 activity following selective activation by small molecule agonist GS15106790A. PLoS One. 2011; 6(2): e16713.

Rajasekar P, Poole DP, Liedtke W, Burnett NW, Veldhuis NA. P2Y1 receptor activation of the TRPV4 ion channel enhances purinergic signaling in satellite glial cells. J Bio Chem. 2015; 290(48): 29051-29062.

Pais R, Zietek T, Hauner H, Daniel H, Skurk T. RANTES (CCL5) reduces glucose-dependent secretion of glucagon-like peptides 1 and 2 and impairs glucose-induced insulin secretion in mice. Am J Physiol-Gastr L. 2014; 307(3): G330-G337.

Zhou D, Samovski D, Okunade AL, Stahl PD, Su X. CD36 level and trafficking are determinants of lipolysis in adipocytes. FASEB J. 2012; 26(11):4733-4742.

Ho KJ, Spieze M, Owens CD, Lancerio H, Kroemer AH, Pande R, et al. Aspirin-triggered lipoxin and resolin E1 attenuates PDGF-induced vascular smooth muscle cell migration via the cyclic adenosine monophosphate/protein kinase A (cAMP/PKA) pathway. PLoS One. 2017; 12(3): e0174936.

Petrini MH, Laguna-Fernandez A, Tseng CN, Hedin U, Ferretti M, Back M. Aspirin-triggered 15-epi-lipoxin A4 signals through FPRI2/ALX in vascular smooth muscle cells and protects against intimal hyperplasia after carotid ligation. Int J Cardiol. 2015; 179: 370-372.

Niu X, Pi SL, Baral S, Xia YP, He QW, Li YN, et al. P2Y12 promotes migration of vascular smooth muscle cells through cofilin dephosphorylation during atherogenesis. Arterioscl Throm Vas. 2017; 37(3): 515-524.